MILK PROTEINS: FROM EXPRESSION TO FOOD
Food Science and Technology
International Series

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Preface

Proteins are vital ingredients for the food industry because they provide all the essential amino acids needed for human health combined with a wide range of dynamic functional properties, such as the capacity to form network structures and stabilize emulsions and foams. The proteins of milk have excellent functional properties and nutritional value, and some have distinctive physiological properties, which are widely exploited in the food industry. Milk proteins have been the subject of intensive research during the last 50 years in an effort to unravel their molecular structures and interactions, relationship between structure and functional attributes, interactions of proteins during processing and, more recently, their physiological functions.

Recent studies on the interactions of milk proteins in complex food systems are leading to a new understanding of the nature of these interactions and their impact on food quality. The knowledge has resulted in the development of several specialized milk protein ingredients tailored to meet specific needs of the food industry. Currently, there is a growing demand by the food industry for milk protein ingredients for specialist high-value applications such as functional foods. In the future, application of novel analytical approaches (genomics, proteomics, nanotechnology) to milk proteins and food materials will provide further understanding of molecular structures and interactions to enable the dairy industry to produce highly functional and healthy protein ingredients for specific applications.

Several books have been published about milk and milk proteins—so why another one? Most of the earlier books have addressed different specialist aspects of dairy science and technology. The primary theme of this book is to present a view along the dairy food chain—starting at the cow (and its mammalian relatives) and finishing with nutritional aspects affecting the consumer, dipping into important current research topics along the way. The molecular structures and interactions of milk proteins under various processing environments are covered most prominently. More importantly, the book also contains a considerable amount of material from dairy industry-based or industry-funded research. Thus, it provides fresh perspectives on milk proteins, from an advanced dairy industry point of view.
The editors particularly thank Fonterra Cooperative Limited for making available time for staff members to contribute their chapters, and for making available hitherto unpublished material. This book is designed to provide an update and call for attention, for industry and academic researchers alike, to important and relevant milk protein science in areas that have the potential to advance the dairy industry.

The overall theme covered in this book was piloted at a meeting organized by the Riddet Center and Fonterra Cooperative Limited in February 2006, with invited presentations from a number of experts in the relevant fields from Australasia, the USA and the UK. This meeting was particularly successful, with a large number of international delegates attending from a broad range of disciplines. This confirmed the growing interest of milk protein scientists in looking beyond the boundaries of their immediate topic area to gain an understanding of how the whole food chain fits together. Such an understanding can help elucidate mechanisms and processes, identify novel research opportunities, and provide additional applications for new developments.

This book includes chapters covering many of the topics addressed at the meeting, as well as some new subjects that we felt were important in order to provide a more complete picture of the journey from expression to food. We would like to thank both the contributors who have been involved from the meeting in 2006 and those who have come on board more recently.

We have chosen to start the book with a comprehensive overview of the biology and chemistry of milk, to set the stage and give a broad underpinning of the later chapters for readers not familiar with this field. Attention is then turned to the biology, and particularly the molecular biology, of lactation, looking first at some “extreme” mammals—the tammar wallaby, which can express two different milk compositions at the same time, and the fur seal, which produces an extremely concentrated milk—to give an idea of the range of biology of milk production. The book thereafter focuses on bovine milk, with mention of the milk from other domestic species as appropriate. This starts with an update on the genomics of bovine milk proteins, and is followed by an overview on post-translational modifications, which completes our view of the biology of milk protein production.

The structural chemistry of milk proteins, including the latest model of casein micelle and molecular structures of whey proteins, is covered in detail. The behavior of milk proteins under a variety of processing regimes, including ultra high pressure, functional systems, drying and storage of powders, is dealt with in a series of chapters. These chapters address our current state of knowledge about existing and emerging processes for the production of milk protein-based food ingredients.

Attention is then turned to the behaviors of milk proteins in real and model food systems, and finally to consumer aspects—the sensory and nutritional/functional food aspects of milk proteins. A wrap-up chapter gives a view on likely issues of future importance for milk proteins, including the emerging area of nutrigenomics.

As with any volume written by a large number of contributors, this contains a variety of styles of presentation. We have made no attempt to homogenize the authors’ styles, but have provided guidance on chapter content to make for best possible continuity.
A volume of this kind always requires a large amount of work by a large number of people. We would like particularly to thank all the contributing authors for their efforts and their expeditious preparation of manuscripts that allowed for the timely publication of this book. We are pleased to acknowledge Claire Woodhall for assisting with the technical editing, and the staff at Elsevier for producing this book.

*Abby Thompson*

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Milk is the characterizing secretion of mammals, of which there are about 4500 species, produced to meet the complete nutritional requirements of the neonate of the species, as well as some defensive and other physiological requirements. The milks of all species are basically similar but there are very significant species-specific differences. In addition to supplying all the nutritional requirements of the neonate, many of the micro-constituents of milk, such as oligosaccharides, immunoglobulins, metal-binding proteins and enzymes, serve protective roles. Milk is an aqueous solution of lactose, inorganic and organic salts, and numerous compounds at trace levels (milk serum), in which are dispersed colloidal particles of three size ranges: whey proteins dissolved at the molecular level, the caseins dispersed as large (50–500 nm) colloidal aggregates (micelles), and lipids emulsified as large (1–20 μm) globules.

The colloidal stability of milk, especially of the casein micelles, is very important from the nutritional and technological viewpoints. The micelles are destabilized and aggregate or gel following limited proteolysis or acidification to ≈pH 4.6. In vivo, aggregation occurs in the stomach of the neonate, thereby slowing transit and improving digestibility. Technologically, destabilization of the micelles can be undesirable or can be exploited in the production of cheese and fermented milk products.

Man has used milk in his diet for about 8000 years and a major industry has developed around the processing of milk of a few species for human foods, especially milk from cattle, buffalo, sheep and goats. Milk processing, which exploits certain physico-chemical properties...
of milk, is practiced worldwide, especially in Europe and North America. Milk is a very flexible raw material, from which a very wide range of different products is produced, including about 1400 varieties of cheese.

In this introductory chapter, the chemical and biological characteristics of milk constituents and the physico-chemical and technological properties of milk will be summarized.

Introduction

Milk is a fluid secreted by the female of all mammalian species, of which there are about 4500 extant species (about 80% of mammalian species are extinct), primarily to meet the complete nutritional requirements of the neonate. The principal requirements are for energy (supplied by lipids and lactose and, when in excess, by proteins), essential amino acids and amino groups for the biosynthesis of non-essential amino acids (supplied by proteins), essential fatty acids, vitamins, inorganic elements and water. Because the nutritional requirements of the neonate depend on its maturity at birth, its growth rate and its energy requirements, which depend mainly on environmental temperature, the gross composition of milk shows large inter-species differences, which reflect these requirements (see Fox and McSweeney, 1998). Milk also serves a number of physiological functions, which are performed mainly by proteins and peptides, including immunoglobulins, enzymes, enzyme inhibitors, growth factors, hormones and antibacterial agents.

Of the 4500 species of mammal, the milk of only about 180 species has been analyzed, and of these the data for only about 50 species are considered to be reliable (i.e. a sufficient number of samples, samples taken properly, representative sampling, adequate coverage of the lactation period). Milk from the commercially important species—cow, goat, sheep, buffalo, yak, horse and pig—are quite well characterized; human milk is also well characterized, as is that of experimental laboratory animals, especially rats and mice. Reviews on non-bovine milks include: general (Evans, 1959; Jenness and Sloan, 1970), buffalo (Laxminarayana and Dastur, 1968), goat (Parkash and Jenness, 1968; Haenlein, 1980; Jenness, 1980), sheep (Bencini and Pulina, 1997), sheep and goats (International Dairy Federation, 1996; Jandal, 1996; Park et al., 2007), camel (Rao et al., 1970; Farah, 1993), horse (Doreau and Boulot, 1989; Solaroli et al., 1993; Doreau, 1994; Park et al., 2006), human (Atkinson and Lonnerdal, 1989; Jensen, 1989, 1995) and sow (Verstegen et al., 1998). The Handbook edited by Park and Haenlein (2006) is a particularly useful source of information on the milk of non-bovine mammals; it includes chapters on goat, sheep, buffalo, mare, camel, yak, reindeer, sow, llama, minor species (moose, musk ox, caribou, alpaca, ass, elk, seal, sea lion and polar bear) and human. Inter-species comparisons of more specific aspects are cited in the appropriate sections.

The milks of certain domesticated animals, and dairy products therefrom, are major components of the human diet in many parts of the world. Domesticated goats, sheep and cattle have been used for milk production since about 8000 BC. Recorded milk production today is about $600 \times 10^6$ tonnes per annum, about 85% of which is bovine, 11% is buffalo and about 2% each is ovine and caprine, with small amounts produced from horses, donkeys, camels, yaks and reindeer. Milk and dairy products
are consumed throughout the world but are particularly important in Europe, the USA, Canada, Argentina, India, Australia and New Zealand. The contribution of milk and dairy products to dietary intake varies widely for different regions of the world, e.g. the kilocalories per day supplied by milk range from 12 in China to 436 in Ireland; in the UK, milk and dairy products supply \( \approx 30\% \) of dietary protein consumed by young children, \( \approx 27\% \) of dietary lipids and \( \approx 65\% \) of calcium (Barker, 2003).

The chemistry and the physico-chemical properties of milk have been studied for about 200 years and are now understood in considerable detail, and described in a voluminous literature. The objectives of this chapter are to provide a summary and overview of the evolution of mammals and lactation and of the principal constituents of milk, especially proteins, which are the subject of this book. Where possible, inter-species comparisons of milk and its constituents are made. Numerous textbooks and review articles are cited and may be consulted for the primary literature.

**Evolution of mammals and lactation**

The secretion of milk is one of the characterizing features of mammals, which evolved from egg-laying, pre-mammalian reptiles, synapsids and cynodonts. Cynodonts are believed to be the ancestors of all mammals and evolved \( \approx 200 \) million years ago (at the end of the Triassic Period). The word “mammal” is derived from “mamma,” which is Latin for breast. Initially, mammals were small shrew-like creatures but they have evolved and diversified to occupy all niches on land, sea and air. They range in size from a few grams (pigmy shrew) to 200 tonnes (blue whale). Their dominance occurred especially after the extinction of the dinosaurs, 60–70 million years ago, at the interface between the Cretaceous and Tertiary Periods (C/T interface). Mammals have been successful: the young of most species are born alive (viviparous) and all are supplied with a specially designed food—milk—for the critical period after birth. No other class of animal is so pampered (for an interesting discussion on this point, see Peaker, 2002). Not surprisingly, the evolution of mammals is a popular subject; reviews include: Crampton and Jenkins (1973), Kemp (1982), Lillegraven et al. (1987), Benton (1999), Easteal (1999), Forsyth (2003), Lillegraven (2004), and Springer et al. (2004).

Mammals are distinguished from other classes of animals by four criteria:

- they secrete milk to nourish their young;
- they are endothermic, i.e. they can control their body temperature;
- they grow body hair or wool for insulation—even aquatic mammals have some hair; and
- they have different types of teeth (flat incisors, conical canines and multi-cusped molars), which allow them to masticate different types of food.

The class *Mammalia* contains two subclasses—*Prototheria* and *Theria* (young born alive).

*Prototheria*. These egg-laying mammals, known as monotremes because they have only one opening for the elimination of waste, mating and egg laying, were the first
mammals, only five species of which survive, i.e. the duck-billed platypus and four species of echidna (also called spiny anteater), which are found only in Australia and New Guinea. Presumably, there were other species of monotreme, which have become extinct.

**Theria.** About 90 million years ago, the *Theria* split into two infra-classes, *Metatheria* and *Eutheria*. However, the fossil of a eutherian mammal, believed to be ≈125 million years old, was discovered recently in north-eastern China; it was named *Eomonia scansoria*, meaning “earliest eutherian mammal with specialized features for climbing” (Ji *et al.*, 2002).

*Metatheria.* Usually called marsupials, of which there are about 330 species. The young are born alive (*viviparous*) but very immature and develop in an abdominal pouch (*marse* = pouch, purse). Marsupials survive mainly in Australia and surrounding islands (>200 species), with several species in South America and one species, the Virginia opossum, in North America; there are none in Europe, Asia or Africa.

*Eutheria.* The fetus develops in utero, where it receives nourishment from the maternal blood via a highly specialized organ called the placenta (these are called placental mammals); ≈95% of all mammals are eutherians.

### Classification of mammals

There are ≈4500 species of mammal, which are classified into 20 orders (see MacDonald, 2004). It is estimated that only about 20% of the species that have evolved over the last 200 million years are still extant. The classification and nomenclature of mammals commenced with the work of Carolus Linnaeus in 1758 and was based initially on morphological characteristics (see MacDonald, 2004). More soundly based classification is now possible, based on DNA sequences (Madsen *et al.*, 2001; Murphy *et al.*, 2001; Springer *et al.*, 2004) and on the primary sequence of certain proteins, e.g. growth hormone and prolactin (Forsyth and Wallis, 2002). It should be possible to classify mammals based on the structure of milk proteins, especially caseins, which are fast-mutating proteins; some preliminary work has been reported (Goldman, 2002; Rijnkels, 2002; Simpson and Nicholas, 2002). However, the sequences of milk proteins from a sufficient number of species have not been determined to make a comprehensive classification scheme possible, but considerable progress on the structure of milk proteins is being made (see Martin *et al.*, 2003) and is discussed further later.

A minor whey protein—whey acidic protein (WAP)—has already been useful for tracing the relationships between mammalian families (Hajjoubi *et al.*, 2006). To date, WAP has been found only in the milk of platypus, echidna, tammar wallaby, opossum, mouse, rat, rabbit, camel and pig. In humans and ruminants, the gene for WAP has been frame shifted and occurs as a pseudogene that is not transcribed. The distribution of WAP suggests that the loss of a functional WAP gene occurred after the divergence of the pig and ruminant lines but before the *Bovidae* diverged from the other ruminants. Analysis of the milks from a wider range of species for WAP should be interesting.
Classification and phylogenetic relationships of the principal dairying species

All of the principal, and many of the minor, dairying species belong to the family Bovidae, a member of the order Artiodactyla (even-toed ungulates [hoofed] mammals, i.e. cloven hoofed). A few minor dairying species (horse and ass) are members of Perissodactyla (odd-toed ungulates). The Bovidae evolved ~18 million years ago; the earliest fossil attributed to the Bovidae is Eotragus, found in 18-million-year-old deposits in Pakistan. The order has three sub-orders: Ruminantia (ruminants; to which all major dairying species belong), Sunia (pigs and related species) and Tylopoda (camels, llama, alpaca and guanaco).

The Ruminantia are classified into six families: Tragulidae (chervrotains), Moschidae (musk deer), Antilocapridae (pronghorns), Giraffidae (giraffes and okapi), Cervidae (deer; 43 species in 16 genera) and Bovidae (137–38 species in 46–7 genera).

The Bovidae are divided into six subfamilies, of which the Bovinae is the most important. The Bovinae are divided into three tribes, of which the Bovini are the most important from our viewpoint. The Bovini are classified into five genera: Bubalus (water buffalo), Bos (cattle), Pseudoryx, Syncerus (African [Cape] buffalo) and Bison (American and European; the latter is also called wisent).

There are seven or eight species of Bos: B. primigenus (aurochs—the ancestor of domestic cattle; they are extinct; the last animal was killed in Poland in 1627), B. javanicus (banteng), B. gaurus (gaur), B. frontalis (gayol), B. mutus (yak), B. sauvali (kouprey), B. taurus (European cattle) and B. indicus (Indian, humped zebu cattle). (B. taurus and B. indicus may be sub-species rather than species.) The phylogenetic relationships of the Bovini have been studied by molecular biology techniques (see Finlay, 2005).

Today, there are about 1.3 × 10^9 cattle worldwide, of which there are two species—B. taurus, of European origin, and B. indicus, which originated in India. B. indicus (zebu) cattle also dominate in Africa but apparently African zebu cattle have some B. taurus genes, probably as a result of cross-breeding many centuries ago. Zebu are less efficient producers of milk or meat than B. taurus but are more resistant to heat stress and various diseases, and therefore dominate in tropical regions.

Since cattle were domesticated ~8000 years ago, they have been bred selectively, especially during the past 200 years, i.e. since Herd books have been kept. These breeding practices have selected for various characteristics, e.g. health, fertility, docility, milk or meat production, or both. Today, there are about 800 breeds of cattle, including dairy, beef or dual-purpose breeds. There are ~200 million dairy cows of many, mainly local, breeds; Holstein-Friesian is the principal breed of dairy cow, representing ~35% of the total (~70 million cows). Other important international dairy breeds are Brown Swiss (~4 million), Jersey (~2 million), Ayrshire, Guernsey and Red Dane.

There are about 160 million buffalo worldwide, of which there are two types—river and swamp—found mainly in South-East Asia, India and Egypt, with smaller numbers in Bulgaria, Italy, Brazil and Australia. Depending largely on the region, buffalo are used for milk, meat or work, or combinations of these. There are no breeds of buffalo, within the meaning applied to cattle; rather, they are named after the area from which they come.
Evolution of the mammary gland

Evolution of the mammary gland is believed to have commenced with the synapsids, ~300 million years ago. These reptiles laid membrane-shelled eggs, which lost water rapidly through evaporation and were kept moist by an aqueous oily secretion from sebaceous apocrine glands on the breast/abdomen of the mother; these were somewhat like the blood patches on the breast of birds. The secretions are believed to have contained a range of bactericidal substances, e.g. oligosaccharides, lysozyme, lactoferrin, transferrin, immunoglobulins and peroxidases, which protected the egg against microbial infection. Presumably, the secretions were licked by the neonate from the mother’s abdomen and served as a source of nutrients.

Considering their importance in the evolution of mammals, including man, the evolution of the mammary gland and the origin of lactation have attracted considerable attention. Reviews on the subject include Pond (1977), Hayssen and Blackburn (1985), Blackburn et al. (1989), Blackburn (1991, 1993), Hayssen (1993) and Oftedal (2000a, 2000b).

Structure of the mammary gland

The microscopic structure of the mammary gland of all species—monotremes, marsupials and eutherians—is basically similar. The structure of the bovine gland from the cellular level to the organ level has been described by Fox and McSweeney (1998). The cells (mammocytes), the structure of which is basically similar to that of all animal cells, are arranged as a monolayer in a pear-shaped organelle, called an alveolus. The alveoli are connected via a system of ducts to a cistern, where the milk is stored until it is expressed from the gland, usually through a teat which is sealed by a sphincter muscle. There is little de novo synthesis in the mammary gland; rather, the major constituents in milk are synthesized from molecules imported from the blood through the basal cell membrane. Within the mammocyte, mainly in the rough endoplasmic reticulum (ER), these molecules are polymerized to lactose, lipids or proteins. The mammocytes are provided with a good blood supply through an extensive system of capillaries and are surrounded by contractile myoepithelial cells which, under the control of the hormones oxytocin and prolactin, contract and express milk from the alveoli through the ducts, and eventually from the gland. The hormonal control of mammary growth and function has been described by Forsyth (1986).

Although, at the microscopic level, the mammary gland is essentially similar across species, the number and the appearance of the gland are characteristic of the species. Monotremes have many glands on the abdomen; the glands do not end in a teat and the milk is licked from the abdomen by the young. Marsupials have two or four glands, which end in a teat, within the pouch. On entering the pouch, the young attaches to a teat and remains attached during the period it spends permanently in the pouch. During this period, an older offspring may use another gland during its visits to the pouch. The two glands secrete milk of very different composition, designed for the specific requirements of the neonate, and the composition changes markedly when the offspring leaves the pouch intermittently. The mammary glands of eutherians are located
externally to the body cavity and end in a teat; their number varies from two (human, goat, sheep, horse, elephant, etc.), four (cattle), 14 (pig) to 24 (some insectivores). The glands are separate anatomically.

The external location of the mammary gland facilitates study of the biosynthesis of milk constituents by isotope dilution techniques, arterio-venus concentration differences, perfusion of the severed gland, tissue slices and cell homogenates.

**Utilization of milk**

As young mammals are born at different stages of maturity and with different nutritional requirements, the milk of each species is designed to meet the requirements of the neonate of that species, i.e. it is species specific. Milk is intended to be consumed unchanged by the young suckling its mother. However, man has consumed the milk of other species for at least 8000 years. Several species have been used for milk production but today cattle, especially *Bos taurus*, is the principal dairying species, accounting for ~85% of total milk production. The other important dairying species are buffalo (*Bubalus bubalis*) (11%), goats and sheep (2% each); other species are significant in certain regions, or for certain purposes, e.g. camel, yak, reindeer, horse and donkey.

Milk is often described as the most “nearly perfect” food; although this is true only for the young of the producing or closely related species, the milk of all species is a nutrient-rich and well-balanced food (Kon, 1959; du Puis, 2002; Patton, 2004). Many of the minor constituents of milk have biological properties, which are described in the appropriate section (see also Korhonen, 2006); these minor constituents have been attracting considerable attention recently.

However, milk is very susceptible to the growth of micro-organisms, which will cause spoilage if the milk is stored. To counteract this, man has developed a range of products that are more stable than milk; some of these date from 4000 BC and have evolved desirable epicurean characteristics, in addition to their nutritional value. Today, several thousand food products are produced from milk; these fall into the following principal groups: liquid/beverage milk (40%), cheese (35%), milk powders (15%), concentrated milks (2%), fermented milk products (2%), butter (30%; some of which is produced from cream/fat obtained as a by-product in the manufacture of other products), ice cream, infant formula, creams, protein-rich products and lactose. Some of these groups are very diverse, e.g. 1400 varieties of cheese have been listed.

**Composition of milk**

Milk is a very complex fluid containing several hundred molecular species (several thousand if all triglycerides are counted individually). The principal constituents are water, lipids, sugar (lactose) and proteins. In addition, there are numerous minor constituents, mostly at trace levels, e.g. minerals, vitamins, hormones, enzymes and miscellaneous compounds. The chemistry of these compounds is generally similar across species but in many cases their structure differs in detail, reflecting evolutionary
changes. The concentration of the principal constituents varies widely among species: lipids, 2–55%; proteins, 1–20%; lactose, 0–10%, reflecting mainly the energy requirements (lipids and lactose) and growth rate (mainly proteins) of the neonate. The concentrations of the minor constituents also vary widely.

Within any species, the composition of milk varies among individual animals, between breeds, with the stage of lactation, feed and health of the animal and with many other factors. The fat content of bovine milk shows large inter-breed differences and within any breed there is a wide range of fat and protein content for individual animals; similar differences occur in the milk of the sheep, goat and buffalo.

Reflecting mainly the nutritional and physiological requirements of the neonate, the composition of milk, and even the profile of constituents therein, changes markedly during lactation. The changes are most marked during the first few days post-partum, especially in the immunoglobulin fraction of proteins. For marsupials, the milk changes from a high-carbohydrate (mainly oligosaccharides) secretion to a high-fat secretion when the neonate begins to leave the pouch, a time that corresponds roughly to the birth of eutherians. The composition of milk remains relatively constant during mid-lactation but changes considerably in late lactation, reflecting the involution of the mammary gland tissue and the greater influx of blood constituents.

**Milk constituents**

In the following sections, the chemistry of milk carbohydrates, lipids, proteins, salts and some minor constituents are described; where possible, inter-species comparisons are made.

**Carbohydrates**

**Lactose**

The principal carbohydrate in the milk of most species is the reducing disaccharide lactose, which is composed of galactose and glucose linked by a β1–4 glycosidic bond. Its concentration varies from 0 to ~10% (Fox and McSweeney, 1998) and milk is the only known source of lactose. Research on lactose commenced with the work of Carl Scheele in about 1780; its chemistry and its important physico-chemical properties have been described very thoroughly. The very extensive literature has been reviewed by Whittier (1925, 1944), Weisberg (1954), Zadow (1984, 1992), Fox (1985, 1997), Fox and McSweeney (1998) and McSweeney and Fox (2008).

Lactose is synthesized in the epithelial mammary cells from two molecules of glucose absorbed from the blood. One molecule of glucose is phosphorylated and converted (epimerized) to galactose-P via the Leloir pathway, which is widespread in animal tissues and bacterial cells. Galactose-P is condensed with a second molecule of glucose through the action of a unique two-component enzyme—lactose synthetase. One component is UDP-galactosyl transferase (EC 2.4.1.22), which transfers galactose from UDP-galactose to any of several acceptor molecules in the biosynthesis of glycoproteins and glycolipids. The specificity of the transferase is controlled and
modified by a-lactalbumin (α-La), one of the principal milk proteins, which reduces
the Michaelis constant ($K_M$) for glucose 1000-fold and, in its presence, most of
the galactose is transferred to glucose, with the synthesis of lactose. There is a positive
correlation between the concentrations of lactose and α-La in milk; the milk of the
Californian sea lion or the hooded seal, which contains no lactose, also lacks α-La.

The synthesis of lactose draws water osmotically into the Golgi vesicles and hence
affects the volume of milk and the concentration of casein, which is packaged in the
Golgi vesicles. There is an inverse correlation between the concentrations of lactose
and casein in milk (Jenness and Holt, 1987).

Lactose serves as a ready source of energy for the neonate, providing 30% of the
calories in bovine milk and acting as an alternative to the more energy-dense lipids.
Milks with a high lactose concentration tend to have lower levels of lipids (Jenness
and Sloan, 1970).

Lactose is also responsible for about 50% of the osmotic pressure of milk, which
is isotonic with blood and hence essentially constant. For milk with a low level of
lactose, the concentration of inorganic salts is high to maintain the osmotic pressure
at the desired level; there is an inverse relationship between the concentrations of
lactose and salts (ash) in milk (Jenness and Sloan, 1970).

During mastitis or in late lactation, the integrity of the mammary cell mem-
branes is damaged and there is an influx of blood constituents into milk; the osmotic
pressure increases and, to adjust this, the concentration of lactose is reduced. This
relationship is expressed as the Koesler Number (% chloride $\times 100 \div %$ lactose),
which is normally <2; a value >3 is considered to be abnormal. Today, the Koesler
Number is rarely used as a diagnostic indicator of mastitis. The electrical conductivity
of milk is commonly used for this purpose, as it depends mainly on the milk salts
and can be measured in-line during milking.

Why milk contains lactose rather than some other sugar(s) is not clear. The pres-
ence of a disaccharide rather than a monosaccharide can be explained on the basis
that twice as much (mass) disaccharide as monosaccharide can be accommodated
for any particular incremental increase in osmotic pressure, which is fixed. Maltose,
which consists of two molecules of glucose, would seem to be the obvious choice of
disaccharide. As energy is expended in converting glucose to galactose, some benefit
must accrue from this conversion; a possible benefit is that galactose, or a derivative
thereof, occurs in some physiologically important lipids and proteins, and a galac-
tose-containing sugar in milk provides the neonate with a ready supply of this impor-
tant monosaccharide.

The properties of lactose are generally similar to those of other sugars, but lactose
differs in some technologically important respects. Some important characteristics of
lactose are as follows:

- Lactose is a reducing sugar, i.e. it has a free, or potentially free, carbonyl group
  (an aldehyde group in the case of lactose).
- Like other reducing sugars, lactose exists partially as an open-chain form with
  an aldehyde group, which can form a hemi-acetal and thus a ring structure.
  Formation of a hemi-acetal creates a new chiral center (asymmetric carbon),
which may exist as two isomers (enantiomorphs), $\alpha$ or $\beta$. By alternately opening and forming the ring structure, the molecule can interchange between the $\alpha$ and $\beta$ isomers, a process referred to as mutarotation.

- The $\alpha$ and $\beta$ isomers of lactose have very different properties, the most important of which are specific rotation, $[\alpha]_{20}^D$ ($+89^\circ$ and $+35^\circ$ for $\alpha$ and $\beta$ respectively), and solubility (70 and 500 g/L for $\alpha$ and $\beta$ respectively).

- Like all reducing sugars, lactose can participate in the Maillard (non-enzymatic) browning reaction, resulting in the production of (off-)flavour compounds and brown polymers. The Maillard reaction contributes positively to the flavor and color of many foods, e.g. the crust of bread, toast and deep-fried products, but the effects in dairy products are negative and must be avoided.

- Redox titration using alkaline CuSO$_4$ (Fehling’s solution) or chloramine-T is the principal standard method for the quantitative determination of lactose. It may also be determined by polarimetry, spectrophotometrically after reaction with phenol or anthrone in strongly acid solution, enzymatically or by high performance liquid chromatography.

- Among sugars, lactose, especially the $\alpha$ enantiomorph, has low solubility in water; however, when in solution, it is difficult to crystallize, which may cause problems in lactose-rich dairy products, e.g. skimmed milk powder and whey powder, unless precautions are taken to induce and control crystallization.

- $\alpha$ and $\beta$ lactose are soluble in water to the extent of about 70 and 500 g/L respectively at 20°C; at equilibrium, the ratio of $\alpha$ to $\beta$ is about 1:2, giving a total solubility of about 180 g/L at 20°C. The solubility of $\alpha$ lactose is more temperature dependent than that of the $\beta$ isomer and $\alpha$ lactose is the more soluble at $>94^\circ$C. Hence, $\alpha$ lactose is the form of lactose that crystallizes at $<94^\circ$C and is the usual commercial form of lactose; $\beta$ lactose may be prepared by crystallization at $>94^\circ$C.

- $\alpha$ lactose crystallizes as a monohydrate whereas $\beta$ lactose forms anhydrous crystals; thus, the yield of $\alpha$ lactose is 5% higher than that of $\beta$ lactose.

- When milk or whey is spray dried, any lactose that has not been pre-crystallized forms an amorphous glass. The amorphous glass is stable if the moisture content of the powder is maintained low but, if the moisture content increases to $>6\%$, the lactose crystallizes as the $\alpha$ hydrate, the crystals of which form interlocking masses and clumps which may render the powder unusable if this is very extensive, i.e. inadequately crystallized powder is hygroscopic. The problem can be avoided by adequate crystallization of lactose before drying or by using effective packaging.

- Interestingly, crystalline lactose has very low hygroscopicity and is used in icing sugar blends.

- Among sugars, lactose has a low level of sweetness; it is only about 16% as sweet as sucrose at 1% in solution and hence has limited value as a sweetening agent, the principal application of sugars in foods. However, it is a useful bulking agent when excessive sweetness is undesirable.

- Lactose is important in the manufacture of fermented dairy products, where it serves as a carbon source for lactic acid bacteria, which produce lactic acid.
Modification of the concentration of lactose in milk through genetic engineering

There has been considerable interest in modifying the lactose content of milk by genetic engineering. As the concentration of lactose is controlled by the concentration of \( \alpha \)-La in the secretory cells, the approach to changing the concentration of lactose involves altering the level of \( \alpha \)-La. There is interest in reducing the level of lactose for a number of reasons, not least because lactose is the least valuable constituent in milk but it costs energy on the part of the animal to synthesis it; therefore, it would be economically advantageous to reduce the lactose content of milk.

As lactose effectively controls the water content of milk, and most dairy processes require the removal of water, it would be advantageous to reduce the amount of water in milk by reducing the level of lactose. However, if the level of lactose is reduced too much, the viscosity of the milk will be too high for easy expression from the mammary gland; the viscosity of mouse milk engineered to contain no lactose was so high that the pups were unable to suckle and died. Obviously, this problem could be overcome by reducing the level of lactose rather than eliminating it. Alternatively, it may be possible to modify the milk secretory mechanism to produce a more useful, or at least a less problematic, sugar than lactose, e.g. glucose, maltose or lactulose (which is a laxative and a prebiotic). It might be possible to increase the concentration of salts in milk.

As discussed below, most adult humans are unable to digest lactose. If the problems arising from high viscosity were resolved, lactose-free or lactose-reduced milk would be nutritionally desirable. The possibility of engineering the mammary cell to secrete \( \beta \)-galactosidase into milk and to hydrolyze lactose in situ has been suggested.

In contrast, there are potentially cases where it would be advantageous to increase the lactose content of milk. The economic benefits of increasing the milk output of sows by increasing its lactose content have been discussed by Wheeler (2003).

Nutritional problems associated with lactose

Mammals cannot absorb disaccharides from the small intestine, where they are hydrolyzed to monosaccharides which are absorbed. Lactose is hydrolyzed by \( \beta \)-galactosidase (lactase) which is secreted by cells in the brush border of the small intestine. The young of most mammalian species secrete an adequate level of \( \beta \)-galactosidase but, as the animal ages, the secretion of \( \beta \)-galactosidase declines and eventually becomes inadequate to hydrolyze undigested lactose, which enters the large intestine into which it draws water, causing diarrhea, and is metabolized by bacteria with the production of gas, which causes cramps and flatulence. In humans, this may occur at 8–10 years of age. These problems cause many individuals to exclude milk from their diet. The problems may be avoided by pre-hydrolyzing the lactose using exogenous \( \beta \)-galactosidase (see Mahoney, 1997). The frequency and the intensity of lactose intolerance/malabsorption vary widely among populations from \( \approx \)100% in South-East Asia to \( \approx \)5% in north-west Europe (Mustapha et al., 1997; Ingram and Swallow, 2008).

Production and utilization of lactose

Previously, whey from cheese or casein production was considered to be a waste material that was fed to farm animals, irrigated on land or disposed of into sewers.
Environmental and economic considerations now dictate that whey be used more efficiently. The principal product lines produced from whey are various whey powders, whey protein products produced by membrane technology and lactose and its derivatives.

Lactose is prepared commercially by crystallization from concentrated whey or ultrafiltrate. The crystals are usually recovered by centrifugation; this process is essentially similar to that used for sucrose or other sugars. About 400,000 tonnes of crystalline lactose is produced annually, compared with ≈100 million tonnes of sucrose.

Because of its relatively low sweetness and low solubility, the applications of lactose are much more limited than those of sucrose or glucose. Its principal application is in the production of “humanized” infant formulas based on cow’s milk (human milk contains ≈7% lactose in comparison with ≈4.6% in bovine milk). The lactose used may be a purified crystalline product or in the form of demineralized whey (for physiological reasons, it is necessary to reduce the concentration of inorganic salts in whey).

Lactose has a number of low-volume, special applications in the food industry, e.g. as a free-flowing or agglomerating agent, to accentuate/enhance the flavor of some foods, to improve the functionality of shortenings, and as a diluent for pigments, flavors or enzymes. It is widely used in the tableting of drugs in the pharmaceutical industry, where low hygroscopicity is a critical property.

Lactose can be converted to several more valuable food-grade derivatives, of which the most significant are: glucose-galactose syrups (≈ three times as sweet as lactose; produced by hydrolysis by β-galactosidase), lactulose (galactose-fructose; a prebiotic and a laxative), lactitol (the alcohol of lactose), lactobionic acid (a sweet-tasting acid, which is a very rare property), tagatose, oligosaccharides (prebiotics) and fermentation products (ethanol, and lactic, acetic and propionic acids).

**Oligosaccharides**

In addition to lactose, a large number of other free saccharides have been found in milk, the concentration, proportions and types of which show large inter-species differences. Oligosaccharides are the most common form, but small amounts of monosaccharides are also present and some milk proteins, especially κ-casein, are glycosylated; there are low levels of highly glycosylated glycoproteins, especially mucins, and glycolipids in the milk fat globule membrane (MFGM).

Almost all of the oligosaccharides have lactose at the reducing end, contain three to eight monosaccharides, may be linear or branched and contain either or both of two unusual monosaccharides—fucose (a 6-deoxyhexose) and N-acetyleneuraminic acid. Fucose occurs quite widely in the tissues of mammals and other animals, where it serves an array of functions (Becker and Lowe, 2003). Its significance in the oligosaccharides in milk is not clear; perhaps it is to supply the neonate with preformed fucose as the concentration of oligosaccharides is higher in colostrum than in milk. General reviews on the oligosaccharides in milk include Newburg and Newbauer (1995) and Urashima et al. (2001, 2007).

The oligosaccharides are synthesized in the mammary gland, catalyzed by special transferases that transfer galactosyl, sialyl, N-acetylglucosaminyl or fucosyl residues from nucleotide sugars to the core structures. These transferases are not affected by
α-La and are probably similar to the transferases that catalyze the glycosylation of lipids and proteins.

The milk of all species examined contains oligosaccharides but the concentration varies markedly. The highest levels are in the milk of monotremes, marsupials, marine mammals, humans, elephants and bears. With the exception of humans, the milk of these species contains little or no lactose and oligosaccharides are the principal carbohydrates. The milk of the echidna contains mainly the trisaccharide fucosylactose whereas that of the platypus contains mainly the tetrasaccharide difucosyllactose. Among marsupials, the best studied is the tammar wallaby; presumably, its lactation pattern and its milk composition are typical of marsupials. A low level of lactose is produced at the start of lactation but, about 7 days after birth, a second galactosyltransferase appears and tri- to pentasaccharides are produced, which by ≈180 days are the principal saccharides; during this period, the saccharide content is high (≈50% of total solids) and the level of lipids is low (≈15% of total solids). At about 180 days, the carbohydrates decrease to a very low level and are mainly monosaccharides, whereas the level of lipids increases to ≥60% of total solids.

Human milk contains ≈130 oligosaccharides, at a total concentration of ≈15 g/L; these are considered to be important for neonatal brain development. Bear milk contains very little lactose but a high level of total sugars (probably mainly oligosaccharides)―1.7 and 28.6 g/kg respectively (Oftedal et al., 1993). Elephant milk contains ≈50 and 12 g/kg of lactose and oligosaccharides respectively a few days postpartum but, as lactation progresses, the concentration of lactose decreases whereas that of oligosaccharides increases, e.g. 12 and 18 g/kg respectively at 47 days (Osthoff et al., 2005). The milk of seals contains both lactose and oligosaccharides but the milks of the Californian sea lion, Northern fur seal and Australian fur seal contain neither, probably because they contain no α-La (Urashima et al., 2001).

Bovine, ovine, caprine and equine milks contain relatively low levels of oligosaccharides, which have been characterized (see Urashima et al., 2001). Caprine milk contains about 10 times as much oligosaccharides as bovine milk and ovine milk, and a process for their isolation by nanofiltration has been reported (Martinez-Ferez et al., 2006). Possible methods for producing oligosaccharides similar to human milk oligosaccharides, by fermentation or by transgenic animals or by recovering oligosaccharides from cow’s milk whey or ultrafiltration permeate, have been discussed by Mehra and Kelly (2006).

As discussed earlier, oligosaccharides with bactericidal properties were probably the saccharides present in the mammary secretions of early mammals, and the high level of oligosaccharides in the milk of monotremes and marsupials conforms with their secretion early in evolution. Messer and Urashima (2002) proposed that the primitive mammary glands of the first common ancestor of mammals produced lysozyme (a predecessor of α-La) and a number of glycosyltransferases but little or no α-La, which resulted in the production of a low level of lactose that was utilized in the synthesis of oligosaccharides and did not accumulate. Initially, the oligosaccharides served mainly as bactericidal agents but later became a source of energy for the neonate and both of these functions persist for monotremes, marsupials and some eutherians, e.g. bears, elephants and marine mammals.
However, most eutherians evolved to secrete predominantly lactose as an energy source, due to the synthesis of an increased level of $\alpha$-La, whereas oligosaccharides continued to play a bactericidal role. Human milk, which contains high levels of both lactose and oligosaccharides, seems to be anomalous. Work on the oligosaccharides of a wider range of species is needed to explain this situation.

The significance of oligosaccharides is not clear but the following aspects may be important. Firstly, for any particular level of energy, they have a smaller impact on osmotic pressure than smaller saccharides. Secondly, they are not hydrolyzed by $\beta$-galactosidase and neither fucosidase nor neuraminidase is secreted in the intestine; hence the oligosaccharides are not hydrolyzed and absorbed in the gastrointestinal tract and function as soluble fiber and prebiotics that affect the microflora of the large intestine. Thirdly, it is claimed that they prevent the adhesion of pathogenic bacteria in the intestine. And finally, galactose and especially N-acetylneuraminic acid are important for the synthesis of glycolipids and glycoproteins, which are important for brain development; hence, it has been suggested that the oligosaccharides are important for brain development (see Kunz and Rudloff, 2006).

**Lipids**

Lipids (commonly called oils or fats, which are liquid or solid respectively at ambient temperature) are those constituents of tissues, biological fluids or foods that are soluble in an apolar solvent, e.g. diethyl ether, chloroform or carbon tetrachloride. Historically, the fat of milk was regarded as its most valuable constituent and, until recently, milk was valued largely or totally on its fat content. This was due at least partially to the development in 1890 of rather simple methods for quantifying the fat content of milk by S. M. Babcock and N. Gerber, long before comparable methods for proteins became available. Milk lipids are very complex chemically and exist as a unique emulsion. Milk lipids have been thoroughly studied and characterized (see Fox, 1983, 1995; Fox and McSweeney, 1998, 2006; and references therein).

The level of fat in milk shows very large inter-species differences, ranging from $\approx 2\%$ to $>50\%$ (see Fox and McSweeney, 1998). Lipids are 2.5 times more energy dense than lactose, so when a highly caloric milk is required (e.g. by animals in a cold environment, such as marine mammals or hibernating bears) this is achieved by increasing the fat content of the milk.

Lipids are commonly divided into three classes.

- **Neutral lipids**: these are esters of glycerol and one, two or three fatty acids for mono-, di- and triglycerides respectively. Neutral lipids are by far the dominant class of lipids in all foods and tissues, representing 98.5% of total milk lipids.
- **Polar lipids** (a complex mixture of fatty acid esters of glycerol or sphingosine): many contain phosphoric acid, a nitrogen-containing compound (choline, ethanolamine or serine) or a sugar/oligosaccharide. Although present at low levels ($\sim 1\%$ of total milk lipids), the polar lipids play critical roles in milk and dairy products. They are very good natural emulsifiers and are concentrated in the MFGM, which maintains the milk lipids as discrete globules and ensures their physical and biochemical stability.
- Miscellaneous lipids: a heterogeneous group of compounds that are unrelated chemically to each other or to neutral or polar lipids. This group includes cholesterol, carotenoids and the fat-soluble vitamins, A, D, E and K. The carotenoids are important for two reasons: firstly, they are natural pigments (yellow, orange, red) and are responsible for the color of butter and cheese; some consumers prefer highly colored cheese, which is obtained by adding a carotenoid-containing extract from annatto beans; secondly, some carotenoids are converted to vitamin A in the liver.

**Fatty acids**

Fatty acids are carboxylic acids with the general formula R-COOH, where the alkyl group (R) is a hydrocarbon chain containing from 3 to 25 carbons (total number of carbons, from 4 to 26), which may be saturated or unsaturated (1 to 6 double bonds), and is usually straight (normal), with small amounts of branched chain, hydroxy and keto (oxo) acids. The vast majority of fatty acids have an even number of carbon atoms because they are synthesized from, and elongated by adding, a two-carbon compound, acetyl CoA, on each cycle of the multi-enzyme fatty acid synthetase (FAS). Although the hydroxy fatty acids are present at low levels, they are important in milk fat because they are converted on heating to lactones that give a desirable flavor to milk fat, which is considered to be the premium cooking fat. Although keto acids are also minor components, they are important flavor precursors because they are converted to highly flavored methyl ketones.

The melting point of fatty acids increases progressively with molecular weight (MW), whereas solubility in water decreases. The melting point decreases with the introduction of double bonds and, for unsaturated fatty acids, the melting point of the cis isomer is lower than that of the trans isomer.

Milk lipids are chemically similar to all other lipids but contain a very wide range of fatty acids (up to 400 fatty acids have been reported in milk lipids, although most of these are present at trace levels). The milk lipids of ruminants are unique in that they are the only natural lipids that contain butyric (butanoic) acid (C\textsubscript{4:0}); they also contain substantial amounts of medium-chain fatty acids (hexanoic [C\textsubscript{6:0}], octanoic [C\textsubscript{8:0}] and decanoic [C\textsubscript{10:0}]), the only other sources of which are coconut oil and palm kernel oil. The short- and medium-chain fats are water soluble and volatile, and have a strong aroma and taste.

The fatty acids in milk fat are obtained from three sources.

- Butanoic acid is produced by reducing β-hydroxybutanoic acid, which is synthesized by bacteria in the rumen; ruminant milk fat is the only natural lipid that contains this fatty acid.
- All hexanoic (C\textsubscript{6:0}) to tetradecanoic (C\textsubscript{14:0}) acids and 50% of hexadecanoic (C\textsubscript{16:0}) acid are synthesized in the mammary gland from acetyl CoA (CH\textsubscript{3}COSCoA); these fatty acids are released from the FAS by chain-length-specific thioesterases, the relative activities of which are responsible for interspecies differences in the proportions of medium-chain fatty acids. Decanoic
acid (C\textsubscript{10:0}) and dodecanoic acid (C\textsubscript{12:0}) are major fatty acids in the milk fats of elephant, horse, donkey, zebra, tapir, rhinoceros, rabbit and hare but these fats contain very little or no butanoic acid (Glass \textit{et al.}, 1967; Glass and Jenness, 1971; Christie, 1995; Osthoff \textit{et al.}, 2005). These species are non-ruminant herbivores with a large caecum, a feature that presumably is somehow responsible for the high levels of C\textsubscript{10:0} and C\textsubscript{12:0}; some of the above species also practice coprophagy.

- All octadecanoic (C\textsubscript{18:0}) acid and 50% of hexadecanoic (C\textsubscript{16:0}) acid are obtained from dietary lipids.

The unsaturated fatty acids are synthesized as follows:

- C\textsubscript{18:1} is produced from C\textsubscript{18:0} in the liver by \(\Delta-9\) desaturase.
- C\textsubscript{18:2} is obtained from the diet, i.e. it is an essential fatty acid.
- The other unsaturated fatty acids are produced from C\textsubscript{18:2} by further desaturation and/or elongation.

Ruminant milk fats contain low levels of polyunsaturated fatty acids (PUFAs) because PUFAs in the diet are hydrogenated by bacteria in the rumen. Biohydrogenation can be prevented by encapsulating dietary PUFAs or PUFA-rich sources in cross-linked (by HCHO) protein or cross-linked crushed oilseeds. PUFA-enriched milk has improved spreadability of butter and perceived improved nutritional qualities.

Incomplete biohydrogenation by the rumen bacterium \textit{Butyrivibrio fibrisolvens} results in the formation of conjugated linoleic acid (CLA; also called rumenic acid), which has potent anti-carcinogenic properties. Eight isomers of CLA are possible but \textit{cis-9, trans-11} is the most biologically active and is the predominant isomer found in bovine milk. The formation of CLA and its nutritional benefits have been the subject of considerable research during the past 15 years; this research has been reviewed by Bauman and Lock (2006) and Parodi (2006).

**Distribution of fatty acids in triglycerides**

As well as the constituent fatty acids, the position of the fatty acids in triglycerides affects their melting point and rheological properties. For these reasons, and to completely characterize the structure of triglycerides, the positions of the fatty acids in milk triglycerides have been determined. An index of this can be obtained by determining the acyl carbon number (ACN) of triglycerides, i.e. the sum of the number of carbons in the three component fatty acids, which can be done by gas chromatography. Probably the first study on this aspect was by Breckenridge and Kuksis (1967), in which the ACNs of the milk triglycerides from seven species were reported. More recent work has been reviewed by Christie (1995) and MacGibbon and Taylor (2006).

The complete structure of triglycerides can be determined by stereospecific analysis, the results of which for milk fat have been described by Christie (1995) and MacGibbon and Taylor (2006). The most notable feature is the almost exclusive esterification of the short-chain fatty acids—C\textsubscript{4:0} and C\textsubscript{6:0}—at the Sn3 position.
Degradation of lipids

Food lipids are susceptible to two forms of deterioration: lipid oxidation, leading to oxidative rancidity, and the hydrolysis of lipids by lipases (lipolysis), leading to hydrolytic rancidity.

Lipid oxidation involves a very complex set of chemical reactions, which have been well characterized; the literature has been comprehensively reviewed by T. Richardson and M. Korycka-Dahl in Fox (1983), by T. P. O’Connor and N. M. O’Brien in Fox (1983), and in Fox and McSweeney (2006).

Milk contains an indigenous lipoprotein lipase (LPL), which is normally inactive because it is separated from the triglyceride substrates by the MFGM. However, if the membrane is damaged, lipolysis and hydrolytic rancidity ensue rapidly. When milk lipids are hydrolyzed by milk LPL, the short- and medium-chain fatty acids are preferentially released. These fatty acids are major contributors to flavor, which may be desirable or undesirable, depending on the product. Hydrolytic rancidity caused by milk LPL is potentially a very serious problem in raw milk and in some dairy products. Lipolysis in milk has been reviewed comprehensively by H. C. Deeth and C. H. FitzGerald in Fox (1983, 1995) and in Fox and McSweeney (2006).

A low level of lipolysis is desirable in all types of cheese, especially in blue cheeses, in which the principal lipases are those secreted by the blue mould Penicillium roqueforti. The free fatty acids are converted to alk-2-ones, the principal flavor compounds in blue cheeses. The characteristic piquant flavor of some cheeses, e.g. Pecorino Romano, is due to short- and medium-chain fatty acids, which are released mainly by an added lipase—pregastric esterase. Other derivatives of fatty acids are alk-2-ols (secondary alcohols), lactones, esters and thio-esters; these are important flavor compounds in cheese.

Milk lipids as an emulsion

Lipids are insoluble in water or aqueous systems. When mixed, a lipid and water (or aqueous solvent) form distinct layers and a force, interfacial tension (γ), exists between the layers. Lipids can be dispersed in water by vigorous agitation (homogenization) but, when agitation ceases, the droplets of lipid coalesce quickly into a single mass (i.e. phase separation). This is driven by the need to reduce the interfacial area and γ to a minimum. If γ is reduced, the droplets of lipid will remain discrete, although they will rise to the surface (i.e. cream) because of the lower density of lipids (0.9) compared with water (1.0). Interfacial tension can be reduced by using a surface-active agent (emulsifier, detergent). Natural emulsifiers include proteins, phospholipids and mono- and diglycerides; there is a wide range of synthetic emulsifiers.

In milk, the lipids are dispersed in the milk serum (specific gravity, 1.036) as globules with a diameter in the range from <1 to ≈20μm (mean 3–4μm). The fatty acids (from the sources described above) and monoglycerides (from blood lipids) are synthesized to triglycerides in the rough endoplasmic reticulum (RER) at the basal end of the epithelial cells. The triglycerides form into globules within the RER and are released into the cell cytoplasm. The globules are stabilized by a complex layer of proteins and phospholipids, known as the MFGM. The inner layer of the MFGM is acquired within the epithelial cell as the fat globules, after release from the RER,
move towards the apical membrane. The outer layer of the MFGM is the apical membrane of the secretory cell through which the lipid globules are pushed as they are expressed from the cell. Because the stability of the milk emulsion is critical in most dairy products, the structure and the stability of the MFGM have been the subject of research for more than 100 years. The MFGM is composed mainly of phospholipids and proteins.

Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) indicates that there are eight main proteins in the MFGM, which have been isolated and characterized (see Mather, 2000). However, SDS-PAGE followed by micro-capillary high performance liquid chromatography-mass spectrometry (HPLC-MS) reveals 120 proteins, of which 23% are involved in protein trafficking, 23% in cell signalling, 21% in unknown functions, 11 in fat trafficking/metabolism, 9% in transport and 7% in protein synthesis/folding, 4% are immune proteins and 2% are contaminating skim milk proteins (Reinhardt and Lippolis, 2006). Many of the 70 indigenous enzymes in milk are concentrated in the MFGM. The very extensive literature on the MFGM has been the subject of many reviews, including Keenan and Mather (2006), who present an up-to-date model of the MFGM.

Some of the MFGM is shed during the aging of milk, specifically if agitated, and forms vesicles (sometimes called microsomes) in the skimmed milk. The MFGM may be damaged by agitation, homogenization, whipping or freezing and may lead to hydrolytic rancidity and non-globular fat which may cause cream plug, oiling-off in coffee and tea, and poor wettability of milk powder (for a review, see Evers, 2004). The MFGM is stripped from the fat globules by extensive agitation (usually of cream), a process referred to as churning; the free fat coalesces and is kneaded (worked) to give a water-in-oil emulsion—butter. The MFGM partitions into the aqueous phase, referred to as buttermilk. The phospholipids in buttermilk give it good emulsifying properties and there is commercial interest in using it as a food ingredient (Singh, 2006). Some of the polar lipids in the MFGM are reported to have desirable nutritional properties (see Ward et al., 2006), but there are conflicting results (see Riccio, 2004).

Presumably, the fat globules in the milk of all species are stabilized by a membrane similar to that in bovine milk but there is very little information on the MFGM in non-bovine milks. Buchheim et al. (1989) and Welsch et al. (1990) studied the glycoproteins in the MFGM of human, rhesus monkey, chimpanzee, dog, sheep, goat, cow, gray seal, camel and alpaca by SDS-PAGE with Periodic Acid Schiff (PAS) staining, Western blotting and lectin biochemistry. Large intra- and interspecies differences were found; very highly glycosylated proteins were found in the MFGM of primates, horse, donkey, camel and dog (see Patton, 1994, 2004, and references therein). Long (0.5–1μm) filamentous structures extend from the surface of the fat globules in equine and human milks; the filaments are composed of mucins (highly glycosylated proteins) which dissociate rapidly from the surface of globules in the bovine milk serum on cooling; they are also lost on heating human milk, e.g. at 80°C for 10 min (see Patton, 2004, and references therein).

The filaments facilitate the adherence of fat globules to the intestinal epithelium and probably improve the digestion of fat. The mucins prevent bacterial adhesion and
may protect mammary tissue against tumors (mammary tumors are very rare in the cow). Why the filaments on bovine milk fat globules are lost much more easily than those in equine milk and human milk is not known; work in this area is warranted. Proteomic methodology is being applied to study the human MFGM and membranes of the mammary epithelial cells (for references, see Reinhardt and Lippolis, 2006).

The fat globules in bovine milk form a cream layer because of the difference in specific gravity between the fat and aqueous phases, but the cream layer is dispersed readily by gentle agitation. The rate of creaming can be calculated from Stokes’ equation:

\[ \nu = 2r^2(\rho_1 - \rho_2)g/9\eta \]

where \( \nu \) is the velocity of creaming, \( r \) is the radius of the fat globules, \( \rho_1 \) and \( \rho_2 \) are the specific gravities of the continuous and dispersed phases respectively, \( g \) is acceleration due to gravity and \( \eta \) is the viscosity of the continuous phase.

Based on the typical values for \( r \), \( \rho_1 \), \( \rho_2 \) and \( \eta \) for milk, one would expect a cream layer to form in milk in about 60 h, but in fact a cream layer forms in about 30 min. The faster than expected rate of creaming is due to the aggregation of fat globules, aided by an immunoglobulin-M-type protein, called cryoglobulin, because it precipitates on to the fat globules when the milk is cooled. The clusters of globules then behave as a unit with a much larger radius. Creaming can be prevented by homogenizing the milk, which reduces the size of the globules and denatures the cryoglobulins. The fat globules in buffalo, ovine, caprine, equine and camel milks do not agglutinate, because these milks lack cryoglobulins.

Previously, the creaming of milk was a very important attribute and was a popular research topic; it has been the subject of several reviews, most recently by Huppertz and Kelly (2006). Traditionally, the fat was removed from milk by natural (gravity) creaming. Gravity creaming is still used to standardize the fat content for some cheese varieties, e.g. Parmigiano Reggiano, but the removal of fat from milk is now usually accomplished by centrifugal separation, in which \( g \) is replaced by \( \omega^2 R \), where \( \omega \) is the centrifugal velocity in radians per second and \( R \) is the radius of the centrifuge bowl. Centrifugal separation is very efficient, essentially instantaneous and continuous.

**Proteins**

The properties of milk and most dairy products are affected more by the proteins they contain than by any other constituent. The milk proteins also have many unique properties; because of this and their technological importance, the milk proteins have been studied extensively and are probably the best characterized food protein system.

Research on milk proteins dates from the early nineteenth century. Pioneering work was reported by J. Berzelius in 1814, by H. Schubler in 1818 on the physico-chemical status of milk proteins, and by H. Braconnot in 1830 who published the first paper in which the word casein was used. A method for the preparation of protein from milk by acid precipitation was described in 1938 by J. G. Mulder, who coined the term “protein.” The acid-precipitated protein was referred to as casein (some early authors called acid-precipitated milk protein caseinogen, which is converted by rennin to
casein, that coagulates in the presence of Ca\(^{2+}\); this situation is analogous to the conversion of fibrinogen in blood by thrombin to fibrin, which coagulates in the presence of Ca\(^{2+}\); about 70 years ago, the term “casein” was universally adopted as the English word for the pH-4.6-insoluble protein in milk. The method for acid (isoelectric) precipitation of casein was refined by Hammarsten (1883) and, consequently, isoelectric casein is frequently referred to as casein nach Hammarsten. An improved method for the isolation of casein was published in 1918 by L. L. van Slyke and J. C. Baker.

The liquid whey remaining after isoelectric precipitation of casein from skim or whole milk is a dilute solution of proteins (whey or serum proteins; \(\approx 0.7\%\) in bovine milk), lactose, organic and inorganic salts, vitamins and several constituents at trace levels. By salting-out with MgSO\(_4\), the whey proteins were fractionated by J. Sebelein, in 1885, into soluble (albumin) and insoluble (globulin) fractions. According to McMeekin (1970), in 1889 A. Wichmann crystallized a protein from the albumin fraction of whey by addition of (NH\(_4\))\(_2\)SO\(_4\) and acidification, a technique used to crystallize blood serum albumin and ovalbumin. Using the techniques available 100 years ago, the whey proteins were found to be generally similar to the corresponding fractions of blood proteins and were considered to have passed directly from blood to milk; consequently, the whey proteins attracted little research effort until the 1930s.

In addition to the caseins and whey proteins, milk contains two other groups of proteinaceous materials—proteose peptones (PPs) and non-protein nitrogen (NPN)—which were recognized in 1938 by S. J. Rowland who observed that, after heating milk fat at 95ºC for 10 min, the whey proteins co-precipitated with the caseins on acidification to pH 4.6. When the pH-4.6-soluble fraction of heated milk was made to 12% trichloroacetic acid (TCA), some nitrogenous compounds precipitated and were designated “proteose peptone”; nitrogenous compounds that remained soluble in 12% TCA were designated NPN. A modified version of S. J. Rowland’s scheme is now used to quantify the principal nitrogenous groups in milk.

Thus, by 1938, the complexity of the milk protein system had been described (i.e. caseins, lactalbumin (now known to consist mainly of \(\alpha\)-lactalbumin, \(\beta\)-lactoglobulin and blood serum albumin), lactoglobulin, PPs and NPN, which represent approximately 78, 12, 5, 2 and 3% respectively of the nitrogen in bovine milk). However, knowledge of the milk protein system was rudimentary and vague at this stage.

Knowledge on the chemistry of milk proteins has advanced steadily during the twentieth century, as can be followed through the progression of textbooks and reviews on dairy chemistry (see Fox, 1992, 2003; Fox and MacSweeney, 2003).

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**Preparation of casein and whey proteins**

The protein fractions may be prepared from whole or skimmed milk, but the latter is usually used because the fat is occluded in isoelectric casein and interferes with further characterization of the proteins. The fat is easily removed from milk by centrifugation (e.g. 3000 \(\times\) g for 30 min) and any remaining fat may be removed by washing the precipitated protein with ether. Isoelectric precipitation is the most widely used method for separating the casein and non-casein fractions of milk protein but several other techniques are used in certain situations (see Fox, 2003).
- Isoelectric precipitation at ≈pH 4.6 at 20°C: the precipitate is recovered by filtration or low-speed centrifugation. Essentially similar methods are used to prepare casein on a laboratory scale or an industrial scale.

- Ultracentrifugation: in milk, the casein exists as large micelles that may be sedimented by centrifugation at 100 000 × g for 1 h; the whey proteins are not sedimentable. The casein pellet can be redispersed in a suitable buffer as micelles with properties similar to those of natural micelles.

- Salting-out methods: casein can be precipitated by any of several salts, usually by (NH₄)₂SO₄ at 260 g/L or saturated NaCl. The immunoglobulins co-precipitate with the caseins.

- Ultrafiltration and microfiltration: all the milk proteins are retained by small-pore, semi-permeable membranes and are separated from lactose and soluble salts. This process, ultrafiltration, is used widely for the industrial-scale production of whey protein concentrates (WPCs) and to a lesser extent for the production of total milk protein. Intermediate-pore membranes are used to separate casein micelles from whey proteins. In microfiltration, using large-pore membranes (0.4 μm), both the caseins and the whey proteins are permeable, but >99.9% of bacteria and other large particles are retained; microfiltration is used for the production of extended-shelf-life beverage milk or cheese milk or to remove lipoprotein particles from whey to improve the functionality of WPC.

- Gel filtration: it is possible to separate the caseins from the whey proteins by permeation chromatography but this method is not used industrially and is rarely used on a laboratory scale.

- Precipitation by ethanol: the caseins are precipitated from milk by ≈40% ethanol, whereas the whey proteins remain soluble; however, precipitation by ethanol is rarely used, on either a laboratory scale or an industrial scale, for the precipitation of casein.

- Cryoprecipitation: caseins, in a micellar form, may be destabilized and precipitated by freezing milk or, preferably, concentrated milk, at about −10°C. Precipitation is caused by a decrease in pH and an increase in Ca²⁺ concentration; the precipitated micelles may be redispersed as micelles by heating to about 55°C. Alternatively, the cryoprecipitated casein may be recovered, washed and dried; it has many interesting properties for food applications, but it is not produced commercially.

- Rennet coagulation: the casein micelles are destabilized by specific, limited proteolysis and coagulate in the presence of Ca²⁺. The properties of rennet-coagulated casein are very different from those of isoelectric casein; rennet-coagulated casein is very suitable for certain food applications, e.g. cheese analogs.

- Caseinates: isoelectric casein is insoluble in water, but may be converted to water-soluble caseinates by dispersion in water and adjusting the pH to ≈6.7 with alkali, usually NaOH to yield sodium caseinate. KOH, NH₄OH and Ca(OH)₂ give the corresponding caseinates, which may be freeze dried or spray dried.
Comparison of key properties of casein and whey proteins

- Solubility at pH 4.6. The caseins are, by definition, insoluble at pH 4.6, whereas the whey proteins are soluble under the ionic conditions of milk. The isoelectric precipitation of casein is exploited in the production of caseins and caseinates, fermented milk products and acid-coagulated cheeses.

- Coagulability following limited proteolysis. The caseins are coagulable following specific, limited proteolysis, whereas the whey proteins are not. This property of the caseins is exploited in the production of rennet-coagulated cheese (≈75% of all cheese) and rennet casein.

- Heat stability. The caseins are very heat stable. Milk at pH 6.7 may be heated at 100°C for 24 h without coagulation and withstands heating at 140°C for up to 20–25 min; aqueous solutions of sodium caseinate may be heated at 140°C for several hours without apparent changes. The heat stability of the whey proteins is typical of globular proteins; they are denatured completely on heating at 90°C for 10 min. The remarkably high heat stability of the caseins, which is probably due to their lack of typical stable secondary and tertiary structures, permits the production of heat-sterilized dairy products with relatively small physical changes.

- Amino acid composition. The caseins contain high levels of proline (17% of all residues in β-casein), which explains their lack of α- and β-structures. The caseins are phosphorylated, whereas the principal whey proteins are not. Whole isoelectric casein contains approximately 0.8% phosphorus, but the degree of phosphorylation varies among the individual caseins. The phosphate is attached to the polypeptides as phosphomonoesters of serine: the presence of phosphate groups has major significance for the properties of the caseins, e.g., (i) molecular charge and related properties and heat stability; and (ii) metal binding which affects their physico-chemical, functional and nutritional properties. Metal binding by casein is regarded as a biological function because it enables a high concentration of calcium phosphate to be carried in milk in a soluble form (to supply the requirements of the neonate); otherwise, calcium phosphate would precipitate in, and block, the ducts of the mammary gland, leading to the death of the gland and perhaps of the animal.

- The caseins are low in sulfur (0.8%), whereas the whey proteins are relatively rich (1.7%). The sulfur in casein is mainly in methionine, with little cystine or cysteine; the principal caseins are devoid of the latter two amino acids. The whey proteins are relatively rich in cysteine and/or cystine, which have major effects on the physico-chemical properties of these proteins and of milk.

- Site of biosynthesis. The caseins are synthesized in the mammary gland and are unique to this organ. Presumably, they are synthesized to meet the amino acid requirements of the neonate and as carriers of important metals required by the neonate. The principal whey proteins are also synthesized in, and are unique to, the mammary gland, but several minor proteins in milk are derived from blood, either by selective transport or due to leakage. Most of the whey proteins have a biological function.

- Physical state in milk. The whey proteins exist in milk as monomers or as small quaternary structures whereas the caseins exist as large aggregates, known
as micelles, with a mass of \( \approx 10^8 \) Da and containing about 5000 molecules. The white color of milk is due largely to the scattering of light by the casein micelles. The structure, properties and stability of the casein micelles are of major significance for the technological properties of milk and have been the subject of intensive research (see below).

**Heterogeneity and fractionation of casein**

O. Hammersten believed that isoelectric casein was a homogeneous protein but, during the early years of the twentieth century, evidence was presented by T. B. Osborne and A. J. Wakeman, and especially by K. Linderstrøm-Lang and collaborators, that it might be heterogeneous (see McMeekin, 1970). By extraction with ethanol-HCl mixtures, K. Linderstrøm-Lang and S. Kodoma obtained three major casein fractions, which contained about 1.0, 0.6 or 0.1% phosphorus, and several minor fractions. The heterogeneity of casein was confirmed by analytical ultracentrifugation and free-boundary electrophoresis by K. O. Pedersen and O. Mellander respectively (see McMeekin, 1970). Electrophoresis resolved isoelectric casein into three proteins, which were named \( \alpha \), \( \beta \) and \( \gamma \) in order of decreasing electrophoretic mobility and represented about 75, 22 and 3% of whole casein respectively.

Following the demonstration of its heterogeneity, several attempts were made to isolate the individual caseins. The first reasonably successful method was developed in 1944 by Warner, who exploited differences in the solubilities of \( \alpha \)- and \( \beta \)-caseins at pH 4.4 and 2ºC. A much more satisfactory fractionation method was developed in 1952 by N. J. Hipp and co-workers based on the differential solubility of \( \alpha \)-, \( \beta \)- and \( \gamma \)-caseins in urea solutions at pH 4.9. This method was widely used for many years until the widespread application of ion-exchange chromatography.

In 1956, \( \alpha \)-casein was resolved by D. F. Waugh and P. H. von Hippel into calcium-sensitive and calcium-insensitive proteins which were called \( \alpha_s \) and \( \kappa \)-caseins respectively. \( \kappa \)-Casein, which represents \( \approx 12\% \) of total casein, is responsible for the formation and stabilization of casein micelles, and affects many technologically important properties of the milk protein system. Numerous chemical methods were soon developed for the isolation of \( \kappa \)-casein (see Fox, 2003). \( \alpha_s \)-Casein, prepared by the method of D. F. Waugh and P. H. von Hippel, contains two proteins, now called \( \alpha_{s1} \) and \( \alpha_{s2} \)-caseins.

Chemical methods for fractionation of the caseins have now been largely superseded by ion-exchange chromatography, which gives superior results when urea and a reducing agent are used (see Strange et al., 1992; Imafidon et al., 1997).

**Application of gel electrophoresis to the study of milk proteins**

Zone electrophoresis on a solid medium, paper or cellulose acetate was introduced in the 1940s. This technique gave good results with many protein systems, but the caseins, because of a very strong tendency to associate hydrophobically, were resolved poorly on these media. Electrophoresis in starch gels (SGE) using discontinuous buffer systems was introduced to general protein chemistry by M. D. Poulak in 1957 and was applied to the study of the caseins by R. G. Wake and R. L. Baldwin
in 1961. The resolving power of SGE was far superior to that of any of its predecessors. When urea (7 M) and a reducing agent, usually 2-mercaptoethanol, were incorporated into the starch gel, isoelectric casein was resolved into about 20 bands, most of which were due to the microheterogeneity of one or more of the caseins.

Electrophoresis on polyacrylamide disk gels (PAGE) was introduced by L. Ornstein in 1964 and was applied to the study of the caseins by R. F. Peterson in 1966. PAGE and SGE give similar results, but PAGE is far easier to use and has become the standard electrophoretic method for the analysis of caseins (and most other protein systems). Gel electrophoretic methods for the analysis of milk proteins have been reviewed by Swaisgood (1975), Strange et al. (1992) and Tremblay et al. (2003).

SDS-PAGE, which resolves proteins mainly on the basis of molecular mass, is very effective for most proteins but, because the masses of the four caseins are quite similar, SDS-PAGE is not very effective. β-Casein, which has very high surface hydrophobicity, binds a disproportionately high amount of SDS and, consequently, has a higher electrophoretic mobility than αs1-casein, although it is a larger molecule. SDS-PAGE is very effective for the resolution of whey proteins and is the method of choice.

Microheterogeneity of the caseins

αs1-, αs2-, β- and κ-caseins represent approximately 38, 10, 35 and 12% respectively of whole bovine casein. However, SGE or PAGE indicates much greater heterogeneity due to small differences in one or more of the caseins, referred to as microheterogeneity, which arises from five factors.

Variability in the degree of phosphorylation

All the caseins are phosphorylated but to a variable degree (αs1-, 8 or 9P; αs2-, 10, 11, 12 or 13P; β-, 4 or 5P; κ-, 1 or 2P per molecule). The number of phosphate residues is indicated thus: αs1-CN 8P, β-CN 5P etc.

Genetic polymorphism

In 1955, R. Aschaffenburg and J. Drewry discovered that β-lactoglobulin exists in two forms (variants, polymorphs) now called A and B, which differ by only two amino acids. The variant in the milk is genetically controlled and the phenomenon is called genetic polymorphism. It was soon shown that all milk proteins exhibit genetic polymorphism and at least 45 polymorphs have been detected by PAGE, which differentiates on the basis of charge and therefore only polymorphs that differ in charge have been detected. It is very likely that only a small proportion of the genetic polymorphs of milk proteins have been detected. The potential of peptide mapping of enzymatic hydrolysates by HPLC-MS has been assessed. The genetic polymorph(s) present is indicated by a Latin letter as follows: β-CN A 5P, αs1-CN B 9P, κ-CN A 1P etc. Genetic polymorphism also occurs in the milk of sheep, goat, buffalo, pig and horse, and probably of all species.

Technologically important properties of milk, e.g. rennetability, heat stability, yield and proportions of milk proteins, are affected by the genetic polymorphs of the milk
proteins present, and work in this area is being expanded and refined. The extensive literature on the genetic polymorphism of milk proteins has been the subject of several reviews, including Ng-Kwai-Hang and Grosclaude (2003).

**Disulfide bonding**

$\alpha_s1$- and $\beta$-caseins lack cysteine and cystine, but both $\alpha_{s2}$- and $\kappa$-caseins contain two half cystine residues which occur as intermolecular disulfide bonds. $\alpha_{s2}$-Casein exists as a disulfide-linked dimer, whereas up to 10 $\kappa$-casein molecules may be linked by disulfide bonds. Inclusion of a reducing agent (usually mercaptoethanol) in SGE or PAGE gels is required for good resolution of $\kappa$-casein; in its absence, $\alpha_{s2}$-casein appears as a dimer (originally called $\alpha_{s5}$-casein).

**Variations in the degree of glycosylation**

$\kappa$-Casein is the only glycosylated casein; it contains galactose, N-acetylgalactosamine and N-acetylneuraminic (sialic) acid, which occur as tri- or tetrasaccharides, the number of which varies from 0 to 4 per molecule of protein (i.e. a total of 9 variants).

**Hydrolysis of the caseins by plasmin**

Milk contains several indigenous proteinases, the principal of which is plasmin, a trypsin-like, serine-type proteinase from blood; it is highly specific for peptide bonds with a lysine or arginine at the P1 position. The preferred casein substrates are $\beta$- and $\alpha_{s2}$-caseins; $\alpha_{s1}$-casein is also hydrolyzed, but $\kappa$-casein is very resistant, as are the whey proteins. All the caseins contain several lysine and arginine residues, but only a few bonds are hydrolyzed rapidly. $\beta$-Casein is hydrolyzed rapidly at the bonds Lys$_{28}$–Lys$_{29}$, Lys$_{105}$–His$_{106}$ and Lys$_{107}$–Glu$_{108}$. The resulting C-terminal peptides are the $\gamma$-caseins ($\gamma^1$: $\beta$-CNf$_{29}$–209; $\gamma^2$: $\beta$-CNf$_{106}$–209; $\gamma^3$: $\beta$-CNf$_{108}$–209), whereas the N-terminal peptides are proteose peptones 5, 8slow and 8fast. The $\gamma$-caseins, which represent ≈3% of total casein, are evident in PAGE gels. Other plasmin-produced peptides are probably present but either are too small to be readily detectable by PAGE or their concentrations are very low relative to those of the principal caseins.

Although $\alpha_{s2}$-casein in solution is also quite susceptible to plasmin, $\alpha_{s2}$-casein-derived peptides have not been identified in milk. $\alpha_{s1}$-Casein in solution is also hydrolyzed readily by plasmin; members of a minor casein fraction, $\lambda$-casein, are N-terminal fragments of $\alpha_{s1}$-casein produced by plasmin (O’Flaherty, 1997).

**Molecular properties of the milk proteins**

The principal, and many of the minor, milk proteins have been very well characterized. The principal properties of the six milk-specific proteins are summarized in Table 1.1. A number of features warrant comment.

The six principal lactoproteins are small molecules, a feature that contributes to their stability. The primary structures of the principal lactoproteins of several species are known, as are the substitutions in the principal genetic variants.

The whey proteins are highly structured, but the four caseins lack stable secondary structures; classical physical measurements indicate that the caseins are unstructured,
but theoretical considerations indicate that, rather than being unstructured, the caseins are very flexible molecules and have been referred to as rheomorphic (Holt and Sawyer, 1993; see also Horne, 2002; Farrell et al., 2006a). The inability of the caseins to form stable structures is due mainly to their high content of the structure-breaking amino acid proline; β-casein is particularly rich in proline, with 35 of the 209 residues. The open, flexible structure of the caseins renders them very susceptible to proteolysis, which facilitates their natural function as a source of amino acids.

In contrast, the native whey proteins, especially β-lactoglobulin, are quite resistant to proteolysis, and at least some are excreted in the feces of infants. This feature is important because most of the whey proteins play a non-nutritional function in the intestine and, therefore, resistance to proteolysis is important.

The caseins are generally regarded as very hydrophobic proteins but, with the exception of β-casein, they are not exceptionally hydrophobic. Because of their lack of stable secondary and tertiary structures, most of their hydrophobic residues are exposed and, consequently, they have a high surface hydrophobicity.

One of the more notable features of the amino acid sequence of the caseins is that the hydrophobic and hydrophilic residues are not distributed uniformly, thereby giving the caseins a distinctly amphipatic structure. This feature, coupled with their open flexible structure, gives the caseins good surface activity, and good foaming and emulsifying properties, making casein the functional protein of choice for many applications. Because of their hydrophobic sequences, the caseins have a propensity to yield bitter hydrolysates.

Also because of their open structure, the caseins have a high specific volume and, consequently, form highly viscous solutions, which is a disadvantage in the production of caseinates. Because of its high viscosity, it is not possible to spray dry sodium caseinate solutions containing >20% protein, thereby increasing the cost of drying and resulting in low-bulk-density powders.

<table>
<thead>
<tr>
<th>Table 1.1 Properties of the principal lactoproteins</th>
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<tr>
<td>Properties</td>
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<tr>
<td>MW (Dalton)</td>
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<tr>
<td>23,612</td>
</tr>
<tr>
<td>Residues</td>
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<tr>
<td>Conc in milk (g/L)</td>
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<tr>
<td>Phosphate residues</td>
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<tr>
<td>½ Cystine</td>
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<tr>
<td>Sugars</td>
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<tr>
<td>Prolyl residue per molecule</td>
</tr>
<tr>
<td>A280, 1% 1 cm</td>
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<tr>
<td>Secondary structure</td>
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<tr>
<td>Hb (ave)</td>
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<tr>
<td>pl</td>
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<tr>
<td>Partial specific volume (ml/g)</td>
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</table>
The lack of stable tertiary structures means that the caseins are not denaturable *stricto sensu* and, consequently, are extremely heat stable; sodium caseinate, at pH 7, can withstand heating at 140°C for several hours without visible change. This very high heat stability makes it possible to produce heat-sterilized dairy products with very little change in physical appearance; other major food systems undergo major physical and sensoric changes on severe heating.

The caseins have a very strong tendency to associate, due mainly to hydrophobic bonding. Even in sodium caseinate, the most soluble form of casein, the molecules form aggregates of 250–500 kDa, i.e. containing 10–20 molecules. This strong tendency to associate makes it difficult to fractionate the caseins, for which a dissociating agent, e.g. urea or SDS, is required. On the other hand, a tendency to associate is important for some functional applications and in the formation and stabilization of casein micelles. In contrast, the whey proteins are molecularly dispersed in solution.

Because of their high content of phosphate groups, which occur in clusters, $\alpha_{s1}^-$, $\alpha_{s2}^-$ and $\beta$-caseins have a strong tendency to bind metal ions, which in the case of milk are mainly Ca$^{2+}$ ions. This property has many major consequences; the most important from a technological viewpoint is that these three proteins, which represent approximately 85% of total casein, are insoluble at calcium concentrations $>\approx 6\text{mM}$ at temperatures $>20^\circ\text{C}$. As bovine milk contains $\approx 30\text{mM}$ calcium, one would expect that the caseins would precipitate under the conditions prevailing in milk. However, $\kappa$-casein, which contains only one organic phosphate group, binds calcium weakly and is soluble at all calcium concentrations found in dairy products.

Furthermore, when mixed with the calcium-sensitive caseins, $\kappa$-casein can stabilize and protect $\approx 10$ times its mass of the former by forming large colloidal particles called casein micelles. The micelles act as carriers of inorganic elements, especially calcium and phosphorus, but also magnesium and zinc, and are, therefore, very important from a nutritional viewpoint. Through the formation of micelles, it is possible to solubilize much higher levels of calcium and phosphate than would otherwise be possible.

**Nomenclature of milk proteins**

During the period of greatest activity on the fractionation of casein (1950–1970), several casein (and whey protein) fractions were prepared that either were similar to proteins already isolated and named, or were artifacts of the isolation procedure. In order to standardize the nomenclature of the milk proteins, the American Dairy Science Association established a Nomenclature Committee in 1955, which has published seven reports, the most recent of which is Farrell *et al.* (2004). In addition to standardizing the nomenclature of the milk proteins, the characteristics of the principal milk proteins are summarized in these articles.

**Whey proteins**

About 20% of the total proteins of bovine milk are whey (serum) proteins. The total whey protein fraction is prepared by any of the methods described for the preparation of casein, i.e. the proteins that are soluble at pH 4.6 or in saturated NaCl or, after
Rennet-induced coagulation of the caseins, are permeable on microfiltration, or not sedimented by ultracentrifugation.

The proteins prepared by these methods differ somewhat: acid whey contains PPs; immunoglobulins are co-precipitated with the caseins by saturated NaCl; rennet whey contains the macropeptides, produced from κ-casein by rennet, plus small amounts of casein; and small casein micelles remain in the ultracentrifugal serum.

On a commercial scale, whey-protein-rich products are prepared by:

- ultrafiltration/diafiltration of casein or rennet whey to remove various amounts of lactose, and spray drying to produce whey protein concentrates (WPCs, 30–85% protein);
- ion-exchange chromatography and spray drying to yield whey protein isolate (WPI), containing ≈95% protein;
- demineralization by electrodialysis or ion exchange, thermal evaporation of water and crystallization of lactose; and
- thermal denaturation, removal of precipitated protein by filtration/centrifugation and spray drying, to yield lactalbumin, which has very low solubility and poor functionality.

Fractionation of whey proteins
It was recognized early that acid whey contains two well-defined groups of proteins: lactalbumins, which are soluble in 50% saturated (NH₄)₂SO₄ or saturated MgSO₄, and lactoglobulins, which are salted out under these conditions. The lactoglobulin fraction contains mainly immunoglobulins. The lactalbumin fraction contains two principal proteins, β-lactoglobulin and α-lactalbumin, and several minor proteins, including blood serum albumin and lactoferrin, which have been isolated by various procedures and crystallized (see Imafidon et al., 1997; Fox, 2003).

There is considerable interest in the production of the major and many minor whey proteins on a commercial scale for nutritional, nutraceutical or functional applications. Several methods have been developed for the industrial-scale production of several whey proteins (see Mulvihill and Ennis, 2003).

Major characteristics of whey proteins

β-Lactoglobulin
β-Lactoglobulin (β-Lg) represents ≈50% of the whey proteins, ≈12% of the total protein, in bovine milk. It is a typical globular protein and has been characterized very well. The extensive literature has been reviewed by, among others, Sawyer (2003).

β-Lg is the principal whey protein in the milk of the cow, buffalo, sheep and goat, although there are slight inter-species differences. Initially, it was considered that β-Lg occurs only in the milk of ruminants, but it is now known that a similar protein occurs in the milk of many other species, including the sow, mare, kangaroo, dolphin and manatee. However, β-Lg does not occur in the milk of human, rat, mouse, guinea pig, camel, llama or alpaca, in which α-La is the principal whey protein.
Bovine β-Lg consists of 162 residues per monomer, with a MW of ≈18 kDa; its amino acid sequence, and that of several other species, has been established. Its isoelectric point is ≈pH 5.2. It contains two intramolecular disulfide bonds and one mole of cysteine per monomer. The cysteine is especially important because it reacts, following thermal denaturation, with the intermolecular disulfide of κ-casein and significantly affects the rennet coagulation and heat stability of milk. It is also responsible for the cooked flavor of heated milk. Some β-Lgs (e.g. porcine) lack a sulfydryl group. Ten genetic variants, A–J, of bovine β-Lg have been identified, the most common being A and B. Genetic polymorphism also occurs in β-Lg of other species.

β-Lg is a highly structured protein: in the pH range 2–6, 10–15% of the molecule exists as α-helices, 43% as β-sheets and 47% as unordered structures, including β-turns; the β-sheets occur in a β-barrel-type calyx. The molecule has a very compact globular structure; each monomer exists almost as a sphere, about 3.6 nm in diameter. β-Lg exists as a dimer, MW ≈36 kDa, in the pH range 5.5–7.5, as a monomer at <pH 3.5 and >pH 7.5, and as a tetramer (MW ≈144 kDa) in the pH range 3.5–5.5. Porcine and other β-Lgs that lack a free thiol do not form dimers, probably not due directly to the absence of a thiol group.

β-Lg is very resistant to proteolysis in its native state; this feature suggests that its primary function is not nutritional. It may have either or both of two biological roles:

- It binds retinol (vitamin A) in a hydrophobic pocket, protects it from oxidation and transports it through the stomach to the small intestine where the retinol is transferred to a retinol-binding protein, which has a similar structure to β-Lg. It is not clear how retinol is transferred from the core of the fat globules, where it occurs in milk, to β-Lg and why some species lack this protein. β-Lg can bind many hydrophobic molecules and hence its ability to bind retinol may be incidental. β-Lg is a member of the lipocalin family, all of which have binding properties (Akerstrom et al., 2000).
- Through its ability to bind fatty acids, β-Lg stimulates lipase activity, which may be its most important physiological function.

β-Lg is the most allergenic protein in bovine milk for human infants and there is interest in producing whey protein products free of β-Lg for use in infant formulas. β-Lg has very good thermo-gelling properties and determines the gelation of WPCs.

α-Lactalbumin
About 20% of the protein of bovine whey (3.5% of total milk protein) is α-La, which is the principal protein in human milk. It is a small protein containing 123 amino acid residues, with a mass of ≈14 kDa, and has been well characterized; the literature has been reviewed by, among others, McKenzie and White (1991) and Brew (2003).

α-La contains four tryptophan residues per mole, giving it a specific absorbance at 280 nm of 20. It contains four intramolecular disulfide bonds per mole but no cysteine, phosphate or carbohydrate. Its isoionic point is ≈pH 4.8. The milk of
Bos taurus breeds contains only one genetic variant of α-La, B, but zebu cattle produce two variants, A and B. α-La has been isolated from the milk of the cow, sheep, goat, sow, human, buffalo, rat, guinea pig, horse and many other species; there are minor inter-species differences in the composition and properties.

The primary structure of α-La is homologous with lysozyme; out of a total of 123 residues in α-La, 54 are identical to corresponding residues in chicken egg white lysozyme and 23 others are structurally similar. α-La is a compact, highly structured globular protein. Because of difficulties in preparing good crystals, its tertiary structure has not been determined but is similar to that of lysozyme, on which a model of the structure of α-La is based (see McKenzie and White, 1991). In evolutionary terms, lysozyme is a very ancient protein; it is believed that α-La evolved from it through gene duplication (see Nitta and Sugai, 1989).

As discussed earlier, α-La is a component of lactose synthetase, the enzyme that catalyzes the final step in the biosynthesis of lactose. There is a direct correlation between the concentrations of α-La and lactose in milk. The milk of some marine mammals contains very little or no α-La.

α-La is a metalloprotein containing one Ca\(^{2+}\) per mole in a pocket containing four Asp residues. The calcium-containing protein is the most heat stable of the principal whey proteins, or, more correctly, the protein renatures following heat denaturation, which occurs at a relatively low temperature, as indicated by differential scanning calorimetry. When the pH is reduced to \(\approx\) pH 5, the Asp residues become protonated and lose their ability to bind Ca\(^{2+}\). The metal-free protein is denatured at quite a low temperature and does not renature on cooling; this characteristic has been exploited to isolate α-La from whey. α-La has poor thermo-gelling properties. Most lysozymes do not bind Ca\(^{2+}\) but equine lysozyme is an exception (Nitta et al., 1987) and seems to be an intermediate in the evolution of lysozyme to α-La (see Nitta and Sugai, 1989).

α-La is synthesized in the mammary gland, but a very low level is transferred, probably via leaky mammocyte junctions, into blood serum, in which the concentration of α-La increases during pregnancy or following administration of steroid hormones to male or female animals (Akers, 2000). The concentration of α-La in blood serum is a reliable, non-invasive indicator of mammary gland development and of the potential of an animal for milk production.

A high MW form of α-La, isolated recently from human acid-precipitated casein, has anti-carcinogenic activity; it was named HAMLET (human α-La made lethal to tumour cells). Bovine α-La can be converted to a form with similar activity, called BAMLET; it is the molten globular state formed from apo-α-La and \(\text{cis}\Delta9\text{-octadecenoic acid}\) (see Chatterton et al., 2006).

**Blood serum albumin**

Normal bovine milk contains 0.1–0.4 g/L of blood serum albumin (BSA; 0.3–1.0% of total nitrogen), presumably as a result of leakage from blood; it has no known biological function in milk. BSA has been studied extensively; for reviews, see Fox (2003). Because of its low concentration in milk, BSA has little effect on the physico-chemical properties of WPC and WPI.
**Immunoglobulins**

Mature bovine milk contains 0.6–1 g immunoglobulins (Igs)/L (≈3% of total nitrogen), but colostrum contains ≈ 10% (w/v) Ig, the level of which decreases rapidly post-partum. IgG1 is the principal Ig in bovine, caprine or ovine milk, with lesser amounts of IgG2, IgA and IgM; IgA is the principal Ig in human milk. The cow, sheep and goat do not transfer Ig to the fetus in utero and the neonate is born without Ig in its blood; consequently, it is very susceptible to bacterial infection with a very high risk of mortality.

The young of these species can absorb Ig from the intestine for several days after birth and thereby acquire passive immunity until they synthesize their own Ig, within a few weeks of birth. The human mother transfers Ig in utero and the offspring are born with a broad spectrum of antibodies. Although the human baby cannot absorb Ig from the intestine, the ingestion of colostrum is still very important because its Igs prevent intestinal infection. Some species, e.g. the horse, transfer Ig both in utero and via colostrum (see Hurley, 2003).

It has been suggested that the neonate secretes chymosin rather than pepsin because the former is weakly proteolytic and does not inactivate Ig. Colostrum contains α2-macroglobulin which may inhibit proteinases in the gastrointestinal tract and protect Ig.

The modern dairy cow produces colostrum far in excess of the requirements of its calf; surplus colostrum is available for the recovery of Ig and other nutraceuticals (Pakkanen and Aalto, 1997). There is considerable interest in hyperimmunizing cows against certain human pathogens, e.g. rota virus, for the production of antibody-rich milk for human consumption, especially by infants; the Ig could be isolated from the milk and presented as a “pharmaceutical” or consumed directly in the milk.

**Whey acidic protein**

WAP was identified first in the mouse milk and has since been found also in rat, rabbit, camel, wallaby, opossum, echidna and platypus milk. As the milks of all of these species lack β-Lg, it was thought that these proteins were mutually exclusive. However, porcine milk, which contains β-Lg, was recently found to contain WAP also. The MW of WAP is 14–30 kDa (the variation may be due to differences in glycosylation) and it contains two (in eutherians) or three (in monotremes and marsupials) four-disulfide domains. As human milk lacks β-Lg, it might be expected to contain WAP but there are no reports to this effect. In humans and ruminants, the WAP gene is frame shifted and is a pseudogene. WAP functions as a proteinase inhibitor, is involved in terminal differentiation in the mammary gland and has antibacterial activity (for reviews, see Simpson and Nicholas, 2002; Hajjoubi et al., 2006).

**Proteose peptone 3**

The PP fraction of milk protein is a very complex mixture of peptides, most of which are produced by the action of indigenous plasmin (see above) but some are indigenous to milk. The fraction has been only partially characterized; the current status has been described by Fox (2003). The PP fraction is of little or no technological significance.
Bovine proteose peptone 3 (PP3) is a heat-stable phosphoglycoprotein that was first identified in the PP (heat-stable, acid-soluble) fraction of milk. Unlike the other peptides in this fraction, PP3 is an indigenous milk protein, synthesized in the mammary gland. Bovine PP3 consists of 135 amino acid residues, with five phosphorylation and three glycosylation sites. When isolated from milk, the PP3 fraction contains at least three components of MW $\approx$ 28, 18 and 11 kDa; the largest of these is PP3, and the smaller components are fragments thereof generated by plasmin (see Girardet and Linden, 1996). PP3 is present mainly in acid whey but some is present in the MFGM. Girardet and Linden (1996) proposed changing the name to lactophorin or lacto-glycophorin; it has also been referred to as the hydophobic fraction of PP.

Because of its strong surfactant properties (Campagna et al., 1998), PP3 can prevent contact between milk lipase and its substrates, thus preventing spontaneous lipolysis. Although its amino acid composition suggests that PP3 is not a hydrophobic protein, it behaves hydrophobically, possibly because of the formation of an amphiphilic $\alpha$-helix, one side of which contains hydrophilic residues whereas the other side is hydrophobic. The biological role of PP3 is unknown.

**Non-protein nitrogen**
The NPN fraction of milk contains those nitrogenous compounds that are soluble in 12% TCA; it represents $\approx$5% of total nitrogen ($\approx$300 mg/L). The principal components are urea, creatine, uric acid and amino acids. Human milk contains a high level of taurine which can be converted to cysteine and may be nutritionally important for infants. Urea, the concentration of which varies considerably, has a significant effect on the heat stability of milk.

**Minor proteins**
Milk contains several proteins at very low or trace levels, many of which are biologically active (see Schrezenmeir et al., 2000); some are regarded as highly significant and have attracted considerable attention as nutraceuticals. When ways of increasing the value of milk proteins are discussed, the focus is usually on these minor proteins but they are, in fact, of little economic value to the overall dairy industry. They are found mainly in the whey but some are also located in the fat globule membrane. Reviews on the minor proteins include Fox and Flynn (1992), Fox and Kelly (2003), and Haggarty (2003).

**Metal-binding proteins**
Milk contains several metal-binding proteins: the caseins (Ca, Mg, Zn), $\alpha$-La (Ca), xanthine oxidase (Mo, Fe), alkaline phosphatase (Zn, Mg), lactoperoxidase (Fe), catalase (Fe), ceruloplasmin (Cu), glutathione peroxidase (Se), lactoferrin (Fe) and transferrin (Fe).

Lactoferrin (Lf), a non-haem iron-binding glycoprotein, is a member of a family of iron-binding proteins, which includes transferrin and ovotransferrin (conalbumin) (see Lonnerdal, 2003). It is present in several body fluids, including saliva, tears, sweat and semen. Lf has several potential biological functions: it improves the bioavailability of iron, is bacteriostatic (by sequestering iron and making it unavailable to intestinal
bacteria) and has antioxidant, antiviral, anti-inflammatory, immunomodulatory and anti-carcinogenic activity.

Human milk contains a very high level of Lf (≈20% of total nitrogen) and therefore there is interest in fortifying bovine-milk-based infant formulas with Lf. The pI of Lf is ≈9.0, i.e. it is cationic at the pH of milk, whereas most milk proteins are anionic, and it can be isolated on an industrial scale by adsorption on a cation-exchange resin. Hydrolysis of Lf by pepsin yields peptides called lactoferricins, which are more bacteriostatic than Lf and their activity is independent of iron status. Milk also contains a low level of serum transferrin.

Milk contains a copper-binding glycoprotein—ceruloplasmin—also known as ferr oxidase (EC 1.16.3.1). Ceruloplasmin is an α2-globulin with a MW of ≈ 126 kDa; it binds six atoms of copper per molecule and may play a role in delivering essential copper to the neonate.

Glutathione peroxidase (GTPase) is a selenium-containing protein. It has been reported that milk contains GTPase and that it binds 30% of the total selenium in milk (see Fox and Kelly, 2006b). GTPase has no known enzymatic function in milk; the activity attributed to GTPase in milk may be due to sulfydryl oxidase.

β2-Microglobulin
β2-Microglobulin, initially called lactollin, was first isolated from bovine acid-precipitated casein in 1963 by M. L. Groves. Lactollin, reported to have a MW of 43kDa, is a tetramer of β2-microglobulin, which consists of 98 amino acids, with a calculated MW of 11 636 Da. β2-Microglobulin is a component of the immune system and is probably produced by proteolysis of a larger protein, mainly within the mammary gland; it has no known significance in milk.

Osteopontin
Osteopontin (OPN) is a highly phosphorylated acidic glycoprotein, consisting of 261 amino acid residues with a calculated MW of 29 283 Da (total MW of the glycoprotein, ≈60 000 Da). OPN has 50 potential calcium-binding sites, about half of which are saturated under normal physiological concentrations of calcium and magnesium.

OPN occurs in bone (it is one of the major non-collagenous proteins in bone), in many other normal and malignant tissues, and in milk and urine; it can bind to many cell types. It is believed to have a diverse range of functions (Bayless et al., 1997) but its role in milk is not clear. A rapid method for the isolation of OPN from milk was reported by Azuma et al. (2006) who showed that it binds Lf, lactoperoxidase and 1gs, and may serve as a carrier for these proteins. An acidic whey protein fraction that contains OPN reduces bone loss in ovariectomized rats (Kruger et al., 2006).

Vitamin-binding proteins
Milk contains binding proteins for at least the following vitamins: retinol (vitamin A, i.e. β-Lg), biotin, folic acid and cobalamine (vitamin B12). The precise role of these proteins is not clear but they probably improve the absorption of vitamins from the intestine or act as antibacterial agents by rendering vitamins unavailable to bacteria. The concentration of these proteins varies during lactation but the influence of other
factors such as individuality, breed and nutritional status is not known. The activity of these proteins is reduced or destroyed on heating at temperatures somewhat higher than high temperature, short time (HTST) pasteurization.

**Angiogenins**
Angiogenins induce the growth of new blood vessels, i.e. angiogenesis. They have high sequence homology with members of the RNase A superfamily of proteins and have RNase activity. Two angiogenins (ANG-1 and ANG-2) have been identified in bovine milk and blood serum; both strongly promote the growth of new blood vessels in a chicken membrane assay. The function(s) of the angiogenins in milk is unknown. They may be part of a repair system to protect either the mammary gland or the intestine of the neonate and/or part of the host-defence system.

**Kininogen**
Two kininogens have been identified in bovine milk, a high (88–129 kDa, depending on the level of glycosylation) and a low (16–17 kDa) MW form. Bradykinin, a biologically active peptide containing nine amino acids, which is released from the high MW kininogen by the action of the enzyme kallikrein, has been detected in the mammary gland, and is secreted into milk, from which it has been isolated. Plasma kininogen is an inhibitor of thiol proteinases and has an important role in blood coagulation. Bradykinin affects smooth muscle contraction and induces hypertension. The biological significance of bradykinin and kininogen in milk is unknown.

**Glycoproteins**
Many of the minor proteins discussed above are glycoproteins; in addition, several other minor glycoproteins have been found in milk and especially in colostrum, the function of which have not been elucidated. One of the high MW glycoproteins in bovine milk is prosaposin, a neurotrophic factor that plays an important role in the development, repair and maintenance of nervous tissue. It is a precursor of saposins A, B, C and D, which have not been detected in milk.

The physiological role of prosaposin in milk is not known, although saposin C, released by digestion, could be important for the growth and development of the young.

**Proteins in the MFGM**
About 1% of the total protein in milk is in the MFGM. Most of the proteins are present at trace levels, including many of the indigenous enzymes in milk. The principal proteins in the MFGM include mucin, adipophilin, butyrophilin and xanthine oxidase (see the section on milk lipids).

**Growth factors**
Milk contains many peptide hormones, including epidermal growth factor, insulin, insulin-like growth factors 1 and 2, three human growth factors (α1, α2 and β), two mammary-derived growth factors (I and II), colony-stimulating factor, nerve growth factor, platelet-derived growth factor and bombasin. It is not clear whether these factors play a role in the development of the neonate or in the development and
functioning of the mammary gland, or both (see Fox and Flynn, 1992; Gauthier et al., 2006; Baumrucker, 2007).

Indigenous milk enzymes
Milk contains about 70 indigenous enzymes, which are minor but very important members of the milk protein system (see Fox and Kelly, 2006a, 2006b). The enzymes originate from the secretory cells or the blood; many are concentrated in the MFGM and originate in the Golgi membranes of the cell or the cell cytoplasm, some of which becomes entrapped as crescents inside the encircling membrane during exocytosis. Plasmin and LPL are associated with the casein micelles and several enzymes are present in the milk serum; many of the latter are derived from the MFGM, which is shed as the milk ages.

The indigenous enzymes are significant for several reasons:

- deterioration of product quality (plasmin, LPL, acid phosphatase, xanthine oxidase);
- bactericidal agents (lactoperoxidase and lysozyme);
- indices of the thermal history of milk (alkaline phosphatase, γ-glutamyltransferase or lactoperoxidase);
- indices of mastitic infection (catalase, acid phosphatase and especially N-acetylglucosaminidase).

The concentrations/activities of the indigenous enzymes in milk show greater inter-species differences than any other constituent, e.g.,

- 3000 times more lysozyme in equine milk and human milk than in bovine milk;
- lactoperoxidase is a major enzyme in bovine milk but is absent from human milk;
- human milk and the milks of a few other species contain bile-salts-stimulated lipase but the milks of most species lack this enzyme;
- the principal lipase in milk is LPL, of which there is <500 times as much in guinea pig milk as in rat milk; and
- bovine milk has a high level of xanthine oxidoreductase (XOR) activity but all other milks that have been studied have low XOR activity because the protein lacks molybdenum (XOR plays a major role in the excretion of fat globules from the mammocyte but in this function it does not act as a non-enzyme).

The reason(s) for these inter-species differences is/are not known but some of them may be significant.

Biologically active cryptic peptides
One of the most exciting recent developments in milk proteins is the discovery that all milk proteins contain sequences that have biological/physiological activities when released by proteolysis. The best studied are phosphopeptides, angiotensin-converting-enzyme inhibitory peptides, platelet-modifying peptides, opiate peptides, immunomodulating peptides and the caseinomacropeptides, which have many biological
properties (see Fox and McSweeney, 2003; Korhonen, 2006; Korhonen and Pihlanto, 2006).

**Casein micelles**

It has been known since the work of H. Schuler in 1818 that the casein in milk exists as large particles, now called casein micelles. The stability of the micelles is critically important for many of the technologically important properties of milk and consequently has been the focus of much research, especially during the past 50 years. Early views and research on casein micelles have been reviewed by Fox and Brodkorb (2008). Although views on the detailed structure of the casein micelle are divided, there is widespread, or unanimous, agreement on their general structure and properties.

Electron microscopy shows that casein micelles are spheres with a diameter in the range 50–500 nm (average ≈120 nm) and a mass ranging from $10^6$ to $3 \times 10^9$ Da (average ≈$10^8$ Da). There are numerous small micelles, but these represent only a small proportion of the mass. There are $10^{14}$–$10^{16}$ micelles/mL of milk, and they are roughly two micelle diameters (≈250 nm) apart. The dry matter of the micelles is ≈94% protein and 6% low-molecular mass species, mainly consisting of calcium phosphate with some magnesium and citrate and trace amounts of other species, referred to collectively as colloidal calcium phosphate (CCP). The micelles bind ≈2.0 g H$_2$O/g protein. They scatter light, and the white color of milk is due largely to light scattering by the casein micelles; the white color is lost if the micelles are disrupted, by dissolving CCP with citrate, EDTA or oxalate, by increasing pH, or by urea (＞5 M) or ethanol (≈35% at 70°C).

**Stability of casein micelles**

The micelles are quite stable to the principal processes to which milk is normally subjected. They are very stable at high temperatures, and withstand heating at 140°C for 15–20 min at pH 6.7. Coagulation is caused by heat-induced changes, e.g. a decrease in pH due to the pyrolysis of lactose to acids, dephosphorylation of casein, cleavage of the carbohydrate-rich moiety of κ-casein, denaturation of the whey proteins and their precipitation on the casein micelles, and precipitation of soluble calcium phosphate on the micelles (see O’Connell and Fox, 2003).

The micelles are stable to compaction (e.g. they can be sedimented by ultracentrifugation and redispersed by mild agitation), to commercial homogenization and to Ca$^{2+}$ concentrations up to at least 200 mM at temperatures up to 50°C. The effects of high pressure (up to 800 MPa) on the casein micelles in bovine, ovine, caprine and buffalo milks have been studied; the size of the micelles increases up to 200–300 MPa but decreases at higher pressure (see Huppertz et al., 2006).

As the pH of milk is reduced, CCP dissolves and is fully soluble at = pH 4.9. Acidification of cold (4°C) milk to pH 4.6, followed by dialysis against bulk milk is a convenient technique for altering the CCP content of milk. If acidified cold milk is readjusted to pH 6.7, the micelles reform, provided that the pH had not been reduced below 5.2. This result seems to suggest that most of the CCP can be dissolved without destroying the structure of the micelles.
Some proteinases, especially chymosin, catalyze a very specific hydrolysis of $\kappa$-casein, as a result of which the casein coagulates in the presence of $\text{Ca}^{2+}$ or other divalent ions. This is the key step in the manufacture of most cheese varieties. The proteinase preparations used for cheesemaking are called rennets.

At room temperature, the casein micelles are destabilized by $\approx40\%$ ethanol at pH 6.7 or by lower concentrations if the pH is reduced. However, if the system is heated to $\geq 70^\circ\text{C}$, the precipitate redissolves and the system becomes translucent. When the system is recooled, the white appearance of milk is restored and a gel is formed if the ethanol-milk mixture is held at 4$^\circ\text{C}$, especially if concentrated milk is used. If the ethanol is removed by evaporation, very large aggregates (average diameter $\approx 3000\,\text{nm}$) are formed. The dissociating effect of ethanol is promoted by increasing the pH (35$\%$ ethanol causes dissociation at $20^\circ\text{C}$ and pH 7.3) or adding NaCl. Methanol and acetone have an effect similar to ethanol, but propanol causes dissociation at $\approx 25^\circ\text{C}$. The mechanism by which ethanol dissociates casein micelles has not been established, but it is not due to the solution of CCP, which is unchanged. The micelles are also reversibly dissociated by urea ($5\,\text{M}$), SDS or raising the pH to $>9$. Under these conditions, the CCP is not dissolved.

The micelles are destabilized by freezing (cryodestabilization) due to a decrease in pH and an increase in the $\text{Ca}^{2+}$ concentration in the unfrozen phase of milk; concentrated milk is very susceptible to cryodestabilization. Cryodestabilized casein can be dispersed by warming the thawed milk to $55^\circ\text{C}$ to give particles with micelle-like properties.

**Micelle structure**

There has been speculation since the beginning of the twentieth century on how the casein particles (micelles) are stabilized (see Fox and Brodkorb, 2008), but no significant progress was possible until the isolation and characterization of $\kappa$-casein by D. F. Waugh and P. H. von Hippel in 1956. The first attempt to describe the structure of the casein micelle was made by Waugh in 1958 and, since then, numerous models have been made and refined. Progress has been reviewed regularly (see Fox, 2003); recent reviews include de Kruif and Holt (2003), Horne (2002, 2003, 2006) and Farrell et al. (2006b).

The principal features that must be met by any micelle model are the following: $\kappa$-casein, which represents $\approx 15\%$ of total casein, must be so located as to be able to stabilize the calcium-sensitive $\alpha_{s1^{-}}$, $\alpha_{s2^{-}}$ and $\beta$-caseins, which represent approximately 85$\%$ of total casein; chymosin and other rennets, which are relatively large molecules (MW $\approx 35\,\text{kDa}$), very rapidly and specifically hydrolyze most of the $\kappa$-casein; when heated in the presence of whey proteins, $\kappa$-casein and $\beta$-Lg (MW $\approx 36\,\text{kDa}$) interact to form a disulfide-linked complex, which modifies the rennet and heat coagulation properties of the micelles.

The arrangement that would best explain these features is a surface layer of $\kappa$-casein surrounding the calcium-sensitive caseins, somewhat analogous to a lipid emulsion in which the triglycerides are surrounded by a thin layer of emulsifier. Most models of the casein micelle propose a surface location for $\kappa$-casein but some early models envisaged $\kappa$-casein serving as nodes in the interior of the micelle (see Fox, 2003).
Removal of CCP causes disintegration of the micelles into particles of MW \( \approx 10^6 \text{Da} \), suggesting that the casein molecules are held together in the micelles by CCP. The properties of the CCP-free system are very different from those of normal milk (e.g. it is precipitated by relatively low levels of Ca\(^{2+}\), it is more stable to heat-induced coagulation and it is not coagulable by rennets). Many of these properties can be restored, at least partially, by an increased concentration of calcium. However, CCP is not the only integrating factor, as indicated by the dissociating effect of urea, SDS, ethanol or alkaline pH. At low temperatures, casein, especially \( \beta \)-casein, dissociates from the micelles.

There has been strong support for the view, first proposed by C. V. Morr in 1967, that the micelles are composed of sub-micelles (\( \approx 10^6 \text{Da} \) and 10–15 nm in diameter) linked together by CCP, giving a micelle with an open, porous structure. On removing CCP (by acidification/dialysis, EDTA, citrate or oxalate), the micelle disintegrates. Disintegration may also be achieved by treatment with urea, SDS, 35% ethanol at 70ºC or pH \( > 9 \). These reagents do not solubilize CCP, suggesting that hydrophobic interactions and hydrogen bonds contribute to micelle structure. Much of the evidence for a sub-micellar structure relies on electron microscopy studies that appear to show variations in electron density, a raspberry-like structure, which was interpreted as indicating sub-micelles.

Views on the proposed structure of the sub-micelles have evolved over the years (see Fox, 2003; Fox and Kelly, 2003). Proposals have included a rosette-type structure similar to that of a classical soap micelle, in which the polar regions of \( \alpha_{s1} \), \( \alpha_{s2} \) and \( \beta \)-caseins are orientated towards the outside of the sub-micelle to reduce electrostatic repulsion between neighboring charged groups; each sub-micelle is considered to be surrounded by a layer (coat) of \( \kappa \)-casein, thus providing a \( \kappa \)-casein coat for the entire micelle.

Several authors have suggested that the sub-micelles are not covered completely by \( \kappa \)-casein and that there are \( \kappa \)-casein-rich, hydrophilic regions and \( \kappa \)-casein-deficient, hydrophobic regions on the surface of each sub-micelle. The latter aggregate via the hydrophobic patches such that the entire micelle has a \( \kappa \)-casein-rich surface layer, but with some of the other caseins on the surface also. In a popular version of this model, it is proposed that the hydrophilic C-terminal region of \( \kappa \)-casein protrudes from the surface, forming a layer 5–10 nm thick and giving the micelles a hairy appearance. This hairy layer, functioning as an ionic brush, is responsible for micelle stability through major contributions to zeta potential (\(-20 \text{mV}\)) and steric stabilization. If the hairy layer is removed through specific hydrolysis of \( \kappa \)-casein or collapsed (e.g. by ethanol), the colloidal stability of the micelles is destroyed, and they aggregate.

A further variant of the sub-unit model envisages two main types of sub-units—one consisting of \( \alpha_{s1} \), \( \alpha_{s2} \) and \( \beta \)-caseins, which are present in the core of the micelle; the other, consisting of \( \alpha_{s1} \) and \( \alpha_{s2} \) and \( \kappa \)-caseins, forms a surface layer. It has also been proposed that \( \beta \)-casein associates to form thread-like structures with which \( \alpha_{s1} \) and \( \alpha_{s2} \)-caseins associate hydrophobically to form the core of the micelle or sub-micelles, which are surrounded by a layer of \( \kappa \)-casein; CCP cements neighboring sub-micelles within the micelle.
Although the sub-micelle model of the casein micelle explains many of the principal features of, and physico-chemical reactions undergone by, the micelles and has been supported widely, it has never enjoyed unanimous support. Indeed new electron microscopy techniques have cast doubts on the authenticity of sub-micelles. Using cryopreparation electron microscopy with stereo-imaging, McMahon and McManus (1998) found no evidence to support the sub-micellar model; they concluded that, if the micelles do consist of sub-micelles, these must be smaller than 2 nm or less densely packed than previously presumed. Like other forms of electron microscopy, field emission scanning electron microscopy showed that casein micelles have an irregular surface, but Dalgleish et al. (2004) concluded that the caseins form tubular structures rather than spherical sub-micelles; in principle, this model seems basically similar to earlier sub-unit models.

Three alternative models have been proposed recently. Visser (1992) proposed that the micelles are spherical conglomerates of randomly aggregated casein molecules held together by amorphous calcium phosphate and hydrophobic bonds, with a surface layer of κ-casein. Holt (1992) considered the casein micelle to be a tangled web of flexible casein molecules forming a gel-like structure in which micro-granules of CCP are an integral feature and from the surface of which the C-terminal region of κ-casein extends, forming a hairy layer. In what he referred to as the dual-binding model, Horne (2002, 2003, 2006) described how casein molecules interact hydrophobically and through calcium phosphate nanoclusters to form micelles. These three models retain the key features of the sub-micellar model, i.e. the cementing role of CCP and the predominantly surface location and micelle-stabilizing role of κ-casein, and differ from it mainly with respect to the internal structure of the micelle.

Inter-species comparison of milk proteins

The milks of the species for which data are available show considerable differences in protein content, from ≈1 to ≈20%. The protein content reflects the growth rate of the neonate of the species, i.e. its requirements for essential amino acids. The milks of all species for which data are available contain two groups of protein—caseins and whey proteins—but the ratio of these varies widely. Both groups show genus- and even species-specific characteristics, which presumably reflect some unique nutritional or physiological requirements of the neonate of the species.

Interestingly, and perhaps significantly, of the milks that have been characterized, human and bovine milks are more or less at opposite ends of the spectrum. Among the general inter-species comparisons of milk proteins are Woodward (1976), Jenness (1973, 1979, 1982), Ginger and Grigor (1999) and Martin et al. (2003); reviews on milk proteins of individual species include: buffalo (Addeo et al., 1977), goat (Trujillo et al., 1977, 2000), sheep (Amigo et al., 2000), camel (Ochirkhuyag et al., 1997; Kappler et al., 1998), yak (Ochirkhuyag et al., 1997), horse (Ochirkhuyag et al., 2000; Park et al., 2006) and sow (Gallagher et al., 1997).

There is considerably more and better information on the inter-species comparison of individual milk proteins than on overall milk composition, probably because only one sample of milk from one animal is sufficient to yield a particular protein for characterization. The two principal milk-specific whey proteins, α-La and β-Lg, from
quite a wide range of species have been characterized and, in general, show a high degree of homology (see Brew, 2003; Sawyer, 2003). The caseins show much greater inter-species diversity, especially in the $\alpha$-casein fraction; all species that have been studied contain a protein with an electrophoretic mobility similar to that of bovine $\beta$-casein, but the $\beta$-caseins that have been sequenced show a low level of homology (Martin et al., 2003).

Sheep’s milk is used mainly for cheese production, with small amounts used for the production of fermented milks; hence the coagulation and gel-forming properties of ovine milk are particularly important. The $\alpha_{s1}$-casein of ovine milk is very heterogeneous—to date, 10 genetic variants have been identified; not only do the properties of the variants differ, but also the concentration of $\alpha_{s1}$-casein varies from 0 to 26% of total casein and, consequently, the total protein content varies considerably and this, in turn, has major effects on the rennet-induced coagulation properties of ovine milk and on the yield and quality of cheese produced therefrom (Amigo et al., 2000; Clark and Sherbon, 2000a, 2000b).

Human $\beta$-casein occurs in multi-phosphorylated form (0–5 mol phosphorus per mol protein; see Atkinson and Lonnerdal, 1989), as does mare’s $\beta$-casein (Ochirkhuyag et al., 2000). Considering the critical role played by $\kappa$-casein, it would be expected that all casein systems contain this protein, but Ochirkhuyag et al. (2000) failed to identify $\kappa$-casein in mare’s milk and suggested that the micelle-stabilizing role was played by $\beta$-casein with zero or a low level of phosphorylation; more recent work has shown that equine milk contains a low level of $\kappa$-casein (Iametti et al., 2001; Egito et al., 2002). Human $\kappa$-casein is very highly glycosylated, containing 40–60% carbohydrate (compared with $\approx 10\%$ in bovine $\kappa$-casein). The $\alpha_s$-casein fraction differs markedly between species; human milk probably lacks an $\alpha_s$-casein whereas the $\alpha$-casein fractions in horse and donkey milks are very heterogeneous. The caseins of only about 10 species have been studied in some detail.

There are very considerable inter-species differences in the minor proteins of milk. The milks of those species that have been studied in sufficient depth contain approximately the same profile of minor proteins, but there are very marked quantitative differences. Most of the minor proteins in milk have some biochemical or physiological function, and the quantitative inter-species differences presumably reflect the requirements of the neonate of the species. The greatest inter-species differences, in some cases 4000-fold, seem to occur in the indigenous enzymes (Fox and Kelly, 2006a, 2006b).

In the milk of all species, the caseins exist as micelles (at least the milks appear white), but the properties of the micelles in the milk of only a few non-bovine species have been studied: caprine (Ono and Creamer, 1986; Ono et al., 1989), ovine (Ono et al., 1989), buffalo (Patel and Mistry, 1997), camel (Attia et al., 2000), mare (Welsch et al., 1988; Ono et al., 1989) and human (Sood et al., 1997, 2002). The appearance and the size of casein micelles in the milk of 19 species—guinea pig, rat, nutria (coypu), dog, cat, grey seal, rabbit, donkey, horse, alpaca, dromedary camel, cow, red deer, sheep, pig, water buffalo, goat, porpoise and man—were studied by Buchheim et al. (1989); the structures of all micelles appeared to be similar on electron microscopy but there were large inter-species differences in size—human
micelles were smallest (64 nm) whereas those of the alpaca, goat, camel and donkey were very large (300–500 nm).

**Salts**

When milk is heated at 500°C for ≈5h, an ash derived mainly from the inorganic salts of milk, and representing ≈0.7% w/w of the milk, remains. However, the elements in the ash are changed from their original forms to oxides or carbonates; the ash contains phosphorus and sulfur derived from caseins, lipids, sugar phosphates or high-energy phosphates. The organic salts, the most important of which is citrate, are oxidized and lost during ashing; some volatile metals, e.g. sodium, are partially lost. Thus, ash does not accurately represent the salts of milk.

However, the principal inorganic and organic ions in milk can be determined directly by potentiometric, spectrophotometric or other methods. The typical concentrations of the principal elements—the macro-elements—are shown in Table 1.2; considerable variability occurs, due, in part, to poor analytical methods and/or to samples from cows in very early or late lactation or suffering from mastitis. Milk also contains 20–25 elements at very low or trace levels. These micro-elements are very important from a nutritional viewpoint: many, e.g. Zn, Fe, Mo, Cu, Ca, Se and Mg, are present in enzymes, and many are concentrated in the MFGM; some micro-elements, e.g. Fe and Cu, are very potent lipid pro-oxidants. Although the salts are relatively minor constituents of milk, they are critically important for many technological and nutritional properties of milk.

Some of the salts in milk are fully soluble but others, especially calcium phosphate, exceed their solubility under the conditions in milk and occur partly in the colloidal state, associated with the casein micelles; these salts are referred to as CCP, although some magnesium, citrate and traces of other elements are also present in the micelles. As discussed earlier, CCP plays a critical role in the structure and stability of the casein micelles. The typical distribution of the principal organic and inorganic ions between the soluble and colloidal phases is summarized in Table 1.2. The principal inorganic and organic ions can be determined or calculated after making certain assumptions; typical values are shown in Table 1.2.

The solubility and the ionization status of many of the principal ionic species are interrelated, especially $\text{H}^+$, $\text{Ca}^{2+}$, $\text{PO}_4^{3-}$ and citrate$^{3-}$. These relationships have major effects on the stability of the caseinate system and consequently on the processing properties of milk. The status of various species in milk can be modified by adding certain salts to milk, e.g. the $\text{Ca}^{2+}$ concentration is reduced by adding $\text{PO}_4^{3-}$ or citrate$^{3-}$; addition of $\text{CaCl}_2$ affects the distribution and ionization status of calcium and phosphate, and the pH of milk.

The precise nature and the structure of CCP are uncertain. It is associated with the caseins, probably via the casein phosphate residues; it probably exists as nanocrystals that include phosphate residues of casein. The simplest stoichiometry is $\text{Ca}_3(\text{PO}_4)_2$ but spectroscopic data suggest that $\text{CaHPO}_4$ is the most likely form.

The distribution of species between the soluble and colloidal phases is strongly affected by pH and temperature. As the pH is reduced, CCP dissolves and is
Milk: an overview

Completely soluble at \(<\approx\text{pH } 4.9\); the reverse occurs when the pH is increased. The solubility of calcium phosphate decreases as the temperature is increased and soluble calcium phosphate is transferred to the colloidal phase, with the release of \(\text{H}^+\) and a decrease in pH:

\[
\text{CaHPO}_4/\text{Ca(H}_2\text{PO}_4)_2 \leftrightarrow \text{Ca}_3(\text{PO}_4)_2 + 3\text{H}^+
\]

These changes are quite substantial and are at least partially reversible on cooling.

As milk is supersaturated with calcium phosphate, concentration of milk by evaporation of water increases the degree of supersaturation and the transfer of soluble calcium phosphate to the colloidal state, with the concomitant release of \(\text{H}^+\). Dilution has the opposite effect.

Milk salts equilibria are also shifted on freezing; as pure water freezes, the concentrations of solutes in unfrozen liquid are increased. Soluble calcium phosphate precipitates as \(\text{Ca}_3(\text{PO}_4)_2\), releasing \(\text{H}^+\) (the pH of the unfrozen liquid may decrease to 5.8). The crystallization of lactose as a monohydrate increases the degree of supersaturation by reducing the amount of solvent water.

There are substantial changes in the concentrations of the macro-elements in milk during lactation, especially at the beginning and end of lactation and during mastitic infection. Changes in the concentration of some of the salts in milk, especially calcium phosphate and citrate, have major effects on the physico-chemical properties of the casein system and on the processability of milk, especially rennet coagulability and related properties and heat stability.

Vitamins

Milk contains all the vitamins in sufficient quantities to enable normal maintenance and growth of the neonate. Cow’s milk is a very significant source of vitamins, especially biotin (B_7), riboflavin (B_2) and cobalamine (B_{12}), in the human diet. For general information on the vitamins and for specific aspects in relation to milk and dairy products, including stability during processing and storage, the reader is referred to a set of articles in Roginski et al. (2003) and in McSweeney and Fox (2007).

In addition to their nutritional significance, four vitamins are significant for other reasons: vitamin A (retinol) and carotenoids are responsible for the yellow-orange color of fat-containing products made from cow’s milk; vitamin E (tocopherols) is a potent antioxidant; vitamin C (ascorbic acid) is an antioxidant or pro-oxidant, depending on its concentration; vitamin B_2 (riboflavin), which is greenish-yellow, is responsible for the color of whey or ultrafiltration permeate, co-crystallizes with lactose and is responsible for its yellowish color, which may be removed by recrystallization or bleached by oxidation. It also acts as a photocatalyst in the development of light-oxidized flavor in milk, which is due to the oxidation of methionine.

Water

Water is the principal constituent in the milk of most species. In addition to meeting the requirement of the neonate for water, the water in milk serves as a solvent for milk salts, lactose and proteins, and affects their properties and stability. It controls the rate of many reactions, e.g. Maillard browning, lipid oxidation, enzyme activity and microbial growth, thus affecting the stability of milk and milk products. The preservation of milk products is effected by reducing the water activity by dehydration or by adding sugar or NaCl; the stability and the quality of most dairy products are strongly affected by their water content and small differences can cause major instability problems.


Conclusions

Milk is a very complex fluid. It contains several hundred molecular species, mostly at trace levels. Most of the micro-constituents are derived from blood or mammary tissue but most of the macro-constituents are synthesized in the mammary gland and are milk specific. The constituents of milk may be in true aqueous solution (e.g. lactose and most inorganic salts), or as a colloidal solution (proteins, which may be present as individual molecules or as large aggregates of several thousand molecules, called micelles) or as an emulsion (lipids). The macro-constituents can be fractionated readily and are used widely as food ingredients. The natural function of milk is
to supply the neonate with its complete nutritional requirements for a period (sometimes several months) after birth and with many physiologically important molecules, including carrier proteins, protective proteins and hormones.

The properties of milk lipids and proteins may be readily modified by biological, biochemical, chemical or physical means and thus converted into novel dairy products.

In this overview, the chemical and physico-chemical properties of milk sugar (lactose), lipids, proteins and inorganic salts were discussed. There is a vast primary literature and a range of specialist textbooks on the technology used to convert milk into a range of food products but these aspects were not considered herein.

References


The comparative genomics of tammar wallaby and Cape fur seal lactation models to examine function of milk proteins

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Abstract

The composition of milk includes nutritional and developmental factors that are crucial both to the function of the mammary gland and to the growth and physiological development of the suckled young. This chapter examines the option of exploiting the comparative biology of species with extreme adaptation to lactation to identify milk protein bioactive components with these functions that are likely to have commercial potential either as a nutraceutical in functional foods or as pharmaceuticals. Increasingly, we find that these molecules are more readily identified in animal models with very different reproductive strategies than by screening...
laboratory and livestock species that give birth to a developed neonate and subsequently provide milk with unchanging composition during lactation.

Two animal models with very different lactation cycles are the tammar wallaby (*Macropus eugenii*) and the Cape fur seal (*Arctocephalus pusillus pusillus*). The tammar wallaby has adopted a reproductive strategy that includes a short gestation (26.5 days), birth of an immature young and a relatively long lactation (300 days). The composition of the milk changes progressively during the lactation cycle and this is controlled by the mother and not by the sucking pattern of the young. The tammar wallaby can practice concurrent asynchronous lactation; the mother provides a concentrated milk for an older animal that is out of the pouch, and a dilute milk from an adjacent mammary gland for a newborn pouch young, demonstrating that the mammary gland is controlled locally. These changes in milk composition control the development of the young and therefore provide new opportunities to identify proteins regulating specific developmental processes in the pouch young.

The second study species, the Cape fur seal, has a lactation that is characterized by a repeated cycle of long at-sea foraging trips (up to 23 days) alternating with short suckling periods of 2–3 days on-shore. Lactation almost ceases while the seal is off-shore but the mammary gland does not progress to apoptosis and involution. It is likely that specific milk proteins have a role both to reduce apoptosis in the mammary gland during foraging and to meet the challenges of growth and fasting physiology of the pup on-shore. Technology platforms using genomics, proteomics and bioinformatics have been used to exploit these models to identify milk bioactive components. In addition, the availability of sequenced marsupial, dog and bovine genomes permits rapid transfer of information to the cow to provide outcomes for the dairy industry.

### Introduction

The composition of milk includes all the factors required to provide appropriate nutrition for growth of the neonate. However, it is now clear that milk also comprises bioactive molecules that play a central role in regulating developmental processes in the young and providing a protective function for both the suckled young and the mammary gland during the time of milk production (Trott *et al*., 2003; Waite *et al*., 2005). Identifying these bioactive components and their physiological function in eutherians can be difficult and requires extensive screening of milk components to meet the demand for foods that improve well-being and options for the prevention and treatment of disease.

In addition, human targets for milk bioactive components are focused to specific and changing health needs during the progression from infant to elderly (Figure 2.1). For example, the need to support growth and development in early life differs from the maintenance of muscle, eyes and brain function in the elderly. Health issues such as obesity, diabetes, hypertension and cancer increase with an aging population. Therefore, functional foods that both improve general nutrition and well-being and target prevention and treatment of health problems are very attractive. New animal models with unique reproductive strategies are now becoming increasingly relevant to search for these factors. This chapter focuses on two animal models that provide new insights into how the lactation cycle is regulated and how, when combined with technology platforms that include genomics, proteomics and bioinformatics, it can be exploited to
identify milk proteins with bioactivity and subsequent application as nutraceuticals and pharmaceuticals.

The lactation cycle is common to all mammals, although marsupials and some pinnipeds have evolved a reproductive strategy that is very different from that of most eutherians. These animal models have an extreme adaptation to lactation and have increased our understanding of the function of milk proteins by revealing mechanisms that are present but not readily apparent in many eutherian mammals (Brennan et al., 2007). Eutherians have a long gestation relative to their lactation period, and the composition of the milk does not change substantially.

In contrast, reproduction in marsupials such as the tammar wallaby (*Macropus eugenii*) is characterized by a short gestation followed by a long lactation, and all the major milk constituents change progressively during lactation (Nicholas, 1988a, 1988b; Tyndale-Biscoe and Janssens, 1988). The tammar wallaby neonate is altricial and remains attached to the teat for the first 100 days of lactation, and may be considered as a fetus maintained in the pouch as opposed to the uterus. The conversion of milk to body mass is very similar to the conversion of precursors to body mass observed in the eutherian fetus in utero (Tyndale-Biscoe and Janssens, 1988). Therefore, examination of the milk provides a unique opportunity to identify specific molecules that play a primary role in regulating the development of the young.

The lactation cycle of the Cape fur seal (*Arctocephalus pusillus pusillus*) comprises maternal foraging and infant nursing that are spatially and temporally separate (Bonner, 1984). Lactating mothers suckle offspring over a period of 4–12 months and females alternate between short periods on-shore suckling their young and longer periods of up to 3–4 weeks foraging at sea (Gentry and Holt, 1986). Cape fur seals provide new opportunities to examine a potential role for milk in protecting the

![Figure 2.1 Nutraceuticals for the lifecycle. Potential life-stage-specific targets for milk bioactive components address the specific health needs associated with the progression from infant to elderly.](image-url)
mammary gland from cell death and infection while the lactating animal is foraging, and to examine a function for milk in the growth and protection of the pup that remains on-shore.

The tammar wallaby (*Macropus eugenii*)

**Milk protein gene expression; naturally occurring gene knockouts used to assess protein function**

Lactation in the tammar wallaby has been divided into phases that are defined by the composition of the milk and the apparent sucking pattern of the young (Figure 2.2; Nicholas *et al.*, 1995, 1997). Phase 1 comprises a 26.5-day pregnancy followed by

**Figure 2.2** The lactation cycle of the tammar wallaby. (a) Development of the pouch young from day 6 to day 220 of age. (b) The lactation cycle in the tammar wallaby has been divided into four phases characterized by changes in milk composition and the sucking pattern of the pouch young. (c) Expression of the major milk protein genes during the lactation cycle. The α-casein, β-casein, α-lactalbumin and β-lactoglobulin genes are induced at parturition and expression remains elevated for the entire lactation. The genes encoding ELP (early lactation protein), WAP (whey acidic protein) and the LLPs (late lactation proteins A and B) are expressed only for specific phases of the lactation cycle.
parturition, and the subsequent 200 days of Phase 2 are characterized by lactogenesis and the secretion of small volumes of dilute milk high in complex carbohydrate and low in fat and protein. The pouch young remains attached to the teat for approximately the first 100 days (Phase 2A), after which it relinquishes permanent attachment to the teat and sucks less frequently while remaining in the pouch (Phase 2B). The onset of Phase 3 of lactation (200–330 days) is characterized by temporary exit from the pouch by the young, a large increase in milk production and a change in the composition of the milk to include elevated levels of protein and lipid, and low levels of carbohydrate.

There is a progressive increase in protein concentration (Nicholas, 1988a, 1988b) and protein production during lactation, and the composition of the proteins changes considerably (Nicholas, 1988a, 1988b; Nicholas et al., 1997; Simpson et al., 2000; Trott et al., 2002). Interestingly, macropodids such as the tammar wallaby can practice concurrent asynchronous lactation; the mother provides a concentrated milk high in protein and fat for an older animal that is out of the pouch, and a dilute milk low in fat and protein but high in complex carbohydrate from an adjacent mammary gland for a newborn pouch young (Nicholas, 1988a, 1988b; Figure 2.3). This phenomenon shows that the mammary gland is controlled locally and it is likely that specific milk factors contribute to this process (Trott et al., 2003).

To provide more detailed information on changes in milk protein during lactation in the tammar wallaby, we recently developed a custom tammar wallaby mammary expressed sequence tag (EST), array printed with 10,000 cDNAs representing genes

**Figure 2.3** Concurrent asynchronous lactation. The pouch has been retracted to expose the four mammary glands. A 6-day-old young is attached to a teat from a mammary gland secreting Phase 2A milk. An older animal at approximately 275 days of age has vacated the pouch and sucks from the elongated teat, which provides Phase 3 milk from the enlarged mammary gland. The remaining two teats are from quiescent mammary glands.
expressed across the lactation cycle to transcript profile the mammary gland at all
the major stages of pregnancy and lactation. A comprehensive set of cDNA libraries
was produced from mammary tissues collected throughout the lactation cycle of the
tammar wallaby and a total of 14,837 ESTs were produced by cDNA sequencing.
A database was established to provide sequence analysis and sequence assembly, pro-
tein and peptide prediction and identification of motifs correlated with bioactivity. The
microarray was used to transcript profile approximately 5000 genes; Figure 2.4 shows
the expression profile of the major milk protein genes during lactation. These micro-
array data confirm earlier published results using Northern blot analysis examining

![Graph showing gene expression over lactation phases](image)

**Figure 2.4** Microarray analysis of genes expressed in the tammar wallaby mammary gland during lactation. The phases of lactation are described in Figure 2.2. (a) The major milk protein genes expressed throughout lactation. (b) The major milk proteins expressed at specific phases of the lactation cycle.
the timing and level of expression of these genes (Bird et al., 1994; Simpson et al., 2000; Trott et al., 2002) at each major phase of the lactation cycle.

There are two temporally different patterns of milk protein gene expression during the lactation cycle: one group of genes is induced to high levels around parturition and is expressed throughout lactation; a second group of genes is expressed highly only during specific phases of lactation (Simpson and Nicholas, 2002). For example, the genes for the whey proteins β-lactoglobulin, trichosurin and α-lactalbumin, and the α-casein and β-casein genes, are induced co-ordinately around the time of parturition, and are expressed for the duration of lactation (Figure 2.4a). In contrast, the early lactation protein (ELP) gene is expressed at very high levels in Phase 2A, the whey acidic protein (WAP) gene is expressed most highly in Phase 2B and a gene coding for an outlier lipocalin protein, referred to as late lactation protein A (LLP-A), is highly expressed in Phase 3 (Figure 2.4b).

A second LLP gene referred to as LLP-B is not on the microarray but is first expressed after 200 days of lactation (Trott et al., 2002). Whereas some milk proteins may be specific to the tammar wallaby, in most cases, orthologues of the proteins are found in other species (Simpson and Nicholas, 2002). Therefore, the tammar wallaby is now emerging as a valuable model to provide a “temporal gene knockout” that allows for a more accurate assessment of the role of the gene product in regulating either a specific stage of development of the young or mammary function during each phase of the lactation cycle.

Changes in milk composition regulate growth of the tammar wallaby pouch young

A study by Trott et al. (2003) examined the hypothesis that the sucking pattern of the pouch young controls the pattern of milk secretion in the tammar wallaby. To test this hypothesis, groups of 60-day-old pouch young were fostered at 2-weekly intervals on to one group of host mothers so that a constant sucking stimulus on the mammary gland was maintained for 56 days. This allowed the lactational stage to progress ahead of the age of the young.

Analysis of the milk in fostered and control groups showed that the timing of changes in the concentration of protein and carbohydrate was unaffected by altering the sucking pattern. However, the rate of growth and development of the foster pouch young was significantly increased relative to that of the control pouch young. This probably resulted from the foster pouch young ingesting more milk of higher energy content and milk with a composition that was approximately 50 days more advanced than the milk normally consumed at their age (Trott et al., 2003). It was concluded from these studies that the lactating tammar wallaby regulates both milk composition and the rate of milk production, and that these determine the rates of pouch young growth and development, irrespective of the age of the pouch young. Therefore, the tammar wallaby allows the correlation of milk with a specific composition, particularly the milk proteins, with defined developmental stages of the suckled young.

More recent experiments (Waite et al., 2005) have extended the studies of Trott et al. (2003) and have examined a role for milk in specific changes in the development
The comparative genomics of tammar wallaby and Cape fur seal lactation models

of the gut of the tammar wallaby pouch young. Parietal cells positive for the alpha sub-unit of the H\(^+\)/K\(^+\)-ATPase pump were present in both the forestomach and the hindstomach of pouch young at 168 days of age (Figure 2.5). However, at 260 days of age, parietal cells were seen in sections from the hindstomach but were no longer detectable in the forestomach. It is likely that this change in the forestomach is essential to establish the appropriate microflora as the pouch young emerges from the pouch and starts to eat herbage.

A similar cross-fostering approach to that used by Trott et al. (2003) accelerated the growth rate of the fostered young but did not accelerate development of the stomach. However, these experiments need to be repeated by fostering 170-day-old pouch young to mothers at day 200 of lactation and then examining the stomach 30 days later. It is conceivable that the milk secreted after day 200 of lactation either contains a factor(s) that leads to a decrease in parietal cells or has lost a factor that is required to maintain parietal cells in the forestomach.

Identification of milk bioactive components in tammar wallaby milk using a functional genomics platform

It is likely that milk bioactive components have a variety of roles in regulating both mammary gland function and development of the suckled young. Depending on their function, some bioactive components will be required continuously whereas
others will have a role at specific stages of lactation and development of the young. Therefore, it is likely that their pattern of secretion can be matched to the developmental profile of the suckled young and may follow similar trends to the major milk proteins described in Figure 2.4.

Generally, two approaches can be used to identify bioactive proteins secreted in milk. The first is a proteomics approach that fractionates milk using a variety of separation technologies and examines fractions for bioactivity. A second approach uses a genomics platform to identify the differentially expressed genes that code for a secreted protein. Adopting the latter approach, the tammar wallaby EST database and the microarray data were used to predict the cDNAs coding for proteins with a signal peptide which represented differentially expressed genes when assessed by microarray analysis (M. Digby, C. Lefevre and K. R. Nicholas, unpublished data). This approach identified 75 genes, excluding all the identified major milk proteins, with the expression profile shown in Figure 2.6.

It is clear that the pattern of expression of these genes is variable and that many proteins are secreted into the milk at specific stages of the lactation cycle. Individual proteins encoded by the cDNA were expressed in vitro and the corresponding proteins were examined for activity in cell-based assays for immune modulation, inflammatory responses, growth and differentiative effects. At least 30 of these proteins had at least one of these bioactivities. In addition, some of the proteins showed a capacity
to alter the development of three-dimensional structures of tammar wallaby mammary epithelial cells in a mammosphere culture model, raising the possibility that these proteins may also contribute to growth and function of the mammary gland during lactation.

A causal relationship between these bioactive components and the development of the pouch young remains to be established. However, bovine orthologues for each of the new tammar wallaby bioactive components have been identified and subsequent research will focus on identifying these factors in bovine milk, determining if they can be recovered from waste streams during milk processing, and assessing market application and potential.

**Fractionation of milk and analysis of bioactive proteins**

The fractionation of milk from tammar wallabies at the major phases of lactation provides an opportunity to identify fractions with bioactivity and to follow changes in activity in specific fractions across the duration of lactation. Temporal changes in activity would probably suggest an association with particular developmental process(es) in the developing young, although a potential autocrine effect on mammary gland function could also be considered. The microarray used to identify differentially expressed genes, described previously, was limited to approximately 5000 genes and it is likely that the number of differentially expressed genes coding for secreted bioactive components will increase when a microarray with whole genome coverage becomes available.

The genome of another marsupial, *Monodelphis domestica*, has been sequenced to eight-fold coverage and the genome of the tammar wallaby has been sequenced to two-fold coverage; this information will probably lead to a marsupial whole transcriptome microarray. The disadvantage of analyzing fractionated whey is that often the sample assayed comprises a mixture of proteins, requiring additional work to identify the specific protein(s) associated with the bioactivity. However, this approach potentially allows identification of proteins and peptides that are not currently represented on the tammar wallaby EST microarray and which may have either undergone post-translational processes or resulted from alternative splicing of genes.

Whey samples collected from tammar wallabies in Phases 2B and 3 were fractionated by reverse phase high performance liquid chromatography (HPLC) as shown in Figure 2.7. Proteins and peptides bound to the column were eluted with a linear acetonitrile gradient and the 60 fractions were analyzed in cell-based assays for potential to stimulate growth and differentiation of cells (ERK activity), pro- and anti-apoptotic assays, and assays for stimulation and inhibition of immune response. Analysis of ERK showed activity in specific fractions (Figure 2.7) in whey from tammar wallabies at Phase 2B. However, activity was evident not only in the same fractions in whey from tammar wallabies in Phase 3 of lactation, but also in an additional set of whey fractions. A specific role for this factor/s is not yet apparent, but it has considerable potential because it is correlated with a dramatic increase in milk production and growth of the young as they emerge from the pouch and begin to eat herbage and consume milk.
WAP; a potential role in development of the young, and local regulation of mammary development in the tammar wallaby

Although increasing attention is directed to identifying the minor protein components of milk that have bioactivity, it is surprising how little is known about the function of the major whey proteins in most species. Identifying the primary function for milk proteins such as β-lactoglobulin and WAP has proved to be challenging. WAP is a major whey protein and is secreted throughout lactation in many eutherian species including rat, mouse, rabbit, camel and pig (Simpson et al., 2000; Simpson and Nicholas, 2002). WAP is secreted in the milk of all marsupials studied to date including the tammar wallaby (Simpson et al., 2000), the red kangaroo (Nicholas et al., 2001), the brush-tailed opossum (Demmer et al., 2001) and the fat-tailed dunnart

Figure 2.7 Fractionation and analysis of tammar wallaby whey. Whey from milk collected in Phase 2B and Phase 3 of lactation was fractionated by reverse phase HPLC. Proteins and peptides bound to the column were eluted with a linear acetonitrile gradient and the 60 fractions were analyzed for potential to stimulate ERK (growth and differentiation of cells) in a cell-based assay.
(De Leo et al., 2006), and WAP mRNA has been detected in the lactating mammary gland of a stripe-faced dunnart and a short-tailed opossum (D. Topcic and K. R. Nicholas, unpublished data).

However, the secretion of the protein is specific to a phase of lactation in all the marsupials mentioned above and therefore the tammar wallaby provides a naturally occurring gene knockout model to assess the role of this protein in regulating specific stages of development of the young, the mammary gland or potentially both. The amino acid sequence of WAP from the two monotremes platypus and echidna has been reported (Simpson et al., 2000; Sharp et al., 2007), but the pattern of secretion in these species is yet to be established.

Alignment of the amino acid sequences of WAPs from marsupial, monotreme and eutherian species shows limited sequence identity (Simpson et al., 2000; Sharp et al., 2007). However, these proteins are recognized by the presence of the WAP motif (KXGXCP) and a domain structure known as the four-disulfide-core (4-DSC) domain, which consists of eight cysteine residues (Ranganathan et al., 1999). The WAPs secreted in the milk of eutherians have two 4-DSC domains (Simpson et al., 2000), whereas the WAPs in all marsupials and monotremes studied to date have three domains (Simpson et al., 2000; Demmer et al., 2001; De Leo et al., 2006; Sharp et al., 2007). The five exons of the marsupial and monotreme WAP genes contrast with the four exons only of the eutherian WAP genes (De Leo et al., 2006; Sharp et al., 2007). The significance of the loss of the third domain in eutherians remains unclear, particularly as convincing evidence for biological function of milk WAP remains to be established.

Recent studies have shown that mouse WAP added to a culture medium of mouse HC-11 cells (a mouse mammary epithelial cell line) is anti-proliferative, and may act by an autocrine or paracrine mechanism (Nukumi et al., 2004). This is consistent with reports showing that over-expression of WAP in transgenic mice inhibits development of the mammary gland and the secretion of milk (Burdon et al., 1991). Studies in our laboratory have used in vitro models to examine the effect of tammar wallaby WAP on proliferation of an epithelial-enriched population of tammar wallaby mammary cells. In contrast to the inhibitory action of mouse WAP on proliferation of HC-11 cells, results show that tammar wallaby WAP added to culture medium stimulates proliferation of mammary epithelial cells and increases expression of the cell cycle gene cyclin D1 (Figure 2.8).

Earlier studies have shown that DNA synthesis in mammary tissue is higher in Phase 2 than in Phase 3 (Nicholas, 1988a), which is consistent with a potential role of tammar wallaby WAP in development of the mammary gland. This is further supported by our studies showing that tammar wallaby WAP stimulates DNA synthesis in primary cultures of mammary epithelial cells (D. Topcic, unpublished data). Interestingly, studies using a WAP gene knockout mouse have shown that mammary development is normal. However, the pups had limited development at the later stages of lactation (Triplett et al., 2005).

As tammar wallaby WAP is a major milk protein that is secreted during the middle third of lactation when the pouch young’s diet comprises only milk, it could be speculated that this protein plays a specific role in the development of both the mammary
gland and the suckled young at this time. The apparent milk WAP genes in human, cow, ewe and goat have accumulated mutations, making them non-functional pseudo-genes (see Hajjoubi et al., 2006). It is likely that a putative function for WAP in mammary development is predominantly found only in marsupials and monotremes, and that the activity has been lost in human, cow, ewe and goat because of a loss of evolutionary pressure on this protein that relates to changes in the reproductive strategy of eutherians. Therefore, the marsupial may be a more appropriate model to explore the potential of other major milk proteins such as β-lactoglobulin which, like WAP, is not

![Graph showing proliferation of Wall-MEC](image)

**Figure 2.8** (a) Proliferation of wallaby mammary epithelial cells (Wall-MEC) cultured in the presence and absence of tammar wallaby WAP (tWAP). The results shown are after day 3 of treatment with (+) and without (−) tWAP. The vertical bars indicate SEM (n = 3), P < 0.05. (b) Expression of the cyclin D1 gene in Wall-MEC grown in the presence and absence of tammar wallaby WAP (day 3 post-treatment). Expression of the cyclin D1 and GAPDH genes was determined by RT-PCR analyses. The relative expression levels of the cyclin D1 gene in Wall-MEC was quantified by National Institutes of Health image software and was presented as a proportion of the housekeeping gene GAPDH. Reprinted from Brennan et al. (2007). URL http://jds.fass.org/, © American Dairy Science Association.

The tammar wallaby (*Macropus eugenii*)
expressed in all eutherians. Resolution of the causes and effects of these comparative differences will provide valuable insights into the roles these molecules perform.

A role for milk in the control of mammary function in the tammar wallaby

The expression of milk protein genes is regulated concurrently by systemic endocrine factors, by paracrine factors such as the extracellular matrix and by autocrine factors secreted in the milk. Previous studies using a tammar wallaby mammary explant culture model (Nicholas and Tyndale-Biscoe, 1985) have shown that different combinations of insulin, cortisol and prolactin are required for expression of the α- and β-casein genes, and whey protein genes including α-lactalbumin and β-lactoglobulin (Simpson and Nicholas, 2002).

Tammar wallaby mammary explants from late pregnant tammar wallabies can be induced with insulin, cortisol, prolactin, thyroid hormone and oestrogen to express the WAP gene (Simpson et al., 2000). Therefore, the inhibition normally observed in vivo during Phase 2A, and the subsequent induction of WAP gene expression around 100 days post-partum, may be hormonally regulated. In addition, the LLP genes can be down-regulated in mammary explants from Phase 3 tammar wallabies and then restimulated with insulin, cortisol and prolactin, but expression of these genes cannot be induced in mammary explants from pregnant tammar wallabies with any hormone combination tested (Trott et al., 2002). Either the appropriate hormonal milieu was not used and additional hormones are required, or the tissue requires additional serum or local mammary factors to express these genes.

This conclusion is supported by an earlier study showing that constructs with up to 1.8 kb of the LLP-A gene promoter did not express a reporter gene after transfection into CHO cells incubated with insulin, cortisol and prolactin, whereas control experiments showed that a rat β-casein gene construct was hormone responsive. In addition, the same construct was not expressed in lactating transgenic mice (Trott et al., 2002).

Recent studies in our laboratory have demonstrated that constructs comprising short-tailed opossum LLP-A and tammar wallaby LLP-B promoters (up to 5 kb of DNA) with a reporter gene were not transcriptionally active following transfection into HC-11 cells, regardless of the hormonal combination in the culture. However, unlike the LLP gene promoters, constructs with various WAP promoter fragments from tammar wallaby, short-tailed opossum and stripe-faced dunnart showed increased transcriptional activity when prolactin was added to the medium (Topcic and Nicholas, unpublished data). This suggests that the mechanisms controlling the expression of these milk protein genes in marsupial species are likely to be different from those in eutherians.

There is increasing evidence to suggest that milk plays an important role in regulating mammary epithelial function and survival, and this is particularly evident during involution (Brennan et al., 2007). Apoptosis was induced preferentially in the sealed teats of lactating mice (Li et al., 1997; Marti et al., 1997), while the litter suckled successfully on the remaining teats, which indicates that cell death is stimulated by an intra-mammary mechanism that is sensitive to milk accumulation (Quarrie et al.,
1995). A protein known as the feedback inhibitor of lactation (FIL) has been suggested as a candidate and is secreted in the milk of the tammar wallaby (Hendry et al. 1998) and other species. It acts specifically through interaction with the apical surface of the mammary epithelial cell to reduce secretion (Wilde et al., 1995).

More recent studies using the tammar wallaby mammary explant culture model (Nicholas and Tyndale-Biscoe, 1985) to examine the process of involution have confirmed the likely role of milk, and particularly putative autocrine factors, for controlling mammary function during involution (Brennan et al., 2007). Mammary explants from pregnant tammar wallabies were cultured for 3 days with insulin, cortisol and prolactin to induce milk protein gene expression. To mimic involution, all hormones were subsequently removed from the culture medium for 10 days to down-regulate expression of the milk protein genes (Figure 2.9). Surprisingly, the explants retained the same level of response during a subsequent challenge with lactogenic hormones.

Previous studies have shown that there is limited secretion of milk proteins from tammar wallaby mammary explants, but it is unlikely that milk constituents accumulated to elevated concentrations (Nicholas and Tyndale-Biscoe, 1985). The maintenance of epithelial cell viability and hormone responsiveness in explants cultured

![Figure 2.9](image-url)  

**Figure 2.9** Northern blot analysis of β-lactoglobulin, α-casein and α-lactalbumin gene expression in tammar wallaby mammary explants. Gene expression is shown in: mammary tissue from day 24 pregnant tammar wallabies prior to culture (t₀); explants cultured in media with insulin, cortisol and prolactin (IFP) for 3 days; explants subsequently cultured in the absence of hormones (NH) for 10 days; and following the re-introduction of IFP for 3 days. The length of culture in days is shown by the subscript. Total RNA (10μg, lower panel) was assayed by Northern blot analysis using [α-32P]dCTP-labelled cDNA probes for the β-lactoglobulin, α-casein and α-lactalbumin genes (upper panels). Arrows indicate transcript size in nucleotides and RNA ribosomal bands.
in the absence of hormones is consistent with a more active mechanism, such as the accumulation of local factors in the milk being the primary stimulus for apoptosis of mammary epithelial cells in the tammar wallaby mammary gland. This model permits the uncoupling of hormone and milk-regulated involution and, clearly, a primary outcome of these studies is evidence for the extraordinary capacity for the survival and maintenance of hormone responsiveness by tammar wallaby mammary epithelial cells cultured in a chemically defined medium with no exogenous hormones and growth factors.

Assuming that bovine mammary epithelial cells show similar characteristics to tammar wallaby mammary epithelial cells, it is likely that milk components will play a major role in the shape of the lactation curve in dairy cattle following peak lactation at approximately 20 weeks of milk production. The lactation cycle in dairy cattle includes a period of increasing milk yield in early lactation followed by a steadily declining yield for the remainder of lactation. The amount of milk produced during lactation is determined by the peak yield and the persistency of lactation (see McFadden, 1997). Milk production is largely a function of the number and the activity of secretory cells in the udder, which decline significantly between the time of peak yield and late lactation. Therefore, it follows that approaches to address the decline in milk yield and lactational persistency after peak lactation must involve changes to the frequency of apoptosis in mammary secretory cells.

Endocrine treatment of cattle with bovine somatotrophin increases milk yield but, in many cases, persistency is not altered. Furthermore, although moderate heritability of persistency suggests that selection for this trait is possible, it is achieved at the cost of milk yield. More recently, increased frequency of milking has been shown to increase milk yield, suggesting that the mammary gland has a local intrinsic resistance to regression. Identification of the milk components that impact significantly on mammary cell fate may provide new approaches for strategies to improve lactational persistency.

The Cape fur seal (*Arctocephalus pusillus pusillus*)

The Cape fur seal suckles its pup for up to 10 months (Gentry and Holt, 1986) and lactation comprises alternate periods of several days on-shore suckling the young and extended periods at sea (Figure 2.10; Bonner, 1984; Ofstedal et al., 1987; Trillmich, 1996). Foraging trips are variable but can extend to 23 days (Gamel et al., 2005). Nursing periods are generally 1–3 days and depletion of the mother’s body reserves necessitates a return to the ocean to forage.

A study by Cane et al. (2005) showed that the mean protein contents of the milks of the Antarctic fur seal (*Arctocephalus gazella*) and the Australian fur seal (*Arctocephalus pusillus doriferus*) were 10.6 and 10.9% respectively, and did not change significantly across the lactation cycle. Electrophoresis of skim milk of the Australian fur seal showed 12 separate bands, representing five caseins and seven whey proteins. The majority of the whey protein was β-lactoglobulin, with possibly
three different isoforms present. However, there was no significant change in the composition of milk proteins during lactation.

More recent studies have shown that milk protein gene expression also decreases during foraging trips as the $\beta$-casein, $\alpha_s2$-casein and $\beta$-lactoglobulin genes are all down-regulated in the mammary gland of the foraging Cape fur seal (*Arctocephalus pusillus pusillus*) (Cane, 2005; Sharp et al., 2006). During time at sea, the lactating mammary gland does not progress to involution but produces less milk compared with that of the on-shore lactating female. For example, milk production in Antarctic fur seals (*A. gazella*) has been shown to continue while the female is foraging at sea but at only 19% of the rate of production on land (Arnould and Boyd, 1995).

**Milk protein gene expression in the lactating Cape fur seal during suckling and foraging**

During lactation on-shore, the mammary alveoli of the Cape fur seal are engorged with milk containing a large amount of lipid (Figures 2.11a, 2.11b; Sharp *et al*., 2006). In contrast, during the mother’s extended foraging trip, the alveoli appear less distended, epithelial cells surrounding the alveoli appear columnar and the lipid component within the milk is decreased. High sequence conservation between the Cape fur seal and the dog, 95% similarity at the DNA level, permits a significant detection rate of measurable hybridization signals between seal cDNA and the Affymetrix canine microarray (Sharp *et al*., 2006).
Figure 2.11  Histological sections of the mammary gland from Cape fur seals: (a) lactating while nursing on-shore and (b) lactating while foraging at sea. Sections are stained with hematoxylin and eosin. Magnification x 100. (c) Milk protein gene expression. β-Casein expression during Cape fur seal lactation cycle. Analysis of expression using canine Affymetrix chips hybridized to cDNA probes generated from RNA from pregnant (placental gestation and non-lactating, n = 2), lactating on-shore (n = 2) and lactating at sea (n = 1) (animals in embryonic diapause) Cape fur seals. (d) Cluster analysis of gene expression profiles from the Cape fur seal mammary gland during different stages of lactation. A total of 1020 Cape fur seal mammary messenger RNA (mRNA) transcripts were identified with expression levels above an intensity of 250 in any sample type. Hierarchical clustering was conducted using Euclidean distance. Pregnant and on-shore lactating data represent an average of two animals. Off-shore data represent a single sample. Reprinted from Current Topics in Developmental Biology, 72, G. P. Schatten, sr ed., J. A. Sharp, K. N. Cane, C. Lefevre, J. P. Y. Arnold, and K. R. Nicholas, Fur Seal Adaptations to Lactation: Insights into Mammary Gland Function, pp. 276–308, New York: Academic Press. Copyright 2006, with permission from Elsevier (see also Plate 2.11).
Expression of the β-casein gene is barely detectable in the mammary gland of pregnant seals whereas the level of expression is significantly elevated in the mammary gland of seals lactating on-shore (Figure 2.11c). However, expression of this gene is also reduced during the foraging trip. Cluster analysis of gene expression profiles in mammary tissue from pregnant, on-shore lactating and off-shore lactating seals revealed that the overall expression profile of the lactating mammary gland of the foraging Cape fur seal is more closely related to the profile of pregnant non-lactating animals (placental gestation) than to the profile obtained from on-shore lactating animals (Figure 2.11d). This result suggests that the interruption of lactation in foraging animals involves a major reprogramming of mammary gland gene expression.

In most mammals during natural weaning, as the alveoli fill with milk due to cessation of suckling, there is a decline in milk protein gene expression in the mammary epithelial cells, and the epithelium regresses and enters involution (Li et al., 1997). This process is characterized by apoptotic cell loss and mammary gland remodelling (Strange et al., 1992; Lund et al., 1996; Metcalfe et al., 1999). Apoptosis associated with involution in the mammary gland of the foraging seal has been analyzed and found to be barely detectable; even after extended periods when there is no sucking stimulus, the gland does not regress (J. A. Sharp et al., unpublished data).

The process by which the lactating seal reduces milk production and avoids entering apoptosis while foraging is unknown. However, it is clear that transcript profiling of the mammary gland of the foraging seal has shown that the acute immune response is associated with involution, presumably to protect the mammary gland from infection during the extended foraging trip while the milk remains in the gland (Sharp et al., 2007). Gene expression profiles associated with survival and preservation of tissue architecture were also either maintained or up-regulated, preventing degradation of mammary tissue. These global gene expression data suggested that the immune and acute phase responses observed in the mouse at involution (Clarkson et al., 2004), and mimicked in the foraging Cape fur seal mammary gland, are independent of the apoptotic phase of involution.

A consequence of reduced nursing is that putative factors responsible for regulating apoptosis are retained in the mammary gland. As discussed previously, it has been shown that the sealing of a single mammary teat of a mouse induces an accumulation of milk, resulting in changes in mammary gene expression and apoptosis within the sealed gland but not of the remaining glands of the same animal (Li et al., 1997; Marti et al., 1997). Studies in lactating animals from a variety of species indicate that regulation of milk secretion may involve an inhibitory factor (Knight et al., 1994; Wilde et al., 1995; Peaker et al., 1998).

As mentioned previously, experiments have identified a small whey protein, termed feedback inhibitor of lactation (FIL) (Wilde et al., 1995), that is synthesized by the secretory epithelial cells of the mammary gland and is secreted into the alveolar lumen along with other milk constituents. It has been proposed that FIL acts on the synthesis and secretory pathway by binding a putative receptor on the apical surface of the epithelial cells (Rennison et al., 1993; Blatchford et al., 1998). Furthermore, FIL may block translation of milk protein transcripts (Rennison et al., 1993) and inhibit secretion of milk constituents.
Preliminary data (Cane, 2005) have demonstrated a FIL-like activity in fractionated seal milk but the level of inhibitory activity measured was similar to that reported for other species (Blatchford et al., 1998). In addition, the activity did not differ in milk collected from seals either arriving on-shore after foraging at sea or after they had been on-shore suckling their pups for 1–2 days. Recently, we showed that mammary-epithelial-cell-enriched fractions from a pregnant Cape fur seal formed three-dimensional structures (mammospheres) in culture (Sharp et al., 2007). The seal cells secreted their own extracellular matrix and formed mammospheres that responded to insulin, cortisol and prolactin in the medium to express milk protein genes. This model will provide new opportunities to assess the role of milk factors on mammary function in the Cape fur seal.

The role of \(\alpha\)-lactalbumin in lactose synthesis is well established. However, studies have suggested that this protein may also be implicated in the process of involution (Hakansson et al., 1995, 1999; Baltzer et al., 2004). Milk from otariid pinnipeds contains little or no lactose (Urashima et al., 2001) and \(\alpha\)-lactalbumin protein has not been detected in otariid milk (Cane et al., 2005) suggesting that this protein is probably absent in these species. It is interesting to speculate that the absence of biologically active \(\alpha\)-lactalbumin in milk may be consistent with the absence of apoptosis in the mammary gland of lactating seals during foraging and that loss of this protein has provided an opportunity to alter the lactational strategy of the otarrid family of seals.

This unique capacity to inhibit apoptosis associated with involution, to maintain mammary epithelial cells in a viable and hormone-responsive state, and to stimulate mammary gland growth to increase milk production during the lactation cycle provides new opportunities to identify the mechanism controlling these events. An understanding of the genetic and/or local factors in the residual milk regulating this process may have application for improving lactational persistency in the dairy cow, either by improved breeding programs or by manipulation of milk factors.

A genomics platform in the Cape fur seal to identify milk bioactive components

The capacity of the Cape fur seal to uncouple involution from the cessation of sucking while foraging at sea is one example of how evolution has adapted this mammal’s lactation cycle. It is likely that the mammary gland also makes other temporal changes such as preventing local infection and reducing inflammatory responses while at sea, and preparing to secrete a milk that delivers protection from infection and the appropriate growth factors to stimulate growth of the gut in the fasting pup that remains on-shore. The reprogramming of this gland offers new opportunities to identify mechanisms and specific mammary factors that are important for the gland to adapt to the stress and challenges during a long foraging trip in the ocean.

To assist in the identification of these molecules, a cDNA library produced from the mammary gland of a lactating off-shore Cape fur seal has been sequenced. Analysis of 10,000 EST sequences, representing approximately 4000 gene transcripts expressed in the lactating off-shore Cape fur seal mammary gland, allows for identification of putative secreted proteins and potentially improves annotation of sequences.
detected by Affymetrix analysis (J. A. Sharp, C. Lefevre and K. R. Nicholas, unpublished data). One interesting observation is the high abundance of lysozyme ESTs in the library, indicating that secretion of this protein in the milk may have a role in protecting the mammary gland of the mother and the gut of the pup from bacterial infection. Further analysis of the EST library has now identified the genes for more than 30 new putative bioactive components and a bovine orthologue for each protein has been identified. These proteins are now being characterized and examined for commercial potential.

Conclusions

This chapter has described two animals with extreme adaptation to lactation and has explored approaches to exploit their unique reproductive strategies to identify new proteins in milk with bioactivity that have the potential to regulate mammary function and growth of the suckled young. In contrast, the lactation cycle in most eutherian mammals is characterized by the initiation of lactation around parturition and the production of milk in which the individual components vary little during the entire period of lactation. For example, in the dairy cow, evolution and selection pressure have led to a lactation cycle that does not include significant changes in the secretory pattern of the milk proteins. Therefore, identification of new protein bioactive molecules in the dairy cow requires a broad screening program and will probably identify molecules in milk that are secreted at low and unchanged levels during lactation.

In contrast, in the tammar wallaby and the Cape fur seal, the temporal pattern of secretion of a milk bioactive component that is correlated with either a specific developmental phase of the suckled young, or altered mammary function, is more likely to indicate a cause-and-effect relationship, and these factors are more likely to have increased commercial value. The increasing availability of sequenced genomes in public databases has underpinned increased interest in comparative genomics and bioinformatics, and our results using the tammar wallaby and Cape fur seal models have led to the identification of many bovine and human orthologues of proteins with bioactivity. Conceptually, it is very likely that the relevance and use of species with an extreme adaptation to reproduction or environmental pressure will become increasingly attractive for improved understanding of the genetic regulation of physiological processes.

References


Significance, origin and function of bovine milk proteins: the biological implications of manipulation or modification


Abstract

The rapid advances in technology for both evaluating and understanding the structure of animal genomes and their functional significance have presented opportunities for scientists to more clearly understand the complexity of bovine milk proteins and the control of their expression. Used in conjunction with proteomic databases, we can start to expand our knowledge of how milk proteins are processed into peptides, which represent much of the biological activity
residing within colostrum and milk. The challenge then remains to translate this information into products that form the basis of a functional foods industry, helping to underpin the commercial viability of the dairy industry. In this chapter, we present a review of the current status of bovine milk genomics and functional genomics, and describe the roles, characteristics and key bioactivities of the major bovine milk proteins and their encrypted peptides. The application of these analytical tools to the full spectrum of lactation strategies adopted by eutherians, marsupials and monotremes to improve our understanding of the milk proteome is discussed.

**Introduction**

As well as being the source of nutrition for the neonates of mammals, milk has long been considered to be a healthy source of dietary proteins and minerals for human consumption, and a global dairy industry has evolved to cater to that need. With competition from other foods and beverages, there has been an impetus for diversification of dairy-based foods, manipulation of milk composition to remove saturated fats, and fortification of milk with fatty acids, minerals and vitamins.

The next evolution of dairy products is to characterize, accentuate and refine naturally occurring biological activities to develop dairy-based functional foods or “nutraceuticals.” Although a significant amount of information in relation to the biological activities present in the major bovine milk proteins is already available, the advances in genomic technologies and the utilization of comparative genomics approaches provide opportunities for further discovery and characterization of the bioactivities in bovine milk.

**Milk genomics: a contemporary approach to milk composition**

Trends in the consumer market are driving an increased focus on the health and dietary functions of food products, and there is an emerging market for milk-derived products. The major focus in dairy is on milk proteins, which are rich in bioactive properties and may therefore be particularly valuable to dairy processors and suppliers providing specialist milk to specification. This area of product development is poised to undergo a significant expansion based on the bovine genome sequencing project.

Milk has been a subject for nutrition and dairy science research over many years, but especially during the post Second World War period, when it received significant support from the British government for its role as a major source of public nutrition. Increasingly, milk is the source of animal protein of choice worldwide because of its ease of handling. Industrial processing of milk into dairy products has been based largely on the insights into milk chemistry that arose from seminal studies conducted during this period. These studies were concerned primarily with nutritional and processing properties of major milk proteins. Recent advances in bovine genomics now extend the capacity to analyze milk at a more detailed level, to provide a basis to understand the properties of milk proteins and, along with parallel advances in
humans and other animals, to open a window on the evolutionary origins and specific benefits of milk. Milk genomics is the study of milk using post-genomic approaches.

**Advances in bovine genome science**

Genomics is concerned with the study of an organism’s gene sequence, gene expression and gene function. Genome science has developed rapidly in the past decade as a result of a vast expansion of genome sequence data, initially arising from the human genome sequencing project, and the concomitant development of applied information technology (bioinformatics). In December 2003, only months after the completion of the human genome sequencing project, an international consortium formally declared the initiation of the bovine genome sequencing project. The human genome sequencing project resulted in the establishment of significant infrastructure, expertise and technological advances. Consequently, sequencing of the bovine genome progressed quickly and was completed in 2005, well ahead of the original schedule.

**DNA sequence**

Bovine genetics, with a focus on the identification of quantitative trait loci (QTL), has been a very active area of study for many years (Khatkar *et al.*, 2004). Using classical genetic and molecular genetic approaches, over 1000 bovine genes had been mapped by the mid 1990s. A phase then followed that was based on candidate gene mapping, and creation and sequencing of bovine expressed sequence tag (EST) libraries. The Institute for Genome Research assembled a bovine gene index that contained information on more than 100 000 sequences. The bovine sequencing strategy employed for the bovine genome sequencing project was a whole genome shotgun approach. An individual animal, a Hereford cow, was chosen as the source of DNA. The total sequencing effort provided approximately 7x coverage of the genome, representing about 2.7 billion base pairs (reviewed in Tellam, 2007). The project complements and in many cases supersedes existing resources for bovine genome science, and provides a platform for developing bovine-specific post-genome research tools and applied industry applications.

**Genome map/structure**

The bovine genome comprises 29 autosomes, plus the X and Y chromosomes. An international bovine bacterial artificial chromosome (BAC) map was produced to assist in assembling the genome, complemented by an integrated mapping strategy using radiation hybrid and existing physical mapping data. A virtual assembly was released in 2005 and the first de novo assembly was released in 2006 (http://genomes.tamu.edu/bovine/). The current genome assembly is version btau_3.1 with a more accurate version 4 due to be released in 2008.

Comparative genetic maps have revealed significant similarity (synteny) between bovine genomes and the genomes of other species. This can be viewed most effectively using the OXGRID display at (http://oxgrid.angis.org.au/). The synteny between
bovine and other species is striking; for example, the mean length of syntenic segments between bovine and human is approximately twice as long as the average length of conserved syntenic segments between human and mouse.

**Polymorphism**

Understanding genome polymorphism within the bovine is a primary goal of the genome sequencing project, and has implications for variants of milk proteins that affect processing characteristics and in some cases consumer health. The sequencing project itself resulted in the discovery of a large number of single nucleotide polymorphisms (SNPs). This effort has been extended to resequence six breeds—Holstein, Angus, Jersey, Brahman, Limousine and Norwegian Red—at low coverage, with the aim of identifying genome-wide SNPs (Tellam, 2007). In addition, mining of existing ESTs has contributed a putative 17344 SNPs (Hawken et al., 2004).

Initial analysis of dairy herds using a subset of SNPs revealed that the predominant cattle breed used internationally for milk production, Holstein Friesian, has low genetic diversity and may be considered to be a single population unit (Zenger et al., 2007). The average minor allele frequency for SNPs was 0.29 based on 9195 SNP genotypes in 1000 bulls (Khatkar et al., 2007). Further, using the same genotyping data, Khatkar and colleagues in the Australian Cooperative Research Center for Innovative Dairy Products determined that the extent of linkage disequilibrium in the Holstein Friesian breed is significantly greater than that reported for humans. They went on to construct the first bovine haplotype map and identified 727 blocks (covering just 2.18% of the genome) with a median length of 2.9 kb.

**Functional genomics**

In addition to increased understanding of dairy herd genetics, the bovine genome sequencing project has led to improved annotation of bovine genes, which underlies comprehensive studies of bovine gene expression. This was enabled by the release of a comprehensive bovine DNA chip. Based on oligonucleotide microarrays and developed using the Affymetrix platform (Dalma-Weiszhausz et al., 2006), the chip contains ≈22000 probesets, representing ≈19000 genes. More recently, a long-oligonucleotide spotted array has been developed by an international consortium of researchers. These tools are now employed for the study of mammary gland gene expression, and consequently lactation functional genomics data are emerging.

Studies reporting lactation profiles using microarrays constructed from a bovine cDNA library or ESTs have already been published (Suchyta et al., 2003). We completed a study of the bovine lactation cycle using the first-release comprehensive bovine Affymetrix DNA chip. This study compared gene expression during pregnancy, lactation and involution phases, and identified highly expressed and differentially expressed genes (unpublished data). As a result, a library of almost 4000 genes that were differentially expressed between phases was created. Utilizing these data, further analysis identified a large group of predicted cow’s milk polypeptides. These peptides are the subject of ongoing validation and development as dairy products.
A complementary approach to gene expression studies of the mammary gland is provided by milk proteomics (Smolenski et al., 2007). Cow’s milk comprises approximately 3.5% protein, 80% of which are caseins; the remainder are predominantly the major whey proteins, but there are also a large number of low-abundant proteins, some of which have significant bioactivity. This area has also benefitted markedly from the bovine genome sequencing project. It is now possible to identify variants of both major and low-abundant proteins in milk based on bovine protein sequences. Together with rapid advances in mass spectrometry, particularly improvements in sensitivity and quantification, and complemented by gene expression data, the protein composition of milk can now be dissected with great accuracy.

Comparative milk genomics

The bovine genome sequencing initiative was not the only project to follow the human genome sequencing effort. As this initial project abated, international consortia formed to sequence the genomes of other organisms, including the chicken, dog, opossum, chimpanzee, macaque and platypus. Currently, there is a major project underway, referred to as the Mammalian Genome Project, to sequence 24 animal genomes. Funded by the National Institutes of Health (NIH), diverse species were included to maximize the branch length of the evolutionary tree. Control of lactation and milk composition vary considerably across different species. The Mammalian Genome Project will contribute enormously to the study of the evolution of milk proteins, and will complement a growing body of work on whole genome expression studies in mammary tissue, now mostly in cattle, humans and mice.

Aligned with the goal to develop a deep understanding of milk proteins, we have developed a comparative approach to assess the genomic basis of diversity in milk composition between species. Experimental models have been chosen on the basis of reproductive/lactation strategy, and have been cross-referenced to the dairy cow. The study is based on gene expression profiles and integration with genomic mapping information and dairy QTL analysis. Within this framework, a specific methodology has been applied to identify those genes that may code for milk proteins that are either inherently active in milk or could be developed as dairy products.

A recent example of how comparative genomics may be exploited to alter milk protein composition is the use of functional genomics to exploit animal models with extreme adaptation to lactation, as an alternative approach to identifying key genes that specifically regulate protein synthesis in the mammary gland. For example, milk protein is the only component of milk that is synthesized de novo in the mammary gland of the Cape fur seal. The protein content of this seal milk is more than double that of bovine milk (Cane et al., 2005). The second model, the tammar wallaby, increases both the milk protein concentration and the total production of milk protein in the latter half of lactation (Nicholas et al., 1997). Changes in global gene expression in mammary tissues from the tammar wallaby at early and late lactation using a cDNA array (see Chapter 2), and in pregnant and lactating seals and dairy cows using an Affymetrix microarray, indicate that folate metabolism, and particularly the
role of the folate receptor, may be a crucial regulatory point of milk protein synthesis in mammary epithelial cells.

Although this analysis is limited because of redundancy on the arrays (number of probes and number of genes are different), non-symmetric mapping between species, dependence upon threshold for mapping and sequence quality and reliance on UniGene gene assemblies, feeding supplement experiments in the dairy cow have indicated a possible role for folate in manipulating milk proteins.

The importance of the folate receptor 1 (or $\alpha$) for cellular uptake of folate was established by the analysis of renal folate handling in mice with targeted gene knock-outs of folate-binding proteins 1 and 2 (folbp1 and folbp2, equivalent to human and cow folr$\alpha$ and folr$\beta$) (Birn, 2006). Molecular studies in human, monkey and mouse cell lines have shown that the folate receptor $\alpha$ mediates cellular uptake of folate and that folate receptor populations are regulated at multiple levels. Folate may either have a direct effect on the mammary gland or may reduce the competition for precursors between gluconeogenesis and methylneogenesis and increase metabolic efficiency of the mammary gland to influence milk protein production. Furthermore, the folate receptor population may play a crucial role in the capacity of mammary epithelial cells to respond and utilize circulating serum folate.

The folate requirements of lactating dairy cows have been extensively reviewed by Girard et al., (2005). Lactation increases the demands both for methylated compounds (synthesis of milk choline, creatine, creatinine and carnitine) and for methionine to support milk protein synthesis (Xue and Snoswell, 1985; Girard and Matte, 2005). The high demand for folate during lactation is demonstrated by the observation that total serum folates decrease by 40% across the lactation period in dairy cows (Girard et al., 1989).

Folate supplement experiments in dairy cows have shown a positive response to milk production and milk protein yields (Girarde and Matte, 1998; Girard et al., 2005; Graulet et al., 2007). This positive response was not consistent amongst all experimental cows in the first two studies, and appeared to be dependent on cow lactation status (premiparous were non-responsive, multiparous were responsive), stage of lactation and serum vitamin B$12$ status.

The more recent study by Graulet et al. (2007), including vitamin B$12$ supplementation, suggested that the increases in milk and milk protein yields as a result of folic acid supplements were not dependent on the supply of vitamin B$12$ and were probably closely related to the role of folic acid in the DNA cycle. However, this does not diminish the importance of the role that folate metabolism plays in milk protein synthesis. This is an example of how the combined approach of comparative genomics and bioinformatics, plus species with extreme adaptation to lactation, may be utilized to identify key genes and cellular processes involved specifically in modifying milk composition and milk protein production.

**Origins of milk proteins**

The study of milk protein genes and encoded milk proteins of domesticated mammals and undomesticated mammals alike has made these data relevant to dairy science.
The data highlight adaptive features and the diversity of milk protein genes over evolutionary time. Evolutionary evidence suggests that complex lactation preceded divergence of the mammalian lineages. Monotreme lactation appears to be the most ancestral form of lactation whereas marsupials and eutherians developed divergent strategies. The study of the evolution of milk proteins assists in determining the significance of maintaining certain proteins while changing or losing others. These differences may reflect divergence of lactation strategies and may provide clues as to why some proteins have become non-essential and are in the process of being lost by some eutherians.

Bioinformatic predictions of genes encoding secreted proteins up-regulated during lactation in the mammary gland suggest that as many as 300 different proteins may be found in milk. However, caseins are the major proteins in milk, representing about 80% of total milk protein content. Expression of a number of casein genes has been confirmed in monotremes (platypus and echidna) and marsupials.

Three evolutionarily related calcium-sensing casein-like genes and the functionally related $\kappa$-casein were identified in monotremes. As in other mammals, a number of casein splice variants could be identified. Interestingly, all these casein genes cluster tightly within a 100 kb region of the platypus genome, a physical linkage that has been observed in the casein locus of all mammalian genomes examined so far (Rijnkels, 2002). The respective organization of the casein locus is also highly conserved in all mammalian lineages. Casein genes occur in the order $\alpha_s1$, $\beta$, $\alpha_s2$- and $\kappa$, where the $\beta$-casein gene is encoded in the opposite direction from the other genes. The close proximity of $\alpha_s1$- and $\beta$-casein genes in an inverted tail—tail orientation and the relative orientation of the more distant $\kappa$-casein genes are similar in all mammalian genome sequences available so far. This observation suggests that the synchronized tissue-specific expression of casein genes in the lactating mammary gland may be controlled, in part, by a common mechanism at this locus. However, genome sequence analysis has also revealed variation in the $\alpha_s2$-casein gene content of mammalian lineages.

Marsupials possess only one copy of $\alpha$-casein corresponding to $\alpha_s1$-casein (Lefevre et al., 2007). In the monotreme lineage, a recent duplication or a gene conversion event involving $\beta$-casein occurred to produce an $\alpha$-like casein gene at a genomic location similar to that of the eutherian $\alpha_s2$-casein. A similar but more ancestral duplication of $\beta$-casein occurred in eutherians to produce $\alpha_s2$-casein and further duplications have also occurred since in human and mouse lineages to produce $\alpha_s2a$- and $\alpha_s2b$-casein genes.

$\beta$-Lactoglobulin ($\beta$-Lg) is the major whey protein in ruminant milk and is present in milk from a variety of species; however, it is absent in rodents, such as the mouse, lagomorphs (such as the rabbit), and humans, although reports to the contrary exist. $\beta$-Lg in human milk probably arises either from spurious antibody cross-reactivity (Brignon et al., 1985) or from the presence of ingested bovine $\beta$-Lg in the mother’s milk (Fukushima et al., 1997). Not all primate milks are devoid of $\beta$-Lg as both the macaque and the baboon have the protein within their milk (Hall et al., 2001).

Because of the absence of $\beta$-Lg in human milk, the dairy industry has developed ways in which to convert the protein composition of bovine milk to something more similar to the protein composition of human milk. Genetic variants have been observed in essentially all of the species from which the protein has been observed. There are several genes encoding $\beta$-Lgs within the different species. Sequence analysis
of species genomes suggests that some species such as the dog and the cat have three \( \beta \)-Lg genes, whereas others such as the horse and the donkey have two \( \beta \)-Lg genes, and that bovine, sheep and goat have one \( \beta \)-Lg gene and bovine has a pseudogene. Marsupials, monotremes and some primates have one \( \beta \)-Lg gene.

Comparison of \( \beta \)-Lg protein sequences shows high conservation of the protein during evolution. Bioactive peptides derived from \( \beta \)-Lg are currently being intensively studied. These peptides are active only once they are released from the precursor protein. Once released, these peptides are suggested to play important roles in human health, including having antimicrobial, antihypertensive and antioxidant activities (Hernandez-Ledesma et al., 2007). The variation of gene number and protein content in milk represents a specific evolution of \( \beta \)-Lg for each lineage. It has not yet been explained how a major protein in cow’s milk such as \( \beta \)-Lg can be completely abolished in human, rat or rabbit milk with little to no consequence to the offspring.

Whey acidic protein (WAP) is a major whey protein that is found in the milk of numerous species such as the mouse (Hennighausen and Sippel, 1982), rat (Campbell et al., 1984), pig (Simpson et al., 1998), rabbit (Devinoy et al., 1988), camel (Beg et al., 2000), wallaby (Simpson et al., 2000) brushtail possum (Demmer et al., 2001), echidna and platypus (Sharp et al., 2007), but is absent in the milk of cows, sheep and humans (Hajjoubi et al., 2006). WAP pseudogenes have been identified in the bovine and the human. A nucleotide deletion at the end of the first exon is reported to have caused a truncation of the bovine WAP protein (Hajjoubi et al., 2006). This deletion is also found in ovine and caprine species. In comparison, it is the absence of an ATG initiation codon in human WAP that is suggested to be the cause of the pseudogene in this species. Interestingly, the polyadenylation signal AATAAA is still present in the ruminant sequence but not in the human sequence.

WAP proteins from different species show moderate sequence similarity; however, they differ in the number of four-disulfide core (4-DSC) motifs between the species. Each specific 4-DCS domain is recognizable by sequence similarity and it is possible to trace the origins of each domain during evolution (Figure 3.1). Although not expressed, both bovine and human pseudogenes have similar gene structure to the eutherian WAPs and it is suggested that these were once functional. As suggested in Chapter 1, the function of the WAP proteins appears to differ with the presence or absence of each of these domains, and it is interesting to speculate why this protein appears to be dispensable in some species.

The apparently dispensable role of proteins such as \( \beta \)-Lg and WAP may be related to the diet of the young, which is being adequately compensated in more domesticated animals or may be linked to improved care and hygiene in domesticated life.

Constraints and opportunities for evolution or manipulation of bovine milk proteins

Although the protein component of bovine milk is only \( \approx 2-4\% \) of the total weight, which typically is marginally less than both total fat and carbohydrate, the population of proteins present in milk is diverse and impacts on the physical properties
of milk, the nutrition of a neonate and the metabolic homeostasis and development of both the dam and the offspring. Approximately 80% of the protein present in milk consists of four structurally and functionally interrelated proteins called the caseins. These include $\alpha_s^1$- and $\alpha_s^2$-caseins as well as $\beta$- and $\kappa$-caseins. Some of these proteins show some conservation of gene and protein sequence and structure and there is similar evidence of relatedness between proteins observed in milks of different species. The remaining 20% of the milk protein population includes the major whey proteins $\beta$-Lg and $\alpha$-lactalbumin ($\alpha$-La) as well as other constituent proteins including immunoglobulins, serum proteins, milk fat globule proteins, transferrin, lactoferrin, $\beta_2$-microglobulin, an array of enzymes and numerous proteolytic products.
The gene and protein structures and sequences of the six major bovine milk proteins (the four caseins, β-Lg and α-La) have been well characterized although the precise secondary and tertiary structures of the caseins have been difficult to demonstrate experimentally as they have proven to be challenging to investigate via X-ray crystallography. The casein phosphoproteins precipitate at pH 4.6, which is a significant characteristic that is utilized in cheese production but, in milk, these proteins aggregate to form protein aggregates or micelles of \( \approx 50-300 \) nm in diameter. \( \kappa \)-Casein stabilizes the surface of the protein micelles with the other caseins forming the core, which is stabilized by the phosphorylation of seryl residues. The organic phosphate moieties in turn enable calcium binding, which also stabilizes the micelle structure and hence, as well as being a significant source of dietary amino acids, the micelle aggregates also form a valuable source of calcium and phosphorus for neonatal nutrition and development.

Apart from the neonatal nutritional and industrial (dairy products) significance of bovine milk proteins, a wide range of biological activities is observed in the neonate as well as the dam; these activities are attributed to both the intact proteins and more importantly their products of \textit{in vivo} digestion (see the review by Meisel, 2005). When considered together, bovine milk proteins, which provide neonatal nutrition, regulate physiological processes, co-ordinate neonatal development and form the basis of an internationally significant agricultural commodity market, are globally important proteins. For this reason, their functional characteristics need to be considered when evaluating the potential for enhancement or modification of their structure or biosynthesis.

The detailed structure of milk proteins and their genes

An understanding of the detailed structure of milk proteins and their genes is critical in the consideration of the impact of modifying bovine milk proteins to enhance productivity, to enhance milk product manufacturing efficiency or to exploit inherent or proposed bioactivity. The physical and structural characteristics of bovine milk proteins have been extensively reviewed (Farrell \textit{et al.}, 2004).

The bovine casein genes are present as single copy genes per chromosome and are clustered together on chromosome 6 in a 200 kb region at position q31–33 in the order \( \alpha_{s1} \), \( \beta \), \( \alpha_{s2} \) and \( \kappa \) (Threadgill and Womack, 1990). The bovine \( \alpha_{s1} \)-casein gene is 17508 bp in length and contains 19 exons. The protein is the most abundant of the bovine caseins, present at 12–15 g/L. There are up to eight reported genetic variants of this protein, with the B variant being the most common in \textit{Bos taurus} species and comprising 199 amino acids with a molecular weight of 23.6 kDa. The \( \alpha_{s1} \)-casein protein may contain up to eight serine monophosphate residues, which cluster in a hydrophilic domain between amino acids 43–80 and, through modelling studies, are thought to be connected to a hydrophobic domain (amino acids 100–199) by helical and sheet secondary structures. The less abundant bovine \( \alpha_{s2} \)-casein gene is 18483 bp in length and contains 18 exons with significant exon duplication (Mercier and Vilotte, 1993). This gene codes for 207 amino acids with a molecular weight of 25.23 kDa. This protein exhibits a number of phosphorylation states as well as four variants that have been observed in common bovine populations.
In contrast, the bovine $\beta$-casein gene is significantly less complex, containing nine exons in 8498 bp. The protein is present at 9–11 g/L in modern dairy cattle, consists of 209 amino acids and has a molecular weight of approximately 24 kDa. Up to 12 genetic variants have been observed and the protein exhibits a flexible structure that can adopt multiple conformations, although a small amount of secondary structure in the form of $\alpha$ helices and $\beta$ sheets has been predicted. There are five phosphoserine residues that reside at the N-terminal domain of the protein.

The bovine $\kappa$-casein gene exhibits fewer structural similarities than the other casein genes and contains five exons and is approximately 13,700 bp in length. It has diverged from the fibrinogen gene family but is more highly conserved between species than the other casein genes. Two common variants are observed in dairy cattle (although several others have been reported), with the B variant containing 169 amino acids with a molecular weight of 19 kDa. Up to six threonine residues may be glycosylated and, similar to the other caseins, $\kappa$-casein has a relatively high proline content that influences the predicted secondary structure of two sets of antiparallel $\beta$ sheet structures exhibiting hydrophobic side chains that may be important for micelle formation. As with the other caseins, the detailed structure of bovine $\kappa$-casein is poorly characterized although the N-terminal domain of amino acids 1–44 has been experimentally shown to contain a helical structure (Bansal et al., 2006).

The bovine $\alpha$-La gene contains four exons and is approximately 3.5 kb in length. It is located on bovine chromosome 5 and exhibits significant interspecies homology in exon 4 (significant for its interaction with galactosyltransferase) and the 5′ untranslated region. Two variants have been characterized for the protein with the most common B variant exhibiting a molecular weight of 14.1 kDa and being present at 0.6–1.7 g/L in skim milk. The 123 amino acids are organized into four helical domains and a small sheet structure that is dependent on pH and metal ion concentrations (Sawyer and Holt, 1993). Different glycosylation states have been observed and structurally bovine $\alpha$-La has some similarity to lysozyme proteins (Brew et al., 1970). This protein, like the caseins, also has calcium-binding properties, as described by Hiraoka et al., (1980).

Bovine $\beta$-Lg is located on chromosome 11, contains four exons in 4723 bp and is well conserved between species (Folch et al., 1994). Structural similarities occur throughout the lipocalin protein family and $\beta$-Lg has been shown to bind to a number of hydrophobic ligands including palmitic acid (Kontopidis et al., 2004; Konuma et al., 2007). There are two common variants of this protein—A and B—with the protein itself containing 162 amino acids, with a molecular weight of 18.3 kDa. It exhibits a number of $\beta$ strands, which form a $\beta$-barrel structure that contains the hydrophobic-ligand-binding site, and a single $\alpha$ helical domain is observed at the C-terminal end of the protein.

The secondary structures of the major bovine milk proteins are diverse, yet are clearly critical for the functions of the proteins including micelle formation, calcium and phosphate binding and other bioactivities. Although there may be opportunities for manipulation of structure and therefore function of these proteins, even modest changes to nucleotide and amino acid sequence may significantly alter protein and peptide functionality.
The function of bovine milk proteins

The structural complexity of these milk proteins suggests that they serve many functions in the neonate in addition to the mere provision of a source of amino acids for protein biosynthesis in the fast-developing neonate. Yet the range of functions is not likely to be as great as that in more altricial species such as the monotremes and marsupials, in which the milk constituents orchestrate almost the entire developmental process from the fetus through to attaining physiological independence (see Chapter 2). The contributions that individual proteins make to this developmental spectrum are poorly understood, although longitudinal studies on gene expression and related changes in milk composition will assist in unravelling this relationship.

The protracted emphasis on selection of cows for milk volume has probably also influenced the balance of biological activities residing within these proteins, because contemporary dairy herds are larger in stature than their predecessors. Thus, developmental activities encoded within the protein fraction may have been altered to emphasize growth promotion rather than other key functions such as the storage of body energy reserves to support reproductive activity and even to promote maturation of the immune system.

The complexity increases further once these proteins are digested enzymatically or are hydrolyzed in the developing gastrointestinal tract and then absorbed to be transported to their site of action. The precise balance between the peptide component that is absorbed as functional entities and the peptide component that remains for further processing into short peptides as a source of dietary amino acids is not well understood. However, the need to conserve a constant supply of developmental molecules is probably more important than the need for protein synthetic substrate in sustaining the development of the neonate.

The observation that over 100 hormones and growth factors have been identified in colostrum or milk (Koldovsky, 1996) would suggest that the milk proteins may play important roles, working in concert with these established endocrine/paracrine signals. Therefore, the sustained selection of animals for milk volume output will have altered the balance of bioactivities in colostrum and milk to account for this response. Thus, the balance of bioactive peptides may differ from those in species in which this selection pressure has been absent. From a commercial point of view, these milk constituents also play a role in determining the sensory properties of milk and its manufactured products. Of course, these same sensory factors probably also serve as attractants to the newborn as it seeks its nutrient supply from the udder.

The extraction of intact proteins of high commercial value from milk has been attractive to dairy farmers tied to tightly regulated commercial milk prices. The profitability of such ventures depends on the ease of extraction, the relative concentration of the protein in milk and the value placed on the bioactivity. A range of these proteins is presented in Table 3.1.

Both α-La and β-Lg have been isolated on a large scale using a range of chromatographic, ultra-high pressure and membrane separation techniques (Korhonen and Pihlanto, 2007). Both of these proteins convey important activities (Table 3.1) that provide the impetus for large-scale production. One of the most important clinical...
findings is the ability of these whey proteins to decrease the incidence of certain cancers. In most studies, crude whey protein mixes are used; these are thought to act through the provision of additional sulfur amino acids that act as substrates for the synthesis of glutathione in many tissues (Bounos and Gold, 1991). This tripeptide is a potent antioxidant, acting through selenium-dependent glutathione peroxidase that contributes to the removal of reactive oxygen species (Parodi, 2007).

Many of the highly expressed proteins clearly have important immunological functions. The passive immunity provided through the immunoglobulin complement of colostrum to the neonate is supplemented by a range of other bioactivities designed to countenance pathogen loads to which the newborn is exposed in its new environment. The activity of these proteins extends to the initiation of mucosal immunity in the gut and an optimal environment for beneficial microflora. Such proteins include immunoglobulins, many chemokines, mucin 1, lactoferrin, cathelicidin 1, lactoperoxidase, S100 calcium-binding proteins, complement proteins and polymeric immunoglobulin receptor.

It is beyond the scope of this review to detail all of the bioactivities identified in milk. However, some specific proteins are illustrative of principles that are found across the milk proteome.

Lactoferrin is one such protein; it has the dual roles of scavenging iron and acting as a bacteriostat. It is found in exocrine secretions such as milk, tears, tubotympanum and nasal exudate, saliva, bronchial mucus, gastrointestinal fluids, cervicovaginal mucus and seminal fluid, all of which require antibiotic activity. It is also found in neutrophils that are attracted to sites of infection where both iron sequestration and antibiotic activity assist in the healing process. However, its biological effects

<table>
<thead>
<tr>
<th>Protein</th>
<th>Concentration in colostrum (g/L)</th>
<th>Concentration in milk (g/L)</th>
<th>Molecular weight</th>
<th>Exploitable bioactivity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caseins (αs1, αs2, β, and κ)</td>
<td>26</td>
<td>28</td>
<td>14000-22000</td>
<td>Ion carriers, bioactive peptide sources, immunomodulators</td>
</tr>
<tr>
<td>β-Lactoglobulin</td>
<td>8.0</td>
<td>3.3</td>
<td>18400</td>
<td>Antioxidant, vitamin-A-binding protein, bioactive peptide source</td>
</tr>
<tr>
<td>α-Lactalbumin</td>
<td>3.0</td>
<td>1.2</td>
<td>14200</td>
<td>Lactose synthesis and milk volume. Calcium binding, immunomodulator, bioactive peptide source</td>
</tr>
<tr>
<td>Immunoglobulins</td>
<td>20-150</td>
<td>0.5-1.0</td>
<td>150000-1000000</td>
<td>Specific immune protection, potential bioactive peptide source</td>
</tr>
<tr>
<td>Glycomacropeptide</td>
<td>2.5</td>
<td>1.2</td>
<td>8000</td>
<td>Antimicrobial, bifidogenic, gastric modulator, anticoagulant factor</td>
</tr>
<tr>
<td>Lactoferrin</td>
<td>1.5</td>
<td>0.1</td>
<td>80000</td>
<td>Antimicrobial, antioxidant, anti-inflammatory, anticarcinogenic, bioactive peptide source, immunomodulator</td>
</tr>
<tr>
<td>Lactoperoxidase</td>
<td>0.02</td>
<td>0.03</td>
<td>78000</td>
<td>Antimicrobial, immunopotentiator</td>
</tr>
<tr>
<td>Lysozyme</td>
<td>0.0004</td>
<td>0.0004</td>
<td>14000</td>
<td>Antimicrobial, immunopotentiator</td>
</tr>
<tr>
<td>Serum albumin</td>
<td>1.3</td>
<td>0.3</td>
<td>66300</td>
<td>Bioactive peptide source</td>
</tr>
</tbody>
</table>

**Table 3.1** Major milk proteins with potential for extraction on a commercial scale to exploit bioactivities (adapted from Korhonen and Pihlanto, 2007)
Significance, origin and function of bovine milk proteins
do not end there. Its commercial utility is increased through its antiviral, antioxidant, antitoxicogenic, antithrombotic and, importantly, anti-carcinogenic actions. However, it is important to note that lactoferrin is just one component of a mix of protective molecules in biological fluids, as lysozyme, beta defensins and the surfactant proteins A and D (SP-A, SP-D) among others also contribute to this important function.

Typically, lactoferrin is found in concentrations of 100 mg/L in colostrum and is extracted by conventional fractionation and chromatography. However, the expression of milk proteins in cereal grains such as rice using recombinant DNA technology (Lonnerdal, 2006) seems to be a more effective way of providing these unique milk proteins in our diet. Similarly, recombinant human lactoferrin given orally has been suggested as a mechanism for the prevention of certain gastrointestinal infections in pre-term infants (Sherman and Petrak, 2005). There is little doubt that this will be a continuing trend, the potential for which was recognized in the last decade (Arakawa et al., 1999); transgenic plants may well supersede the cow as a source of this and other unique milk proteins.

Bioactive peptides sequestered within milk proteins
Milk proteins contain latent biofunctional peptide sequences within their primary structures that exert physiological responses in vivo. They remain latent until released through the processes of enzymatic processing in the gastrointestinal tract. A large range of these peptides has been identified, including opioid, antimicrobial, immunomodulatory, mineral-transporting, growth-promoting, anticancer, proteinase and angiotensin-converting enzyme (ACE) inhibitory peptides (Shah, 2000; Ferranti et al., 2004). In an evolutionary sense, the cow is targeting peptides to specific effector sites through this process. Presumably, if the availability of metabolic substrates to support maximal biological response is inadequate, then the animal has the ability to limit the quantities of peptides released by suppressing the release of rate-limiting enzymes.

The biofunctional peptides currently most studied in food proteins appear to be those that inhibit ACE. This enzyme plays a central role in the regulation of blood pressure through the production of the potent vasoconstrictor angiotensin (Ang) II and the degradation of the vasodilator bradykinin. ACE inhibitory peptides may therefore have the ability to lower blood pressure in vivo by limiting the vasoconstrictory effects of Ang II and by potentiating the vasodilatory effects of bradykinin (Murray and Fitzgerald, 2007). However, this peptide is just one of a number that contribute to the regulation of this important physiological parameter; in addition to the renin–angiotensin system, endothelins and their converting enzymes, the kinin nitric oxide pathway and the neutral endopeptidases all play a role.

The development of functional foods with antihypertensive properties provides an attractive and potentially commercially lucrative range of products. Although this enzyme is found widely through metabolic tissues, it is largely associated with the vascular epithelium (Ondetti and Cushman, 1982). These ACE inhibitory peptides can be enzymatically released from intact proteins in vitro and in vivo during food processing and gastrointestinal digestion respectively. ACE inhibitory peptides may
be generated in or incorporated into functional foods in the development of “natural” beneficial health products (Murray and Fitzgerald, 2007). In view of the extensive range of peptides derived from various precursor proteins that are involved in the regulation of blood pressure, it may be too simplistic to expect a functional food containing just one or two of these peptides to decrease suppressor activity in the long term. However, several products are currently being evaluated as beneficial functional foods/food ingredients.

One major success story in deriving a commercial product from milk protein tryptic digests is the casein phosphopeptides used in the chewing gum “Recaldent” (Cross et al., 2007). These peptides, containing the sequence -Pse-Pse-Pse-Glu-Glu-, where Pse is a phosphoseryl residue, stabilize calcium and phosphate ions in aqueous solution and make these essential nutrients bioavailable. Investigations of the chemistry of these intriguing peptides have shown that they can be altered to form casein phosphopeptide–amorphous calcium phosphate which in turn is capable of forming a calcium fluoride. The practical outcome from these outstanding discoveries is a product that is capable of remineralizing carious lesions in dental enamel and that can be incorporated into dental care products and foodstuffs (Cross et al., 2007).

The potential for the discovery of new peptides is high and it would seem reasonable to suspect that careful screening of digesta from various segments of the gastrointestinal tract may prove to be a valuable approach for this discovery process.

Existing variation in bovine milk proteins and the impact on expression, function and milk quality

As described above, a number of genetic variants of the major bovine milk proteins have been observed and characterized. These variations range from minor amino acid substitutions to larger deletions. There are many studies that suggest that the presence or absence of a particular variant expressed within an individual may be associated with altered production or processing quality of milk (see the reviews of Jakob and Puhan 1992; Martin et al., 2002). Many of these studies are based on statistical associations but others have been well characterized to elucidate cause–effect relationships. These naturally occurring variations in milk protein amino acid sequences serve as examples for the breadth of potential that modification of milk proteins may have on milk production and quality.

The relationship between genetic variants of bovine milk proteins and productive characteristics has been extensively reviewed. The characteristics observed to be associated with milk protein polymorphism include first lactation milk production, protein production and fat percentage in Californian Jersey cows (Ojala et al., 1997); milk and protein yield and fat content in Finnish Ayrshire cows (Ikonen et al., 2001); milk, protein and fat yield in Holstein and Ayrshire cows (Lin et al., 1986); the effect of nutritional regimens on milk composition (Mackle et al., 1999); somatic cell count (Ng-Kwai-Hang et al., 1987); total solids and milk protein profile (McLean et al., 1984); and lifetime performance (Lin et al., 1989).

Yet, despite this body of evidence, milk protein genotype is not currently used in selection to enhance genetic gain for milk production characteristics, possibly
because of the variable and, on occasion, conflicting observations or the potential cost relative to the sometimes modest production differential. Understandably, many of the genotypic variants associated with increases in milk yield are generally already present at a high frequency in modern dairy populations.

The B variant of bovine $\alpha_s$1-casein is the most abundant variant in western dairy populations, possibly because of indirect selection as other variants may in some way be associated with decreased $\alpha_s$1-casein synthesis, such as the A variant which may be subject to a potential decrease in translational efficiency because of observed polymorphism at a polyadenylation signal site (McKnight et al., 1989) or because of promoter polymorphisms that may alter transcriptional efficiency (Prinzenberg et al., 2003). Similarly, animals that are observed to have a lower bovine $\alpha_s$1-casein protein concentration in milk (heterozygous for the G allele) exhibit a lower casein content and a slower clot formation time, yet display a faster curd-firming time and a higher curd firmness (Mariani et al., 2001). In contrast to bovine $\alpha_s$1-casein, although there are four known variants of the bovine $\alpha_s$2-casein protein, few studies have shown any relationships between the genotype of this protein and milk production or quality traits.

The A1 and A2 variants of bovine $\beta$-casein are present in modern dairy populations at high frequencies and the A2 variant has been shown in defined populations of Danish dairy cattle breeds to be associated with higher milk, fat and protein yields when homozygous (Bech and Kristiansen, 1990).

Some controversy surrounds the A2 protein genotype, as it has been suggested that human populations who consume milk containing higher concentrations of the A2 variant exhibit a lower incidence of cardiovascular disease, a lower incidence of type 1 diabetes and decreased severity of symptoms of neurological diseases because the A2 variant does not liberate the opioid bioactive peptide casomorphin 7 upon proteolytic digestion (Laugesen and Elliott, 2003; Tailford et al., 2003; Bell et al., 2006; Kaminski et al., 2007).

In a review of evidence, however, Truswell (2005) suggested that there was limited evidence to suggest that milk containing the A2 variant of $\beta$-casein had any significant human health advantage and, indeed, human trials suggested that there was no differential effect between dietary products containing either the A1 variant or the A2 variant on human plasma cholesterol concentrations (Venn et al., 2006) or cardiovascular health (Chin-Dusting et al., 2006).

With the significant structural role of bovine $\kappa$-casein in the casein micelle, it is not surprising that there are only two common variants—A and B—in modern dairy populations, although up to nine other variants have been characterized in bovine species (Farrell et al., 2004). A number of studies have been conducted to investigate associations between $\kappa$-casein protein variants and milk production and quality characteristics. Of all the genetic polymorphisms of dairy cattle affecting milk composition, the $\kappa$-casein genotype is one of the more significant, with association differences identified between genotype and lifetime production (Lin et al., 1989), concentrations of individual milk proteins (McLean et al., 1984), protein yield and percentage (Tsiaras et al., 2005) as well as cheese production characteristics including rennet clotting time, curd formation and coagulation strength (Pagnacco and Caroli, 1987).
Of the 12 identified bovine β-Lg variants, only the A and B variants occur at high frequencies in dairy cow populations. The B variant has been associated with higher casein percentage (Lundén et al., 1997), higher total solids (McLean et al., 1984) and higher milk yield, fat yield, fat percentage and lactose yield (Tsiaras et al., 2005), whereas the A variant is associated with higher whey protein content and lower casein content (Auldist et al., 2000), possibly suggesting that it may be expressed at higher concentrations than the B variant. The B variant association with the casein composition of milk therefore has implications for milk quality for cheese production. Of the three bovine α-La variants, the B variant is the most common in modern dairy populations, yet the A and B variants differ by a single amino acid residue. The role of this protein in lactose biosynthesis is such that there seems to be little genetic divergence and little function or associated differences observed between the common variants.

**Experimental modifications of bovine milk proteins**

The experimental manipulation of patterns of expression or characteristics of bovine milk proteins has previously been reviewed (Bremel et al., 1989; Yom and Bremel, 1993; Clark, 1996). Both over-expression and impairment or “knockout” of milk proteins have been conducted predominantly in rodent models, presumably because of the expense and technical challenges of conducting studies in bovine or other ruminant species. Although observations from experiments conducted in model species must be considered in context, valuable insight into the potential for future manipulation of bovine milk proteins in ruminant species has been gained.

Although a number of experiments have been conducted to either over-express or inhibit expression of milk proteins in mice and other rodents, only a few have further characterized the milk to assess the impact of manipulation of expression profiles on the quality of milk for dairy product manufacture. A transgenic mouse model over-expressing bovine κ-casein (Gutiérrez-Adán et al., 1989) exhibited no changes in milk protein concentration, yet milk from these mice exhibited a smaller micelle size and a tendency toward stronger curd characteristics. Conversely, a mouse model in which the endogenous κ-casein was suppressed exhibited a loss in micelle stability, resulting in casein precipitation in alveolar lumens, which prevented lactation (Shekar et al., 2006).

These investigations highlight the significance of the role of κ-casein in the tertiary structure of protein in milk regardless of species. In comparison, the manipulation of β-casein expression results in less dramatic consequences. The over-expression of bovine β-casein in a mouse model (Hitchin et al., 1996) had little observed effect on lactation although the processing quality of milk was not assessed in this study. In another study, the inhibition of endogenous murine β-casein (Kumar et al., 1994) changed micelle size and reduced pup growth.

Similarly, in cloned transgenic cattle in which both bovine β-casein and κ-casein were over-expressed (Brophy et al., 2003), milk levels of both proteins were increased, demonstrating that milk protein composition can be altered in large ruminants using recombinant gene technologies. Although the processing characteristics of the milk from these animals were not established, the authors stated that the fat, lactose and mineral contents of the transgenic milk were within normal ranges.
The functionality of caseins can be altered through enzymatic modification of the proteins, e.g. through dephosphorylation. This results in increased pepsin hydrolysis, which in turn alters digestibility (Li-Chan and Nakai, 1989) and potentially the release of bioactive peptides. Importantly, none of the studies discussed identified defects or significant changes to mammary development or structure, suggesting that manipulation of milk proteins may not have undesirable effects on the mammary gland itself, compromising either the potential to lactate or animal health.

The whey proteins have also been investigated for manipulation to alter processing characteristics of milk. Because of its role in the synthesis of lactose and the implications for manipulation of osmotic potential of milk, $\alpha$-La has been a target for manipulation. Mice transgenic for expression of bovine $\alpha$-La exhibited increased lactose concentration in milk, and a slight increase in pup growth rate (Boston et al., 2001).

Similarly, transgenic pigs over-expressing bovine $\alpha$-La produced milk with lower total solids, higher milk yield in early lactation, higher lactose concentration and increased litter growth rates (Noble et al., 2002). In contrast, a mouse model of $\alpha$-La deficiency resulted in milk that was so viscous that pups were not able to suckle effectively (Stinnakre et al., 1994). Bovine $\alpha$-La itself has also been modified to alter its characteristics and various mutations have resulted in changes in affinity for galactosyltransferase, increased glucose binding (Grobler et al., 1994) and changes in molten globule conformation (Uchiyama et al., 1995).

The $\beta$-Lg protein has also been evaluated experimentally. Over-expression in transgenic mice resulted in normal mammary physiology and therefore normal milk secretion and pup growth (Hyttinen et al., 1998; Gutiérrez-Adán et al., 1999). However, a similar study in which ovine $\beta$-Lg was over-expressed reported an increase in total protein content (Simons et al., 1987). Mutants of bovine $\beta$-Lg have also been studied, with variation in thermal stability (Cho et al., 1994), rate of secretion (Katakura et al., 1999) and rate of denaturation and digestion (Jayat et al., 2004) being reported.

**Adding value to milk through the use of milk protein genomics**

The potential value of milk as a source of animal protein varying in characteristics that promote human health has been demonstrated by the range of bioactivities residing within the mammary protein phenome. The ease with which dairy products are distributed to consumers irrespective of their socio-economic status also makes this an ideal vehicle for the administration of therapeutics. The hyperimmune nature of colostrum, containing up to 40% by weight of immunoglobulin, suggests that immunization of cows against specific pathogens may provide a rich source of therapeutic antibodies. As human colostrum contains antibodies reactive to colonization factors 1 and 2 of enterotoxigenic *Escherichia coli* (Correa et al., 2006), thereby preventing diarrhea in infants, the production of similar antibodies in cows seemed to be a logical commercial target. The Australian commercial biotechnology company Anadis has exploited this technology in producing an antibody-enriched colostrum tablet to inhibit travellers’ diarrhea (www.anadis.com).

However, perhaps the greatest potential that milk proteins and peptides have for human health is through the addition of whey protein and casein peptides to foods.
to increase their functionality. Estimates of world whey protein production exceed 0.5 million tonnes, which provides a resource that should be used more effectively than by simply disposing of it as a liquid waste or as an animal feed. Considerable progress has been made in the development of methods for separation and then purification of specific proteins from the whey fraction. Native immunoglobulins, lactoferrin, lactoperoxidase, α-La and β-Lg have all been recovered in industrial quantities. Yet the major advances in recombinant technologies may ultimately provide a cost-competitive alternative supply of these proteins.

The development of transgenic dairy herds expressing altered protein composition and yield has been muted for the past decade. It was thought that the use of yeast and bacterial artificial chromosomes capable of conveying gene constructs of an Mb may be useful in supporting the expression of the 200 kb casein locus (with all four caseins) including the regulatory elements co-ordinating their expression (Zuelke, 1998). To date, the only report of substance is that of Brophy et al. (2003), cited above, in which both β- and κ-casein were over-expressed. Given the intricacy of the ultrastructure of the micelle, and its critical nature in providing calcium and phosphorus in the correct form for absorption by the calf, it is not surprising that any genetic manipulation of protein composition is going to upset this equilibrium, which has been refined through millions of years of evolution. However, our ability to manipulate milk composition through manufacturing technologies has effectively removed the need for such animals. Growing animal welfare concerns must also be considered when embarking on such ambitious projects.

Nevertheless, the development of somatic nuclear transfer techniques that facilitate targeted genetic modifications has driven transgenic research over the past 5 years. Small interfering RNA techniques and lentiviral vectors have also contributed to this cause, as have modified episomal vectors designed to promote high levels of expression of therapeutic genes (Manzini et al., 2006). The use of transgenic animals as bioreactors to synthesize valuable proteins in large quantities is probably the major application for new transgenic animals. Goats and rabbits are the most popular target species, in addition to the cow, for producing a range of proteins including enzymes such as alpha-glucosidase, hormones such as human growth hormone and large proteins such as lactoferrin, albumin, collagen and vaccines (e.g. for malaria). The development of a recombinant human antithrombin III expressed in goat’s milk is the closest of these products to commercial release (Niemannn and Kues, 2007).

Ideally, a milk enriched in peptides promoting immune function, controlling blood pressure, acting as a bacteriostat and minimizing oxidative stress and cancer risk, while at the same time relieving depression and preventing dental caries, would seem to have the makings of a highly valuable functional food. Combining this with an enrichment with n-3 fatty acids thought to increase insulin sensitivity and therefore prevent diabetes, together with certain milk carbohydrates capable of improving cognition, adds greatly to a product that already acts as a rich source of amino acids and energy to promote normal growth processes. Manipulation of these proteins in milk will inevitably occur in the factory and potentially in the cow. The challenge remains to turn this speculation into commercial reality for the benefit of societies in both the developed world and the developing world.
Conclusions

The bovine milk proteins form the basis for a global industry in dairy products, and as such have been intensively studied by both classical deconstructive observation and in dynamic whole animal systems utilizing post-genomic or functional genomics approaches. From these studies we have a greater understanding of the roles of these proteins in milk, their biological significance in the neonate and the biochemical characteristics of bovine milk proteins that affect efficiencies of manufacturing processes.

The emerging interest in elucidating the bioactivities inherent in bovine milk proteins is both challenging and potentially economically and socially rewarding. Initially the benefits of this research will be realized by dairy manufacturers promoting brand differentiation following from marketing trends in milk fortification yet the far greater potential of enhancing these bioactivities of native proteins into complementary health products or nutraceuticals will represent a major repositioning of dairy products in consumer consciousness and result in the re-evaluation of the value of dairy products.

The enthusiasm for exploitation of these bioactivities in milk and colostrum should also be tempered by our knowledge of the biological interrelatedness of the major milk proteins and the bioactivities of trace proteins and elements in milk. These products should be produced through a means that is acceptable to consumers to allay fears regarding food safety and concerns for animal welfare. Approaches ranging from exploitation of existing variation within dairy populations, manipulation during processing or even transgenic approaches may be desirable depending on the value of the end product to human health and nutrition.

References


References


Post-translational modifications of caseins

John W. Holland

Abstract

The caseins exhibit a high degree of heterogeneity as a result of post-translational modifications (PTMs). Phosphorylation of the α- and β-caseins and glycosylation of κ-casein are the best-known modifications and are critical for the formation and stability of casein micelles. κ-Casein, in particular, has long been known to exhibit a high degree of variability in glycosylation. It is somewhat surprising to see so much variability in such important structural features. In recent years, the adoption of proteomic techniques has greatly enhanced our ability to unravel heterogeneity in proteins arising from complex and variable patterns of PTMs. In this chapter, a summary of our knowledge of the PTMs of caseins is attempted, with a particular emphasis on κ-casein and the implications that variations in PTMs have for dairy processors.

Introduction

The caseins are phosphoproteins and constitute about 80% of the protein in milk (Swaisgood, 2003; Farrell et al., 2004). They are assembled in a colloidal complex with calcium phosphate and small amounts of other minerals. Although obviously important for the provision of amino acids, calcium and phosphorus for infant nutrition, the casein micelle structure is also critical in determining the physical properties of milk. The stability of the micelle, or its controlled destabilization in the case of cheese and yoghurt manufacture, is of primary concern to the dairy industry.

A number of reviews of micelle structure have been published in recent years (Rollem, 1992; Holt and Horne, 1996; Horne, 1998; Walstra, 1999; Horne, 2002;
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Chapter 5 in this volume). In simple terms, the micelle is a network of protein molecules held together by a combination of hydrophobic interactions between protein molecules and electrostatic interactions between phosphoserine-rich regions of the \( \alpha_{s1} \), \( \alpha_{s2} \) and \( \beta \)-caseins and micellar calcium phosphate. Whereas the internal structure is still the subject of debate (Horne, 1998; Walstra, 1999), there is general acceptance of the “hairy micelle” concept, in which the hydrophilic C-terminal portion of \( \kappa \)-casein extends from the surface, providing steric and electrostatic repulsion, which prevents micelle aggregation.

A critical factor in micelle formation and stability is the presence of post-translational modifications (PTMs) such as phosphorylation on the \( \alpha_{s1} \), \( \alpha_{s2} \) and \( \beta \)-caseins and glycosylation on \( \kappa \)-casein. PTMs on secreted proteins such as the caseins occur in the endoplasmic reticulum and/or golgi complex after synthesis of the polypeptide chain. As such, they are not encoded by the genes per se but may be dependent on protein (and hence gene) sequence motifs that are necessary, but of themselves not sufficient, for modification to take place. There are a number of other factors that determine whether or not a PTM occurs, including expression of the genes encoding the enzymes necessary for the modification, the availability of their substrates and the accessibility of the modification site on the protein, especially after folding. Therefore, although it may be possible to predict the theoretical sites of modification on proteins, determination of the actual sites and the degree to which they are modified requires considerable experimental characterization.

Advances in our understanding of complex systems such as the caseins micelle are frequently preceded by advances in technology. In recent years, the development of proteomic technologies has greatly enhanced our ability to analyze milk proteins, particularly with respect to PTMs. Two-dimensional electrophoresis (2-DE), in particular, provides a high-resolution methodology for displaying the heterogeneity of the major milk proteins. As can be seen in Figure 4.1, genetic variants, phospho-variants and glyco-variants of the caseins can be resolved on a single 2-D gel. Advances in mass spectrometry (MS) have enhanced our ability to analyze the proteins arrayed on 2-D gels. Therefore, not only is it possible to resolve many proteins and their isoforms but also it is possible to characterize them, particularly with respect to the many PTMs that affect their electrophoretic mobility.

As will be seen below, the four caseins are present in many diverse forms as a result of differential PTMs. The biological reasons behind the diversity are not clear. However, what is clear is that the PTMs of the caseins are critical for their function in micelle formation and stability. The first part of this chapter summarizes what is known about the PTMs of the \( \alpha_{s1} \), \( \alpha_{s2} \) and \( \beta \)-caseins. The second part focuses on \( \kappa \)-casein, which has been the subject of recent proteomic studies, and includes an extended discussion on the functional significance of \( \kappa \)-casein heterogeneity.

**Bovine casein**

The \( \alpha_{s1} \), \( \alpha_{s2} \) and \( \beta \)-caseins are phosphoproteins that are generally well characterized in terms of their PTMs and have been the subject of a number of reviews (Mercier, 1981;
Ginger and Grigor, 1999; Farrell et al., 2004). In fact, the $\alpha$- and $\beta$-caseins have been used as model phosphoproteins in the development of proteomic techniques to examine global phosphorylation patterns (e.g. Cox et al., 2005; Kapkova et al., 2006; Sweet et al., 2006; Zhou et al., 2006; Wu et al., 2007). The multitude of techniques developed for the analysis of phosphorylation is beyond the scope of this chapter but has been reviewed recently (Bodenmiller et al., 2007; Collins et al., 2007). The PTMs of the $\alpha_s1$-, $\alpha_s2$- and $\beta$-caseins are summarized in the sections below.

An important note concerns the numbering of the amino acids in the casein sequences described in subsequent sections. The numbering of residues is based on the Swiss-Prot database entry for the relevant protein (see Figures 4.2 and 4.3) and includes the signal peptide which is normally removed during processing to generate the mature protein. This differs from the numbering in most of the dairy literature, in which the N-terminal amino acid of the mature protein is numbered 1.

$\alpha_s1$-Casein

The predominant form of $\alpha_s1$-casein in bovine milk contains eight phosphate groups. The phosphates are attached to hydroxyamino acids occurring in the sequence motif Ser/Thr-Xxx-Glu/Asp/pSer (Mercier, 1981). However, the vast majority of casein phosphorylation sites, including the eight sites in the major form of $\alpha_s1$-casein, occur in the more restricted Ser-Xxx-Glu/pSer motif. Phosphorylation of threonine or of serine in the Ser-Xxx-Asp motif is relatively uncommon. A minor form of $\alpha_s1$-casein with nine phosphates, originally called $\alpha_s0$-casein, also occurs in bovine milk. It contains one extra phosphate on Ser$^{56}$, which occurs in a Ser-Xxx-Asp motif (Manson et al., 1977).
Post-translational modifications of caseins

P02662 | CASA1_BOVIN Alpha-s1-casein—Bos taurus (Bovine).

1 MKLLILTCLV AVALARPKHPIKHHQGLPQEVLNENLLRFFVAPFPEVFGKE 50
51 KVNELSKDIGSSEDESTQAMIDEIKQMEAESISSSEIVPNVSEQIKHDKED 100
101 VPSERLYGLIEQLLRLKYYKVPQLEIVPNAEBERLHSMKEGIHAQQKEMP 150
151 IGVQNRELAYIFYEPLFRQFYQLDAYPSGAWYVYPLGTQYTDAPSFSDPNP 200
201 IGSENSEKTTPMLW 214

P02663 | CASA2_BOVIN Alpha-s2-casein—Bos taurus (Bovine).

1 MKFFIFTCLV AVALAKNTMEHVSSSEESIIQQETYKQKMNAINPSKENL 50
51 CSTFCKEVRVANEEESIGSSEESAEPATEEEVITVDDKHYQKALNEI 100
101 NQFYQKFQYQLYQGGPILNPWMVTQKRNAPVITPMLNREQLSTSEENS 150
151 KTVQDMESTEVFTKTKLTEEENRLNFLKISQRYQKFLLPQYLKTVYQ 200
201 HQKAMKFWIQPKTVIPYVRYL 224

P02666 | CASB_BOVIN Beta-casein—Bos taurus (Bovine).

1 MKVLILALCLVALALAREELELNVGPEIVELSSSEESITRINKKIEKFQSS 50
51 EBQQQTDELQDKIMFQAQTQLVVPFPFGIPvPLQTNPIQPPLVQTVVVP 100
101 PFLQPEVMGVSVKKEAMAPKHKMEMPPKYPVESPTEQSLTLTDVENLHL 150
151 PLPLLQSWHMPHQQLPLPTVMPFQSVSLSLSQSKLPVPQKAVPYPQRDM 200
201 PIQAFLLLYQEPYLGVPVRGPFPIIIV 224

Figure 4.2  Amino acid sequences of the bovine α- and β-caseins. Swiss-Prot accession numbers, entry names, protein names, species and sequences for the α- and β-caseins. The database sequences are for the B variant of αs1-casein, the A variant of αs2-casein and the A2 variant of β-casein. Potential phosphorylation sites are shown in bold type and those that have been experimentally confirmed are underlined. The signal peptides are shown in italics.

↓

1 MMKSSFLVVTILALTLPFLGAEQENQEOPIRECEKDERFFSKIAKYIPIQ 50
51 YVLRSRPYQGLNYYQKQPVALINQQLPPYPPYAKPAAVSPAOQILQWQLV 100
101 SNTVPK继承AQPTTMARHPHPLSPMAIPPKKNQDKTEIPINTIASEG 150
151 PTSTPTTEAVESTNTELEDSPSVIESPEINTVQTVSTAV 190

Figure 4.3  Amino acid sequence of κ-casein. Amino acid sequence of κ-casein A (Swiss-Prot accession number P02668). The N-terminal signal sequence (1–21) is shown in italics. The arrow indicates the amino-terminus of the mature protein. Recognized sites of potential phosphorylation and glycosylation are indicated in bold, underlined text. Amino acid substitutions distinguishing the B variant are shown above the main sequence.

The reference protein for αs1-casein is αs1-CN B-8P, where B-8P signifies the B genetic variant with eight phosphates (Farrell et al., 2004).

A number of genetic variants of αs1-casein have been described (Farrell et al., 2004). Only the less common forms, D and F, are likely to have altered phosphorylation profiles. Variant D has Ala68 substituted with Thr, which generates a phosphorylation site of the form Thr-Xxx-Glu. Phosphorylation of this residue was detected
when the variant was first identified (Grosclaude et al., 1972a). Variant F has Ser$^{81}$ substituted with Leu, which disrupts the serine cluster—Ser$^{79}$-Ile-Ser-Ser-Ser-Glu-Glu$^{85}$—and eliminates the secondary phosphorylation sites at Ser$^{79}$ and Ser$^{81}$. The amino acid sequence and the modifications of $\alpha_{s1}$-casein are shown in Figure 4.2.

$\alpha_{s2}$-Casein

The $\alpha_{s2}$-casein component of bovine milk is more varied than the $\alpha_{s1}$-casein component. It generally presents as a mixture of four phosphoforms with 10–13 phosphates. The reference protein for $\alpha_{s2}$-casein is $\alpha_{s2}$-CN A-11P (Farrell et al., 2004). The A variant has 12 serine residues in Ser-Xxx-Glu/pSer motifs and four threonine residues in Thr-Xxx-Glu motifs (Figure 4.2). Consequently, up to 16 phosphates are theoretically possible. Presumably, the 12 serine residues are the first to be phosphorylated. However, it is not known whether specific residues remain unphosphorylated in the different forms but, given the consistent appearance of the forms in milk, it is likely to be the case. Unfortunately, it is not possible to draw any conclusions with regard to phosphorylation site occupation until the individual phosphoforms are analyzed. This should be possible using gel-based proteomic techniques.

Only four genetic variants of $\alpha_{s2}$-casein have been described (Farrell et al., 2004) and, in each case, an altered phosphorylation profile would be expected. In variant B, Ser$^{23}$ is changed to Phe as a result of a single nucleotide substitution (Ibeagha-Awemu et al., 2007). This causes loss of a phosphorylation site in the first phosphoserine cluster. Variant C has three amino acid changes: Glu$^{48}$ is changed to Gly, with loss of the phosphorylation site at Ser$^{46}$; Ala$^{62}$ is changed to Thr, creating a potential site with the motif Thr$^{62}$-Xxx-Glu$^{64}$; Thr$^{145}$ is changed to Ile, with loss of the potential site at Thr$^{145}$.

Variant D has a deletion of nine amino acids as a result of skipping exon VIII (Bouniol et al., 1993). This results in loss of the first three serines from the second phosphoserine cluster. A second PTM on $\alpha_{s2}$-casein is the formation of an intramolecular disulfide bond between the two cysteine residues in the protein (Rasmussen et al., 1994). The functional role of disulfide bonding is not clear at this stage, but it may contribute to micelle stability and is discussed further in the section on $\kappa$-casein.

$\beta$-Casein

Bovine $\beta$-casein is usually present as a single form with five phosphates, indicating that all five Ser-Xxx-Glu/pSer sites in the sequence are constitutively phosphorylated. The reference protein is $\beta$-CN A$^{2}$-5P (Farrell et al., 2004). Some 12 genetic variants of $\beta$-casein have been characterized, but only two variants appear to have altered phosphorylation profiles. Variant C has a Glu to Lys substitution at residue 52, which removes the phosphorylation site at Ser$^{50}$. Variant D has a Ser to Lys substitution at residue 33, which removes the primary phosphorylation site at Ser$^{33}$. Variation in $\beta$-casein arises primarily as a result of proteolysis rather than PTMs. The sequence and the phosphorylation sites of $\beta$-casein are summarized in Figure 4.2.

Although much of the focus has been on bovine milk, other species have not been entirely neglected (Ginger and Grigor, 1999). It is apparent that considerable variations
Post-translational modifications of caseins occur between different species. For example, the β-casein of human milk exists as six different forms with 0 to 5 phosphates (Greenberg et al., 1984). Equine β-casein also shows variation in phosphorylation, with typically 3 to 7 phosphates on full-length β-casein (Girardet et al., 2006) and 1 to 7 phosphates on a low-molecular-weight form that arises from an internal deletion (Miclo et al., 2007). Ovine β-casein has also been reported to be variably phosphorylated, with 0 to 7 phosphates (Ferranti et al., 2001), although it is not clear where the seventh phosphorylation site is. Caprine β-casein appears to be more like bovine β-casein with the same five phosphorylation sites and an additional site on Thr27 (Neveu et al., 2002).

κ-casein is not part of the micelle structure. Its major feature is a variable degree of glycosylation. Its major feature is a variable degree of glycosylation. The keen interest in κ-casein arises largely from its key role as a stabilizer of the micelle structure. In mice, the absence of κ-casein causes a failure of lactation, as the lumina of the mammary gland become clogged with aggregated caseins (Shekar et al., 2006). The PTMs of κ-casein have been the subject of more recent research and are covered here in much greater detail than those of the other caseins.

The full amino acid sequence of bovine κ-casein was first reported in 1973 (Mercier et al., 1973). The mature protein consists of a single chain of 169 amino acids (Figure 4.3) and has a theoretical molecular weight of 18974 Da and a theoretical pl of 5.93 (A variant). The amino terminal glutamine is cyclized to form a pyrrolidone glutamic acid residue. The bovine κ-casein gene sequence was published in 1988 (Alexander et al., 1988). It consists of five exons spread over about 14 kilobases, with most of the protein-coding region located in exon 4. A cleavable amino terminal signal sequence of 21 amino acids directs secretion of the mature protein.

A number of polymorphisms in the κ-casein gene have been identified, resulting in one or more amino acid substitutions in the mature protein. The most common variants are the A and B variants, which differ by two amino acids (Asp169/Thr157 in variant A and Ala169/Ile157 in variant B). The genetic variants of κ-casein are summarized in Table 4.1. A number of polymorphisms in the non-coding region have also been identified (Schild et al., 1994; Keating et al., 2007). Although these do not affect the amino acid sequence, they have the potential to affect expression levels. The full amino acid sequences of κ-casein from 18 species are currently in the Swiss-Prot database, with another 24 entries covering sub-species and incomplete sequences (Table 4.2). The reference protein for κ-casein is κ-CN A-1P, UniProt P02668 (Farrell et al., 2004).

Phosphorylation

κ-casein appears to be constitutively phosphorylated at Ser170 and only partially phosphorylated at Ser148 (Talbot and Waugh, 1970; Mercier et al., 1973). A minor tri-phosphorylated form has also been detected (Vreeman et al., 1986; Molle and Leonil, 1995). Other studies have managed to detect only mono-phosphorylated
Table 4.1 Genetic variants of bovine κ-casein

<table>
<thead>
<tr>
<th>Variant</th>
<th>Amino acid changes (relative to A variant)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td></td>
<td>(Grosclaude et al., 1972)</td>
</tr>
<tr>
<td>B</td>
<td>Thr\textsuperscript{157} to Ile, Asp\textsuperscript{169} to Ala, Thr\textsuperscript{157} to Ile</td>
<td>(Mercier et al., 1973)</td>
</tr>
<tr>
<td>B\textsuperscript{2}</td>
<td>Asp\textsuperscript{169} to Ala, Ile\textsuperscript{174} to Thr, Arg\textsuperscript{136} to His</td>
<td>(Gorodetskii et al., 1983)</td>
</tr>
<tr>
<td>C</td>
<td>Thr\textsuperscript{157} to Ile, Asp\textsuperscript{169} to Ala</td>
<td>(Miranda et al., 1993)</td>
</tr>
<tr>
<td>E</td>
<td>Ser\textsuperscript{176} to Gly, Asp\textsuperscript{169} to Val</td>
<td>(Schlieben et al., 1991)</td>
</tr>
<tr>
<td>F\textsuperscript{1}</td>
<td>Asp\textsuperscript{169} to Val</td>
<td>(Sulimova et al., 1992)</td>
</tr>
<tr>
<td>F\textsuperscript{2}</td>
<td>Arg\textsuperscript{31} to His, Arg\textsuperscript{118} to Cys</td>
<td>(Prinzenberg et al., 1996)</td>
</tr>
<tr>
<td>G\textsuperscript{2}</td>
<td>Asp\textsuperscript{169} to Ala</td>
<td>(Sulimova et al., 1996)</td>
</tr>
<tr>
<td>H</td>
<td>Thr\textsuperscript{156} to Ile, Ser\textsuperscript{125} to Ala, Thr\textsuperscript{157} to Ile</td>
<td>(Prinzenberg et al., 1999)</td>
</tr>
<tr>
<td>I</td>
<td>Ser\textsuperscript{125} to Ala, Arg\textsuperscript{31} to His</td>
<td>(Prinzenberg et al., 1999)</td>
</tr>
<tr>
<td>J</td>
<td>Asp\textsuperscript{169} to Ala, Ser\textsuperscript{176} to Arg</td>
<td>(Mahe et al., 1999)</td>
</tr>
<tr>
<td>A(1)</td>
<td>Silent (Pro\textsuperscript{150}, CCA to CCG)</td>
<td>(Prinzenberg et al., 1999)</td>
</tr>
</tbody>
</table>

Notes: Other variants have been described but not confirmed or have proven to be identical to those above.

forms (Rasmussen et al., 1997; Riggs et al., 2001) although, in one of these (Riggs et al., 2001), the phosphorylation site appears to have been identified incorrectly.

Phosphorylation has also been examined by MS of intact κ-casein extracted from 2-D gels. Both mono- and di-phosphorylated forms were observed and phosphorylation at Ser\textsuperscript{170} was confirmed by MS/MS (Claverol et al., 2003). However, the electrophoretic mobility of some phospho-forms was not consistent with the MS analysis and probably reflected artifactual modification (e.g. deamidation) during purification. Using 2-DE with isoelectric focusing as the first dimension, phosphorylation variants in whole milk can be easily resolved because of the pI shifts caused by the acidic phosphate groups (Holland et al., 2004).

The two main phosphorylation sites have been confirmed by tandem MS sequencing of peptic peptides released from protein forms separated by 2-DE. Tri-phosphorylated forms of both the A and B variants have been observed and the third phosphorylation site has been identified recently as Thr\textsuperscript{166} (Holland et al., 2006). This site is consistent with the general observation of casein phosphorylation on the Ser/Thr-Xxx-Glu/pSer motif, with only relative low levels of phosphorylation on threonine residues.

**Glycosylation**

Whereas about 40% of κ-casein has been estimated to be non-glycosylated, the remaining 60% has up to six glycans attached (Vreeman et al., 1986). The presence
of sugars on $\kappa$-casein was recognized as long ago as 1961 (Alais and Jolles, 1961) and, during the 1970s and 1980s, a large number of studies were directed at elucidating the sugar composition, sequence and sites of attachment to the protein. The major glycan is a tetrasaccharide composed of galactose (Gal), N-acetylgalactosamine (GalNAc) and sialic or neuraminic acid (NeuAc) of the form

<table>
<thead>
<tr>
<th>Name (accession number)</th>
<th>Full length or fragment</th>
<th>Species (common name)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CASK_BALPH (Q27952)</td>
<td>(Fragment)</td>
<td><em>Balaeoptera physalus</em> (Finback whale) (Common rorqual)</td>
</tr>
<tr>
<td>CASK_BISBO (P42155)</td>
<td>(Fragment)</td>
<td><em>Bison bonasus</em> (European bison)</td>
</tr>
<tr>
<td>CASK_BOVIN (P02668)</td>
<td>precursor</td>
<td><em>Bos taurus</em> (Bovine)</td>
</tr>
<tr>
<td>CASK_BUBBU (P11840)</td>
<td>precursor</td>
<td><em>Bubalus bubalis</em> (Domestic water buffalo)</td>
</tr>
<tr>
<td>CASK_CAMDR (P79139)</td>
<td>precursor</td>
<td><em>Camelus dromedarius</em> (Dromedary) (Arabian camel)</td>
</tr>
<tr>
<td>CASK_CAPCA (Q95146)</td>
<td>(Fragment)</td>
<td><em>Capreolus capreolus</em> (Roe deer)</td>
</tr>
<tr>
<td>CASK_CAPCR (P42156)</td>
<td>precursor</td>
<td><em>Capricornis crispus</em> (Japanese serow)</td>
</tr>
<tr>
<td>CASK_CAPHI (P02670)</td>
<td>precursor</td>
<td><em>Capra hircus</em> (Goat)</td>
</tr>
<tr>
<td>CASK_CAPSU (P50420)</td>
<td>precursor</td>
<td><em>Capricornis sumatrensis</em></td>
</tr>
<tr>
<td>CASK_CAPSW (P50421)</td>
<td>precursor</td>
<td><em>Capricornis swinhoei</em></td>
</tr>
<tr>
<td>CASK_CAVPO (P19442)</td>
<td>precursor</td>
<td><em>Cavia porcellus</em> (Guinea pig)</td>
</tr>
<tr>
<td>CASK_CERDU (Q95147)</td>
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<td><em>Cervus duvauceli</em> (Swamp deer)</td>
</tr>
<tr>
<td>CASK_CEREL (Q95149)</td>
<td>(Fragment)</td>
<td><em>Cervus elaphus</em> (Red deer)</td>
</tr>
<tr>
<td>CASK_CERNI (P42157)</td>
<td>precursor</td>
<td><em>Cervus nippon</em> (Sika deer)</td>
</tr>
<tr>
<td>CASK_CERUN (Q95177)</td>
<td>(Fragment)</td>
<td><em>Cervus unicolor</em> (Sambur)</td>
</tr>
<tr>
<td>CASK_ELADA (Q95184)</td>
<td>(Fragment)</td>
<td><em>Elaphurus davidianus</em> (Pere David’s deer)</td>
</tr>
<tr>
<td>CASK_EQUGR (Q28400)</td>
<td>(Fragment)</td>
<td><em>Equus grevyi</em> (Grevy’s zebra)</td>
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<td>CASK_GIRCA (Q28417)</td>
<td>(Fragment)</td>
<td><em>Giraffa camelopardalis</em> (Giraffe)</td>
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<td><em>Hippopotamus amphibius</em> (Hippopotamus)</td>
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<td>precursor</td>
<td><em>Equus caballus</em> (Horse)</td>
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<td>precursor</td>
<td><em>Homo sapiens</em> (Human)</td>
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</tr>
<tr>
<td>CASK_OVIDA (Q95224)</td>
<td>(Fragment)</td>
<td><em>Ovis dalli</em> (Dall sheep)</td>
</tr>
<tr>
<td>CASK_OVIMO (Q95227)</td>
<td>(Fragment)</td>
<td><em>Ovis moschatus</em> (Muskox)</td>
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<td>CASK_PIG (P11841)</td>
<td>precursor</td>
<td><em>Sus scrofa</em> (Pig)</td>
</tr>
<tr>
<td>CASK_RABIT (P33618)</td>
<td>precursor</td>
<td><em>Oryctolagus cuniculus</em> (Rabbit)</td>
</tr>
<tr>
<td>CASK_RANTA (Q95239)</td>
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<td><em>Rangifer tarandus</em> (Reindeer) (Caribou)</td>
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<td>CASK_RAT (P04468)</td>
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<td><em>Rattus norvegicus</em> (Rat)</td>
</tr>
<tr>
<td>CASK_RUPRU (P50424)</td>
<td>precursor</td>
<td><em>Rupicapra rupicapra</em> (Chamois)</td>
</tr>
<tr>
<td>CASK_SAITA (P50425)</td>
<td>precursor</td>
<td><em>Saiga tatarica</em></td>
</tr>
<tr>
<td>CASK_SHEEP (P02669)</td>
<td>precursor</td>
<td><em>Ovis aries</em> (Sheep)</td>
</tr>
<tr>
<td>CASK_TAPI (Q29135)</td>
<td>(Fragment)</td>
<td><em>Tapirus indicus</em> (Asiatic tapir) (Malayan tapir)</td>
</tr>
<tr>
<td>CASK_TAYTA (Q28794)</td>
<td>(Fragment)</td>
<td><em>Tayassu tajacu</em> (Collared peccary) (Pecari tajacu)</td>
</tr>
<tr>
<td>CASK_TRAJ (Q29137)</td>
<td>(Fragment)</td>
<td><em>Tragulus javanicus</em> (Lesser Malay chevrotain)</td>
</tr>
<tr>
<td>CASK_UNCUN (Q29150)</td>
<td>(Fragment)</td>
<td><em>Uncia uncia</em> (Snow leopard) (Panthera uncia)</td>
</tr>
</tbody>
</table>
NeuAca(2-3)Galβ(1-3)[NeuAca(2-6)]GalNAc, but monosaccharide (GalNAc), disaccharide (Galβ(1-3)GalNAc) and trisaccharide (NeuAca(2-3)Galβ(1-3)GalNAc or Galβ(1-3)[NeuAca(2-6)]GalNAc) are also found (Fournet et al., 1975, 1979; van Halbeek et al., 1980; Fiat et al., 1988; Saito and Itoh, 1992).

The relative amounts have been determined by high-performance liquid chromatography (HPLC) as 56.0% tetrasaccharide, 36.9% trisaccharide (18.4% linear and 18.5% branched), 6.3% disaccharide and 0.8% monosaccharide (Saito and Itoh, 1992). It is not known whether the minor forms arise from incomplete synthesis of the tetrasaccharide in mammary epithelial cells or are products of degradation after synthesis and/or secretion of κ-casein into the lumen of the mammary gland.

Establishment of the attachment site(s) of the glycans has been more controversial. On the basis of Edman sequencing of short glycopeptides obtained by enzymatic digestion, Jolles et al. (1973) proposed Thr^{152} or Thr^{154} as the glycan attachment site. Kanamori et al. (1980) proposed Thr^{152}, Thr^{154} and Thr^{156} (or Thr^{157}) after analyzing a glycopeptide that was derived from κ-casein and that contained three GalNAc residues. Work from the same laboratory on bovine κ-casein from colostrum also indicated glycosylation at Thr^{152}, Thr^{154} and Thr^{156} (Doi et al., 1980). Subsequently, using a different peptide fraction prepared from κ-casein of normal milk, glycosylation at Thr^{154} and Ser^{162} was reported (Kanamori et al., 1981). Meanwhile, further work from Jolles’ laboratory identified Thr^{152} as the glycan attachment site on κ-casein from normal milk and Thr^{152} and Thr^{162} as the attachment sites on κ-casein from colostrum (Fiat et al., 1981). Zevaco and Ribadeau-Dumas (1984) suggested that glycans could be attached to any of the previously identified sites (Thr^{152}, Thr^{154}, Thr^{156}, Thr^{157}, Ser^{162} or Thr^{163}) but their published study contained no conclusive evidence for any site.

All these studies were limited by the technology available at the time. In normal Edman sequencing, glycosylated serine or threonine residues are not detected and their presence is inferred from a blank in the sequencing cycle where serine or threonine is expected (for a more detailed discussion, see Pisano et al., 1994). This is further complicated by the fact that serine and threonine are themselves low-yield amino acids. Thus, in assigning O-glycosylation sites, Edman sequencing data can easily be misinterpreted.

Conclusive identification of glycosylation sites in κ-casein was achieved using solid-phase Edman sequencing, which allows the direct detection of glycosylated serine and threonine residues (Pisano et al., 1994). Variable levels of glycosylation at Thr^{142}, Thr^{152}, Thr^{154}, Thr^{157} (A variant only), Thr^{163} and Thr^{186} were detected and no evidence of glycosylation at any serine residue was obtained. However, even this study did not give the full picture of κ-casein glycosylation, as it could detect only average glycosylation site occupancy in a crude mixture of glycoforms.

We have shown that κ-casein glycoforms can be separated by 2-DE (Holland et al., 2004) and the resolution obtained is shown in Figure 4.4. When the glycosylation site occupancy of individual glycoforms was investigated using tandem MS sequencing of chemically tagged peptides, an interesting pattern was observed. The monoglycoform was glycosylated exclusively at Thr^{152}, the di-glycoform was glycosylated at Thr^{152} and Thr^{163} and the tri-glycoform was glycosylated at Thr^{152}, Thr^{154} and Thr^{163} (Holland et al., 2005). Further studies using enriched fractions of κ-casein...
Post-translational modifications of caseins

Separated on 2-D gels showed up to six glycans on κ-casein B where only five sites had been previously identified. Tandem MS analysis provided evidence for glycosylation at Thr\(^{166}\) on the tetra-glycoform (Holland et al., 2006). The remaining two glycosylation sites were not confirmed but were presumably on Thr\(^{142}\) and Thr\(^{186}\), as proposed earlier (Pisano et al., 1994). This pattern is illustrated in Figure 4.5.

As Thr\(^{166}\) can be phosphorylated or glycosylated, there is potential for competition at this site. However, as both the tri-phosphate and tetra-glycoforms are very minor forms, it may be of little significance. Overall, the glycosylation of κ-casein in the mammary epithelial cells appears to take place in a highly controlled manner and this suggests that it is a rather important process with considerable functional significance.

**Disulfide bonding**

κ-Casein purified from bovine milk occurs as both monomeric forms and oligomeric forms with up to eight or more monomers linked by disulfide bonds (Swaisgood et al., 1964; Talbot and Waugh, 1970). A more recent study has shown that reduced and carboxymethylated κ-casein can form large fibrillar structures, although these do not occur in milk (Farrell et al., 2003). The nature of the disulfide-linked complexes has been examined by a number of authors (Groves et al., 1992, 1998; Rasmussen et al., 1992, 1994, 1999; Farrell et al., 2003).

There are only two cysteine residues (Cys\(^{32}\) and Cys\(^{109}\)) in bovine κ-casein (Mercier et al., 1973) and they appear to be randomly linked in disulfide bonds in oligomeric forms (Rasmussen et al., 1992). In monomeric κ-casein, the two cysteines form an intramolecular disulfide bond (Rasmussen et al., 1994). As can be seen

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**Figure 4.4** Heterogeneity of κ-casein in cow’s milk. 2-D gel showing multiple forms of κ-casein. The genetic variant, number of phosphate residues and glycosylation state are indicated. Numbers in brackets indicate the extra negatively charged residues relative to κ-casein B-1P. Adapted from Holland et al. (2004).
in Figure 4.6, disulfide-bonded monomers, dimers and trimers can be resolved on 2-D gels of whole milk when reducing agents are omitted (Holland et al., 2008). It is not clear whether these higher order complexes of \( \kappa \)-casein have any importance in micellar structure but it would be expected that they would be less likely to dissociate from the micelles.

The cysteine residues are not well conserved across species, with human, porcine and rodent \( \kappa \)-caseins containing only a single cysteine, precluding the formation of disulfide-linked oligomers larger than dimers (Rasmussen et al., 1999; Bouguyon et al., 2006). However, the ability of \( \kappa \)-casein to form disulfide-linked complexes with itself or with other proteins during heat treatment is relevant to dairy processing (see below). The combined PTMs of bovine \( \kappa \)-casein are summarized in Figure 4.7.

**Sources and functional significance of \( \kappa \)-casein heterogeneity**

Although the heterogeneity of \( \kappa \)-casein has been recognized for many decades and the structural elements are now fairly well defined, the source of the heterogeneity,
particularly in glycosylation, and its functional role(s) are not known. Early studies on the influence of the glycosylation of κ-casein on its biological properties have been reviewed previously (Dziuba and Minikiewicz, 1996). They highlighted a number of studies addressing factors that could influence the degree of glycosylation
and what influence glycosylation might have on micelle stability. The sections below cover some of those studies again and highlight more recent work related to the sources and functional significance of κ-casein heterogeneity.

**Sources of heterogeneity**

A large number of studies have examined the influence of milk protein polymorphism on milk composition and yield, and these have been extensively reviewed (e.g. Ng-Kwai-Hang, 1997; Martin et al., 2002). However, in many cases, the results have been inconsistent, which is probably a reflection of the multi-factorial nature of milk production. It is difficult to isolate the effects of a single protein polymorphism from those of the other major milk proteins, especially as there appears to be a substantial degree of co-ordination of their expression. There are also a number of environmental or cow factors such as feed type and lactational stage that are frequently interrelated, as they all vary with the seasonal changes in dairy farming. Studies on specific effects of κ-casein variants have largely focused on the common A and B variants and there appears to be a consensus that milk from B variant cows contains more fat, protein, casein and κ-casein than milk from A variant cows (Bovenhuis et al., 1992; Ng-Kwai-Hang, 1997; Bobe et al., 1999).

Studies relating to the glycosylation status of κ-casein are more limited. Robitaille et al. (1991a) identified a number of factors that appeared to contribute to variation in the NeuAc content of bovine κ-casein. The NeuAc content was higher in cows with the κ-casein AB phenotype than in cows with the AA phenotype; it decreased with increasing parity and varied over the course of lactation, dropping to a minimum at 2–3 months after calving before increasing over the next 9–10 months. They also examined the association between κ-casein glycosylation and milk production/composition (Robitaille et al., 1991b).

Although there appeared to be a statistically significant association between the NeuAc content of κ-casein and milk yield, the most striking result of these investigations was the variability of NeuAc/κ-casein measurements (mean, $64\pm21\,\mu g/mg$; range, $23–166\,\mu g/mg$), which suggests that other factors could have had a large impact on glycosylation or that the inherent variability in the assay masked any true associations. Limited 2-D gel analyses suggest that the pattern of glycosylation is far more consistent than these measurements indicate (Holland et al., 2004, 2005).
It would be very informative to examine some of these supposed extremes in NeuAc and hence the glycosylation level of κ-casein on 2-D gels.

Significant differences in the content of non-glycosylated κ-casein in milk have been reported for cows of different κ-casein genotypes (Lodes et al., 1996). Non-glycosylated κ-casein levels were higher (as a percentage of total protein) in milk from cows with the B variant than in milk from cows with the A variant. The rarer variants, C and E, were generally associated with lower levels. However, as no measurements of glycosylated κ-casein levels were reported, no effect of genetic variant on glycosylation can be inferred from this report.

Electrophoretic and chromatographic techniques have been used to profile the κ-casein macropeptide from cows of the AA and BB phenotypes (Coolbear et al., 1996). They found that the B variant macropeptide was more highly glycosylated than the A variant, with an increased content of both hexosamine (i.e. GalNAc) and sialic acid (i.e. NeuAc). After anion-exchange HPLC on a MonoQ column, the elution profile of the B variant contained more peaks, suggesting that an increased number of oligosaccharide chains were attached. These results were consistent with earlier studies suggesting more extensive glycosylation of the B variant (Vreeman et al., 1986) compared with the A variant (Molle and Leonil, 1995), despite the fact that the A variant contains an extra (potential) glycosylation site (Pisano et al., 1994). From these and other results, Coolbear et al. (1996) suggested that there were generally consistent patterns of glycosylation for the genetic variants but that the overall extent of glycosylation could vary.

Variations in κ-casein glycosylation during pregnancy and lactation have been touched on above. Early studies indicated a higher degree of glycosylation of κ-casein in colostrum than in mature milk as well as the presence of an additional sugar moiety, N-acetylglucosamine (GlcNAc) (Guerin et al., 1974; Fournet et al., 1975). Subsequently, a number of studies addressed the structure of the oligosaccharides attached to colostral κ-casein and how they varied with time after parturition (Saito et al., 1981a, 1981b, 1982; van Halbeek et al., 1981; Fiat et al., 1988).

As well as the structures already identified above in normal milk, the following structures have been reported: the acidic hexasaccharide, NeuAca(2-3)Galβ(1-3)[NeuAca(2-3)Galβ(1-4)GlcNAcβ(1-6)]GalNAc; the acidic pentasaccharide, NeuAca(2-3)Galβ(1-3)[Galβ(1-4)GlcNAcβ(1-6)]GalNAc; the acidic tetrasaccharide, GlcNAcβ(1-3)Galβ(1-3)[NeuAca(2-6)]GalNAc; the neutral pentasaccharide Galβ(1-3)[Galβ(1-4){Fuca(1-3)}GlcNAcβ(1-6)]GalNAc; the neutral tetrasaccharide, Galβ(1-3)[Galβ(1-4)GlcNAcβ(1-6)]GalNAc; and the neutral trisaccharide, Galβ(1-3)[GlcNAcβ(1-6)]GalNAc.

This extra complexity is already observable 15 min after parturition but decreases to normal over about 66 h (Fiat et al., 1988). These results suggest changes in the expression profiles of the glycosyltransferases responsible for assembling the O-linked glycans on κ-casein. The initial step of attachment of GalNAc to a threonine residue is catalyzed by UDP-GalNAc:polypeptide N-acetylgalactosaminyltransferase (GalNAcT) and, although not much is known about the bovine enzymes, there are at least 12 mammalian GalNAcTs that have been cloned and functionally expressed (Ten Hagen et al., 2003). In mice, the expression of several isoforms changes markedly during pregnancy and lactation (Young et al., 2003). It is likely
that expression of the other required glycosyltransferases also varies, presumably under control of lactogenic hormones.

Other seasonal factors related to climate, such as heat stress, drought and nutrition (e.g. pasture versus fodder), can have an impact on milk production and composition. However, we are not aware of any specific studies on their effect on κ-casein glycosylation.

Functional significance

κ-Casein plays a key role in micelle stability by acting as a hairy layer that provides both steric and electrostatic repulsion between micelles, preventing aggregation. Glycosylation of κ-casein increases both the size of the hydrophilic C-terminal “hairs” and their charge—because of the bulk of the hydrophilic sugar residues with their hydration shells and the negative charge of the neuraminic acid groups respectively. Theoretically, the higher the degree of glycosylation of κ-casein, the greater its stabilizing ability should be. As such, it might be expected that the degree of glycosylation of κ-casein would have a marked effect on both the size and the stability of the casein micelles.

However, whereas the size of the casein micelles has been shown to be inversely related to their κ-casein content (relative to total casein) by a number of authors (McGann et al., 1980; Davies and Law, 1983; Donnelly et al., 1984; Dalgleish et al., 1989; O’Connell and Fox, 2000), there is no clear correlation between micelle size and degree of glycosylation. Slattery (1978) found an apparent inverse relationship between the proportion of glycosylated κ-casein and micelle size but it did not apply to all of the size fractions isolated.

In contrast, Dalgleish (1985, 1986) found that the proportions of glycosylated and non-glycosylated κ-casein did not vary with micelle size. More recently, O’Connell and Fox (2000) showed an apparent increase in κ-casein glycosylation with increasing micelle size. Some of these discrepancies are probably the result of the different experimental approaches adopted. Currently, there is no conclusive evidence for a distinct relationship between micelle size and κ-casein glycosylation.

As stated above, micellar stability, or controlled destabilization in the case of cheese and yoghurt manufacture, is of key importance in dairy manufacturing. A number of authors have looked for effects of κ-casein heterogeneity on micellar stability and the processing properties of milk. Takeuchi et al. (1985) used ion-exchange chromatography to prepare nine fractions of κ-casein A-1P that varied in the level of glycosylation. The ability of these sub-fractions to stabilize αs1-casein was shown to increase with increasing carbohydrate content. In cheese manufacture, the initial step is the chymosin (rennet-)catalyzed cleavage of the Phe126-Met127 bond in κ-casein, resulting in release of the hydrophilic glycomacropeptide from the micelle surface, which leads to micellar aggregation or clotting.

Doi et al. (1979) examined the susceptibility to chymosin action of κ-casein preparations with different degrees of glycosylation. They found that more highly glycosylated forms were less susceptible to hydrolysis not only by chymosin but also by other proteases. Others have also found an inverse relationship between glycosylation and
chymosin susceptibility with purified \( \kappa \)-casein fractions (Addeo et al., 1984; Vreeman et al., 1986) and in model systems (Addeo et al., 1984; Leaver and Horne, 1996).

In milk, the relationship is not so clear. Chaplin and Green (1980) claimed that all \( \kappa \)-casein molecules were hydrolyzed with equal efficiency, whereas van Hooydonk et al. (1984) found that the rate of chymosin-catalyzed hydrolysis decreased with increasing glycosylation. Again, differences in experimental approach were probably responsible for at least some of the apparent discrepancy. Rennet clotting time (RCT), rate of curd firming and curd firmness have been measured to assess the effect of \( \kappa \)-casein glycosylation on the coagulation properties of milk (Robitaille et al., 1993). Whereas no effect on RCT was observed, the rate of curd firming decreased and the curd firmness increased at higher glycosylation levels.

Differences in rennet coagulation properties have also been observed for genetic variants of \( \kappa \)-casein. Shorter RCTs, higher curd-firming rates and higher curd firmness have been reported for milk from cows with the BB variant than for milk from cows with the AA variant (Walsh et al., 1998; and references therein). These differences between A and B variant milks were maintained after heat treatments of up to 80°C for 2 min, despite an overall deterioration of the coagulation properties at elevated temperatures (Choi and Ng-Kwai-Hang, 2003). Milks containing the rarer \( \kappa \)-casein C variant form rennet gels even more slowly than the A or B variant milks, possibly because of the substitution of histidine for arginine at residue 118, which may affect chymosin binding (Smith et al., 1997). Similar results have been observed for the \( \kappa \)-casein G variant, which has cysteine at residue 118 (Erhardt et al., 1997), and a similar explanation has been proposed (Smith et al., 1997).

Coagulation of milk can also be induced by acid, as is the case in yoghurt manufacture. There are fewer studies on the effect of the glycosylation of \( \kappa \)-casein on acid coagulability. Cases et al. (2003) found that partial deglycosylation with neuraminidase had little effect on micellar surface charge and solvation but caused a decrease in acid gelation time, a higher rate of gel firming and a higher final firmness.

Heat treatment of milk can also destabilize the casein micelle structure. The heat-induced coagulation of milk is a very complex process that is affected by many parameters (O’Connell and Fox, 2003). A number of studies have examined the influence of genetic variants of \( \kappa \)-casein on heat stability parameters and it is generally accepted that B variant milks are more stable than A variant milks (FitzGerald and Hill, 1997). The reason may be more related to the effects on \( \kappa \)-casein concentration and micelle size mentioned above than to the structural differences between the variants (Smith et al., 2002). Again, there are fewer studies related to the influence of the glycosylation of \( \kappa \)-casein on heat stability. Using a model system composed of casein micelles in simulated milk ultrafiltrate, Minkiewicz et al. (1993) showed that enzymatic removal of neuraminic acid using neuraminidase caused a decrease in heat stability.

However, Robitaille and Ayers (1995), using whole milk, could not find a significant effect of neuraminidase treatment on heat stability. When milk is heated above 65°C, \( \beta \)-lactoglobulin denatures, exposing a previously buried sulfhydryl group that can participate in disulfide exchange reactions with other cysteine-containing proteins including \( \kappa \)-casein. This interaction has been recognized for many decades (Sawyer, 1969) and has been the subject of numerous investigations and reviews over
the years, but a detailed analysis is beyond the scope of this review (for an extensive review, see O’Connell and Fox, 2003). Recent studies have addressed both the mechanism of formation (Guyomarc’h et al., 2003) and the impact on product quality (Vasbinder et al., 2003) of disulfide-linked complexes. Despite the vast amount of literature on this topic, there do not appear to be any studies that have addressed the impact of the variable glycosylation of κ-casein on its ability to form disulfide-linked complexes either with itself or with β-lactoglobulin.

Heat-induced changes in micelle structure are particularly relevant for UHT milk production and storage. The extremes of heat treatment (of the order of 140–145°C for 4–10 s) produce a number of changes in the milk, not least of which is the formation of κ-casein-β-lactoglobulin complexes. On storage, UHT-treated milks show a variable tendency to form gels and this phenomenon, known as age gelation, affects product shelf life (for a review, see Datta and Deeth, 2001). Again, despite extensive studies over many years on UHT processing and product performance, the influence of κ-casein heterogeneity, particularly heterogeneity with respect to glycosylation, has not been addressed.

From a theoretical perspective, higher initial levels of glycosylation may act to temper the deleterious effects of heat treatment through effects on micellar size, micellar stability and the formation of disulfide-linked complexes. The heat treatment itself may affect the glycosylation level at the surface of the micelle either indirectly, through loss of κ-casein in complex formation with β-lactoglobulin, or directly, through degradation of glycosidic residues (van Hooydonk et al., 1987; as quoted in Dziuba and Minikiewicz, 1996). Subsequent changes in the glycosylation level during storage could be mediated by the action of heat-stable glycosidases originating from psychrotrophic bacteria present in the raw milk (Marin et al., 1984).

Release of monosaccharides during the storage of UHT milk has been observed (Recio et al., 1998; Belloque et al., 2001). Thus, both the initial glycosylation level of the κ-casein and the residual amount after UHT treatment may affect the storage properties of UHT-treated milk. As the actions of heat-resistant proteases can contribute to the age gelation of UHT milk, the inhibitory effects of glycosylation on the activity of proteases such as plasmin (Doi et al., 1979) may be important for prolonging shelf life. Unravelling specific effects will require the application of modern proteomic technologies for κ-casein analysis (Claverol et al., 2003; Holland et al., 2004, 2005, 2006; O’Donnell et al., 2004). Using these technologies, it will be possible to elaborate the heterogeneous glycoforms of κ-casein in raw milk, after pre-treatment(s), after UHT processing and during storage leading up to gelation. This will allow a definitive assessment of the functional significance of κ-casein glycosylation.

One aspect of κ-casein heterogeneity that has not been considered above is its influence on the biological properties of milk. This area has been reviewed extensively (Dziuba and Minikiewicz, 1996). There are two main areas to consider. Firstly, there is the nutritional contribution of the carbohydrate residues in κ-casein, particularly NeuAc. The importance of NeuAc and its roles in numerous biological functions have been reviewed recently (Schauer, 2000). NeuAc is commonly found as the terminal sugar residue on mammalian glycoproteins. Although mammals can synthesize NeuAc, the high levels in milk and especially colostrum may be related to a high demand for neonatal growth and development. The normal glycans on κ-casein
are part of a class known as the Thomsen-Friedenreich-related antigens (Dall’Olio and Chiricolo, 2001). The terminal NeuAc residues may play a key role in preventing colonization of the gut by pathogenic organisms by providing alternative binding sites that minimize binding to the normal gut epithelium.

The second aspect relates to the enormous interest in bioactive peptides derived from milk proteins (Clare and Swaisgood, 2000; Kilara and Panyam, 2003). Numerous in vitro activities have been ascribed to κ-casein, its glycomacropeptide or peptides derived from them (Dziuba and Minikiewicz, 1996; Brody, 2000). Some of these activities appear to be associated with particular forms of κ-casein (Malkoski et al., 2001) and can be glycosylation dependent (Li and Mine, 2004).

Whether or not the same activities occur in vivo is not always clear because it requires both generation and absorption of the active component during digestion and this is not easy to detect. In vivo production of glycomacropeptide is known to occur after milk ingestion (Ledoux et al., 1999; and references therein), and has been detected in the plasma of infants after the ingestion of milk (Chabance et al., 1995). Any naturally occurring bioactivity of glycomacropeptide-derived peptides could be strongly influenced by the glycosylation status of κ-casein either directly, by modifying the activity of the peptide, or indirectly, by affecting proteolysis of κ-casein and hence release of the peptide.

Conclusions

PTMs such as phosphorylation, glycosylation and perhaps disulfide bond formation play a critical role in casein micelle formation and stability. It seems somewhat surprising then that so much variability occurs in these PTMs on the caseins. Whereas significant functional differences in milk properties have been consistently reported for milks with different genetic variants of the caseins, the effects reported for variable PTMs have been limited and not always consistent. Undoubtedly, a major contributing factor has been the lack of adequate methodology to definitively assess the heterogeneous casein variants present within a milk sample and the way these change as a result of treatment. High-resolution proteomic techniques may well hold the key to advancing our knowledge of milk protein PTMs and their influence on milk quality, processability and storage stability.

References


Casein micelle structure and stability

David S. Horne

Abstract

The physico-chemical properties of the casein proteins are reviewed, highlighting the factors controlling the strength of those interactions most important to the assembly and structure of the casein micelle, namely electrostatic repulsion and hydrophobic attraction, with particular emphasis on their magnitude and range. The various strands are drawn together step-by-step to develop the dual-binding model of casein micelle assembly and structure as a polymerization of a block copolymer system. The model is then used to predict the behavioral properties of the micelle in fluid milk where, by considering the rheology of high concentration milks, the hard sphere colloidal approach is shown to be a special case limited to milk of normal concentration and pH. The necessity for a dual-binding approach is then forcefully demonstrated in its ability to provide full mechanistic explanations of observed behavior in the renneting, acid gelation and alcohol-induced destabilization of skim milk. Bond mobility is identified as a crucial factor but also important is bond location and whether bonding can be extended beyond the protective shell of the micelle—its hairy layer.

Introduction

The caseins are a family of phosphoproteins found in the milks of all mammals. They exist in these milks generally as complex aggregates or micelles of the proteins and mineral calcium phosphate. Because the caseins utilize the same calcium-sequestering mechanism to regulate the calcium phosphate concentration of their environment, they...
have recently been identified as members of a wider family of secretory calcium-binding phosphoproteins descended from a common ancestor gene (Kawasaki and Weiss, 2003, 2006). These secretory phosphoproteins include enamel matrix proteins, dentine, salivary proteins, bone extracellular matrix proteins and the caseins, amongst others. All are descended from early primordial genes by duplication and divergence to serve their specialized adaptive functions. Their genes retain common functional and sequence features even after this extensive divergence. It is thought that primordial calcium-sensitive casein genes diverged from enamel matrix protein genes before the appearance of monotremes in the Jurassic era (Kawasaki and Weiss, 2003).

Casein makes milk supersaturated with calcium phosphate. Essentially, it transports safely through the mammary gland the mineral calcium phosphate that is essential for the development of bones and teeth in the suckling infant, through the medium of the casein micelle. Locking up the calcium phosphate in this package is one aspect of the biological function of the micelle. Ensuring release of this same calcium phosphate in the gastric destination of the milk is a property of the micelle that is not generally given great consideration. More research effort has been put into trying to give mechanistic understanding to the technological behavior of the micelle, understanding necessary to achieve efficient conversion of milk into products such as cheese and yoghurt, or the behavior of milk components in emulsions or reconstituted dairy products.

Many of the physical and technological properties of the casein micelle (diffusion, viscosity, light scattering) can be described by treating the casein micelles as colloidal hard spheres (Alexander et al., 2002). The initial stages, up to the onset of instability, in processes such as renneting, acid-induced gelation and flocculation in the presence of ethanol can apparently be well described by allowing these micellar hard spheres to become adhesive, essentially, as described later, treating their \( \kappa \)-casein outer layer as a salted polyelectrolyte brush (De Kruif and Zhulina, 1996; De Kruif and Holt, 2003). Beyond the critical point in all three processes, however, changes in micellar integrity and internal structure render the simple colloidal particle approach inadequate (Horne, 2003a, 2003b; Choi et al., 2007a).

Neither is the adhesive sphere approach at all helpful in furthering our understanding of the processes of micellar assembly, the pathways to dissociation or the maintenance of micellar integrity. For that, we must turn to a structural model of the casein micelle, bearing in mind that, notwithstanding the inadequacy of the adhesive sphere approach, our micelle model has also to be adaptable enough to explain why that primitive approach has been so successful within its limitations.

Various models of casein micelle assembly and structure have appeared over the years and have been subjected to regular review and appraisal, most recently by Fox (2003). The three principal contenders are (i) the sub-micelle model of Slattery and Evard (Slattery and Evard, 1973; Slattery, 1977), subsequently elaborated by Schmidt (1980); (ii) the nanocluster model of Holt (Holt, 1992, 2004; De Kruif and Holt, 2003); (iii) the dual-binding model proposed by Horne (1998). There has been considerable debate in more recent years over the suitability and success of these models (Farrell et al., 2006; Horne, 2006). It is not our intention to rehash those arguments here but rather to highlight the necessity for a dual-binding model by demonstrating its usefulness.
in providing mechanistic understanding of the production of renneted and acidified milk gels, of micellar assembly and dissociation, and of the physical properties of micellar suspensions including their rheological behavior at high concentration. We first summarize the physico-chemical properties and interactions of the caseins and show how these lead naturally to the dual-binding picture.

**Casein primary structure and interactions**

Just as casein micelles are aggregates of all of the casein proteins and micellar calcium phosphate, so does the dual-binding model involve the properties and interactions of all of the caseins. Central to this argument are those features of the proteins that are conserved across species and through millennia. The caseins were identified as members of the wider secretory calcium phosphate binding family by their possession of functional and sequence features common to that family (Kawasaki and Weiss, 2003, 2006). Among the conserved motifs is the SXE peptide (Ser-X_{aa}-Glu) where X_{aa} may be any amino acid. In the caseins, this peptide provides a recognition template for post-translational phosphorylation of the serine in the mammary gland by a casein kinase (Mercier, 1981).

Moreover, in the caseins, the serine residues are often found clustered in groups of two, three or four. Such clusters in the αs- and β-caseins are highly conserved (Martin et al., 2003) and their numbers attest to the significance of the calcium phosphate requirement for post-natal growth in mammals; even more so when it is noted by reference to their sequences (Swaisgood, 2003) that the α-caseins, e.g. the αs1- and αs2-caseins of bovine milk, themselves possess two or more such clusters. From now on, discussion will be confined largely to the behavior of bovine caseins because it is for these that the largest body of research data is available. However, the extensions to the behavior in the micelles of other milks will be obvious.

These clusters of phosphoserine residues and the necessary glutamic acid residues templating their existence give rise to massive downward spikes in the hydrophobicity profiles of the bovine αs1- and β-caseins (Figure 5.1). Associated with these are significantly high densities of negative charge at normal milk pH. There is a charge density of $-9e$ within the span of residues 65–72 of αs1-casein and a further $-6e$ along the sequence 48–53 of the same protein. A similarly high charge density of $-9e$ is found between residues 16–23 of β-casein, encompassing the phosphoserine cluster there. Similar high densities are found around the phosphoserine clusters of αs2-casein. All of these estimates of charge density assume a contribution of $1.5e$ from each phosphoserine residue at or near the natural pH of milk, 6.7.

Away from the phosphoserine clusters, the casein molecules are distinctly hydrophilic. This segregation of hydrophilic and hydrophobic residues confers on the caseins a definite amphipathic nature, which contributes to their ability to function successfully as stabilizers in oil-in-water emulsions. The topography of β-casein adsorbed at the oil–water interface was probed by testing the accessibility of the reactive sites to the proteolytic enzyme trypsin (Leaver and Dalgleish, 1990). In aqueous solution, the reactive sites of β-casein were attacked randomly at no preferential rate.
Figure 5.1 Hydrophobicity plots of (a) $\alpha_{s1}$-casein, (b) $\alpha_{s2}$-casein and (c) $\beta$-casein calculated as a moving average (window $n = 3$) of amino acid hydrophobicities taken from the consensus scale used by Horne (1988). Asterisks denote centers of electrostatic repulsion arising from phosphoserine cluster motifs, the size indicating the number of negative charges associated with each, as listed in the text.

With $\beta$-casein-stabilized emulsions, however, the peptides released showed the lysines at positions 25 and 28 of the sequence to be readily accessible to trypsin, whereas all other possible attack sites were less so (Leaver and Dalgleish, 1990). These residues lie in the center of the highly charged, hydrophilic N-terminal region containing the
four-phosphoserine cluster in β-casein. Measured by dynamic light scattering, a decrease of approximately 13 nm in hydrodynamic radius of the emulsion droplets also accompanied the scission of these peptides, indicating the extent to which they stretched out into the aqueous phase from the emulsion droplet surface (Dalgleish and Leaver, 1991).

The remaining hydrophobic portion of the molecule was speculated to lie along the droplet surface, shielded from trypsin attack. Similar changes in hydrodynamic radius were observed when β-casein was adsorbed from aqueous buffers onto the surface of polystyrene latex particles, indicating a similar adsorption pattern (Dalgleish, 1990; Brooksbank et al., 1993), a pattern that was replicated at the air–water interface, as observed by neutron reflectivity (Dickinson et al., 1993).

The combined experimental evidence was therefore consistent with the view that much of the hydrophobic end of the adsorbed β-casein was directly associated with the hydrophobic interface, with the hydrophilic N-terminal tail extending significantly out into the aqueous phase. Self-consistent-field calculations of the conformation of β-casein adsorbed at a planar hydrophobic interface confirmed this picture of a tail–train structure, and also predicted a train–loop–train structure for adsorbed αs1-casein with anchor points at both ends of the molecule (Leermakers et al., 1996; Dickinson et al., 1997a, 1997b). Schematic representations of these structures were drawn by Horne (1998) and were used in depicting assembly of the casein micelle via the dual-binding model. Perhaps rather than depicting the hydrophobic regions of the molecules as rectangular bars, it would have been more realistic to depict these regions as puckered, as in Figure 5.2, because not all amino acids therein are equally hydrophobic, as the profile plots of Figure 5.1 demonstrate.

Such representations were also used by Horne (1998) to picture the aggregates produced by self-association of β-casein or αs1-casein (Figure 5.3). Thus β-casein was envisaged as a hedgehog-like micelle subject to a monomer/micelle equilibrium,
Figure 5.3  Diagramatic representations of the polymeric structures generated when the hydrophobic chains of the caseins interact: the worm-like chain of $\alpha_s$-casein; the micelle of $\beta$-casein, where only two molecules have been included to simplify the diagram.

a mechanism proposed by Payens and co-workers (Payens and van Markwijk, 1963; Payens et al., 1969). In this picture, the hydrophobic trains are buried inside and the charged hydrophilic tails extend from the surface into solution.

More recently the self-association of $\beta$-casein has been revisited by De Kruif and collaborators (De Kruif and Grinberg, 2002; Mikheeva et al., 2003; O’Connell et al., 2003), who applied high sensitivity differential scanning calorimetry and static and dynamic light scattering techniques to the problem. These experiments again concluded that a micelle-like structure was adopted but, rather than being formed by the highly co-operative monomer/micelle equilibrium suggested previously, instead the micellization took place as a series of consecutive additions of monomer to a growing micelle, as suggested in the shell model of Kegeles (1979). In a like fashion, with its hydrophobic chains at opposite ends of the molecule and a central section containing the highly charged phosphoserine clusters, $\alpha_s$-casein self-associates to produce a worm-like chain polymer (Payens and Schmidt, 1966; Schmidt, 1970a, 1970b).

Though driven by hydrophobic interactions, electrostatic repulsive interactions are also very important in the self-association of these caseins. In particular, note how the equilibrium structures adopted by their polymers place the centers of charge as far apart as possible while still permitting the self-association to take place. Compared with hydrophobic interaction, electrostatic repulsion is a long-range force, an important factor now being recognized in studies of protein–protein interactions (Kegel and Van der Schoot, 2004; Piazza, 2004; Stradner et al., 2004), and certainly manifesting itself in these reactions of the caseins. These electrostatic interactions define the degree of polymerization and limit further growth. Thus, increasing the pH, which increases the protein charge, decreases the polymer size in both $\alpha_s$-casein and $\beta$-casein solutions, whereas increasing the ionic strength, which decreases the range of the electrostatic repulsion component, allows the formation of larger polymers for both casein species (Payens et al., 1969; Schmidt, 1970a, 1970b).
The importance of charge in controlling the extent of aggregation of the caseins cannot be stressed too highly. Precipitation of the caseins can be achieved by lowering the pH and titrating away sufficient of the charge of the phosphoseryl and carboxyl groups to reach the isoelectric points of the proteins. $\alpha_{s2}$-Casein, $\alpha_{s1}$-casein and $\beta$-casein are termed the calcium-sensitive caseins because they can be precipitated in the presence of ionic calcium, the order of sensitivity being as given (Swaisgood, 2003).

The most extensive data set is available for $\alpha_{s1}$-casein. Here the aggregation shows a lag phase with little change in molecular weight with time until a critical time, beyond which rapid aggregation occurs. Horne and Dalgleish (1980) demonstrated that the logarithm of this critical coagulation time was a linear function of $Q^2$, where $Q$ is the net negative charge of the protein. Thus $Q$ is the algebraic sum of the negative and positive charges of the protein, reduced by twice the number of calcium ions bound to the protein, each calcium carrying two positive charges. Furthermore, this relationship held when changes in the net negative charge were produced by chemical modification, whether by conversion of positively charged lysine residues to neutral or negatively charged derivatives, or even by introduction of new negatively charged sites by iodination of tyrosine residues to the di-iodo form (Horne, 1983; Horne and Moir, 1984). Each of these modifications effectively increases the net negative charge of the protein, thereby reducing its propensity for calcium-induced precipitation and slowing down the rate of aggregation.

However, once the protein charge is corrected for the measured extent of modification, the logarithm of the rate of precipitation has been shown to remain linear in $Q^2$, all points lying on the same line as those obtained with the unmodified protein, all other reaction conditions being the same (Horne, 1983; Horne and Moir, 1984). The net charge of the protein therefore dominates its precipitation behavior. Farrell et al. (2006) have suggested that positively charged residues in the N-terminal hydrophobic chain of $\alpha_{s1}$-casein could participate in binding to the phosphate groups of phosphoseryl residues, but such $+/-$ bridging does nothing to reduce the net negative charge of the protein having already been accounted for in the algebraic summation leading to $Q$, which, as we have demonstrated, controls the level of aggregation in these proteins.

It is only when the local balance of electrostatic repulsion and hydrophobic attractive interaction is in favor of attraction that hydrophobic bonds are formed. Sequentially, the major centers of electrostatic repulsion—the phosphoserine clusters—are remote from the hydrophobic regions in the trains of Figure 5.2, though, depending on the adopted conformation, they may not be remote spatially. Figure 5.2 is a depiction of a possible conformation of each protein at a hydrophobic interface, the puckering displaying the tendency to form a multitude of weak, short-range bonds by the hydrophobic chain. Lowering the temperature weakens hydrophobic bonds; hence the tendency to find monomeric $\beta$-casein at low temperatures but a micellar aggregate at room temperature and above.

The aggregation of $\beta$-casein induced by calcium also shows a marked temperature dependence with no precipitation observed at 4°C (Parker and Dalgleish, 1981). At higher temperature, these hydrophobic bonds are in general stronger but, because they are relatively weak overall, statistically individual bonds are readily ruptured by the increased thermal energy available, leading to more mobile labile interactions between molecules.
Casein micelle properties

Almost all of the casein proteins present in bovine milk are incorporated into the casein micelles, together with a high proportion of the available calcium and inorganic phosphate. The calcium and phosphate within the micelle form low molecular mass species collectively known variously as colloidal calcium phosphate, micellar calcium phosphate and latterly calcium phosphate nanoclusters. The micelles are very open, highly hydrated structures, with typical hydration values of 2–3 g H$_2$O/g protein, depending on the method of measurement.

Electron microscopy shows that casein micelles are generally spherical in shape, with diameters ranging from 50 to 500 nm (average ≈ 150 nm) and a molecular weight ranging from $10^6$ to $>10^9$ Da (average ≈ $10^8$ Da) (Fox, 2003). For a casein content of 2.5 g/100 mL milk, there are some $10^{14}$–$10^{16}$ micelles/mL milk, which implies a relatively close packing with inter-surface separations less than one micelle diameter.

Milk is white largely because the colloidal dimensions of the casein micelles are such that they scatter significant amounts of light, an effect compounded by their high number density. Scattering of shorter wavelength radiation (neutrons and X-rays) reveals the internal structure to be heterogeneous, with a correlation length of variations in scattering length density within the particle of approximately 18 nm. This scattering behavior has been interpreted by Stothart and Cebula (1982) as due to a structure composed of closely packed spherical sub-units of this diameter, a picture that mirrors the raspberry-like appearance in early electron micrographs of the casein micelle (Schmidt, 1982).

More recent electron microscopy studies (McMahon and McManus, 1998) have suggested that these well-defined structures are likely to be artifacts of the fixation process, although micrographs recently obtained by field emission scanning electron microscopy (SEM) show a complex surface structure of cylindrical or tubular, but not spherical, protrusions between 10 and 20 nm in diameter, extending from the surface of the micelle (Dalgleish et al., 2004). These samples were not metal coated, although they were of necessity subjected to a fixation and dehydration process, which might have introduced some collapse of more loosely bound protein on to a denser skeleton.

In a careful study using cryo-transmission electron microscopy (cryo-TEM) and small angle X-ray scattering (SAXS), Marchin et al. (2007) have extended previous structural studies to elucidate greater detail of micellar fine structure. Their cryo-TEM pictures show small regions of high electron density, approximately 2.5 nm in diameter, uniformly distributed in a homogeneous web of protein, giving the micelles a granular aspect that diminishes when the pH is reduced from 6.7 to 5.2. Paralleling this change in appearance, the SAXS scattering profile loses its characteristic shoulder at high wave vector when the pH is reduced. This clearly demonstrates that the shoulder is directly linked to the presence of micellar calcium phosphate, and its dissociation from the micelle on acidification to pH 5.2.

The results also demonstrate that this micellar calcium phosphate is uniformly distributed through the micelle in small particles, approximately 2.5 nm in diameter and that, after their removal, other forces/interactions must contrive to maintain micellar structural integrity. Their loss on acidification does not result in micellar disruption.
Any structural model of casein micelle assembly would have to lead to a structure that could reproduce both the cryo-TEM pictures and the field emission SEM pictures, allowing for possible changes in appearance brought about by techniques of sample preparation.

Casein micelle structure is not fixed, but dynamic. In various ways, it responds to changes in micellar environment, temperature and pressure. Cooling milk on release from the udder at 37°C to storage at refrigeration temperatures brings about significant solubilization of $\beta$-casein, some $\kappa$-casein and much lower amounts of $\alpha_s1$- and $\alpha_s2$-casein from the micelles (Dalgleish and Law, 1988). Raising the temperature back to 37°C reverses the process. None of this movement of $\beta$-casein does anything to disrupt the internal structure of the micelle, as observed by cryo-TEM and SAXS (Marchin et al., 2007). Almost complete disruption of the micelles, manifested by a loss of their scattering power and removal of the white color of milk, can be achieved by:

- addition of a strong calcium sequestrant such as ethylene diamine tetraacetic acid (EDTA) (Griffin et al., 1988);
- addition of urea (McGann and Fox, 1974);
- dialysis against a phosphate-free buffer (Holt et al., 1986);
- increasing the pH, by exposure to high pressure (Huppertz et al., 2006); or
- addition of ethanol at $\approx70^\circ$C (O’Connell et al., 2001).

Significantly, the colloidal calcium phosphate can also be solubilized by lowering the pH but, as confirmed by Marchin et al. (2007), without substantial disruption of the micelle structure.

Fractionation of the casein micelles according to size can be realized by a stepwise centrifugation protocol. The proportions of $\alpha_s1$- and $\alpha_s2$-casein remain constant with micelle size but $\kappa$-casein content increases inversely with that size (Donnelly et al., 1984; Dalgleish et al., 1989). For a solid sphere, the surface-to-volume ratio is inversely proportional to the radius of the sphere and these results imply that $\kappa$-casein resides on the micellar surface, where its content controls the micellar total surface area and hence the micelle size.

A surface location for the $\kappa$-casein component may also be inferred from the requirement that this protein be readily accessible for rapid and specific hydrolysis by chymosin and similar proteases, a reaction that destabilizes the micelles and leads to clot formation, which is exploited in cheese manufacture. A surface location is also required to enable the $\kappa$-casein to interact with $\beta$-lactoglobulin in milk to form a complex on heating, the formation of which modifies the rennet and acid coagulation properties of the micelles. It is evident that a principal requirement, which must be met by any micelle model, is that it should generate a surface location for $\kappa$-casein.

Models of casein micelle structure

Casein micelle structure and casein micelle models have been extensively reviewed (Schmidt, 1982; Walstra, 1990, 1998; Holt, 1992; Rollema, 1992; Horne, 1992,
Casein micelle structure and stability

1998, 2006; Fox, 2003; Farrell et al., 2006). As mentioned previously, based on the biochemical and physical properties of the micelles and the casein proteins outlined above, three main models have been proposed: the sub-micelle model (Slattery and Evard, 1973; Schmidt, 1982; Walstra, 1998), the nanocluster model of Holt (Holt, 1992; De Kruijff and Holt, 2003) and the dual-binding model (Horne, 1998, 2002).

In the first model, the casein micelles are composed of smaller proteinaceous sub-units—the sub-micelles—linked together via colloidal calcium phosphate. In the second model, the nanoclusters of colloidal calcium phosphate are randomly distributed, cross-linking a three-dimensional web of casein molecules. Both of these models have been severely criticized (Farrell et al., 2006; Horne, 2006), and the dual-binding model arose first as an attempt to overcome their deficiencies. Rather than reiterate those arguments, we present first a summary of the dual-binding model as providing a rational mechanism for micelle assembly and structure, and demonstrate how this model may be exploited to explain various observations of micellar properties and behavior.

The dual-binding model for micelle assembly and structure

In the dual-binding model, micellar assembly and growth take place by a polymerization process involving, as the name suggests, two distinct forms of bonding, namely, cross-linking through hydrophobic regions of the caseins and bridging across calcium phosphate nanoclusters. Central to the model is the concept that bond formation is facilitated, and hence micellar integrity and stability are maintained, by a local excess of hydrophobic attraction over electrostatic repulsion, bearing in mind the quite different ranges of these interaction components. The individual casein molecules behave and interact as they do in their self-association equilibria, as described previously.

Each casein molecule effectively functions as a block copolymer, as detailed in Figure 5.2, with the hydrophobic region(s) offering the opportunity for a multitude of individual, weak, hydrophobic interactions. The hydrophilic regions of the casein molecules contain the phosphoserine cluster (or clusters), with the exception of $\kappa$-casein, which has no such cluster, each offering multiple functionality for cross-linking. Thus, as we have seen, $\alpha_{s1}$-casein can polymerize (self-associate) through the hydrophobic blocks, giving the worm-like chain of Figure 5.3. Further growth is limited by the strong electrostatic repulsion of the hydrophilic regions but, in the casein micelle situation, the negative charges of the phosphoserine clusters are neutralized by intercalating their phosphate groups into a facet of the calcium phosphate nanocluster.

This has two very important implications for the micelle. Firstly, by removal of a major electrostatic repulsion component, it increases the propensity for hydrophobic bonding upstream and downstream of the nanocluster link. It effectively permits and strengthens those bonds. Secondly, it allows for multiple protein binding to each nanocluster, allowing a different network to be built up. $\beta$-Casein, with only two blocks—a hydrophilic region containing its phosphoserine cluster and the hydrophobic C-terminal tail—can form polymer links into the network through both, allowing further chain extension through both. $\alpha_{s2}$-Casein is envisaged in this model as having two of each

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1 The description here largely follows that found in Horne (2002) with minor refinements highlighted.
block, two (possibly three, see below) phosphoserine clusters and two hydrophobic regions. It is only a small fraction of the total bovine casein but, by being able to sustain growth through all its blocks, it is likely to be bound tightly into the network.

κ-Casein is the most important of the caseins in the dual-binding model of micellar assembly and structure. It can link into the growing chains through its hydrophobic N-terminal block but its C-terminal block is hydrophilic and cannot sustain growth by linking hydrophobically to another casein molecule. Neither does κ-casein possess a phosphoserine cluster and therefore it cannot extend the polymer cluster through a nanocluster link. Thus, chain and network growth are terminated wherever κ-casein joins the chain. This leaves the network with an outer layer of κ-casein, satisfying the prime requirement recognized earlier.

The nanocluster bridging pathway through the phosphoserine clusters is the only pathway allowed in the nanocluster micelle model of Holt (Holt, 1992; De Kruif and Holt, 2003), where around 50 casein molecules are considered to link into each calcium phosphate nanocluster. Horne et al. (Horne, 2006; Horne et al., 2007a) have argued on the basis of mineral content and stoichiometry that the functionality of these nanoclusters will be much lower and is more likely to be four to six phosphoserine clusters, these not necessarily originating from different casein molecules.

Some consideration has also to be given to what constitutes a phosphoserine cluster capable of linking into the calcium phosphate nanocluster. Aoki et al. (1992) suggested a minimum of three phosphoserine residues but De Kruif and Holt (2003) argued that two might be sufficient. This would allow the phosphoserine pair at positions 46 and 48 of αs1-casein, or those at positions 129 and 131 of αs2-casein, to function as nanocluster linkage sites, particularly if the carboxyls of the neighboring glutamate residues acted as pseudo-phosphate groups. This would give αs1-casein two linkage sites and αs2-casein three linkage sites.

This level of functionality in these caseins is absolutely essential to the Holt model to build the required three-dimensional network as, without them, an αs2-casein molecule with only two linkage sites and with such a low percentage of the total casein would probably prove to be insufficient. Although they are not essential to the dual-binding model, these mini-clusters of pairs of phosphoserines may provide for a weaker bridging link to the calcium phosphate nanocluster, allowing a range of nanocluster bond strengths to prevail.

**Calcium phosphate equilibria in the dual-binding model**

From the mineral viewpoint, casein micelle assembly is a frustrated crystallization of calcium phosphate. Milk is supersaturated in calcium and phosphate and, were it not for the presence and intervention of the highly phosphorylated caseins, a precipitation of calcium phosphate and potentially painful calcification of the mammary gland duct system would occur. The phosphoseryl clusters on the casein have the potential to act as a template for the crystal formation but can also act as a cap, cutting off growth at a particular facet. In this way, the overall total number of phosphoseryl clusters in a millilitre of milk controls the size and the number of nanoclusters present in the milk (Horne et al., 2007a). This can happen only if all (or close to all) of the phosphoserine cluster motifs are involved in nanocluster stabilization.
In turn, this must mean that the binding of a serine phosphate group into the nanocluster must present a bonding advantage thermodynamically over that of a free phosphate group going into a growing calcium phosphate crystal in an equilibrium situation. It is a more favorable outcome thermodynamically.

But, just as the calcium phosphate crystal in equilibrium in solution with free calcium and phosphate is subject to environmental constraints shifting that equilibrium, so too is the micellar nanocluster species. The two equilibria may exist in parallel though when micelles are present in normal milk conditions, the nanocluster form is the favored option. However, conditions may change where “solution” crystal growth is favored, and we try to explore these speculations in some of the discussions following. The possibility of shifting calcium phosphate equilibria is an aspect of casein micelle structural behavior that has not been considered previously but, as the nanoclusters are involved, may offer alternative avenues to explore in addressing problems of micellar behavior associated with the application of high pressure, or of the addition of ethanol.

**Application of the dual-binding model**

**Predictions of casein micelle properties**

**Size and appearance**

A major failing of the earlier micelle models was their lack of a plausible mechanism for assembly, growth and, more importantly, termination of growth. All such elements are in place in the dual-binding model. Furthermore, the product of the dual-binding model satisfactorily represents the appearance and scattering behavior of the native casein micelle. Network growth is envisaged as a random process and its termination along any particular pathway depends on the serendipitous arrival of a $\kappa$-casein molecule. Micelle size will therefore depend on the proportion of $\kappa$-casein in the mix, but will also present a range of sizes dependent as it is on random events. The model also reproduces the heterogeneity in structure required by the X-ray and neutron scattering data. The dense calcium phosphate nanoclusters will be rather homogeneously distributed through the matrix and will give rise to the structures observed by cryo-TEM and inferred from SAXS (Marchin *et al.*, 2007).

The dual-binding model presents the casein micelle as a dynamic, “living” entity. The hydrophobic interactions are individually weak and capable of breaking and recombining on an almost continuous basis. To some extent, the molecular movements this allows will be restricted by the stronger nanocluster linkages and the low probability of rupturing simultaneously all hydrophobic bonds involving any particular molecule. Molecular movement has several consequences, however. The micelle may have an outer layer of individual $\kappa$-casein molecules when initially constructed but conditions within the Golgi vesicle are suitable for disulfide bond formation; otherwise $\beta$-lactoglobulin and the other whey proteins would not fold properly. Movements of that outer “hairy” layer may bring those $\kappa$-caseins into proximity and allow their polymerization through disulfide bridging, the size of the polymer depending on when the chain closes into a loop, but giving rise to the polymeric $\kappa$-casein entities observed on micellar dissociation.
Another consequence of the “living” nature of these hydrophobic bonds is that the dehydration of the micelle required in the preparation of a sample for electron microscopy would also tend to be accompanied by the collapse of the more mobile, weaker and less multitudinously bonded regions on to those more strongly cross-linked; hence, perhaps giving rise to the raspberry-like (Schmidt, 1982) or tubular (Dalgleish et al., 2004) structures seen in some electron micrographs. Even the putative caps suggested for those tubules by Dalgleish et al. (2004) can be provided by the dual-binding model, as the disulfide bridging between the \( \kappa \)-casein molecules would enhance the Velcro effect of the weak hydrophobic bonding of an individual molecule to many such molecules in the chain.

**Effects of urea, pH, sequestrants, temperature**

The concept of a local excess of hydrophobic attraction over electrostatic repulsion, as well as permitting the visualization of micellar growth, successfully accommodates the response of the micelle to changes in pH, temperature, urea addition or removal of calcium phosphate by sequestrants, all in accordance with experimental observations.

Urea disrupts hydrophobic bonds and high concentrations will bring about micellar disintegration. In some regions of the micelle, this may be only partial because the nanocluster cross-links through the phosphoserines remain, unaffected by this reagent. Micellar fragments in a range of sizes may be produced, even some as large as some of the original micelles, though perhaps more open and swollen from their own starting state before urea treatment. Extensive disruption does occur, however, as is observed by the loss of the white appearance of skim milk (McGann and Fox, 1974). The dual-binding model fully accounts for these observations.

Removal of calcium from the calcium phosphate nanocluster by sequestrant addition, whether EDTA, citrate or oxalate, restores the negative charge of the hydrophilic region, if the pH is maintained at the native milk pH. This shifts the hydrophobic attraction/electrostatic repulsion balance in favor of repulsion and the micelle breaks up. Decreasing the milk pH solubilizes the colloidal calcium phosphate (Dalgleish and Law, 1989), but the negative charges associated with the cross-linking phosphoseryl groups are also titrated away. The strength of the hydrophobic bonds remains unaffected or may be enhanced if other carboxyl charges are also titrated away. The integrity of the micelles is maintained but their scattering behavior and their appearance in cryo-TEM micrographs reflect the loss of the nanoclusters (Marchin et al., 2007).

Increasing the pH may be expected to be the reverse of the dissolution process and to favor the formation of calcium phosphate species. Fox (2003) noted that raising the pH to \( >9.0 \) does not dissolve colloidal calcium phosphate but rather increases its level. However, increasing the milk pH to these levels does lead to dissociation of the micelles and creation of a translucent solution. Whether this is due to conversion of the phosphoserine residues from singly to doubly charged units, which are no longer capable of binding to the calcium phosphate nanoclusters, and hence allowing greater amounts of colloidal calcium phosphate, or whether the increase in charge is sufficient to upset the balance of electrostatic repulsion and hydrophobic attraction in upstream and downstream hydrophobic bonds, it is not possible to say—both routes leading to micellar disruption.
Decreasing the temperature is known to decrease the strength of hydrophobic attraction and to shift the monomer/micelle equilibrium in β-casein solutions towards the monomer side at temperatures below 15°C (De Kruif and Grinberg, 2002). Lowering the temperature of milk to refrigeration levels also brings about dissociation of a large fraction of the β-casein from the casein micelle (Dalgleish and Law, 1988), possibly some of which is not bound into the micellar matrix through its phosphoserine cluster. Raising the temperature back to its initial value reverses the process and the β-casein is reincorporated into the micelle. It is to be anticipated that, even at room temperature, the ongoing equilibrium would allow exchange of micellar casein, albeit with low levels of β-casein in the serum phase.

There are also shifts in the calcium phosphate equilibria in milk associated with temperature change. Ultrafiltration permeate is a clear, straw-yellow liquid when prepared at 4°C but becomes turbid when heated to room temperature and above because of the precipitation of calcium phosphate. Even permeate collected at room temperature clouds on heating but reverts to clarity on cooling. Hilgemann and Jenness (1951) noted that calcium phosphate also precipitates in milk. However, the calcium phosphate precipitate was only slowly resolubilized (Jenness and Patton, 1959). Weakening the calcium phosphate "solution" equilibrium would favor preservation of the nanoclusters but anything that pushes that "solution" equilibrium to the solid side could have an effect on the continuing existence of the nanoclusters. There are indications that heating milks in the temperature range 50–90°C brings about increasing mobility in the micelle (Rollemo and Branches, 1989), which would be in line with partial disruption. The behavior of casein micelles in this temperature range merits further scrutiny, particularly as so many processes in the dairy industry are conducted just in this range.

The dual-binding model and micellar interactions

The ideas outlined earlier in this chapter allow us to schematically describe in Figure 5.4 how the casein micelle might appear as an interacting species at the various pH values indicated.

Internally, at pH 6.7 (Figure 5.4a), the micellar matrix is closely interlinked through a combination of nanocluster bridging bonds (the small black circles) and hydrophobic interactions, occurring randomly along any selected polymer chain. The hydrophobic interactions at this pH (indicated as crossover points in the tangled protein network in the diagrams in Figure 5.4) are many but relatively weak, being counterbalanced by the negative charges present on ionized carboxyl groups, dispersed along the chains and throughout the network. The micellar outer reaches are mainly κ-casein molecules, which have terminated polymer extension and limited micellar growth in the dual-binding model. The negative charges from the ionized carboxyls and sialic acid groups on the κ-casein macropeptides provide the electrostatic repulsion component in the inter-micellar interaction potential, which inhibits micellar aggregation. Its longer range, illustrated by the thickness of the shell around the micelle, prevents close approach of the hydrophobic regions buried beneath
the shell, and amply fulfills the requirements of a hard sphere model colloid at this pH, 6.7.

At the lower intermediate pH of 5.6 (Figure 5.4b), the same shell continues to prevent close approach of the micelles. The pK values of the acidic groups giving rise to the negative charge are generally lower than 5.5 and have yet to be titrated away. Internally though, most of the micellar calcium phosphate nanoclusters have been solubilized and the bridges between phosphoserine cluster motifs have been lost, weakening the overall network structure of the micelle. The bond strengths of hydrophobic interactions remain relatively weak, still being counterbalanced by ionized carboxyl groups dispersed through the micelle.

By pH 5.1 (Figure 5.4c), the surface charges are being titrated away, the shell in Figures 5.4a and b is much thinner and aggregation begins. Internally, the hydrophobic interactions are effectively being strengthened because the counterbalancing electrostatic repulsions are also being removed from the equation, leading to reduced mobility within the micellar particles.

**Figure 5.4** Representations of casein micelle structures at various pH values as indicated. The pale chains indicate protein molecules, where they cross being a hydrophobic interaction junction. The small black circles are the calcium phosphate nanoclusters that are solubilized when the pH is lowered. The outer circle is indicative of the range of steric repulsion generated between micelles and preventing interaction of the surface protein chains (see also Plate 5.4).
Rheology of micellar dispersions

In milk as produced from the cow at its natural pH of 6.7 and temperatures from ambient to blood heat, casein micelles closely follow the behavior of hard sphere colloids (De Kruif, 1998; Alexander et al., 2002). Justification for this assertion comes from studies utilizing light and neutron scattering to measure micelle size and polydispersity (Hansen et al., 1996), from sedimentation behavior (De Kruif, 1998) and from measurements of micellar voluminosity (De Kruif, 1998), diffusivity (De Kruif, 1992) and viscosity of micellar suspensions (Griffin et al., 1989).

Paralleling colloidal hard sphere behavior holds only for a limited range of concentrations and, above a critical concentration, micellar suspensions show strong deviations from expected hard sphere behavior (Mezzenga et al., 2005). The viscosity continues to increase but at a slower rate than that expected for hard spheres. This is accompanied by a transition from Newtonian viscosity behavior at natural milk concentration to non-Newtonian viscoelastic behavior in the high concentration regime. More enlightening demonstrations of the departure from hard sphere behavior come from studies of the rheology of high concentration micellar suspensions produced by ultrafiltration (Karlsson et al., 2005), by evaporation to 45% total solids (Bienvenue et al., 2003) and by centrifugal sedimentation and pelleting (Horne, 1998). In these instances, concentrated micellar suspensions are close packed and show a gel-like behavior, which can be interpreted with the assistance of the dual-binding model.

Karlsson et al. (2005) concentrated skim milk by ultrafiltration to produce a micellar suspension with 19.5% casein and studied the effects of pH and ionic strength on its viscoelastic properties. Their suspensions exhibited Newtonian viscosities at very low (Brownian) and very high (hydrodynamic) shear rates, with shear thinning at intermediate shear rates and stresses. The concentration of the micelles by ultrafiltration forced the micelles to interact, jamming them together at this high volume fraction and producing a honeycomb-like structure in freeze-fracture electron micrographs. The elastic modulus of these gels decreased as the pH was lowered from the value achieved in the ultrafiltration retentate. Addition of NaCl at levels of 0.33 and 0.66 mol/kg prior to ultrafiltration increased the elasticity of the gels but shifted their pHs to more acid values. Thereafter in the salt-added systems, lowering the pH produced a decrease in elasticity that paralleled the untreated suspension behavior, the higher salt level giving the greater elasticity throughout.

Karlsson et al. (2005) also measured the phase angle—the partitioning between viscous and elastic components in these gels as the pH was reduced. In the no-added-salt system, they found the phase angle to increase through a maximum close to 45° and thereafter decrease with decreasing pH. In the presence of added salt, the maximum in phase angle was again observed but shifted to much lower pH values: in the case of the higher salt level, to a pH value lower than for acid gel formation in milk of normal concentration and, in both cases, where elasticity had been observed to increase again in these salted concentrated suspensions.

The dual-binding model explains this behavior with reference to the schematic of the micellar interaction potential depicted in Figure 5.5. The increase in micellar concentration in the no-added-salt case forces the micelles together and into the secondary minimum generated by hydrophobic interactions. This is the source of the
attractive interaction giving rise to the viscoelasticity observed. The micelles are also in a jammed structure and their internal bonding contributes to the measured elasticity. On lowering the pH, the loss of the calcium phosphate nanocluster bridges weakens this structure and the elasticity decreases, as observed. The bonding due to hydrophobic interactions is relatively weak and the loss of the nanocluster bridges further contributes to the mobility in the gel, as evidenced by the observed increase in phase angle. Dropping the pH further titrates away carboxyl groups; however, it reduces the counterbalancing electrostatic component and thereby strengthens hydrophobic bonds in the matrix, reducing mobility and producing the subsequent drop in phase angle.

The major effect of the addition of salt is to reduce the Debye-Huckel parameter and shorten the range of the electrostatic repulsion between micelles. This makes it easier to enter the secondary minimum in the interaction potential and increases the gel elasticity, as observed, with more salt producing the stronger gel. Again, though, the calcium phosphate nanocluster bridges contribute stress-carrying bonds and their removal by lowering the pH leads to the observed decrease in the elasticity of the gel. Throughout this titration, the bonds in the system are relatively stronger than in the no-salt case—the salt also contributes to decreasing the effectiveness of intramicellar electrostatic repulsion—and the phase angles are lower in comparison.

Karlsson et al. (2005) suggested that a significant effect of the salt addition is to exchange bound calcium within the micelle for monovalent ions, which would imply no nanoclusters in the system to be solubilized on decreasing the pH and thereby voiding the above explanation for the decrease in elasticity with pH. Huppertz and Fox (2006) did indeed find increased levels of calcium in serum when 600 mM NaCl was added to a two-times-concentrated milk but they found no increase in serum inorganic phosphate, suggesting that the increase in calcium came from displacement.

Figure 5.5  Repulsive inter-micellar interaction potential with inner hydrophobic interaction minimum. The dashed line shows the effect of salt addition on the range of the electrostatic repulsion component.
of casein-bound calcium rather than a salt-induced dissociation of the calcium phosphate nanoclusters, leaving these to be solubilized on acidification.

The evaporated milks produced by Bienvenue et al. (2003) had 45% total solids or were approximately concentrated from normal by a factor of four, rather than the eight times concentration of the micelles in the ultrafiltration retentates of Karlsson et al. (2005). The milks of Bienvenue et al. (2003) increased in viscosity on storage at 50°C, with salt addition accelerating the increase. Such behavior is in line with the predictions of the dual-binding model outlined above. The collision rate increased by concentration will be further increased by raising the temperature, bringing about a higher frequency of micelles attempting to enter the secondary minimum. A higher success rate and flocculation due to more thermal energy will give rise to the observed increase in viscosity. The weak flocs can be disrupted by higher shear stresses, giving the observed shear-thinning behavior. The effect of salt, as above, would be to render it easier to enter the secondary minimum and promote the flocculation reaction.

Finally, the casein micelle pellets produced by the centrifugation of skim milk at 19,000 g for 60 min had protein concentrations of approximately 20% (Horne, 1998). At high temperatures (40°C), this pellet flowed freely. Its viscosity was Newtonian, independent of shear rate or frequency. At low temperature (5°C), however, this micellar suspension exhibited all the properties of a classical viscoelastic gel, with elastic moduli independent of frequency and phase angles less than 45°. At intermediate temperatures, there was a crossover between viscous and elastic behavior. The behavior here is dominated by that of the hydrophobic interactions.

At low temperatures, the strength of these interactions is low. Both β-casein and κ-casein are known to depart from the micelle under such conditions (Dalgleish and Law, 1988) but, in the close-packed conditions prevailing in the pellet, they are liable to migrate or link to neighboring micelles or to become entangled with proteins loosened from those micelles, leading to the gel-like behavior. As the temperature is increased, the strength of the hydrophobic interaction increases but the ability to break bonds is also enhanced and more mobility is allowed. The strengthening of the bonding may also lead to a tightening up of the micelles and their becoming more compact may allow the suspension to flow more freely.

The dual-binding model and micellar destabilization

The concept of the casein micelle electrosterically stabilized by a “hairy layer” coat of κ-casein appears to enjoy universal acceptance (Holt, 1975; Walstra, 1979; Holt and Horne, 1996).

Because the dual-binding model of the casein micelle naturally provides a surface location for κ-casein in a growth-limiting role, it readily explains the destabilization of the casein micelle system on the proteolysis of κ-casein by chymosin and the loss of the steric-stabilizing hairs. Such proteolysis also leads to a significant drop in the micellar zeta potential (Dalgleish, 1984), and consequent reduction in the electrostatic repulsion between micelles. Further confirmation of the importance of electrostatic repulsion in inter-micellar interactions is evinced by the necessary presence of ionic calcium to bring about/promote the aggregation of the chymosin-treated micelles.
Notwithstanding the importance of electrostatics, hydrophobic interactions also play an important part, as evidenced by the fact that fully renneted micelles show no signs of aggregation at low temperatures (\(<10^\circ C\)) (Dalgleish, 1983) or that rennet gels increase in elasticity as the incubation temperature is raised (Horne, 1998).

Similarly, the action of ethanol in collapsing the hairs and inducing micellar aggregation is a major coup for the “hairy micelle” model, translated into the adhesive sphere picture of De Kruijff and Holt (2003). As the $\kappa$-casein hairs are also negatively charged, their neutralization on acidifying milk would also remove a component of the stabilizing barrier and induce aggregation (De Kruijff and Holt, 2003). Attractive as these scenarios are for explaining these three routes to micellar destabilization, in none of them does the adhesive sphere/hairy micelle approach tell the whole story. To achieve this, the influence of reaction conditions on micellar integrity has to be considered and it is here that the full power of the dual-binding model comes into play.

**Dual-binding model and rennet curd formation**

The aggregation and gelation of casein micelles induced by the chymosin proteolysis of $\kappa$-casein is the reaction that comes closest to the colloidal aggregation model, especially in strictly controlled laboratory studies, many of which maintained the pH of the milk at its natural value of 6.7. It is under such conditions that the casein micelle exhibits most closely the properties of a colloidal hard sphere and, importantly for the aggregation observed, where the internal integrity of the micelle is undisturbed. It seems that the internal binding through calcium phosphate nanocluster bridges limits extensive rearrangements and constrains excursions of hydrophobic regions so that only those close to the surface behind the barrier of the charged macropeptide can take part in micellar aggregation once that barrier is removed. This explains the success of the reaction schemes for the initial stages of aggregate growth based on the particle model (reviewed by Hyslop, 2003, and previously by Dalgleish, 1992). It also explains the success of the gel strength model described by Horne (1995, 1996), with the particles remaining largely unchanged through the gel formation process.

However, cheeses are seldom manufactured at the natural pH of milk, because this is not the optimum pH for enzyme action (Dalgleish, 1992). Generally, some acidification of the milk is applied and this modifies the internal integrity of the micelles with consequent effects on the rennet coagulation and curd properties. Some of these properties, and their influence on the cheeses produced from such curds, have been studied by Choi et al. (2007, 2008), by varying the milk pH or by adding EDTA at a fixed milk pH of 6.0, the objective being to examine in detail the impact of removing micellar calcium phosphate.

In all samples, the elastic modulus of the rennet gel passed through a maximum as the gel was formed, declining during longer reaction times. Rennet gels produced at pH 6.4 had the highest maximum, probably due to a lower electrostatic repulsion because only low levels of micellar calcium phosphate were solubilized at this pH. The maximum elasticity in the gelation profile thereafter decreased with the decreasing pH of the preparation from 6.4 to 5.4. This is explained in the dual-binding model by the decline in the number of nanocluster bridges within the micelle that contribute to the overall strength of the gel matrix. There was also a decrease in the maximum gel elasticity.
with an increase in the added concentration of EDTA, which removed nanocluster links by sequestration of calcium. In both cases, pH adjustment and EDTA addition, the decrease in the maximum elasticity in the gel curds was accompanied by an increase in the rheological loss tangent. The removal of the nanocluster bridges was permitting greater mobility in these gels; a weaker, more flexible network was being produced.

The microstructure of these rennet-induced gels was also examined, near the point of their maximum elasticity and again some 2–10h later using fluorescence microscopy (Choi et al., 2007). When elasticity was at its maximum, the gels obtained at pH 6.4 manifested more branched, interconnected networks than those obtained at pH 5.4 where the strands/clusters were larger with more obvious open regions between. In all cases, there was a decrease in apparent interconnectivity between strands in the gel microstructure during aging, which agreed with the decrease in elasticity beyond the maximum.

It is apparent that gel strength is a function not only of the number and strength of potential bonds in a system but also of their spatial distribution. Here, the loss of the nanocluster bridges weakens the network but also introduces into it greater mobility. Rearrangements occur at a rate governed by that mobility and apparently towards a more compact clustering of the casein proteins, which weakens the matrix structure. Predicting the relationship between matrix morphology, bond strength and bond number is one of the challenges yet to be addressed in food materials science.

These changes in rennet gel matrix structure have a direct impact on the functional properties of the cheeses made from the gels. Choi et al. (2008) demonstrated that removal of micellar calcium phosphate contributed to a greater softening and ease of flow of these cheeses at higher temperature. However, they found that there was an optimum pH of preparation, below which the increasing influence of attractive hydrophobic interactions in the balance of forces reduced bond lability and inhibited curd stretching.

Dual-binding model and ethanol stability
Although studies of the response of dilute suspensions of casein micelles to the addition of ethanol constituted one of the greater successes of the hairy micelle model, they also provided pointers to the failings of the adhesive sphere concept developed from that model (Horne, 2003a).

Dynamic light scattering studies (Horne, 1984, 1986; Horne and Davidson, 1986) demonstrated the collapse of the hairy layer and the consequent loss of the steric-stabilizing component with sub-critical concentrations of ethanol, but attempts to measure layer thickness as a function of buffer pH or ionic calcium concentration were confounded by the observation that initial micelle size was a function of these parameters. Raising the pH or decreasing the ionic strength produced an increase in the hydrodynamic size of the micelle, presumably due to a loosening up of the micelle as either treatment increased the effective electrostatic repulsion between the hairs.

More importantly for the ethanol-induced collapse of the hairs, this loosening of the structure extended deeper into the micelle and greater apparent layer thickness was shown as the micelle structure was caused to expand. This behavior is accommodated within the dual-binding model as a manifestation of the control of internal...
binding by the balance of hydrophobic attraction and electrostatic repulsion, and also by the presence of the calcium phosphate nanocluster bridges, which can be lost on acidification. The model allows for changes in the rigidity of the micelle structure, particularly in the surface layers, which would then control the apparent size of the micelle when the steric-stabilizing hairs are collapsed by the non-solvent—ethanol.

The experiments described above confirm yet again the contribution of steric stabilization to micelle stability but relate to the behavior of the casein micelle in a highly dilute suspension in an artificial environment, devoid of inorganic phosphate. In milk, the results of the alcohol stability test, and particularly the behavior of the alcohol stability/pH profile as a function of mineral content, demand consideration of a different mechanism of destabilization (Horne, 1987), but a mechanism wholly consistent with the dual-binding model, although initially proposed well in advance of that model.

The mechanism, proposed by Horne (1987), suggests that there are two competing effects of ethanol on the micellar system: destabilization through loss of the hairy layer, and shifts in the calcium phosphate equilibria, first noted by Pierre (1985). The behavior of calcium phosphate and its colloidal form has been at the center of much discussion in this chapter. If ethanol promotes the precipitation of calcium phosphate external to the micelle, it would first of all reduce the concentration of free calcium, reduce the level of caseinate-bound calcium and disrupt the binding through calcium phosphate nanoclusters. Moderate losses would increase the negative charge of the caseins and increase the thickness of the steric-stabilizing layer. The higher the alcohol level, the faster and more extensive would be the precipitation of calcium phosphate. The ensuing adjustment in protein charge and conformation, although relatively rapid, still requires a finite response time.

Countering these changes are the effects of ethanol as a non-solvent for the proteins, promoting cross-linking and collapse of the hairy layer. When the coagulation reaction occurs faster than the adjustment of charge and conformation resulting from shifts in the calcium phosphate equilibria, or the extent of the latter is limited by insufficient ethanol, the aggregation reaction dominates and precipitation of micelles follows.

The origin of the sigmoidal ethanol stability/pH profile can also be explained through the effect of pH on calcium phosphate precipitation. Increasing the pH brings about increased calcium phosphate precipitation, possibly further enhanced by the ethanol, which means that more ethanol is required to precipitate the protein, i.e. to overcome the increased energy barrier being erected following the transfer of calcium phosphate from the nanocluster state. Conversely, decreasing the pH acts to diminish the influence of ethanol-induced precipitation of calcium phosphate by titrating away negative charge and reducing electrostatic repulsion between protein species. Other effects of milk serum composition, of forewarming the milk and of modifying the milk concentration and ionic strength, can all be explained in a similar fashion (Horne, 2002).

Finally, what is effectively a competition between mineral precipitation and protein aggregation explains the anomalous destabilization of milk by trifluoroethanol (TFE) (Horne and Davidson, 1987) and the behavior of ethanol in milks at high temperature (~70°C) (O’Connell et al., 2001). With TFE, it was found that, after passing through a critical range of TFE concentrations, which caused protein precipitation, higher levels gave rise to micellar dissociation and produced translucent suspensions.
When ethanol/milk mixtures were heated by O’Connell et al. (2001), these too became translucent as the micelles dissociated. In both instances in the mechanism proposed here, this behavior would be seen as the result of the calcium phosphate precipitation proceeding so fast or to such an extent that the micelle would disintegrate before the micellar aggregation reaction could occur—any protein aggregates remaining small and giving rise to insignificant turbidity.

Dual-binding model and acid gel formation

It is in trying to describe milk acid gel formation in terms of molecular events that the dual-binding model is most useful. Superficially, the initial stages of acid-induced aggregation of casein micelles can be accommodated by the adhesive sphere model, where titration of micellar charge collapses the “hairy layer” and allows aggregation to proceed (De Kruif and Holt, 2003), but closer study, particularly of the kinetics of gel formation using glucono-d-lactone (gdl) as acidulant, reveals anomalies (Horne, 2003b).

Firstly, at any given temperature, increasing the quantity of gdl used, and hence the rate of acidification of the milk, leads to stiffer gels. There is no mechanism for this in the adhesive sphere model, which has pH as the only variable but, in the dual-binding model, we postulate that titration of the negative charges of the caseins will also affect internal bonds in the micelle, and that reduction of electrostatic repulsion will deepen the attraction between the molecules. If the acidification is proceeding slowly, then this may allow equilibration and rearrangement into localized denser structures with few linkages between, giving rise to weaker gels. More rapid drops in pH may lock the protein into a more dispersed structure with greater density of possibly stronger strands, as was observed microscopically when similar trends in elasticity were detected in rennet gels (Choi et al., 2007a).

Secondly, if the skim milk is heat treated at 90°C for 10 min, a process practiced by the dairy industry and known as forewarming, prior to acid-induced gelation by gdl at 40°C, not only is the critical coagulation pH shifted to a higher pH value and a much stiffer gel produced, but also a distinct step is introduced on to the gelation profile. This is not an artifact introduced by slippage in the rheometer. It is fully reproducible and its presence is subject to the reaction conditions applied. It can be made to disappear with forewarmed milk, for example, by lowering the incubation temperature to 30°C and below (D S Horne, unpublished observations).

A similar step can be introduced into the profile of a non-heat-treated milk by raising the incubation temperature to 40°C or above and employing a high concentration of gdl. Such steps in the profiles were also seen following the acidification of milks in the presence of xanthan (Aichinger, 2005), where their magnitude and definition were dependent on the level of xanthan introduced. With heat-treated milks, the presence, the magnitude and the definition of the stepped gelation profile change in response to the temperature and duration of the preheating (Figure 5.6).

The critical pH for gelation and the maximum in complex modulus are monotonic functions of the level of denaturation of the whey proteins brought about by the prior heat treatment (Figure 5.7). Stepped gelation profiles like this are also seen during the formation of fermented milk gels, again with forewarmed milks, where
the step can be removed by suitable modest additions of trisodium citrate (TSC) (Oczan-Yilsay et al., 2007). The critical pH and the loss tangent remained unaffected by additions of TSC up to 10 mM although the gel elasticity recorded at pH 4.6 was enhanced and the gelation profile lost its step and assumed a monotonic increase in $G'$ with decreasing pH.

All of the above observations can be reconciled within the following picture, which relies heavily on the precepts of the dual-binding model. Aggregation, and thereafter
gelation, of the casein micelles on acidification begins only when the electrostatic repulsion component is reduced below a critical level, i.e. when a critical pH is reached. The rate of aggregation is a function of a number of factors: the nature of the surface, the energy (or temperature) involved and the collision rate. Modifying the surface by partially or wholly coating it with denatured whey proteins through formation of a β-lactoglobulin/κ-casein complex, raising the incubation temperature and increasing the micellar concentration by imposing a phase separation through addition of incompatible polysaccharide will all increase the rate of aggregation and shift the critical pH. These are all operational in the above examples, and in all cases we are forcing the micelles to interact, probing further into the inner reaches of the interaction potential with greater frequency and accessing the hydrophobic minimum there.

So far, this is no more than another adhesive hard sphere description but, during acidification, the internal integrity of the casein micelles is also being compromised through the solubilization of the micellar calcium phosphate nanoclusters. Pushing the critical gelation pH to higher values means that more nanoclusters are still present in the aggregating micelles, which are in consequence more rigid than at lower pH. Because the hydrophobic interactions are counterbalanced by any remaining electrostatic repulsion, these hydrophobic bonds are weaker the higher is the coagulation pH. Rearrangements and bond breakage are relatively rapid, as evidenced by the higher value of the phase angle at this point. As the pH continues to drop, the loss of the remaining nanoclusters allows more mobility in the gel, giving the increasing phase angle, but also permitting stronger strands to be formed in the rearrangements, and hence the greater elasticity in the gel.

However, the elasticity is not increasing so rapidly that it cannot be overtaken by its diminution on the loss of the nanocluster bridges; hence the maximum or inflexion point in $G'$ with system pH. Eventually the titration of the carboxyl groups wins out and the increasing contribution from attractive hydrophobic interactions produces increasing elasticity in the gels. Compared with the gel produced in a reaction regime with a low critical gelation pH, this “hi-pH” gel is anticipated to have a more uniform distribution of contributing strands, as has been observed in yoghurt studies (Oczan-Tilsay et al., 2007). When the critical pH is low, as for an unheated milk for example, the reactions described above relating to the behavior on the loss of the remaining calcium phosphate nanoclusters still occur but are now confined within the micellar particle, which remains unaggregated because the inter-micellar potential is still repulsive.

Mobility is introduced and bonds are strengthened but only within the micelle so that, when eventually aggregation and gelation do take place, fewer inter-cluster bonds are formed and a more open gel structure is obtained. Location and confinement are major factors in defining the final architecture and strength of the gels formed, and the dual-binding model helps us to get to grips with understanding what is going on.

Conclusions

This chapter has reviewed the physico-chemical properties and interactions of the caseins and blended them on to the construction of a mechanistic framework for a
working model of the casein micelle—the dual-binding model. The various properties of the casein micelle have been rationalized in terms predicted by the model. Particular attention has been paid to those treatments leading to destabilization and gel formation, where the necessity for the dual-binding model to explain fully the behavior in the developing gels has been forcefully demonstrated.

Constraints of space have limited the number of cases we have been able to consider. A major omission has been explaining the dissociation and reassociation behavior of the casein micelle under high pressure, but this has recently been covered by Huppertz et al. (2006) in terms that can be aligned with little difficulty with the dual-binding model. Neither have we extensively mentioned the application of this model to explaining the behavior of non-bovine milks, although a recent study by Horne et al. (2007b) has demonstrated its usefulness in this context for marsupial milks. Our objective has not been to be exhaustive but to provide examples of how the model can be made to work in various situations. Every reader could come up with other examples where the model might prove to be useful. Testing it to failure can only result in superseding it with a better model.

References


Structure and stability of whey proteins

Patrick B. Edwards, Lawrence K. Creamer and Geoffrey B. Jameson

Abstract

The chemical and physical stability of the more common proteins of bovine and, where available, ovine, caprine and equine whey (β-lactoglobulin, α-lactalbumin, serum albumin, immunoglobulins and lactoferrin) is reviewed with regard to their molecular structures and dynamics. The behavior of the proteins separately and in combination, to temperature, pressure, pH, denaturants (such as guanidinium chloride and urea) and stabilizers (such as fatty acids, and metal ions) has been considered. Particular emphasis has been placed on studies that have utilized X-ray, NMR, fluorescence and circular dichroism techniques. Attention is directed to the role of cysteines and disulfide bridges with regard to chemical stability. Whereas there is considerable knowledge of structure–function relationships of individual proteins, there is a dearth of three-dimensional structural knowledge of combinations of proteins, despite clear importance of such knowledge to functionality, especially with regard to food processes.

Introduction

Information regarding whey protein structure and stability has great potential to facilitate knowledge-based product design. Recent reviews highlight the importance of knowledge of structure and stability, including the effects of pressure, temperature and...
chemical denaturants (Chatterton et al., 2006; Lopez-Fandino, 2006a). In this chapter, we discuss the structures of the whey proteins shown in Table 6.1 under quiescent and destabilizing conditions (change in pH, temperature and pressure and addition of chaotropes) and in the presence or absence of small-molecule ligands. Particular emphasis is placed on information that has been obtained via high-resolution X-ray crystallographic and high-field nuclear magnetic resonance (NMR) studies. The results of these studies have been applied to many projects reported in the present volume (e.g. Chapters 7, 8, 9, 10 and 13).

Table 6.1 Typical protein composition of whey (based on Farrell et al., 2004)

<table>
<thead>
<tr>
<th>Protein</th>
<th>Proportion by mass (%)</th>
<th>No. amino acids</th>
<th>Molecular mass (Da)</th>
<th>Iso-ionic point</th>
<th>Disulfide bonds/thiols</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>β-Lactoglobulin (β-Lg)</td>
<td>60</td>
<td>162</td>
<td>18 363^a</td>
<td>5.35</td>
<td>2/1</td>
<td>Two common variants, A and B</td>
</tr>
<tr>
<td>α-Lactalbumin (α-La)</td>
<td>20</td>
<td>123</td>
<td>14 178</td>
<td>4.80</td>
<td>4</td>
<td>About 10% of molecules are glycosylated</td>
</tr>
<tr>
<td>Bovine serum albumin (BSA)</td>
<td>3</td>
<td>583</td>
<td>66 399</td>
<td>17/1</td>
<td></td>
<td>Also present in blood serum</td>
</tr>
<tr>
<td>Immunoglobulin G (IgG)</td>
<td>10</td>
<td>&gt;500</td>
<td>161 000 (G1)^b</td>
<td>Many isoforms</td>
<td></td>
<td>Passive transfer of immunities</td>
</tr>
<tr>
<td>Lactoferrin (Lf)</td>
<td>&lt;0.1</td>
<td>689</td>
<td>76 110</td>
<td>8.95</td>
<td>17</td>
<td>Bacteriostatic role; glycoprotein</td>
</tr>
</tbody>
</table>

^a Molar mass for the A variant.
^b G1 is the major immunoglobulin; two other classes, IgM and IgA, are present in much lower abundance.

Bovine β-lactoglobulin

β-Lg contains 162 amino acids and has a molecular weight of 18.3 kDa (Hambling et al., 1992). It is a member of the lipocalin (a contraction of the Greek lipos, “fat, grease” and calyx, “cup”) family of proteins (Banaszak et al., 1994; Flower, 1996), so called because of their ability to bind small hydrophobic molecules into a hydrophobic cavity. This led to the proposal that β-Lg functions as a transport protein for retinoid species, such as vitamin A (Papiz et al., 1986).

β-Lg is the most abundant whey protein in the milk of most mammals (≈10% of total protein or ≈50% of whey protein), but has not been detected in the milk of humans, rodents or lagomorphs. In the case of human milk, α-lactalbumin (see below) is the dominant whey protein. Bovine β-Lg is the most commonly studied milk protein.

There are ten known genetic variants of bovine β-Lg. The most abundant variants are labelled β-Lg A and β-Lg B (Farrell et al., 2004) and differ by two amino acid substitutions, Asp64Gly and Val118Ala respectively. The quaternary structure of the protein varies among monomers, dimers or oligomers depending on the pH, temperature and ionic strength, with the dimer being the prevalent form under physiological
conditions (Kumosinski and Timasheff, 1966; McKenzie and Sawyer, 1967; Gottschalk et al., 2003). This variable state of association is likely to be the result of a delicate balance among hydrophobic, electrostatic and hydrogen-bond interactions (Sakurai et al., 2001; Sakurai and Goto, 2002).

**Molecular structure of bovine β-Lg**

β-Lg was an early target of X-ray diffraction as newly applied at the Royal Institution to protein crystals. This was due to its high abundance and relatively easy purification from milk, and its propensity to form suitable crystals. In retrospect, this was a very ambitious project because β-Lg was not the easiest protein to analyze (Green et al., 1979), partly because of the multiple crystal forms. Nevertheless, this study established that the protein monomer was near spherical with a block of electron density with a rod-like structure across one face.

The next attempt (Creamer et al., 1983) to determine the structure was by calculation using sequence data and structural probabilities to estimate which portions of the amino acid sequence might form into the helices, strands and sheets. The secondary structure of β-Lg was predicted to comprise 15% α-helix, 50% β-sheet and 15–20% reverse turn (Creamer et al., 1983). It is interesting to note that many of the residues that reside in the extended structures of the native protein have been shown to have a nascent propensity to form α-helical structures in the presence of trifluoroethanol or amphiphiles (Hamada et al., 1995; Kuroda et al., 1996; Chamani et al., 2006).

In 1986, the first medium-resolution structure of β-Lg was published (Papiz et al., 1986). Structural similarity to a seemingly different type of protein, plasma retinol-binding protein, has given rise to much speculation as to the role of β-Lg in bovine milk. Higher resolution structures subsequently revealed the now familiar eight-stranded β-barrel (calyx), flanked by a three-turn α-helix. A final ninth strand forms the greater part of the dimer interface at neutral pH (Papiz et al., 1986; Bewley et al., 1997; Brownlow et al., 1997). The β-barrel is formed by two β-sheets, where strands A to D form one sheet and strands E to H form the other (with some participation from strand A, facilitated by a 90° bend at Ser21). Two disulfide bonds link Cys66 on loop CD (which, as its name suggests, connects strands C and D) with Cys160 near the C-terminus, and Cys106 on strand G with Cys119 on strand H, leaving Cys121 as a free, but unexposed, thiol. The loops connecting strands BC, DE and FG are relatively short whereas those at the open end of the barrel, strands AB, CD, EF and GH, are longer and more flexible. These features are illustrated in Figure 6.1.

The structures of the A and B variants are very similar. However, the Asp64Gly substitution results in the CD loop adopting different conformations (Qin et al., 1999). The Val118Ala substitution causes no detectable change to the structures, but the void created by substituting the bulky isopropyl substituent with the smaller methyl group results in the hydrophobic core of the B variant being less well packed, and may account for the lower thermal stability of the B variant (Qin et al., 1999).

Very careful titrimetric and thermodynamic measurements in the late 1950s (Tanford and Nozaki, 1959; Tanford et al., 1959) established the presence of a carboxylate residue with an anomalously high pKₐ value of 7.3. This was attributed to
a pH-dependent conformational change, a conclusion that rationalized earlier measurements of pH-dependent sedimentation coefficients (Pedersen, 1936) and specific optical rotation data (Groves et al., 1951). Much later, X-ray structure analyzes (Qin et al., 1998a) at pH values above and below this so-named Tanford transition established that, at pH 6.2, the EF loop is closed over the top of the barrel, burying Glu89 (the carboxylic acid with the anomalous $pK_a$) inside the calyx. At pH 8.1, this loop is articulated away from the barrel such that the formerly buried glutamic acid becomes exposed in the carboxylate form (Qin et al., 1998a).

An early structure of bovine $\beta$-Lg crystallized in the presence of retinol appeared to show retinol bound externally to the protein (Monaco et al., 1987), apparently later confirmed by a body of fluorescence data (Dufour et al., 1994; Lange et al., 1998; Narayan and Berliner, 1998). However, subsequent structural analyzes have shown that fatty acids, retinoid species (including vitamin A) and cholesterol (including vitamin D) all bind inside the calyx (Kontopidis et al., 2004). Induced circular dichroism (CD) measurements and NMR measurements confirm the X-ray crystallographic observations. Ligand binding is discussed in more detail below, because it relates to the probable physiological function of $\beta$-Lg as well as to the stability of this molecule and current technological interest in the role of protein–ligand interactions in flavor perception (see Chapter 13). At this stage, there is no evidence for the binding of fatty acids or retinoid species outside the calyx. Except for very bulky ligands (see below), ligands bind inside the calyx of $\beta$-Lg at pH $\approx 7$.

At about the same time as the Tanford transition and ligand-binding modes were elucidated by high-resolution X-ray crystallography (Qin et al., 1998a, 1998b), NMR
studies of β-Lg structure in low-pH solutions, where the protein is monomeric, were initiated (Ragona et al., 1997; Fogolari et al., 1998; Kuwata et al., 1998; Uhrínová et al., 1998). These NMR studies, described below, have provided proof of persistence of the tertiary structure down to pH 2 and have yielded a depth of insight into structural stability and protein dynamics that is not possible by standard X-ray crystallographic techniques.

**Structure of bovine β-Lg in aqueous solution**

NMR spectroscopy is used to obtain protein structures in solution (Cavanagh et al., 1995). The technique is best suited to monomeric proteins with molecular weights $<\approx 25$ kDa and usually requires recombinant singly ($^{15}$N) or doubly ($^{15}$N/$^{13}$C) labeled material for molecules with molecular weights $>\approx 8$ kDa. Therefore, most NMR studies of bovine β-Lg have been at a pH of between 2 and 3 where the molecule is monomeric. Early studies using wild-type B-variant protein (Fogolari et al., 1998) confirmed the presence of the eight-stranded β-barrel. However, the full structure (of the A variant) (Kuwata et al., 1999; Uhrínová et al., 2000) required the use of isotopically labeled recombinant material (Kim et al., 1997; Denton et al., 1998). As the full structure was determined by NMR techniques independently and near simultaneously by two groups from Tokyo and Edinburgh, this has provided objective and very useful comparisons (Jameson et al., 2002a).

The solution structure was shown by both Kuwata et al. (1999) and Uhrínová et al. (2000) to have an overall similarity to that established earlier by X-ray crystallography at pH 6.2, despite the considerably lower pH (and concomitant increase in the protein’s surface charge) necessary to obtain usable NMR spectra. The EF loop is firmly closed over the open end of the β-barrel at this pH and the side chain of the Glu89 “latch” is buried, as in the X-ray structures. The biggest difference, when compared with the Z lattice X-ray structure at pH 6.2 (Qin et al., 1998a), is that the three-turn α-helix adopts a different position with respect to the β-barrel, possibly because of the pH-induced increase in positive charge on this part of the protein’s surface (Uhrínová et al., 2000). The lower pH was also found to move the conformation of the AB loop by up to 3.5 Å, which may be significant for the disruption of the dimer interface.

Further differences were found at the N- and C-termini, but these can be ascribed to limitations imposed by the use of recombinant protein with a non-native N-terminus for the NMR structure and possible crystal-packing effects at the C-terminus for the X-ray structure. In some crystal forms, much of the C-terminus from residues $\approx 152$ to 162 is not observed or is very poorly defined in electron density maps.

**Studies of bovine β-Lg by NMR at neutral pH**

The large size of the bovine β-Lg dimer at pH 7 is expected to cause some broadening of the peaks in its $^{1}$H NMR spectrum because of slower molecular reorientation. However, this problem is exacerbated by chemical exchange broadening of peaks in the vicinity of the dimer interface by the dynamic equilibrium of molecules in the
Structure and stability of whey proteins

associated or unassociated state. These factors render the resulting spectra unsuitable for structure determination. Several methods have been employed to allow NMR studies at neutral pH. The most straightforward of these has been to use a non-ruminant \( \beta \)-Lg that is intrinsically monomeric, yet with the same overall tertiary structure as the bovine protein, in this case equine \( \beta \)-Lg (Kobayashi et al., 2000). Alternatively, the dimer interface may be disrupted by producing bovine \( \beta \)-Lg mutants with amino acid substitutions carefully chosen to disrupt the intermolecular interactions between either the I strands or the AB loops (Sakurai and Goto, 2002) (see Figure 6.1).

It is worth noting that an attempt to form dimeric equine \( \beta \)-Lg by producing a mutant with amino acid substitutions chosen to mimic those of the bovine protein at the interface was not successful (Kobayashi et al., 2002), indicating that subtle features in \( \beta \)-Lg conformation remote from the interface have an impact on successful dimer formation. Indeed, reaction of the free thiol of Cys121 (located away from the interface in the H strand and covered by the main \( \alpha \)-helix, Asp129–Lys141; see Figure 6.2) with 2-nitro-5-thiobenzoic acid produces a monomeric species with native structure at pH 2 and a monomeric but unfolded structure at pH 7 (Sakai et al., 2000). The configuration of the \( \alpha \)-helix is known to change with pH (Uhrinová et al., 2000) and this may therefore also have an important influence on both the protein’s stability and its quaternary state.

The third approach to overcome the problems of the rate constants for the dissociation/reassociation equilibrium of the bovine \( \beta \)-Lg dimer being in the intermediate exchange regime has been to covalently bond two monomers via an Ala34Cys mutant (Sakurai and Goto, 2006). This variant was used to study the dynamics of the EF loop across the Tanford transition (see the following section) and, more recently, to examine the nature of ligand binding to \( \beta \)-Lg (Konuma et al., 2007). Although no full structure determination was reported, the amide chemical shifts of the mutant were within 0.1 ppm of those from monomeric \( \beta \)-Lg (except for seven residues that encompassed the substitution site). This fact, combined with the similarity of the mutant and wild-type \( \beta \)-Lg CD spectra, indicated that the tertiary structures of the mutant and wild-type proteins were similar.

Bovine \( \beta \)-Lg dynamics

Crystallographic atomic displacement parameters, often loosely referred to as temperature factors or just \( B \) factors, describe the spread of an atom’s electron density in space and can therefore be used to infer residue-specific mobility. However, for surface residues, the \( B \) factors of both main-chain and side-chain atoms are highly sensitive to intermolecular crystal-packing contacts. Moreover, except where data to ultra-high resolution (better than 1.0 Å, which is not yet the case for any \( \beta \)-Lg) are available, similarity restraints are imposed on \( B \) values of adjacent atoms and residues along the polypeptide chain to ensure stable refinement.

Nonetheless, in the case of isomorphous structures at similar resolution (where structures share the same average \( B \) value, the same space group, very similar unit cell parameters and, hence, very similar intermolecular contacts), or in regions where non-isomorphous structures lack intermolecular contacts, some meaning can be
Figure 6.2  Ligand-binding sites on β-Lg as inferred from NMR measurements of binding of small (<12 atoms) ligands to β-Lg at acidic pH (Luebke et al., 2002). The binding site of 12-bromododecanoic acid is shown for reference (Qin et al., 1998b). (a) View into the calyx showing the primary binding site of fatty acids at pH ≈ 7 and of flavor components at pH ≈ 2, highlighted in yellow. (b) Secondary binding site for flavor components at pH 2 at N-terminal ends of strands A, B, C and D, and the C-terminal strand, highlighted in pink. (c) Secondary binding site for flavor components at pH 2 adjacent to the three-turn helix and strand G, highlighted in cyan. To show more clearly the attachment of side chains to the main chain, loops and strands have not been smoothed. The pH-sensitive EF loop is colored in magenta. Figure drawn with PyMOL (Delano, 2002) using co-ordinates with PDB code 1bso (see also Plate 6.2).

placed on differences observed in B factors. These differences can be both within a particular structure and between structures determined, for example, at different pH or in the presence/absence of added ligands. High B factors, indicating apparent high mobility, can also arise from a distribution of slightly different yet immobile conformations or from errors in model building. For these reasons, it is advantageous to study dynamics of the protein (particularly those of the backbone) by NMR techniques, using uniformly 15N-labeled protein.
Flexibility on the nanosecond timescale can be inferred from low $^{15}$N steady-state nuclear Overhauser effect values. As might be expected, mobile residues for $\beta$-Lg tend to have highly accessible surface areas (and such residues identified in NMR studies correlate in general with those that have relatively high $B$ factors; Kuwata et al., 1999). Slower conformational exchange processes can be indicated by large values for the ratio of the $T_1$ (spin–lattice) and $T_2$ (spin–spin) relaxation times of $^{15}$N nuclei. Such residues include Ser21 at the midpoint (kink) of the A strand, possibly caused by fluctuations of the barrel, and residues 61 and 66 at either end of the CD loop, consistent with a slow segmental or hinging motion of this loop (Uhrínová et al., 2000).

All three sets of relaxation parameters can be analyzed in concert using the extended model-free formalism to give amplitudes and timescales for the internal motion of the backbone N–H bond vectors (Lipari and Szabo, 1982; Clore et al., 1990).

When applied to variant A of $\beta$-Lg, this method confirms the above observations, but also identifies a number of residues in the EF loop undergoing substantial conformational change (Edwards et al., 2003). Preliminary results also suggest that the B variant is more mobile relative to the A variant at the Asp64Gly (Gly in variant B) substitution site, whereas the dynamics at the Val118Ala (Ala in variant B) substitution site are very similar (Edwards et al., 2003).

NMR measurements of the dynamics of the covalently bonded Ala34Cys mutant dimer have recently given complementary information regarding the structural changes associated with the Tanford transition established previously using X-ray crystallography (Qin et al., 1998a). The $^{15}$N dynamics of the EF loop measured either side of the transition indicate a three-step process. With increasing pH, the first event is a conformational change to the GH loop. This is followed by the breaking of hydrogen bonds at the hinges of the EF loop followed by the subsequent articulation of the EF loop away from the calyx (Sakurai and Goto, 2006). The dynamic flexibility of the EF loop at pH ≈ 2 is important, as it means that, at low pH, neither ingress into nor egress from the hydrophobic pocket is kinetically prevented.

**Structures of $\beta$-Lgs from other species**

Equine (horse) $\beta$-Lg, which shares 58% identity with bovine $\beta$-Lg, has been shown to be monomeric over a wide pH range, whereas porcine (pig) $\beta$-Lg, which shares 63% identity with bovine $\beta$-Lg, is dimeric below pH 5 and monomeric at pH 5 and above (in contrast to bovine $\beta$-Lg). At pH 7, both equine $\beta$-Lg and porcine $\beta$-Lg are monomeric and therefore amenable to NMR study. Equine $\beta$-Lg has been extensively studied by NMR with regard to denaturation processes, but a full structural characterization by either NMR or X-ray methods has yet to be published.

The X-ray crystal structure of porcine $\beta$-Lg at pH 3.2 clearly revealed a dimeric structure formed by domain swapping of N-terminal regions, a quaternary structure quite different from that observed for bovine $\beta$-Lg (Hoedemaeker et al., 2002). The EF loop adopts the closed conformation over the calyx, as found also for bovine $\beta$-Lg at acidic pH, consistent with the notion that this loop acts as a lid to the calyx. However, the porcine protein is much less conformationally stable at acidic pH than its bovine counterpart (Burova et al., 2002; Invernizzi et al., 2006), which has led to
questioning of the role of \( \beta \)-Lg as a transporter of hydrophobic molecules through the acidic environment of the gut (Burova et al., 2002). Despite 63% identity between bovine \( \beta \)-Lg and porcine \( \beta \)-Lg, the RMS difference in \( \mathrm{C}_\alpha \) positions between these two structures is remarkably high at 2.8 Å, although inspection of the two structures shows that the core \( \beta \)-barrel structure superimposes closely and that these differences are concentrated in the flexible loop regions, which comprise nearly a third of the structure.

The 2.1 Å resolution structure of rangiferine (reindeer) \( \beta \)-Lg at pH \approx 6.5 was published recently (Oksanen et al., 2006). Both the monomeric tertiary structure and the dimeric quaternary structure are very similar to those of bovine \( \beta \)-Lg, which is not unexpected as polypeptide lengths are identical and sequence identity is greater than 94%. At pH \approx 6.5, the EF loop is observed to be in the closed position. There are few structural data on ovine (sheep) and caprine (goat) \( \beta \)-Lgs, despite the commercial importance of their milk.

Although \( \beta \)-Lg is absent from human milk, two other secreted lipocalins share limited sequence identity (but close structural similarity) with bovine \( \beta \)-Lg. Tear lipocalin is the major protein in tears, and structural and functional studies indicate that it, like \( \beta \)-Lg, binds a broad range of hydrophobic molecules (Glasgow et al., 1995). Glycodelin, a heavily glycosylated lipocalin, is found in the human endometrium in early pregnancy where its function remains unclear. It has been suggested (Kontopidis et al., 2004) that \( \beta \)-Lg arose by gene duplication of glycodelin and exists in milk solely for nutritive value. However, there appears to have been considerable selection pressure to retain not only the Glu89 in all sequences of the \( \beta \)-Lgs found to date but also the pH-sensitive conformational switch for the EF loop of all \( \beta \)-Lgs (and indeed for tear lipocalin) that have been functionally characterized. Such preservation is not consistent with a solely nutritive role for \( \beta \)-Lg, although the origin of the \( \beta \)-Lg gene from glycodelin remains an intriguing possibility.

**Ligand binding to \( \beta \)-Lg**

\( \beta \)-Lg has the ability to bind a large number of small molecules (for a review, see Sawyer et al., 1998), although the location of the bound ligand continues to generate controversy (Dufour et al., 1994; Lange et al., 1998; Narayan and Berliner, 1998). However, in recent years, NMR techniques have been used to map ligand-binding sites on the protein through changes in chemical shift and relaxation times of protein residues.

Another technique that provides reliable evidence of binding to the protein is induced CD, where an achiral chromophore “lights up” in the CD spectrum on being placed into a chiral environment on or in the protein. This technique was applied to bovine \( \beta \)-Lg by us (Creamer et al., 2000) to show unequivocally that retinol and fatty-acid binding (e.g. palmitic and \textit{cis}-parinaric acids) is competitive. The binding of these and other chirally active ligands, including \textit{trans}-parinaric acid and retinoic acid, was explored over a range of pH and with non-chromophoric fatty-acid ligands to gain an understanding of the parameters surrounding the Tanford transition. More recently, induced CD has been used to study the binding of these (Zsila et al., 2002)
and other ligands (Zsila, 2003; Zsila et al., 2005), including piperine, to β-Lg and to other lipocalins.

Ligand binding has also been extensively studied by fluorescence spectroscopy in which, typically, the fluorescence from tryptophan residues is monitored for changes, which may be positive or negative, that are interpreted as being due to ligand binding. For bovine β-Lg, one tryptophan (Trp19) is buried and is part of the highly conserved lipocalin Gly–X–Trp motif at the beginning of strand A (see Figure 6.2b), whereas the other (Trp61) is largely exposed and is part of the mobile CD loop. Fluorescence by both tryptophans is sensitive to small changes in the positions of nearby quenchers, respectively a nearby charged side chain of Arg124 and the Cys66–Cys160 disulfide bond. NMR spectroscopy, induced CD and X-ray results clearly indicate that interpretation of fluorescence measurements for evidence of ligand binding poses hazards.

Ragona et al. (2003) used a combination of electrostatic calculations, docking simulations and NMR measurements to suggest that the pH-dependent conformational change of the EF loop triggered by the protonation of Glu89 is common to all β-Lgs and that ligand binding (of palmitic acid) is determined by the opening of this loop. In earlier work using 13C-labelled palmitic acid, this group had shown that the ligand also undergoes conformational change with increasing pH (Ragona et al., 2000).

Recent NMR studies of the binding of palmitic acid to the “NMR friendly” Ala34Cys mutant dimer of bovine β-Lg have indicated that, although a rigid connection is made by the protein with the ligand at the bottom of the calyx, the interaction at the open end of the calyx is more dynamic (Konuma et al., 2007). These observations complement those of Ragona et al. (2003) and also the X-ray studies on the binding of fatty acids (Qin et al., 1998b; Wu et al., 1999), which showed the carboxylate head group to be substantially less well ordered than the hydrophobic tail. The results of the study with the Ala34Cys mutant suggest that it is the plasticity of the D strand and the EF and GH loops that allows β-Lg to accommodate such a wide range of ligands (Konuma et al., 2007). With the exception of changes in conformations of the side chains of Phe105 and Met107, NMR and X-ray studies show that the core lipocalin structure remains invariant upon ligand binding.

Crystallographic data clearly show that both fatty acids and retinol bind in the calyx (Qin et al., 1998b; Sawyer et al., 1998; Wu et al., 1999). Although all crystallographic studies of ligand binding have been under conditions of high ionic strength, congruence of these data with NMR and induced CD data collected under conditions of low ionic strength indicate that the X-ray results are not an artifact of ionic strength. The preservation of the structure of β-Lg, in particular the hydrophobic cavity, at conditions of near-zero ionic strength at the pI of the protein (≈5.3) (Adams et al., 2006) further demonstrates that the primary, and possibly only, ligand-binding site at ≈pH 7 is inside the calyx.

Although ligands such as palmitic acid appear to be released at acid pH (Ragona et al., 2000), NMR evidence (based on perturbations of backbone chemical shifts) for the binding of the flavor compounds γ-decalactone and β-ionone at pH 2 has been reported (Luebke et al., 2002; Tromelin and Guichard, 2006). Therefore, there is evidence for three binding sites to β-Lg: the canonical site inside the calyx;
a second site involving perturbation of residues Trp19, Tyr20, Glu44, Gln59, Gln68, Leu156, Glu157, Glu158 and His161; and a third site involving residues Tyr102, Leu104 and Asp129. These binding sites are illustrated in Figure 6.2.

Initially, it was thought that porcine $\beta$-Lg did not bind fatty acids (Frapin et al., 1993). However, recent NMR studies have shown that the pH for 50% uptake of ligand has shifted by nearly 4 pH units from $\approx 5.8$ for bovine $\beta$-Lg to 9.7 for porcine $\beta$-Lg, whereupon the EF loop undergoes a structural change analogous to that of its bovine counterpart (Ragona et al., 2003).

**Effect of temperature on bovine $\beta$-Lg**

The thermal properties of $\beta$-Lg variants are of considerable commercial relevance because of their role in the fouling of processing equipment as well as the functional qualities that can be imparted to dairy products by thermally induced $\beta$-Lg aggregation. Consequently, this aspect of the protein’s behavior has been the focus of extensive experimental work.

At neutral pH, the midpoint of the thermal unfolding transitions, as determined by differential scanning calorimetry (DSC), is $\approx 70^\circ C$ (de Wit and Swinkels, 1980), whereupon the protein dimer dissociates and the constituent molecules begin to unfold. This reveals the free thiol of Cys121 (located at the C-terminal end of the H strand—see Figure 6.1) and a patch of hydrophobic residues, leading to the possibility of both covalent and hydrophobic intermolecular association (Qi et al., 1995; Iametti et al., 1996). The ensuing disulfide interchange reactions lead to the formation of a variety of mixed disulfide-bonded polymeric species (Creamer et al., 2004).

Genetically engineered mutants with an extra cysteine positioned to allow a third disulfide bond to be formed to Cys121 have been shown both to retard thermal denaturation by 8–10°C and to resist heat-induced aggregation (Cho et al., 1994). In mixtures of bovine $\beta$-Lg, $\alpha$-lactalbumin ($\alpha$-La) and bovine serum albumin (BSA), or of $\beta$-Lg and one or other of $\alpha$-La and BSA at pH 6.8 subjected to high temperatures, homo- and heteropolymeric disulfide-bridged species were observed (Havea et al., 2001). The formation of $\alpha$-La–$\alpha$-La disulfide links ($\alpha$-La has no free cysteine; see below) is attributed to catalysis by BSA or $\beta$-Lg (Havea et al., 2001). At low pH, where the protein is monomeric, denaturation is largely reversible at temperatures below 70°C (Pace and Tanford, 1968; Alexander and Pace, 1971; Mills, 1976; Edwards et al., 2002). Heating above this temperature leads to the formation of large aggregates, but, in contrast to the behavior at neutral pH, the species are predominantly non-covalently bonded (Schokker et al., 2000).

The precise denaturation process is complex and is influenced by factors such as pH, protein concentration, ionic environment, genetic variant and presence of ligands. Both lowering the pH (Kella and Kinsella, 1988; Relkin et al., 1992) and adding calyx-bound ligands (Puyol et al., 1994; Considine et al., 2005a; Busti et al., 2006) make the protein more resistant to thermal unfolding. The stability of the genetic variants (at pH 6.7) appears to decrease in the order C $>$ A $>$ B, with the A variant showing the least co-operative unfolding transition (Manderson et al., 1997). The protein’s susceptibility to thermal denaturation at pH 6.7–8 is strongly concentration dependent
up to about 6 mM, being most susceptible to unfolding at a concentration of \( \approx 1.4 \) mM (Qi et al., 1995). It is possible that, at high protein concentration (\( \approx 6 \) mM), tertiary structure is lost directly from the native dimer state (Qi et al., 1995).

There is some evidence that the thermal unfolding occurs in more than one step. Kaminogawa et al. (1989) used antibody binding affinities to propose that thermal unfolding of variant A of \( \beta \)-Lg occurs in at least two stages, starting with conformational changes near the N-terminus followed by changes in the region of the three-turn \( \alpha \)-helix. Fourier-transform infrared (FT-IR) measurements by Casal et al. (1988) have also indicated a loss of helical content early in the denaturation process (using variant B of \( \beta \)-Lg in 50 mM phosphate buffer at pH 7). Qi et al. (1997) used FT-IR and CD measurements to propose that variant A of \( \beta \)-Lg forms a molten globule with reduced \( \beta \) structure when heated above 65°C in 30–60 mM NaCl at pH 6.5. NMR studies, observing hydrogen/deuterium (H/D) exchange of the backbone amide protons of \( \beta \)-Lg A at pH 2–3, have revealed a stable core comprising the FG and H strands, possibly stabilized by the Cys106–Cys119 disulfide bond between strands F and G (Belloque and Smith, 1998; Edwards et al., 2002). Significant secondary structure even at a temperature as high as 90°C has been reported (Casal et al., 1988; Qi et al., 1997; Bhattacharjee et al., 2005).

Effect of pressure on bovine \( \beta \)-Lg

High-pressure treatment of food is of increasing commercial importance because of increasing consumer demand for products that have been subjected to minimal processing damage. Pressure treatment as part of the processing regime has the potential to produce dairy products with improved functional and organoleptic properties compared with those produced by thermal treatment alone (Messens et al., 2003).

Of the major whey proteins, \( \beta \)-Lg is the most susceptible to pressure-induced change (Stapelfeldt et al., 1996; Patel et al., 2005). Presumably this is due to its relatively inefficient packing, caused by the presence of the \( \beta \)-barrel with its large solvent-exposed hydrophobic pocket, and the lower number of disulfide bonds (two compared with four in, for example, the similar-sized \( \alpha \)-La). A reduction in the molar volume of bovine \( \beta \)-Lg has been detected at pressures as low as 10 MPa, possibly because of a contraction of the calyx (Vant et al., 2002). A number of studies have shown that \( \beta \)-Lg becomes more susceptible to enzymatic cleavage when exposed to pressure, possibly because of pressure-induced conformational change. The free cysteine has been shown to become exposed at between 50 and 100 MPa (Stapelfeldt et al., 1996; Tanaka and Kunugi, 1996; Moller et al., 1998).

Exposure of the protein to pressures in excess of \( \approx 300 \) MPa causes irreversible changes to \( \beta \)-Lg’s tertiary and quaternary structure. A combination of CD and fluorescence spectroscopy of \( \beta \)-Lg at neutral pH exposed to pressures as high as 900 MPa indicated that pressure induces monomer formation with subsequent aggregation, but with only small irreversible effects on \( \beta \)-Lg tertiary structure (Iametti et al., 1997). However, more recent results from tryptic hydrolysis suggest that, whereas exposure to pressures below 150 MPa has no detectable permanent effect on \( \beta \)-Lg A’s conformation, pressures above 300 MPa lead to the detachment of strands D and G from
the β-barrel together with the formation of disulfide-bonded oligomers (Knudsen et al., 2002).

In mixtures of bovine β-Lg with either α-La or BSA at pH 6.6 subjected to high pressures, intermolecular disulfide-bridged aggregates form only between β-Lg and itself. No β-Lg–α-La or β-Lg–BSA disulfide-bridged species are detected (Patel et al., 2005), in contrast to heat-treated mixtures where such species are observed (Havea et al., 2001).

In order to correlate the pressure-induced conformational changes with the protein’s primary sequence, Belloque et al. (2000) made NMR amide H/D exchange observations of β-Lg A and B following exposure of solutions at neutral pH to pressures of up to 400 MPa. Little H/D exchange was reported at 100 MPa, which indicated that any conformational change that occurred did not increase the exposure of most amide protons to the solvent compared with their exposure in the native conformation at ambient pressure. A large increase in the extent of H/D substitution at 200 MPa and above indicated increased conformational flexibility, but the similarity of the spectra of control samples recorded in H₂O rather than D₂O before and after pressurization demonstrated that any pressure-induced conformational changes were largely reversible up to 400 MPa. The authors proposed that the structure of the A variant was more sensitive to changes in pressure than that of the B variant and that the F, G and H strands of the protein’s β-barrel were the most resistant to conformational change, the latter conclusion paralleling the effects of temperature (Belloque and Smith, 1998; Edwards et al., 2002).

FT-IR and small-angle X-ray scattering experiments suggest that, even at 1 GPa, the unfolded state contains significant secondary structure (Panick et al., 1999). Combined application of pressure and heat has shown that changing the temperature over the range from 5 to 37°C has negligible effect on the susceptibility of β-Lg to pressures up to 200 MPa (Skibsted et al., 2007). However, combined application of pressure and moderate temperature at 600 MPa/50°C (Yang et al., 2001) and 294 MPa/62°C (Aouzelleg et al., 2004) has indicated the formation of a molten globule with an α-helical structure on the basis of results obtained from CD spectroscopy.

It should be noted therefore that the potential for temperature increases induced by rapid pressurization of the sample needs to be considered when studying the effects of pressure on protein conformation and stability.

Enzymatic proteolysis observations indicate that β-Lg is less susceptible to pressure-induced change at acidic pH (Dufour et al., 1995) than at neutral or basic pH. Nevertheless, NMR measurements of monomeric β-Lg at pH 2 while under pressure at up to 200 MPa have shown that the two β-sheets unfold independently to form two intermediates in an unfolded state that still appears to contain significant secondary structure (Kuwata et al., 2001).

A three-step mechanism has been proposed for β-Lg denaturation at neutral pH and ambient temperature, which broadly encompasses the above observations: a pressure of 50 MPa causes partial collapse of the calyx (with concomitant reduction in ligand-binding capacity) together with exposure of Cys121. Increasing the pressure to 200 MPa causes further (partially reversible) disruption to the hydrophobic structure.
together with a decrease in the molecular volume. Higher pressures cause irreversible aggregation reactions involving disulfide interchange reactions (Stapelfeldt and Skibsted, 1999; Considine *et al.*, 2005b).

**Effect of chemical denaturants on bovine β-Lg**

Chemical denaturants are often used to unfold proteins and to characterize mechanisms and transition states of protein-folding processes. Commonly used denaturants include alcohols, particularly 2,2,2-trifluoroethanol (TFE), urea and guanidinium chloride (GdmCl).

Theoretical calculations predict a significantly higher amount of α-helical secondary structure than is actually observed in native β-Lg (Creamer *et al.*, 1983; Nishikawa and Noguchi, 1991). That is, the native structure is the result of competition between α-helix-favoring local interactions and β-sheet-forming long-range interactions. However, addition of alcohols such as TFE can disturb this balance by weakening the hydrophobic interactions and strengthening the helical propensity of the peptide chain (Thomas and Dill, 1993). The ability to increase the α-helical content of bovine β-Lg by the addition of alcohols (ethanol, 1-propanol, 2-chloroethanol) was first demonstrated by Tanford *et al.* (1960) using optical rotary dispersion measurements.

Contemporary studies tend to favor the use of TFE, where the β-Lg β-sheet-to-α-helix transition has been shown to be highly co-operative, occurring over the range ≈15–20% v/v of cosolvent (Shiraki *et al.*, 1995; Hamada and Goto, 1997; Kuwata *et al.*, 1998). The higher proportion of α-helical structure in the so-called TFE-state is found in the N-terminal half of the molecule (Kuwata *et al.*, 1998). Magnetic relaxation dispersion measurements of the solvent nuclei have shown that this state is an open, solvent-permeated structure (unlike the collapsed state of a molten globule) and that its formation is accompanied by a progressive swelling of the protein with increasing TFE concentration (Kumar *et al.*, 2003). High protein and TFE concentration (8% v/w and 50% v/v respectively) can lead to fibrillar aggregation and gel formation of bovine β-Lg at both acid and neutral pH (Gosal *et al.*, 2002).

The ability of urea to induce protein unfolding is thought to be via a combination of hydrogen-bond formation with the protein backbone and a reduction in the magnitude of the hydrophobic effect (Bennion and Daggett, 2003). Therefore, in contrast to TFE, both the helical propensity and the hydrophobic effect are reduced. Urea-induced unfolding of bovine β-Lg at acidic pH was first reported as a two-state process (Pace and Tanford, 1968). Subsequent NMR H/D exchange measurements of bovine β-Lg B at pH 2.1 also allowed the urea-induced unfolding to be well approximated as a two-state transition between folded protein and the unfolded state via a co-operative unfolding of the β-barrel and the C-terminus of the major α-helix (Ragona *et al.*, 1999). However, Dar *et al.* (2007) have recently provided evidence that urea also causes unfolding via an intermediate, albeit with structural properties between those of the native and unfolded states. Addition of anionic amphiphiles, sodium dodecyl sulfate (SDS) or palmitate, causes β-Lg to resist urea-induced unfolding because of binding inside the calyx (Creamer, 1995).
GdmCl is often used as an alternative to urea in studies of protein stability. At the neutral or acidic pH of most stability studies, GdmCl will be fully dissociated. At low GdmCl concentration (< ≈ 1 M), chloride ions screen the electrostatic repulsion between positively charged groups of the protein (Hagihara et al., 1993). The result is that the additional electrostatic interactions of GdmCl compared with the neutral urea molecule have the potential to both stabilize and destabilize protein structure depending on the concentration of GdmCl (Hagihara et al., 1993).

D’Alfonso et al. (2002) have compared the denaturations of bovine β-Lg B with both GdmCl and urea between pH 2 and 8, as monitored by CD, UV differential absorption and fluorescence measurements. Discrepancies between unfolding free energies obtained using the two denaturants could be reconciled if GdmCl denaturation was assumed to occur via an intermediate state. The secondary structure of this state is similar to that of the native protein, but with greater rigidity in the vicinity of the tryptophan residues, consistent with the screening of electrostatic repulsion between charged residues (D’Alfonso et al., 2002). The GdmCl-induced unfolding intermediate of bovine β-Lg A at pH 2 has been reported to have increased α-helical structure (Dar et al., 2007).

Porcine β-Lg has also been shown to unfold via an intermediate state on addition of GdmCl. The stability of the porcine protein was lower than that of its bovine counterpart and the intermediate state was richer in α-helical structure. Most of the hydrophobic–hydrophobic interactions of the buried core of the native state are conserved between bovine β-Lg and porcine β-Lg. However, four pairwise interactions of the Phe105 side chain of bovine β-Lg are lost on the change to Leu in the porcine protein. This indicates that the presence of the aromatic residue may play an important role in the increased stability of the bovine protein (D’Alfonso et al., 2004). It is interesting to note that this residue is particularly resistant to H/D exchange in heated β-Lg solutions (Edwards et al., 2002).

**α-Lactalbumin**

α-La is a 123-amino-acid, 14.2-kDa globular protein that is found in the milk of all mammals. The bovine protein binds Ca\(^{2+}\), with the holo form being the more abundant form in milk. Within the Golgi apparatus of the mammary epithelial cell, α-La is the regulatory component of the lactose synthase complex (in which it combines with N-acetyl lactosamine synthase, now named β-1,4-galactosyltransferase-I), the role of which is to transfer galactose from UDP-galactose to glucose (Brew, 2003). In the absence of α-La and in the presence of a transition metal ion such as manganese(II), the catalytic domain of bovine β-1,4-galactosyltransferase-1 (residues 130–402) transfers galactose (Gal) to N-acetylgalactosamine (GlcNAc), which may be either free or linked to an oligosaccharide, generating a disaccharide unit, Gal-β-1,4-GlcNAc (N-acetyl lactosamine). The Ca\(^{2+}\) ion binding site is remote from the active site of the α-La–β-1,4-galactosyltransferase-I complex (Brew, 2003). α-La has been studied extensively, largely because of its formation of a molten globule state under mild denaturing conditions (Dolgikh et al., 1981).
Molecular structure of bovine $\alpha$-La

The tertiary structure of bovine $\alpha$-La is typical of that of the protein from other mammalian species (Acharya et al., 1991; Calderone et al., 1996; Pike et al., 1996) and is similar to that of lysozyme, with which it shares significant homology. As illustrated in Figure 6.3a, $\alpha$-La is made up of two lobes: the $\alpha$-lobe contains residues 1–34 and 86–123; and the smaller $\beta$-lobe spans residues 35–85. The $\alpha$-lobe contains three $\alpha$-helices (residues 5–11, 23–34 and 86–98) and two short $3_{10}$-helices (residues 18–20 and 115–118). A small, three-stranded $\beta$-sheet (residues 41–44, 47–50 and

![Figure 6.3](image-url)

Figure 6.3  (a) Structure of bovine $\alpha$-La showing the Ca$^{2+}$ ion binding site (PDB code: 2yfd). The peptide chain is rainbow colored, beginning at the N-terminus in blue and progressing to the C-terminus in red, in order to show the assembly of the sub-domains. The Ca$^{2+}$ ion is five co-ordinate. Loop 79–84 provides three ligands, two from main-chain carbonyl oxygen atoms of Lys79 and Asp84 and one from the side chain of Asp82. Co-ordination about the Ca$^{2+}$ ion is completed by carboxylate oxygen atoms from Asp87 and Asp88 at the N-terminal end of the main four-turn helix and by two water molecules. The four disulfide bonds are shown in ball-and-stick representation (one in the helical domain is obscured and the two linking the helical domain and the Ca$^{2+}$ ion binding loop to the $\beta$ domain are on the left half of the panel). (b) The lactose synthase complex formed from bovine $\alpha$-La (yellow) with $\beta$-1,4-galactosyltransferase (gray) (PDB code: 1f6s). Several substrate molecules are observed, together with the cleaved nucleotide sugar moiety (cyan sticks). The Mn$^{2+}$ ion is shown as a pink sphere and the Ca$^{2+}$ ion is shown as a gray sphere. For clarity, loop regions are given a smoothed representation. Figure drawn with PyMOL (Delano, 2002) (see also Plate 6.3).
The structure is stabilized by four disulfide bonds (Cys6–Cys120 and Cys28–Cys111 in the α-lobe, Cys60–Cys77 in the β-sheet and Cys73–Cys90 tethering the two lobes together) (Brew, 2003).

Unlike β-Lg, there is no free thiol. A Ca\(^{2+}\) ion binds with a sub-micromolar dissociation constant at the so-called binding elbow formed by residues 79–88 located in a cleft between the two lobes, with the metal ion co-ordinated in a distorted pentagonal bipyramidal configuration by the side-chain carboxylate groups of Asp82, Asp87 and Asp88, the carbonyl oxygens of Lys79 and Asp84 and the oxygen atoms of two water molecules (Pike et al., 1996).

The structure of the apo form of bovine α-La is similar to that of the holo protein; the largest changes involve the movement of the Tyr103 side chain in the inter-lobe cleft with little change in the vicinity of the Ca\(^{2+}\) ion binding site (Chrysina et al., 2000). The salient features of the structure of the holo protein are depicted in Figure 6.3, together with its complex with β-1,4-galactosyltransferase-I with bound substrates and, interestingly, a trapped intermediate species.

The structures of α-La from several other species, including baboon, human, guinea pig and buffalo, have also been characterized by X-ray diffraction methods. Consistent with high sequence identity, there are no significant differences among these structures, except for a flexible loop at residues 105–110 implicated in the formation of the lactose synthase complex (Acharya et al., 1989; Calderone et al., 1996; Pike et al., 1996). It is worth noting that the recombinant goat protein, which has an added methionine at the N-terminus, is markedly less stable, by \(-14\) kJ/mol, than the native protein, mostly the result of an increased rate of unfolding (but a preserved rate of refolding) (Chaudhuri et al., 1999). Similar observations have been made on recombinant bovine α-La (Acharya et al., 1989). Thus, native protein functionality and stability should not in general be inferred from measurements of recombinant proteins heterologously expressed in bacterial systems (which generally add an N-terminal methionine residue).

**Effect of temperature on bovine α-La**

In general, holo-α-La undergoes a thermal unfolding at a lower temperature than does β-Lg (Ruegg et al., 1977). The role of bound Ca\(^{2+}\) ions appears to be to confer stability to the tertiary structure: with less than equimolar amounts of bound calcium, the thermal unfolding transition is lowered substantially, decreasing to about 35°C for the apo form (Relkin, 1996; Ishikawa et al., 1998). The presence of Ca\(^{2+}\) ions also accelerates the rate of refolding of α-La by more than two orders of magnitude (Wehbi et al., 2005). The presence of Ca\(^{2+}\) ions also aids in the refolding and the formation of the correct disulfide linkages of the denatured reduced protein (Belloque et al., 2000). The structurally closely related, but functionally unrelated, enzyme lysozyme can be subdivided into two classes: a non-calcium-binding subclass, typified by egg-white lysozyme (Grobler et al., 1994; Steinrauf, 1998), and a calcium-binding subclass, including equine and echidna lysozyme (Tsuge et al., 1992; Guss et al., 1997).
The thermal denaturation behavior of bovine $\alpha$-La from three different sources has been studied; significant differences have been reported (McGuffey et al., 2005), which has provided some resolution of the apparently discordant denaturation data from different groups. In the presence of $\beta$-Lg or BSA, each of which has an unpaired cysteine, $\beta$-Lg--$\alpha$-La, $\alpha$-La--BSA and even $\alpha$-La--$\alpha$-La oligomers form at high temperature. Because $\alpha$-La (which lacks a free thiol) by itself fails under similar conditions to form disulfide-linked oligomers, intermolecular disulfide–sulfydryl interchange reactions appear to play a role in forming $\alpha$-La--$\alpha$-La oligomers (Havea et al., 2001; Hong and Creamer, 2002).

**Effect of pressure on bovine $\alpha$-La**

Again, the absence of free thiol groups renders $\alpha$-La intrinsically less susceptible to irreversible structural and functional change induced by high pressure. Reversible unfolding to a molten globule state begins at 200 MPa and loss of native structure becomes irreversible beyond 400 MPa (McGuffey et al., 2005) (corresponding figures for $\beta$-Lg are 50 and 150 MPa [Stapelfeldt and Skibsted, 1999; Considine et al., 2005b]). In the presence of Ca$^{2+}$ ions, the denaturation pressure increases by 200 MPa for $\alpha$-La (Dzwolak et al., 1999). Only in the presence of thiol reducers does oligomerization of $\alpha$-La occur at high pressures (Jegouic et al., 1996).

**Effect of chemical denaturants on bovine $\alpha$-La**

At neutral pH, the calcium-depleted, or apo, form of $\alpha$-La reversibly denatures to a variety of partially folded or molten globule states upon moderate heating (45°C), or, at room temperature, by dissolving the protein in aqueous TFE (15% TFE) or by adding oleic acid (7.5 equivalents) (Svensson et al., 2000; de Laureto et al., 2002). Under these various conditions, the UV-CD spectra of apo-$\alpha$-La are essentially identical to those of the most studied molten globule form of $\alpha$-La—the A-state found at pH 2.0 (Kuwajima, 1996). At 4°C and pH 8.3, proteolysis of apo-$\alpha$-La by proteinase-K occurs slowly and non-specifically, leading to small peptides only. In contrast, at 37°C, preferential cleavage by proteinase-K is observed at peptide bonds located in loop regions of the $\beta$-sheet sub-domain of the $\beta$-domain of the protein (residues 35–85), creating peptides in which disulfide bridges link N-terminal residues 1–34 to C-terminal fragments, residues 54–123 or 57–123.

Preferential cleavage at similar sites and similar disulfide-bridged fragments have also been observed for proteolysis of the molten globule states induced by TFE and oleic acid. Polypeptides formed from the molten globule A-state of $\alpha$-La comprise, therefore, a well-structured native-like conformation of the $\alpha$-domain and a disordered conformation of the $\beta$-sub-domain, residues 34–57 (de Laureto et al., 2002).

Oleic acid treatment leads to a kinetically trapped folding variant of the protein, which can also bind Ca$^{2+}$ ions, called HAMLET (human $\alpha$-lactalbumin made lethal to tumor cells, and its bovine analog BAMLET), which has been shown to induce apoptosis in tumor cells (Svensson et al., 2000). Under conditions where thermal denaturation of $\alpha$-La is reversible, thermal denaturation of HAMLET is irreversible, with respect to loss of its apoptotic effect on tumor cells (Fast et al., 2005).
Human $\alpha$-La and bovine $\alpha$-La also weakly bind a second $\text{Ca}^{2+}$ ion, which has been structurally characterized for human $\alpha$-La (Chandra et al., 1998). In addition, $\text{Zn}^{2+}$ ion binding to possibly structurally inequivalent sites has been characterized for human $\alpha$-La and bovine $\alpha$-La by fluorescence spectroscopic techniques (Permyakov and Berliner, 2000). The binding of $\text{Zn}^{2+}$ ions to calcium-loaded $\alpha$-La has been shown to destabilize the native structure to heat denaturation (Permyakov and Berliner, 2000). However, the weak binding of zinc (sub-millimolarity dissociation constant) means that it is probably not physiologically relevant.

### Serum albumin

Serum albumin (SA) is an approximately 580-residue protein that is found in both the blood serum and the milk of all mammals and appears to function as a promiscuous transporter of hydrophobic molecules. However, as with many proteins, this transport role appears not to be the only physiological function for SAs. The structure of human serum albumin (HSA), described in more detail in the next sub-section, is notable also for the number of disulfide bridges, 17 in total. There is one unpaired cysteine, Cys34 in HSA (highlighted in Figure 6.4), and also Cys34 in BSA. This cysteine is part of a highly conserved QQCP(F/Y) motif. It is susceptible to various oxidations, including a two-electron oxidation to sulfenic acid (−SOH) (Carballal et al., 2007).

Recently, evidence that, at least in blood serum, this cysteine is involved in HSA’s role in the control of redox properties has been found (Kawakami et al., 2006). A similar role can be postulated for BSA in blood serum but, in both human milk and bovine milk, this redox role has not been established (or even investigated). In terms of milk flavor and the flavor of milk products, control of the redox states of milk components is obviously of importance. It appears also that, in blood serum, where HSA is the major protein component, present at a concentration of 0.6 mM, HSA is the first line of defence against radicals, including reactive oxygen species and nitric oxide.

*In vitro* studies showing the reactivity of Cys34 in HSA have been complemented by *in vivo* studies that show that, in primary nephrotic syndrome, Cys34 is oxidized to sulfonate, $-\text{SO}_3^-$ (Musante et al., 2006). Again, in milk, defences against reactive oxygen species are essential to preserve milk quality, but HSA and BSA are at much lower concentrations in milk than in blood. HSA has also been shown to have esterase activity (Sakurai et al., 2004). Whether this is physiologically important in blood (or in milk) has not been established for either BSA or HSA. Finally, an active role for HSA in the transport of fatty acids across membranes has been characterized (Cupp et al., 2004). It is in this process that SAs are introduced into mammalian milks.

### Structure of SAs

Although the three-dimensional structure of BSA has not been determined, the structure of HSA, with which BSA shares 75% sequence identity, has been well characterized for
the apo protein as well as for a variety of complexes with a variety of long-chain fatty acids and other more compact hydrophobic molecules. The structure of HSA complexed with the anaesthetic halothane (C$_2$F$_3$Cl$_2$Br) and myristic acid (CH$_3$(CH$_2$)$_{12}$COOH) is shown in Figure 6.4.

The structure of HSA comprises three structurally homologous domains, each of just under 200 residues, denoted I–III (Curry et al., 1998) and involving residues 5–196, 197–383 and 384–582 respectively. Each domain has two sub-domains, each of ≈100 residues, denoted A and B. The structure lacks β-strands and is predominantly (68%) α-helical, with several lengthy loops connecting the A and B sub-domains. On ligand binding, there is substantial movement of the domains with respect to each other, but the tertiary structure of each domain undergoes only small changes (Curry et al., 1998). Medium- and long-chain fatty acids occupy five distinct

![Figure 6.4](image-url)  
*Figure 6.4*  Structure of HSA complexed with halothane (slate/purple) partially occupying seven distinct sites and myristic acid (yellow/red) fully occupying five distinct sites (PDB code: 1e7c). Domain IA (residues 5–107) is shown in blue; domain IB (residues 108–196) is shown in light blue; domain IIA (residues 197–297) is shown in green; domain IIB (residues 297–383) is shown in light green; domain IIIA (residues 384–497) is shown in red; and domain IIIB (residues 498–582) is shown in light red. The single cysteine, Cys34, is labeled (Bhattacharya et al., 2000). The 17 disulfide bonds, which tie together individual sub-domains, are represented in stick format. Figure drawn with PyMOL (Delano, 2002) (see also Plate 6.4).
sites (dissociation constants 0.05–1 μM) (Spector, 1975), one in domain I, a second between domains I and II and the remaining three in domain III, as characterized by X-ray techniques for HSA. In the case of halothane binding, two sites are located in domain I and five are located in domain II.

A comprehensive study of the binding of 17 distinct drugs to HSA, in the presence and absence of myristate, has been published recently (Ghuman et al., 2005). Whereas the binding of steroids to BSA is influenced by the binding of fatty acids, for HSA, there is much less influence (Watanabe and Sato, 1996). NMR titrations have shown that BSA, like HSA, binds five myristates; four of the five sites appear to be structurally homologous to those identified crystallographically for HSA (Hamilton et al., 1984, 1991; Cistola et al., 1987; Simard et al., 2005). The X-ray structure of equine serum albumin (ESA) is very similar to that of HSA, consistent with ≈75% sequence identity between these two proteins (Ho et al., 1993).

**Effect of temperature on SAs**

Careful DSC measurements of defatted HSA and its binding to short- to medium-chain fatty acids have been made. In the absence of fatty acids, a single sharp endotherm is observed, yielding a midpoint temperature for denaturation, $T_m$, of 64.7°C, consistent with a concerted unfolding. In the presence of fatty acids, the endotherm broadens and there is a steady increase in $T_m$ as the chain length of the fatty acid increases from $n$-butanoate ($T_m = 77.6°C$) to $n$-octanoate ($T_m = 87.2°C$); $T_m$s for $n$-nonanoate and $n$-decanoate are very similar to that for $n$-octanoate.

The short-chain fatty acids formate, acetate and $n$-propionate show evidence for inducing increased stability in HSA through binding of the fatty acids at secondary sites that are inaccessible to the longer-chain fatty acids. For a 30 mg/mL ($≈0.5$ mM) solution of HSA at pH 7.0, the concentration of fatty acid at which maximum stability of HSA is achieved decreases from more than 2900 mM for formate to less than 15 mM for $n$-dodecanoate (Shrake et al., 2006).

For the native protein, DSC data could be fitted to a two-state model with about seven more or less equivalent binding sites for $n$-decanoate. This number may be contrasted with the value of five observed by X-ray and NMR methods for the binding of myristate (tetradecanoate acid) to HSA and also to BSA (see above) (Simard et al., 2005).

**Effect of pressure on SAs**

BSA, despite the unpaired cysteine, is relatively stable to high pressures (800 MPa) (Lopez-Fandino, 2006b). BSA undergoes substantial secondary structure changes but, unlike β-Lg, these changes are reversible. It appears that the large number of disulfide bonds protects the hydrophobic core of the protein, including the largely buried Cys34 in sub-domain IA, which is held together by three disulfide bonds (Lopez-Fandino, 2006b). The effects of binding partners, such as fatty acids or other whey components, such as β-Lg, on the structure and stability of BSA at high pressure remain uncharacterized.
Combined pressure/temperature infrared studies of the amide vibrational modes of ESA have shown very recently that high pressure (400 MPa) can convert an intermolecular β-sheet aggregate formed by heating ESA to 60°C at 0.1 MPa (i.e. ambient pressure) to a disordered structure, which reverts to the native structure upon the release of pressure. The activation volume of $\approx +92$ mL/mol and the partial molar volume difference between the native and heat-denatured states ($\Delta V_{N-HA} = +32$ mL/mol) are consistent with decreasing stability of the heat-denatured intermolecular β-sheet with increasing pressure (Okuno et al., 2007).

**Effect of chemical denaturants on SAs**

The denaturation of BSA and HSA by urea and GdmCl has been extensively studied (e.g. Lapanje and Skerjanc, 1974; Khan et al., 1987; Guo and Qu, 2006). In the presence of fatty acids and other molecules, especially molecules that bind to domain III (e.g. diazepam), denaturation of BSA by urea changes from a three-step process to a two-step process, indicating that the initial denaturation involves changes in the tertiary and secondary structure of domain III (Ahmad and Qasim, 1995; Tayyab et al., 2000). For HSA, denaturation appears to be an intrinsically two-step process with fatty acids converting denaturation to a one-step process (Muzammil et al., 2000; Shrake et al., 2006). In both cases, binding of ligands to BSA or HSA stabilizes the protein against urea-induced denaturation. There is evidence that the urea-induced denatured state and that induced by high pressure are similar, at least for HSA (Tanaka et al., 1997).

In the presence of cations, such as guanidinium and cetylpyridinium salts, domain III of SA is again the most susceptible to denaturation (Ahmad et al., 2005; Sun et al., 2005). Consistent with the high helical content of SA, perfluorinated alcohols and alcohols with bulky hydrophobic heads stabilize HSA (and presumably BSA) against denaturation by both urea and GdmCl (Kumar et al., 2005).

The stability of HSA in the presence of polyethylene glycols (PEGs) has also been examined (Farruggia et al., 1999); low-molecular-weight PEG affects ionization of surface tyrosines and high-molecular-weight PEGs lower the thermal transition temperatures.

**Immunoglobulins**

The immunoglobulin (Ig) proteins form a diverse family whose members, when in milk, protect the gut mucosa against pathogenic micro-organisms. In bovine milk, the predominant species of Ig proteins are members of the IgG subfamily, in particular IgG1. Colostrum contains 40–300 times the concentration of IgG proteins than milk does; their role is to confer passive immunity to the neonate while its own immune system is developing (Gapper et al., 2007).

IgG proteins have multiple functions, including complement activation, bacterial opsonization (rendering bacterial cells susceptible to immune response) and agglutination. They inactivate bacteria by binding to specific sites on the bacterial surface.
Given the significance attributed to bovine milk and milk products in human nutrition and health, it is important to note that there are significant differences in the levels of the various subfamilies of Igs in milks from different species. Human colostrum and milk contain relatively low levels of the IgG subfamily compared with bovine milk; the reverse occurs for the IgA subfamily. The properties and the accurate quantitation of bovine Ig proteins have been reviewed recently and in detail (Gapper et al., 2007).

Structure of IgG

The structure of IgG is illustrated schematically in Figure 6.5a and as revealed by X-ray techniques in Figure 6.5b. Both the heavy chain and the light chain are predominantly β-sheet structures. Disulfide bridges link pairs of molecules, as well as the heavy chain to the light chain. The protein is generally glycosylated at a number of sites. However, the actual structure is much less tidy than the schematic Y-shaped figure; in particular, the disulfide bonds are at the base of the light chain–heavy chain associations.

Effects of temperature, pressure and chemical denaturants on Ig structure and stability

The response of bovine IgG (isoform not specified) to temperature and chemical denaturants, urea and GdmCl, has been reported (Ye et al., 2005). Thermal denaturation and thermal denaturation in the presence of denaturants was irreversible, producing, via a series of steps, an incompletely unfolded aggregate. Isothermal chemical denaturation produced, also by a series of steps, a completely unfolded random coil state (Ye et al., 2005). The response of IgG to high pressure (200–700 MPa) in the presence of the kosmotrope sucrose has also been reported (Zhang et al., 1998).

A comparative study of the thermal denaturation of bovine IgG, IgA and IgM has been published, with stability in the order just given (Mainer et al., 1997). As retention of immunological properties under standard milk processing conditions was of interest, the activity of the heat-treated protein was determined by an immunological assay using antibodies raised against these Ig proteins. Relatively recently, the response of IgG to pulsed electric fields (and to heat) was reported (Li et al., 2005). Little change in secondary structure or in immunoactivity was reported for samples subjected to a pulsed electric field of ≈41 kV/cm.

Lactoferrin

Lactoferrin (Lf) is a monomeric, globular, Fe³⁺-binding glycoprotein comprising ≈680 amino acids, giving a molecular mass of ≈80 kDa. It is a member of the transferrin family but, unlike the eponymous protein, to date there is no strong evidence
Figure 6.5  (a) Schematic of the general structure of Igs. Reproduced from Gapper et al. (2007) with permission. The different classes are distinguished by the constant or Fc regions of the heavy and light chains. (b) The X-ray structure of the human IgG1 molecule (PDB code: 1hzh). The heavy chains are in blue and green; the light chains are in magenta and pink. The asparagine N-linked glycan is shown in stick representation. The lack of two-fold symmetry indicates the extreme flexibility of the domains with respect to one another and the consequent sensitivity to crystal-packing effects. Disulfide bridges are shown as spheres. Each domain has a disulfide bridge joining the two sheets; additional disulfide bridges link the two heavy chains and the light chains to the heavy chains. The N-terminus of each chain is at the top left and top right of the diagram; the C-termini of the heavy chains are at the base of the molecule. Structures (e.g. PDB code: 1wej, 3hfm) where antigens are bound at the light chain–heavy chain interface indicate that binding of antigen occurs across the top of the molecule with little embedding of antigen between the domains, contrary to the mode implied by (a). Figure drawn with PyMOL (Delano, 2002) (see also Plate 6.5).
that Lf is involved in iron transport or metabolism under normal circumstances. Indeed, the protein is only lightly loaded with iron(III) in milk, allowing it to perform a major bacteriostatic role by sequestering iron(III) despite bacteria producing iron(III)-sequestering agents (siderophores) that have affinities for iron(III) many orders of magnitude higher than that of Lf. However, Lf is a multifunctional protein, with evidence to suggest that it, and especially its N-terminal arginine-rich fragments, called lactoferricin(s), obtained by pepsin hydrolysis of the entire protein, have active antimicrobial activity as well as activity as antiviral and antiparasitic agents (Strom et al., 2000, 2002).

Accordingly, commercial applications utilizing bovine Lf and its partially digested peptides are appearing as nutraceuticals in infant formulas, health supplements, oral care products and animal feeds. In addition, reputed antioxidant properties are being utilized in cosmetics (van Hooijdonk and Steijns, 2002). For a review on the remarkable properties of Lf and their commercial applications, see Brock (2002). The protein is synthesized in the mammary gland, but is also found in other exocrine fluids besides milk. Bovine Lf is sometimes used as a supplement in bovine-milk-based infant formulas (to offset the lower abundance relative to that found in human milk) and is potentially useful as a constituent in functional foods, with marked effects on bone-cell activity (Cornish et al., 2004).

**Structure of bovine Lf**

Bovine Lf has a tertiary structure (Moore et al., 1997) that is very similar to that of the Lfs of other species determined so far: human (Anderson et al., 1987, 1989; Haridas et al., 1995), buffalo (Karthikeyan et al., 1999) and horse (Sharma et al., 1999). All Lfs and transferrins contain two lobes, which share internal homology (≈40% sequence identity between the N- and C-terminal lobes) and a common fold (see Figure 6.6). The homology between lobes for a given species is less than that between corresponding lobes from different species, indicating that the gene duplication event is of ancient origin. Each lobe contains two α/β domains divided by a cleft that incorporates an iron-binding site. Huge structural changes accompany iron binding (each lobe closes over the iron(III), encapsulating a synergistic bicarbonate anion).

Notwithstanding these structural changes, for the apo protein, at least for human Lf, there is little difference in free energy between the open and closed forms (indeed the structure of human apo-Lf has one lobe open and the other lobe closed). This delicate balance is achieved by both open and closed conformations of the protein having, remarkably, the same surface area exposed to solvent. The same applies, but by a different structural mechanism, to the open and closed forms of the N-terminal recombinantly produced half molecule of human Lf (Jameson et al., 1998, 1999, 2002b). Like β-Lg, holo-Lf undergoes a pH-dependent conformational change (in this case at as low as pH ≈ 3) that releases the bound ferric ion; the structurally related, but genetically and functionally distinct, serum transferrin releases its cargo at significantly higher pH (pH ≈ 5.5) (Baker and Baker, 2004).
Figure 6.6  Structure of Lf. (a) Human apo-Lf (PDB code: 1cb6), showing the domain structure. The N lobe (blue for sub-domain N-I\(_N\) and cyan for sub-domain N-I\(_C\)) is in the open conformation; the C-lobe (magenta for sub-domain C-I\(_N\) and pink for sub-domain C-I\(_C\)), despite no metal ion present, is in the closed conformation. The helix connecting the two lobes is shown in yellow. Metal-binding ligands are shown as spheres. (b) Bovine holo-(Fe(III))-Lf (PDB code: 1blf; the human analog, 1b0l, is structurally very similar). The polypeptide is rainbow colored, blue at the N-terminus to red at the C-terminus, to highlight the manner in which the sub-domains N-I\(_N\) and C-I\(_N\) are formed from residues \(\approx 1-90\) and \(\approx 250-320\) (N-I\(_N\)) and \(\approx 350-440\) and \(\approx 600-680\) (C-I\(_N\)); for reference, these sub-domains are shown in approximately the same orientation in frames (a) and (b). The iron-binding ligands, including the synergistic bicarbonate ion in (b), are shown in stick form, the iron atom is shown as a red sphere and the cysteines are shown as orange spheres (16 Cys forming eight disulfides for human Lf and 34 Cys forming 17 disulfides for bovine Lf). Loops are not smoothed. Figure drawn with PyMOL (Delano, 2002) (see also Plate 6.6).

Effects of temperature, pressure and chemical denaturants on Lf structure and stability

As a minor component of milk proteins, Lf seems largely to have escaped detailed study of its intrinsic response to temperature, pressure and chemical denaturants or of its influence, if any, on the stability of either itself, or other whey proteins, in the presence of other whey proteins. Calcium ions have been reported to bind to bovine...
Lf, probably to the sialic acid groups of the asparagine N-linked glycan, with micromolar dissociation constants; both the apo and holo forms of calcium-bound bovine Lf are more stable to heat and chemical denaturants (Rossi et al., 2002). The calcium-bound forms appear to reduce the Lf-induced release of lipopolysaccharide moieties from bacterial membranes (Rossi et al., 2002).

The heat stability of bovine Lf in isolation (Sánchez et al., 1992) and in association with the major bovine whey proteins, β-Lg (weak association), α-La (no detectable association) and BSA (weak 1:1 association) (Lampreave et al., 1990) has also been studied. The stability of bovine Lf is such that standard pasteurization conditions (but not UHT treatment) are likely to have little effect on its structure and properties, especially its immunological properties (Oria et al., 1993).

Conclusions

Three-dimensional structure determines physiological function and also functionality of whey proteins in their varied applications, both in whole milk and in whey itself. In this review, we have tried to look beyond individual proteins to see what structural features may be important to protein–protein interactions under stresses of temperature, pressure and chemical denaturants. In particular, we have attempted to uncover more recent results that challenge existing paradigms or offer new perspectives. In general, changes that are brought about as a consequence of pH, heat, pressure, chaotropes, etc., shift equilibrium points (e.g. between monomer and dimer, native and unfolded states, etc.) and it is only when a new covalent bond is formed (or removed) that there is irreversible change.

One key interaction between like and unlike proteins arises from disulfide bond interchange, generally facilitated by a single cysteine residue. For this reason, we have focused, especially in the figures, on the observed locations of cysteines. However, detailed structural information, at a level comparable with that of individual partners, is generally lacking on intermolecular assemblies, even when only pairwise associations are formed and isolated.

Whereas considerable attention has been directed to understanding interactions among components of milk and of whey, relatively little attention has been directed to date on the redox state and redox changes during the storage and processing of milk proteins and their effects on milk processing and the flavors of milk-derived products. This remark originates from recent studies on blood sera, where SAs play key roles in redox states and protection against reactive oxygen species. It is important, then, that these structures and functional states are characterized and understood, especially if milk processing, particularly relatively new methods such as pressure treatment, results in refolded proteins that may have highly undesirable properties, such as amyloidogenesis.

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Structure and stability of whey proteins


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Abstract

High-pressure processing (HPP) has been reported to bring about particular changes in the molecular structure of proteins and thus gives rise to new properties that are inaccessible via conventional methods of protein modification. The aim of this chapter is to provide relevant current knowledge, mainly focusing on how HPP can affect the structural conformational, unfolding and aggregation of whey proteins and their interactions with other proteins. The denaturation and aggregation pathways and mechanisms underlying pressure-induced denaturation and aggregation of β-lactoglobulin, α-lactalbumin and bovine serum albumin and their mixtures in different systems are explained. Heat and HPP have been reported to have some similar effects and some different effects on proteins. In some instances, pressure-induced changes in various proteins are compared and contrasted with the heat-induced changes in milk proteins.

Introduction

Processing treatments such as heat and high pressure are normally applied in the food industry for the purpose of microbial destruction or shelf-life extension, or to achieve
High-pressure-induced interactions involving whey proteins

It is well known that the heat-induced interactions of milk proteins have a marked impact on the functionality of the final products and such interactions are of considerable commercial importance in the dairy and food industries. Therefore, considerable research efforts have been directed to studying the detailed pathways/mechanisms of heat-induced functionality and the effects of heat treatments on milk proteins (denaturation, aggregation and gelation of whey proteins) and protein–protein interactions, including interactions of caseins and whey proteins, have been studied in great detail over 5–6 decades (e.g. Haque et al., 1987; Haque and Kinsella, 1988; Hill, 1989; Noh et al., 1989; Matsudomi et al., 1992, 1993, 1994; Hines and Foegeding, 1993; McSwiney et al., 1994a, 1994b; Gezimati et al., 1996, 1997; Havea et al., 1998, 2000, 2001, 2004; Manderson et al., 1998; Schokker et al., 1999, 2000; Hong and Creamer, 2002; Anema and Li, 2003a, 2003b; Cho et al., 2003; Livney et al., 2003; Livney and Dalgleish, 2004; Patel et al., 2004, 2006, 2008) and the subject has often been reviewed (e.g. Mulvihill and Donovan, 1987; de Wit, 1990; Singh and Creamer, 1992; Jelen and Rattray, 1995; Singh, 1995; de la Fuente et al., 2002; O’Connell and Fox, 2003; Singh and Havea, 2003; Singh, 2004; Condé et al., 2007b).

Although thermal processing is effective, economical and readily available, in many cases, it has undesirable effects on the sensory and nutritional qualities of food (Balny and Masson, 1993; Trujillo et al., 2002). More than a century ago, pioneering work by Hite (1899) showed the potential of high-pressure processing (HPP) as a non-thermal (alternative) preservation process and Bridgman (1914) demonstrated the effects of HPP on the denaturation and functional properties of egg proteins. However, it was not until 1990 that equipment advances and growing consumer demand for minimally processed, high-quality, nutritious and safe foods led to considerable research interest in HPP technology (Hayashi, 1988; Farr, 1990; Hoover, 1993; Datta and Deeth, 1999, 2003; Needs et al., 2000a, 2000b; Velazquez de la Cruz et al., 2002).

The recent interest in the HPP of food materials as an alternative to or in addition to temperature treatment led to many fundamental studies on the pressure behavior of proteins. It is also evident from the literature that, in many cases, heat treatment and high-pressure treatment have different effects on different milk proteins, suggesting that HPP has the potential for both preservation and modification of the structure of proteins, alteration to their functional properties and the creation of value-added products (see Ohmiya et al., 1989; López-Fandiño et al., 1996; García-Risco et al., 1998, 2000; Datta and Deeth, 1999, 2003; Patel et al., 2004, 2005, 2006; López-Fandiño, 2006a, 2006b).

Today, HPP is considered to be a possible alternative to heat treatment in many cases and has reached the consumer in a variety of products, such as high-pressure-treated fresh fruit jams, jellies, juices, salad dressings, rice, cakes and guacamole (Farr, 1990; Cheftel, 1992; Earnshaw, 1992; Hoover, 1993; López-Fandiño, 2006a, 2006b), but no high-pressure-treated commercial dairy products are available as yet. However, as there is increased interest in the high-pressure treatment of dairy products, it is currently a major focus of investigation and this subject has been extensively reviewed (see Heremans, 1982; Weber and Drickamer, 1983; Balny et al., 1989, 1992; Masson, 1992; Balny and Masson, 1993; Mozhaev et al., 1994; Johnston, 1995; Heremans et al., 1997;
Messens et al., 1997; Balci and Wilbey, 1999; Datta and Deeth, 1999, 2003; Farkas and Hoover, 2000; Boonyaratankornkit et al., 2002; Huppertz et al., 2002; Lullien-Pellerin and Balny, 2002; Royer, 2002; Trujillo et al., 2002; Claeyes et al., 2003; Huppertz et al., 2006a, 2006b; López-Fandiño, 2006a, 2006b; Considine et al., 2007b; Rastogi et al., 2007; Patel et al., 2008).

As whey proteins are widely used as nutritional and functional ingredients in the food industry, there is obvious interest in studying the effects of HPP on whey proteins and their interactions with other proteins. The specific detailed structural aspects and some fundamental aspects related to the structural stability of β-lactoglobulin (β-LG), α-lactalbumin (α-LA), bovine serum albumin (BSA), immunoglobulins (Igs) and lactoferrin (LF) have been dealt with in Chapter 6. The present chapter builds on these fundamental aspects and extends the discussion to pathways and mechanisms of pressure-induced denaturation, aggregation and interactions of individual whey proteins and their mixtures in various systems.

**Characterization of heat- and pressure-induced changes to proteins**

Several different methods have been used in the study and characterization of protein denaturation and aggregation, including X-ray crystallography, far-UV circular dichroism (CD), Fourier transform infrared (FTIR) spectroscopy, Fourier transform of Raman spectra and hydrogen/deuterium (H/D) exchange using a nuclear magnetic resonance (NMR) method for determining changes in the secondary structure (Tanaka and Kunugi, 1996; Subirade et al., 1998; Panick et al., 1999; Belloque et al., 2000; Edwards et al., 2002; Ngarize et al., 2004). 1H NMR is a technique that is very sensitive to structural changes and can give structural and dynamic information at an atomic level.

Methods such as intrinsic and extrinsic fluorescence and induced CD have been widely used for tertiary structure determination (Pearce, 1975; Masson, 1992; Dufour et al., 1994; Heremans et al., 1997; Narayan and Berliner, 1997; Kelly and Price, 2000; Collino et al., 2003; Kontopidis et al., 2004). Scattering methods such as small-angle X-ray scattering (SAXS) and light scattering (LS) have also been used recently (Pessen et al., 1989; Holt et al., 2003). In particular, SEC–MALLS (size exclusion chromatography coupled to multi-angle laser light scattering) can provide excellent information on aggregate size (Schokker et al., 1999). SEC–MALLS combined with electrophoretic techniques is a powerful tool for characterizing intermediates and aggregates formed during heating (Schokker et al., 1999, 2000). The colorimetric methods used in the determination of protein sulfhydryl groups in milk (Owusu-Apenten, 2005) and other methods used in the study of protein aggregation have been reviewed (Wang, 1999, 2005; De la Fuente et al., 2002; Considine et al. 2007b).

It is evident that less commonly used polyacrylamide gel electrophoresis (PAGE) separation in different environments (e.g. sodium dodecyl sulfate [SDS] solution, in the presence or absence of a disulfide bond reducing agent) allows the separation and differentiation of the various polypeptide chains. SDS-PAGE dissociates the processing-induced non-covalent bonds and leaves the covalent bonds intact, whereas
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native- or alkaline-PAGE generally separates the whey proteins with hydrophobic and disulfide bonds intact; therefore, it is possible to show whether the processing-induced protein aggregates are hydrophobically linked aggregates, reducible disulfide-cross-linked aggregates etc. in heated solutions of pure β-LG (Manderson et al., 1998) or commercial whey protein concentrate (WPC) solutions (Havea et al., 1998, 2000, 2001). The same techniques have been used successfully to examine the pressure-induced aggregation of various dairy proteins, revealing the subtle and not-so-subtle differences (Patel et al., 2004, 2005, 2006), which are discussed in detail in the later part of this chapter.

When heat-treated or pressure-treated samples were analyzed using various PAGE techniques, many changes in the PAGE patterns of these samples were noted (McSwiney et al., 1994b; Gezimati et al., 1996, 1997; Havea, 1998; Havea et al., 1998, 2000, 2001, 2002, 2004; Manderson et al., 1998; Cho et al., 2003; Patel et al., 2004, 2005, 2006; Considine et al., 2005a, 2005b, 2007a). It became necessary to identify each of these changes and aggregates observed in different PAGE environments. For simplification, some specific nomenclature had to be formulated, e.g. “native monomer,” “native dimer,” “SDS monomer,” “SDS dimer,” etc. (see Table 7.1).

Two-dimensional (2D) PAGE (2D native- and then SDS-PAGE, and 2D SDS- and then reduced SDS-PAGE) can be applied to further characterize various intermediate species of protein aggregates and the high-molecular-weight aggregates formed as a consequence of heat or pressure treatment. A combination of two PAGE techniques can be an even more powerful technique; it can determine the composition of the protein aggregates and/or the types of bonds by which the protein aggregates are held together in the heat-treated (Havea et al., 1998, 2000, 2001; Manderson et al., 1998) or pressure-treated (Patel et al., 2004, 2005, 2006) samples.

In the 2D native and then SDS-PAGE procedure, a sample containing protein complexes is analyzed using one-dimensional (1D) PAGE in a Tris HCl buffer at pH 8.7,

Table 7.1 Nomenclature of different forms of proteins and protein aggregates analyzed using various electrophoretic techniques

<table>
<thead>
<tr>
<th>Nomenclature</th>
<th>Applicable to</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Native</td>
<td>All</td>
<td>Has all the characteristics of native proteins</td>
</tr>
<tr>
<td>Native dimer</td>
<td>β-LG</td>
<td>Normal state of native β-LG between pH 4 and pH 7 (at 30°C and at 0.05 M NaCl)</td>
</tr>
<tr>
<td>Native-like</td>
<td>All</td>
<td>Behaves like native protein on alkaline (native)-PAGE</td>
</tr>
<tr>
<td>Non-native monomer</td>
<td>All</td>
<td>Behaves like native protein on SDS-PAGE, but not on alkaline-PAGE</td>
</tr>
<tr>
<td>SDS monomer</td>
<td>All</td>
<td>Has mobility on SDS-PAGE close to that expected from molecular monomer</td>
</tr>
<tr>
<td>SDS dimer, trimer, etc.</td>
<td>All</td>
<td>Has mobility on SDS-PAGE close to that expected from molecular monomer</td>
</tr>
<tr>
<td>Apo</td>
<td>α-LA</td>
<td>Deficient in calcium (or other divalent cation)</td>
</tr>
<tr>
<td>Holo</td>
<td>α-LA</td>
<td>Same as native α-LA</td>
</tr>
</tbody>
</table>
called alkaline- or native-PAGE. The proteins or protein complexes separated on native-PAGE consist of all complexes including covalent (e.g. disulfide bonds) and non-covalent (e.g. hydrophobic)-bonded aggregates. The 1D native-PAGE gel strip with its separated protein bands is transferred to SDS-PAGE in the second dimension (i.e. transferred into a dissociating environment). Once the proteins in the strip are partially equilibrated with the SDS to form SDS-protein complexes, they are elec
trophoresed into a new (SDS) environment in a second dimension SDS-PAGE. SDS-
PAGE in the second dimension dissociates non-covalent bonds (mainly hydrophobic aggregates) from the non-reduced native-PAGE gel strip, whereas the covalent bonds (disulfide bonds) remain unaffected.

The other type of 2D PAGE is SDS- and then reduced SDS-PAGE. This type of 2D PAGE technique separates the initial mixture of proteins using SDS-PAGE with all the native and process-induced disulfide bonds intact. These separated proteins and protein aggregates are treated with a disulfide bond reducing agent, 2-mercaptothanol (2-ME), while still in the gel strip. This gel strip is then used as the sample source for the second dimension PAGE analysis. SDS-PAGE in the second dimension separates the proteins as the reduced SDS-protein species. Thus the components of each of the various disulfide-bonded aggregates can be identified (for example, see Figures 7.3 and 7.4; refer to Patel et al., 2004, 2005, 2006 and Considine et al., 2007b for detailed descriptions of these PAGE methods).

**Effects of high pressure on milk proteins**

The native three-dimensional structure of a protein is maintained by a variety of non-
covalent interactions (such as hydrogen bonding, electrostatic, van der Waals’ and hydrophobic interactions) between amino acid residues within the polypeptide chain and between residues and solvent molecules (Singh, 1995). Three-dimensional structure has a very important role to play in the stability and functional properties of a protein. Excellent reviews describing the molecular basis of whey protein functionalities (Holt, 2000), the structure–function relationship (Qi et al., 2001) and the possible usefulness of knowledge of the three-dimensional structure of milk proteins (Sawyer et al., 2002) are available. The number and the position of free thiols and disulfide bonds present in the structure of different proteins are greatly significant to their stability, denaturation, interactions and functional properties (Holt, 2000; Sawyer et al., 2002).

Applications of high pressure cause proteins to lose their native three-dimensional structure and lead to denaturation and aggregation of whey proteins and/or their interactions with each other (whey protein–whey protein interactions) or with the caseins (casein–whey protein interactions). According to Le Chatelier-Braun’s principle, under pressure, reactions with a negative volume change are enhanced and reactions with a positive volume change are suppressed (Buchheim and Prokopek, 1992; Johnston, 1995; Balci and Wilbey, 1999). Thus, the basic thermodynamic approach to pressure-induced changes to proteins is based on the compressibility of molecules and on changes in volume (ΔV) during pressure treatment as, unlike temperature,
High-pressure-induced interactions involving whey proteins

High-pressure-induced interactions involving whey proteins (Balny and Masson, 1993; Gross and Jaenicke, 1994; Heremans and Smeller, 1998; Lullien-Pellerin and Balny, 2002). Changes in the solvation volume are caused mainly by pressure-induced ionization, changes in solvent exposure of amino acid side chains and peptide bonds, and diffusion of water into cavities located in the hydrophobic core of the protein (Heremans, 1982, 1992; Balny and Masson, 1993; Messens et al., 1997; Hendrickx et al., 1998; Claeys et al., 2003).

In contrast to heat treatments, where covalent bonds and non-covalent bonds are affected, it has been reported that HPP at room temperature (=20°C) disrupts only relatively weak bonding such as intramolecular hydrophobic and electrostatic interactions (Balny and Masson, 1993; Silva and Weber, 1993), whereas hydrogen bonds are relatively insensitive to pressure, suggesting that high pressure affects the tertiary (three-dimensional configuration held together mainly by hydrophobic and ionic interactions) and quaternary (the spatial arrangement by non-covalent interactions into a multimeric protein) structures of globular proteins and has little effect on their secondary structure. There are views that covalent bonds are largely insensitive to pressure treatment at relatively low temperature (Hayakawa et al., 1996), which means that the primary protein structure (the amino acid sequence) will remain intact. This partly explains why HPP has been reported to have slightly different effects on protein structure compared with heat treatments.

The effects of high pressure on denaturation, aggregation and interactions of whey proteins have been studied extensively under various conditions and in different systems such as milk (see Felipe et al., 1997; Law et al., 1998; Arias et al., 2000; Hinrichs, 2000; Scollard et al., 2000; Huppertz et al., 2002, 2004a, 2004b; Anema et al., 2005a), WPC (see Van Camp et al., 1997a, 1997b) and whey protein isolate (WPI) (Hinrichs et al., 1996a, 1996b; Hinrichs, 2000; Michel et al., 2001), and using pure whey proteins (Dumay et al., 1994, 1998; Funtenberger et al., 1995; Galazka et al., 1996a; Jegouic et al., 1996; Olsen et al., 1999).

It has been reported that pressure-induced reactions of whey proteins lead to unfolding of monomeric proteins, aggregation and gelation (Zipp and Kautzmann, 1973; Heremans, 1982; Weber and Drickamer, 1983; Silva and Weber, 1993; Van Camp and Huyghebaert, 1995a, 1995b; Van Camp et al., 1997a, 1997b; Balci and Wilbey, 1999; Tedford et al., 1999a, 1999b; Huppertz et al., 2002; Fertsch et al., 2003), by reformation of intra- and intermolecular bonds within or between the molecules, linked by hydrophobic interactions and disulfide bridges (see Cheftel, 1992; Masson, 1992; Hoover, 1993; Galazka et al., 1996a; Messens et al., 1997; Trujillo et al., 2002), depending on the type of protein, protein concentration, pH, ionic strength, applied pressure, pressurizing temperature and duration of the pressure treatment, etc. (Masson, 1992; Messens et al., 1997; Fertsch et al., 2003; Huppertz et al., 2004a).

Also, different pressurizing temperatures may have different effects on the denaturation and aggregation of proteins (see Huppertz et al., 2004a; Patel, 2007), because of combined effects of pressure and temperature, which can have different effects on the interactions that maintain protein structures. Many studies related to the combined effects of pressure and temperature on milk proteins have been published (e.g. Tedford et al., 1999b; Huppertz et al., 2004a; Patel, 2007). However, the present
review focuses mainly on the HPP-induced changes in milk proteins that occur at ambient temperature, unless specified otherwise.

Selected reports on the effects of high pressure on β-LG, α-LA, BSA or their combinations in various systems are summarized in the following sections.

Denaturation and aggregation of pure whey proteins in model systems

β-Lactoglobulin (β-LG)

Many reports have suggested that β-LG is the most sensitive of the major whey proteins to high pressure and that it dominates the pressure-induced denaturation, aggregation and gelation of the whey protein system (Van Camp and Huyghebaert, 1995a, 1995b; Stapelfeldt et al., 1996; Van Camp et al., 1996, 1997a, 1997b; Kanno et al., 1998; Belloque et al., 2000; Patel et al., 2004, 2005; López-Fandiño, 2006a, 2006b; Considine et al., 2007b). Therefore, the majority of studies have concentrated on the effects of high pressure on β-LG in order to gain insight into the mechanisms of unfolding and aggregation that occur during pressurization or after pressure release.

Different effects of high pressure on proteins have been observed when the samples are analyzed under high pressure ("in situ" analysis) and when analyzed after pressure release. With increasing pressure, protein molecules undergo a sequence of conformational changes because of alterations in stabilizing interactions (Johnston et al., 1992). During the pressure release phase and after pressure treatment, new intermolecular interactions are formed and the proteins may be newly structured (Fertsch et al., 2003). The majority of the reports included in the present review deal with the interactions of proteins after pressure release.

At relatively low pressures (50 MPa), analysis of thiol reactivity (Møller et al., 1998; Stapelfeldt et al., 1999) and NMR studies (Tanaka and Kunugi, 1996) suggested the existence of a "pre-denatured" state of β-LG. This "pre-denatured" form corresponded to a not-completelyunfolded structure, which preceded reversible denaturation. Belloque et al. (2000), using 1H NMR, showed that the degree of deuterium exchange was very small at 100 MPa and that there were no variations in the resonances belonging to the strongly bonded "core" of β-LG.

These observations might suggest that the regions of β-LG affected by the "pre-denatured" state are likely to be different from the "core," as the core was still tight and remained unaltered at 100 MPa. Whereas pressures ranging between 0 and 140 MPa did not affect β-sheets (Subirade et al., 1998), the reactivity of the free sulfydryl group of β-LG increased with pressure up to 150 MPa (Tanaka et al., 1996a, 1996b). These results suggested that, in spite of having a similar overall conformation, the architectures of β-LG before and after dynamic high pressure were stabilized by slightly different interactions (Subirade et al., 1998).

Pressure-induced unfolding and refolding as studied by deuterium exchange (Belloque et al., 2000) further demonstrated that the conformational flexibility of β-LG increased at 200 MPa. It was reported that, even though the core of β-LG
was highly flexible at 400 MPa, its structure was found to be identical to the native structure after equilibration back to atmospheric pressure. It was also suggested that pressure-induced aggregates are formed by β-LG molecules maintaining most of their structure, and that the intermolecular disulfide bonds, formed by sulfydryl–disulfide exchange reactions, are likely to involve Cys66–Cys160 rather than Cys106–Cys119 (Belloque et al., 2000), which is different from the possibility of thiol–disulfide interchange reactions as discussed by Considine et al. (2007b).

In addition, it was reported that the β-LG variants A and B could be distinguished in a $^1$H NMR spectrum of a solution made with the AB mixed variant by the differences in chemical shifts of Met107 and Cys106 and that, under pressure, the core of β-LG A seemed to unfold faster than that of β-LG B and the structural recovery of the core was full for both variants (Belloque et al., 2000). These results are somewhat in agreement with those of Iametti et al. (1997), who found that only 10% of the structure was lost at 600 or 900 MPa.

The pressure denaturation of β-LG was believed to be a simple two-step mechanism until Jonas and Jonas (1994) reported pressure-induced pre-denaturation transitions and thus demonstrated that the pressure denaturation of β-LG could be a stepwise process. A few years later, Stapelfeldt and Skibsted (1999) proposed a three-step denaturation process and recently Considine et al. (2005b) published a three-stage model of the pressure denaturation of β-LG (Figure 7.1; for a detailed description, refer to Considine et al., 2007b). It has also been reported that addition of hydrophobic ligands such as all-trans-retinol, palmitic acid, SDS and 8-anilino-1-naphthalenesulfonate (ANS) to β-LG solution before pressure treatment (see Figure 7.1) affects the pathways of denaturation (Considine et al., 2005b, 2007b).

The overall major changes that occur to the structure of β-LG include monomerization of the dimeric state (Iametti et al., 1997), a decrease in α-helix and β-sheet content (Hayakawa et al., 1996; Panick et al., 1999) and irreversible changes involving the formation of intermolecular disulfide bonds (Funtenberger et al., 1997; Iametti et al., 1997; López-Fandiño et al., 1997; Møller et al., 1998). In the pressure-induced mechanism proposed (Iametti et al., 1997), release of monomers represents one of the earliest events, whereas association of transiently modified monomers stabilizes the denatured forms of the protein. In addition, it has been reported that inter- and intramolecular reactions of sulfydryl groups can occur (Tanaka et al., 1996a, 1996b),

![Figure 7.1](image-url)
leading to the formation of new disulfide bonds through sulfhydryl-disulfide inter-
change reactions (Funtenberger et al., 1997) when samples are pressure treated at 
450 MPa. At 800 MPa, most of the β-LG present in the system becomes involved in 
hydrophobic and disulfide-linked aggregates (Patel, 2007); this behavior is quite simi-

Moreover, it has been reported that factors such as protein concentration (Dumay et al., 
1994), pH, ionic strength, type and molarity of the buffer used for preparation of 
the protein solutions (Funtenberger et al., 1995; Cheftel and Dumay, 1996), pressure 
intensity, pressurizing time and pressurizing temperature (Yang et al., 2001; 
Patel, 2007), and binding of hydrophobic ligands (Considine et al., 2005b, 2007a) 
and small molecules such as sucrose (Dumay et al., 1994) can affect the pressure-induced 
denaturation and aggregation of β-LG. It has been found that high-pressure-
induced denaturation is partially reversible at lower (2.5%) protein concentration but 
that the denaturation is irreversible and that aggregation occurs at higher (5.0%) con-
centration (Dumay et al., 1994).

The progressive formation of intermolecular disulfide-bonded dimers to hex-
amers or higher polymers of β-LG (pH 7.0) has been reported to be a function of 
the pressure level and of the buffer type and molarity (Funtenberger et al., 1995). 
It was suggested that high pressure induced the formation of intermolecular disulfide 
bonds, especially at neutral pH. When the combined effects of pressure, temperature 
and time were evaluated, the pressure intensity was found to have major effects on 
the structure of β-LG (Aouzelleg et al., 2004), which is somewhat different from the 
finding that the combined effects of pressure intensity and temperature have the most 
effect on the denaturation of β-LG (Patel, 2007).

Different aspects of the pressure-induced unfolding and aggregation of β-LG have 
also been reviewed in detail (see López-Fandiño, 2006b; Considine et al., 2007b), 
including the effects of high pressure on the functional properties of β-LG (López-
Fandiño, 2006b).

**α-Lactalbumin (α-LA)**

Several studies reported that, compared with β-LG, α-LA is resistant to pressure 
denaturation (López-Fandiño et al., 1996; Tanaka and Kunugi, 1996; Scollard et al., 
2000; Huppertz et al., 2004a; Patel et al., 2004, 2006). A comparison of pressure-
induced changes of two major whey proteins, α-LA and β-LG, at neutral pH showed 
that the reversible unfolding to a molten globule state of α-LA begins at 200 MPa 
and loss of native structure becomes irreversible only beyond 400 MPa, as compared 
with 50 and 150 MPa respectively for β-LG (Tanaka and Kunugi, 1996; Tanaka et al., 
1996c; Stapelfeldt and Skibsted, 1999; McGuffey et al., 2005).

These different behaviors of the proteins were variously explained, including the dif-
fferences in their secondary structures (which lead to a higher effective hydrophobicity 
in β-LG) and/or in the number of disulfide bonds (four in α-LA and two in β-LG) 
and also the Ca$^{2+}$ binding sites (Tanaka and Kunugi, 1996). In fact, the binding of 
calcium is reported to remarkably stabilize α-LA to pressure, by a 200 MPa increase 
in the pressure value at which denaturation occurs (Dzwolak et al., 1999; Hosseini-nia
et al., 2002). This observation is partly supported by the finding that there was a difference in $\Delta V$ values between apo- and holo-$\alpha$-LA (Kobashigawa et al., 1999).

Fluorescent measurement of dansylated (prepared at atmospheric pressure) proteins, especially the energy transfer from the intrinsic tryptophan residue to the dansyl group, showed that the protein structure was deformed by pressure and that the energy transfer mechanisms of the two proteins were differently affected by high pressure, probably reflecting the degree of compactness of their pressure-perturbed structures (Tanaka et al., 1996c). It has been reported that $\alpha$-LA is present in a molten globule state beyond 200 MPa and up to 400 MPa (Jonas, 2002) and that $\alpha$-LA changes its conformation from the molten globule state to the unfolded state without volume changes (Kobashigawa et al., 1999). The volume of $\alpha$-LA changes only at the transition from the native state to the molten globule state (Kobashigawa et al., 1999). Lasselle et al. (2003) reported that heat and high pressure had similar effects, supporting the view that the molten globule state is stabilized by hydrophobic interactions.

In samples of severely heated solutions of $\alpha$-LA, dimers and larger aggregates of $\alpha$-LA were formed (Lyster, 1970; Havea et al., 2001). However, no effects on monomeric $\alpha$-LA were noticeable when pure $\alpha$-LA was pressure treated at 800 MPa (Patel, 2007), except that some changes in the structure of $\alpha$-LA were found when $\alpha$-LA samples were pressure treated at 1000 MPa (Jegouic et al., 1996).

**Bovine serum albumin (BSA)**

BSA has been found to be quite resistant to pressure treatment up to 400 MPa (Hayakawa et al., 1992; López-Fandiño et al., 1996; Patel et al., 2004, 2005, 2006; López-Fandiño, 2006b). Several reports explaining the pressure stability of BSA are available. There are views that pressure-induced changes in the secondary structure of BSA are mainly reversible (Hosseini-nia et al., 2002) and that the greater stability of BSA is probably related to the fact that this molecule, through its 17 intramolecular disulfide bonds and the presence of several separate domains, has an extremely rigid structure (Hayakawa et al., 1992; López-Fandiño et al., 1996; López-Fandiño, 2006b). It is possible that the relatively high number of disulfide linkages in BSA may impede pressure-induced aggregation by protecting the hydrophobic core/groups present inside the molecule from exposure to the solvent (Hosseini-nia et al., 2002).

Ceolín (2000) studied the hydrodynamic behavior of BSA, using a perturbed angular correlation technique, as a function of high pressure up to 410 MPa. It was reported that, at moderate pressure ($\approx 150$ MPa), the BSA molecule suffers structural modifications that produce an increase in the molecular volume and the rotational correlation time of the molecule. However, it may be possible that, unlike $\beta$-LG, the changes in the secondary structure of BSA are largely reversible (López-Fandiño, 2006b). However, processing at 800 MPa was reported to have a substantial effect on the secondary structure of BSA, and BSA was polymerized through disulfide bonding involving the free sulphydryl residue (Galazka et al., 1996b, 1997; Patel, 2007).
**Immunoglobulins (Igs) and lactoferrin (LF)**

Both IgG and LF are more stable under pressure than under heat (Patel et al., 2005, 2006; Carroll et al., 2006; Palmano et al., 2006; Indyk et al., 2008). This finding has great commercial significance for using HPP in the manufacture of nutritional products containing IgG and LF. However, little attention has been paid to this subject. It was reported that the pressure stability of IgG was better in colostrum solutions than in pure IgG solutions (Indyk et al., 2008), suggesting that some other colostrum milk components had protective effects on the denaturation of IgG.

A study on the response of IgG to high pressure (200–700 MPa) in the presence of the cosmotrope sucrose has been reported (Zhang et al., 1998). Recently, Brisson et al. (2007) studied the effects of iron saturation on the thermal aggregation of LF at neutral pH and found that iron saturation markedly increased the thermal stability of LF and decreased aggregation. A similar observation was made by Palmano et al. (2006) for pressure-treated iron-saturated LF solutions.

**Mixtures of α-LA and β-LG**

α-LA does not form aggregates when pressure treated alone at 800 MPa (Patel, 2007), but forms high-molecular-weight disulfide-bonded oligomers at high pressure only in the presence of thiol reducers (Jegouic et al., 1996) and the oligomers of α-LA are stabilized mainly by non-native interchain disulfide bridges. As reported for heat-treated mixtures of α-LA and β-LG (see Havea et al., 2001; Hong and Creamer, 2002), mixed aggregates of denatured α-LA and β-LG were readily formed in pressure-treated whey protein solutions (Jegouic et al., 1997). This observation supports the view that the presence of reactive thiol groups is a prerequisite for pressure-induced denaturation and aggregation of α-LA (Jegouic et al., 1996, 1997). Therefore, in mixtures of α-LA and β-LG, because of its free sulfydryl group, β-LG can induce the oligomerization of α-LA, resulting in the formation of a large heterogeneous population of oligomers (Jegouic et al., 1997; Grinberg and Haertlé, 2000).

Yet another possibility is that the interactions of α-LA and β-LG occur in a hydrophobic environment. However, it has been reported that, under pressure, the volume change of α-LA is much less (Lassalle et al., 2003) than that of β-LG (Royer, 2002) and therefore there is little possibility for such interactions to take place in a hydrophobic environment. This partly explains why α-LA retains most of its structure under high pressure. At very high pressure (e.g. 800 MPa), the irreversible denaturation of α-LA was much less than that of β-LG, which was assigned to the difference in the number of bonds stabilizing the structure of each protein (Hinrichs et al., 1996b; Messens et al., 1997).

**Mixtures of β-LG, α-LA and BSA**

Little has been reported on the effects of high pressure on mixtures of β-LG, α-LA and BSA in pure protein systems. Recently, Patel (2007) reported that, when mixtures of β-LG, α-LA and BSA were pressure treated, there was a somewhat similar
aggregation trend to that reported for heat-treated mixtures of β-LG, α-LA and BSA (Gezimati et al., 1996, 1997; Havea et al., 2001). However, in the pressure-treated samples, it appeared that β-LG, being the most pressure-sensitive whey protein, formed early aggregates, prior to the unfolding of either α-LA or BSA.

Pressure treatment of a ternary mixture of BSA, β-LG and α-LA generated aggregates comprising a mixture of hydrophobically linked and disulfide-linked aggregates (Patel, 2007), whereas when pure BSA solutions or combinations/mixtures of BSA and other whey proteins (e.g. a binary mixture of BSA and α-LA) were pressure treated, almost all the aggregates were disulfide linked and only a small proportion of the aggregates were hydrophobically linked (Patel, 2007). This may be due to structural differences in each of these proteins, and/or the effects of high-pressure treatment on the structure of each of these proteins may be responsible for such differences.

Commercial whey protein solutions

In addition to the above studies using pure protein systems, several studies conducted on heat- and pressure-induced denaturation, aggregation and gelation of whey proteins using commercial whey protein ingredients such as WPC or WPI are available.

Characterization of pressure-treated WPC solutions using 2D PAGE (Patel et al., 2004, 2005) suggested that HPP generated both hydrophobically bonded and disulfide-bonded aggregates consisting of all whey proteins including β-LG, IgG, LF, BSA and α-LA (Figure 7.2), similar to those reported by Havea et al. (1998) for heat-treated WPC solutions. Heated WPC solutions contained 1:1 disulfide-bonded adducts of α-LA and β-LG, which were more obvious at low concentrations. Almost all of the β-LG was incorporated into the aggregates via disulfide bonds and to a lesser extent via hydrophobic interactions (Havea et al., 1998).

However, when similar samples were pressure treated, the β-LG dimer was predominant (Patel et al., 2004, 2005). The detailed characterization and identification of the disulfide-linked aggregates formed in pressure-treated WPC solutions is shown in Figure 7.3, which clearly shows that severe pressure treatment of WPC solutions generated disulfide-bonded dimer, trimer, tetramer, etc. as well as 1:1 complexes of β-LG:α-LA and that higher molecular weight disulfide-linked aggregates consisting BSA, LF, Ig, β-LG and α-LA were formed.

It was found that the sensitivities of each of the whey proteins to heat treatment (Ig > LF > BSA > β-LG B > β-LG A > α-LA) and pressure treatment (β-LG B > β-LG A > IgG > LF > BSA > α-LA) were considerably different (Patel et al., 2004, 2005). Also, high-pressure treatment generated a comparatively greater proportion of smaller aggregates than did heat treatment (Patel et al., 2004). These results confirm and support the view that there are some similarities and some differences between the heat- and high-pressure-induced aggregations of whey proteins. Similar differences were found when whey protein gel formation was induced by either heat treatment or pressure treatment (Van Camp and Huyghebaert, 1995a, 1995b; Van Camp et al., 1996; Dumay et al., 1998).
Figure 7.2 2D PAGE patterns of control and pressure-treated WPC solutions (12% w/v). Native- and then non-reduced SDS-PAGE patterns of (a) the control and (b) a sample pressure treated for 20 min at 800 MPa. Similarly, SDS- and then reduced SDS-PAGE patterns of (c) the control and (d) a sample pressure treated for 20 min at 800 MPa. Gel strips marked as a' and a" represent the sample strip and the stained strip respectively. X2 and X3 are dimer and trimer of β-LG respectively and X4, X5 and X6 are high-molecular-weight aggregates, which were caught up at the beginning of the resolving gel, caught up within the stacking gel and could not enter the gel respectively. For a detailed description, refer Patel et al. 2005. (Reproduced with permission from Patel et al. (2005), copyright 2005 American Chemical Society.)

At high protein concentration (10%), intermolecular interactions and irreversible aggregation are favored (Wong and Heremans, 1988; Dumay et al., 1994). High-pressure treatment of concentrated (80–160 g/kg) β-LG isolate solutions (pH 7.0) prepared in water or various buffers induces β-LG gelation at low temperature (Zasypkin et al., 1996; Dumay et al., 1998). The decreasing solubility (in various dissociating media) of the protein constituents of pressure-induced gels as a function of storage time after pressure release suggests that the aggregation and gelation result from hydrophobic interactions, and also disulfide bonds, and that a progressive build-up of these interactions takes place after pressure release (Dumay et al., 1998).
Hinrichs et al. (1996b) determined orders of reaction of $n = 2.0$ for α-LA and $n = 2.5$ for β-LG in a WPI solution. These reaction rate constants were found to vary slightly at higher protein concentrations (Keim and Hinrichs, 2004). β-LG, α-LA and BSA participate in pressure-induced aggregation and gelation through disulfide bonding. Moreover, it has been reported that the number of stabilizing disulfide bonds directly influences the texture properties of pressure-induced whey protein gels (Keim and Hinrichs, 2004).

The effects of protein concentration, intensity of pressure treatment, holding time and pressurizing temperature on whey protein aggregation in WPC solutions have been investigated (Patel, 2007). The rate of aggregation of the whey proteins increased with an increase in the concentration of protein in the WPC solution and the pressurizing temperature. The combination of low protein concentration, mild pressure treatment (200 MPa) and low pressurizing temperature (20°C) led to minimal loss of native-like and SDS-monomeric β-LG, whereas the combination of high protein concentration, severe pressure treatment (600 MPa) and higher pressurizing temperature (40°C and higher) led to significant loss of both native-like and SDS-monomeric β-LG. The sensitivity of the pressure-resistant whey proteins, such as α-LA and BSA, to aggregation was significantly increased at pressurizing temperatures of 40°C and higher. Self-supporting gels were formed when 8 or 12% (w/v) WPC solutions were pressure treated at 600–800 MPa and 20°C.
Pressure-induced gelation of whey proteins

At protein concentrations sufficiently high for gel formation, WPC was found to produce pressure-induced gels in the pressure range 200–400 MPa (Van Camp and Huyghebaert, 1995a, 1995b; Van Camp et al., 1996). In addition to protein concentration, applied pressure, holding time and pressurizing temperature were found to be major factors in the formation of pressure-induced gels and the properties of the gels (Van Camp and Huyghebaert, 1995a).

As discussed earlier, significant differences in protein denaturation and aggregation induced by heat compared with high pressure have been demonstrated (Heremans et al., 1997; Patel et al., 2004, 2005), which might suggest that the gels produced from whey proteins by high-pressure treatment may have different properties from those made by heat treatment. For example, it was found that heat-set gels, for equal protein concentrations, were firmer than pressure-induced gels (Van Camp and Huyghebaert, 1995a; Zasypkin et al., 1996). High-pressure treatment generated gels with a more porous structure and lower firmness (Van Camp and Huyghebaert, 1995b; Zasypkin et al., 1996; Dumay et al., 1998) that were weaker, less elastic and more exudative than heat-induced gels (Cheftel and Dumay, 1996; Dumay et al., 1998).

In contrast to heat-induced gels, pressure-induced gels of β-LG underwent mechanical and protein solubility changes when stored at 4°C following pressure release, clearly indicating a time-dependent strengthening of protein–protein interactions, probably because the primary aggregates of β-LG further aggregated during storage through hydrophobic interactions and disulfide bonds (Dumay et al., 1998). Also, the WPC gels produced by high pressure (400 MPa for 30 min) at protein concentrations ranging from 110 g/L up to 183 g/L differed significantly from heat-induced protein gels (80°C for 30 min) with respect to gel strength and appearance (Van Camp and Huyghebaert, 1995a, 1995b).

The gel strength (Van Camp and Huyghebaert, 1995a; Van Camp et al., 1996; Kanno et al., 1998; Fertsch et al., 2003), the storage modulus (G') and the loss modulus (G'') (Van Camp and Huyghebaert, 1995b) and the breaking stress (Kanno et al., 1998) increased with increasing protein concentration and pressure intensity because of increased probability of interactions between denatured proteins (Van Camp et al., 1996; Dumay et al., 1998). The ratio G'/G'' decreased with increasing protein concentration and the gels became more elastic (Van Camp and Huyghebaert, 1995a). In addition, the gel strength increased with increasing pressure and prolonged holding time (Van Camp and Huyghebaert, 1995a; Van Camp et al., 1996; Kanno et al., 1998). Longer pressure holding times improve the strength of the gel network, stimulating the formation of more intensive intermolecular interactions (Van Camp and Huyghebaert, 1995a).

Many such studies on the microstructure and rheological analysis of pressure-induced gels have been reported. β-LG gels made by pressure treatment also exhibited different rheological properties from those made by heating (Dumay et al., 1998). Electron microscopy suggested a higher level of cross-links in the heat-induced gels; high pressure generated a more porous network with a lower level of intermolecular cross-links (Van Camp and Huyghebaert, 1995a, 1995b; Van Camp et al., 1996).
Pressurization of β-LG isolate solutions (70 g/kg protein, pH 7.0) at 450 MPa and 25°C for 30 min formed gels with a sponge-like texture and a porous microstructure that was prone to exudation (Zasypkin et al., 1996). The rigidity and the elasticity of pressure-induced gels increased with increasing β-LG concentration, but remained lower than those of heat-induced gels (87°C for 40–45 min) at the same protein concentration (Zasypkin et al., 1996). These heat-induced gels also displayed a finely stranded network and high water retention. Some of the previous studies also suggested that, at neutral pH, β-LG forms transparent, fine-stranded heat-induced gels (Paulsson, 1990; Stading and Hermansson, 1991; Langton and Hermansson, 1992; Stading et al., 1993; Foegeding et al., 1995). These results suggested that comparatively weaker intermolecular or interparticulate forces are formed by pressure treatment.

A recent study characterized the interactions of whey proteins during pressure-induced gel formation using combinations of techniques such as transmission electron microscopy (TEM), SEC and 1D and 2D PAGE (Patel et al., 2006). Using SEC and 1D PAGE, the pressure-treated samples showed a time-dependent loss of native whey proteins, and a corresponding increase in non-native proteins and protein aggregates of different sizes. These aggregates altered the viscosity and opacity of the samples and were shown, using 1D PAGE (native, SDS and SDS reduced PAGE) and 2D PAGE (native then SDS and then reduced SDS PAGE), to be cross-linked by intermolecular disulfide bonds and by non-covalent interactions. It was concluded that the large internal hydrophobic cavity of β-LG may have been partially responsible for its sensitivity to high-pressure treatment. Conversely, α-LA responds to pressure by modifying its structure to be more molten globule-like and does not fully unfold at very high pressures.

Various possible hypotheses in support of pressure-induced gel formation have been discussed (Patel et al., 2006). It seems likely that, at 800 MPa, the formation of a β-LG disulfide-bonded network precedes the formation of disulfide bonds between α-LA or BSA and β-LG to form multi-protein aggregates, possibly because the disulfide bonds of α-LA and BSA are less exposed than those of β-LG either during or after pressure treatment. It may be possible that intermolecular disulfide bond formation occurs at high pressure and that hydrophobic association becomes important after the high-pressure treatment, i.e. a novel pathway of whey protein gel formation using high pressure.

It was postulated that β-LG plays a major role in the aggregation and gel formation of WPC under pressure (Van Camp et al., 1997a, 1997b; Patel et al., 2006), which suggested that the major whey protein component in WPC primarily determines its functional behavior under high pressure. However, it was suggested that some additional studies will be needed to confirm this hypothesis, as well as to deduce the role of other whey proteins (i.e. α-LA, BSA and Ig) in gel formation (Van Camp and Huyghebaert, 1995b; Van Camp et al., 1996).

Similar to heat-induced gelation of whey proteins, it has been reported that factors such as combinations of pressure and different temperatures (Walkenström and Hermansson, 1997), pHs (Van Camp and Huyghebaert, 1995b; Arias et al., 2000) and calcium contents (Van Camp et al., 1997b) affect the aggregation behavior, pressure-induced
functionality such as gel formation, and physical, rheological and microstructural properties of whey proteins. Protein–protein interactions are favored near the isoelectric point of the whey proteins, and neutral and alkaline pHs stimulate the formation of intermolecular disulfide bonds (Van Camp and Huyghebaert, 1995b).

Pressure-induced $\beta$-LG denaturation increases considerably at alkaline pH and decreases at acidic pH (Arias et al., 2000). Further, it has been reported that the role of calcium in the aggregation and gelation of whey proteins under pressure may be explained in a similar manner to the heat-induced effects on whey proteins (Mulvihill and Kinsella, 1988; Kinsella and Whitehead, 1989; Van Camp et al., 1997b). A combination of pressure and higher pressurizing temperature (up to 70°C) has been recommended for inactivating microbial spores and therefore it is important to determine its effects on the proteins in food systems. However, little information on the effects of pressurizing temperature on whey protein aggregation and gelation is available.

**HPP-induced changes in milk**

Studies on the effects of high pressure on milk can be broadly grouped into several topics, including the effects of high pressure on casein micelle size and its dissociation, changes in the appearance of pressure-treated milks, denaturation of whey proteins and their interaction with the casein micelles in milk, and effects of high pressure on milk from various species. Some of these topics have been reviewed recently (Huppertz et al., 2002, 2006a, 2006b; López-Fandiño, 2006a, 2006b; Considine et al., 2007b), including the effects of HPP on technological properties including rennet coagulation and cheesemaking properties, acid coagulation properties, etc. This section covers the main aspects of the pressure-induced denaturation and aggregation of whey proteins and their interactions with casein in the milk system.

**Denaturation of whey proteins in the milk system**

Compared with heat treatment, fewer studies on whey protein denaturation and aggregation in pressure-treated milk samples have been reported.

Like heat-induced denaturation, the pressure intensity and the holding time have been reported to affect the level of denaturation of whey proteins in milk (López-Fandiño et al., 1996; López-Fandiño and Olano, 1998a, 1998b; Huppertz et al., 2004a; Anema et al., 2005b; Hinrichs and Rademacher, 2005). However, considerable differences in the sensitivities of the different proteins to heat ($\text{LF} > \text{Ig} > \text{BSA} > \beta$-$\text{LG} > \alpha$-$\text{LA}$) and pressure ($\beta$-$\text{LG} > \text{LF} > \text{Ig} > \text{BSA} > \alpha$-$\text{LA}$) have been reported (Patel et al., 2006), showing that $\beta$-$\text{LG}$ is the most pressure sensitive among all the whey proteins. About 70–80% denaturation of $\beta$-LG occurs at 400 MPa (López-Fandiño et al., 1996; López-Fandiño and Olano, 1998a; Arias et al., 2000; García-Risco et al., 2000; Scollard et al., 2000). Relatively little further denaturation of $\beta$-LG occurs at 400–800 MPa (Scollard et al., 2000). Compared with $\beta$-LG, $\alpha$-LA is stable to pressures up to about 400–500 MPa.
in the milk environment at ambient temperature (Hinrichs et al., 1996a, 1996b; López-Fandiño et al., 1996; Felipe et al., 1997; Gaucheron et al., 1997; López-Fandiño and Olano, 1998a; Arias et al., 2000; García-Risco et al., 2000; Needs et al., 2000a; Scollard et al., 2000; Huppertz et al., 2002, 2004b).

Various studies have reported different extents of denaturation of β-LG following high-pressure treatment at 600 MPa of pasteurized milk (Needs et al., 2000a) or reconstituted skim milk powder (Gaucheron et al., 1997); this may be attributed to the level of denaturation caused by treatments before pressurization, which may influence the amount of denaturation measured afterwards. The reaction order of pressure-induced denaturation of β-LG is 2.5 (Hinrichs et al., 1996b), indicating that the denaturation process is concentration dependent and that a lower initial concentration of native β-LG should reduce the extent of denaturation of β-LG under pressure. Also, β-LG and α-LA are reported to be comparatively more pressure resistant in whey than in milk, which may be attributed to the absence of casein micelles and colloidal calcium phosphate in whey (Huppertz et al., 2004b).

Differences in the pressure stabilities of α-LA and β-LG may be linked to the more rigid molecular structure of the former (López-Fandiño et al., 1996; Gaucheron et al., 1997), caused probably by differences in the secondary structure and in the number of disulfide bonds and Ca^{2+} binding sites. The pressure resistance of α-LA is partially caused by the different numbers of intramolecular disulfide bonds in the two proteins (Hinrichs et al., 1996a, 1996b; Gaucheron et al., 1997) or by the lack of a free sulfydryl group in α-LA (López-Fandiño et al., 1996; Funtenberger et al., 1997).

It has also been reported that the molecular structure of α-LA is more stable than that of β-LG, and that oligomerization takes place only if, during unfolding, free sulfydryl groups from other molecules are available (Hinrichs et al., 1996b; López-Fandiño et al., 1996; Gaucheron et al., 1997; Jegouic et al., 1997). This difference in pressure sensitivity can also be explained by the types of bonds stabilizing the conformational structures of β-LG and α-LA (Hinrichs et al., 1996b; Messens et al., 1997).

BSA has also been found to be resistant to pressures up to 400 MPa in raw milk (Hinrichs et al., 1996b; López-Fandiño et al., 1996) or 600 MPa (Hayakawa et al., 1992). The high stability of BSA can be explained by the fact that BSA carries one sulfydryl group and 17 disulfide bonds. The energy received under pressure treatment is too small to disrupt all the disulfide bonds and to change the molecular structure of BSA. IgG in caprine milk (Felipe et al., 1997) and bovine milk (Carroll et al., 2006; Patel et al., 2006) has been reported to be more resistant to pressure denaturation than to heat denaturation.

**Interactions of whey proteins with casein micelles**

One of the major reactions of interest is the interaction between the denatured whey proteins and the casein micelles, particularly the interactions of denatured β-LG with κ-casein (κ-CN) at the micelle surface. There has been considerable research on the specific interactions that occur, the composition of the interaction products and the sequence of events involved in this reaction in heated milks (see Chapter 8), but little information on such interactions on effects of HPP treatments is available.
On high-pressure treatment of milk at 300–600 MPa, β-LG may form small aggregates (Felipe et al., 1997) or may interact with the casein micelles (Needs et al., 2000a; Scollard et al., 2000). It was reported that, when mixtures of κ-CN and β-LG were pressure treated at 400 MPa, the presence of β-LG reduced the susceptibility of κ-CN to subsequent hydrolysis by chymosin, indicating interactions between the proteins (López-Fandiño et al., 1997). SDS-PAGE studies of pressure-treated and untreated milks or solutions containing κ-CN or β-LG or both in the presence or absence of denaturing agents showed evidence of the formation of aggregates linked by intermolecular disulfide bonds (López-Fandiño et al., 1997). Although αs2-casein (αs2-CN) occurs at the same concentration as κ-CN and has one disulfide bond, it has not been reported to interact with β-LG in either heat- or pressure-treated milk systems.

Recently, using modified 2D SDS- and then reduced SDS-PAGE, Patel et al. (2006) showed that the effects of heat treatment and high-pressure treatment on the interactions of the caseins and whey proteins in milk were significantly different. The 2D SDS- and then reduced SDS-PAGE patterns (Figure 7.4) showed that pressure treatment of milk at 200 MPa (Figure 7.4D) caused β-LG to form disulfide-bonded dimers and incorporated β-LG into aggregates, probably disulfide bonded to κ-CN, suggesting that preferential reaction occurred at this pressure (Patel et al., 2006). The other whey proteins appeared to be less affected at 200 MPa.

In contrast, pressure treatment at 800 MPa incorporated β-LG and most of the minor whey proteins (including Ig and LF), as well as κ-CN and much of the αs2-CN, into large aggregates (Figure 7.4E). However, only a proportion of the α-LA was denatured or incorporated into the large aggregates. The relatively lower degree of α-LA reactivity at high pressures is probably related to the relative stability of this protein compared with β-LG, as discussed earlier, and is based on the unusual pressure-dependent behavior of α-LA (Kuwajima et al., 1990; Kobashigawa et al., 1999; Lasalle et al., 2003). These and other results show that the differences between the stabilities of the proteins and the accessibilities of the disulfide bonds of the proteins at high temperature or pressure affect the formation pathways that result in differences among the compositions of resultant aggregation or interaction products (including their sizes) that ultimately may affect product functionalities.

The formation of disulfide-linked complexes involving αs2-CN, κ-CN and whey proteins in heat- and pressure-treated milk has been demonstrated (see Figure 7.4) and has been explained by possible proposed reactions of the caseins and whey proteins in heat- and pressure-treated milk (Figure 7.5; see also Patel et al., 2006).

The virtual absence of αs2-CN from the heat-induced aggregates formed at 85–90°C in milk, as reported in previous studies, might be because αs2-CN is not a surface component of the micelle and therefore its disulfide bond(s) are inaccessible to the denaturing or denatured β-LG. On the other hand, κ-CN is on the surface of the micelles, and its disulfide bond(s) could be readily accessible to a thiol group of β-LG. Based on this fact, a diagrammatic representation of the consequences of the various inter-protein reactions that might take place during the heat treatment of milk at about 90°C has been reported (Patel et al., 2006). However, it has been reported
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α\textsubscript{S2}-CN
κ-CN
β-CN
α-LA
γ-CN

Reduced α\textsubscript{S2}-CN dimer

BSA
IgH

α\textsubscript{S2}-dimer

Control

Reduced sample

LF
BSA
IgH
α\textsubscript{S2}-CN
κ-CN
β-CN
κ-CN
β-LG
α-LA
γ-CN

Disulphide-linked complexes of casein and whey proteins resolved from region X4, X5 and X6 are high-molecular-weight aggregates, which were caught up at the beginning of the resolving gel, caught up within the stacking gel and could not enter the gel respectively. For a detailed description, refer Patel et al. 2006. (Reproduced in part with the permission of Patel et al., 2006, copyright 2006 American Chemical Society.)

Figure 7.4 2D SDS- and then reduced SDS-PAGE patterns of the control sample (a) and samples heat treated at 72°C for 15 s (b) and 140°C for 5 s (c). Similarly, 2D PAGE patterns of samples pressure treated at 200 MPa for 30 min (d) and 800 MPa for 30 min (e). Gel strips marked as a’ and a” represent the sample strip and the stained strip respectively and X4, X5 and X6 are high-molecular-weight aggregates, which were caught up at the beginning of the resolving gel, caught up within the stacking gel and could not enter the gel respectively. For a detailed description, refer Patel et al. 2006. (Reproduced in part with the permission of Patel et al., 2006, copyright 2006 American Chemical Society.)

(Patel et al., 2006) that α\textsubscript{S2}-CN is apparent in the large aggregates in milk samples heated at temperatures above 100°C, supporting an earlier report by Snoeren and Van der Spek (1977), who reported that processing at severe temperatures can affect the proteins in a qualitatively different way.
In contrast, the effects of the pressure treatment of milk (Figure 7.5) are different from the effects of the heat treatment of milk. It can be postulated that both the casein micelles and the whey proteins are compressed at pressures up to about 150 MPa (Anema et al., 2005a, 2005b). Thereafter, the micelles swell as the pressure
is increased up to 400 MPa, and hydrogen bonds and hydrophobic associations are diminished and colloidal calcium phosphate dissolves. As a consequence, the micelles absorb more water, swell and subsequently disperse (Anema et al., 2005a, 2005b). At 200 MPa, β-LG does not appear to become involved in interactions but is constrained to forming a stable, inactive, disulfide-bonded dimer (Figure 7.4) at <400 MPa, supporting the results of Considine et al. (2007b).

At higher pressures (>400 MPa), the polymerization of β-LG becomes the norm and pressure-induced β-LG aggregation becomes similar to heat-induced β-LG aggregation (Figures 7.4 and 7.5). The β-LG in WPC or in milk is not significantly modified by the other components, i.e. β-LG dominates the denaturation and aggregation pathway during pressure (>400 MPa) treatment, as it has been shown to dominate the reaction at high-temperature heat treatments.

Moreover, it has been reported that large quantities of very large aggregates that cannot enter the gel are present to a greater extent in heat-treated milk than in pressure-treated milk (Figure 7.4; Patel et al., 2006), indicating that the sizes of the aggregates are comparatively smaller in pressure-treated milks than in heat-treated milks. Such differences can be attributed to different effects of heat treatment and pressure treatment on the structure of the proteins, which may ultimately lead to different textures of the final products.

Unlike studies on the effects of heat treatment on casein–whey protein interactions and the distribution of casein–whey protein complexes between the colloidal and serum phases as a function of pH, only a limited number of studies have examined the effects of high-pressure treatment on casein–whey protein interactions as a function of pH (Arias et al., 2000; Huppertz et al., 2004a).

Conclusions

HPP is a rapidly growing non-thermal preservation technology that can potentially be used to create novel protein structures by bringing about particular changes in the molecular structure of proteins and thus may give rise to innovative, new generation value-added food products and new properties that are inaccessible via conventional methods of protein modification. In order to realize the full potential of HPP technology, it will be useful to have comparable levels of understanding of the science underlying the differences in the effects of heat treatment and pressure treatment on denaturation, aggregation and interaction of the different milk proteins.

In this chapter, we attempted to review the mechanisms and pathways of pressure-induced denaturation and aggregation, and of interactions of whey proteins in various systems including simple model systems using purified milk proteins and complex systems such as WPC and milk. The selected examples of differences in the heat- and pressure-induced interactions of whey proteins were discussed based on recent research findings. Based on the current literature, it appears that more focus on identifying the specific pressure-induced interactions of milk proteins in actual systems is needed. The effects of a range of processing conditions and the detailed mechanisms of possible synergistic interactions also need to be addressed appropriately.
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High-pressure-induced interactions involving whey proteins


Abstract
This chapter reviews the literature on the denaturation of the whey proteins in milk, their interaction reactions with other milk protein components, and provides some examples on the relationships between denaturation/interaction reactions of the whey proteins and the functional behavior of the milk in selected applications. Early studies on whey protein denaturation in milk were aimed at developing methods to assess denaturation levels and determining the relationships between the denaturation of the whey proteins and the functional behavior of milk products in bakery and other applications. Subsequent studies were directed towards modeling the whey protein denaturation processes through kinetic and thermodynamic evaluations and determining the role of various milk components on these denaturation processes.
Although the denaturation of whey proteins is critical in modifying the functional behavior of dairy products, it has become increasing apparent that a measure of the denaturation level of the whey proteins was not in itself a good predictor of functional performance. Therefore further studies have investigated the interactions of the denatured whey proteins with other milk protein components, in particular the interactions between the denatured whey proteins and the casein micelles (including identification of the specific disulfide bonds involved in complex formation between the denatured β-lactoglobulin and κ-casein). A limited number of recent studies have indicated that manipulation of the interactions of the denatured whey proteins with the other milk protein components may provide a significant tool for modifying or controlling the functional performance of milk protein products in some applications.

Introduction

Milk is produced in the mammary gland of female mammals and is intended for the feeding of the neonate from birth to weaning. Milk is a highly nutritious readily digested food, rich in protein, minerals and energy in an aqueous solution. It also provides the neonate with many other essential compounds such as protective agents, hormones and growth factors. Milk is a highly perishable fluid and was intended by nature to be consumed soon after production. However, humans have used milk and dairy-derived foods to supplement their diet for centuries and dairy products are still a major food source. Because of the commercial and nutritional significance of dairy products, manufacturing processes to preserve the food value of milk long after its initial production have been developed.

Over the last century, modern dairy milk processing has been transformed from an art into a science. Traditional products such as cheeses and yoghurts combine centuries-old knowledge with modern science, technology and processing techniques; in contrast, more recently developed products (such as spray-dried milk products, milk protein concentrates and whey protein concentrates) have been based on modern technologies of the time. Internationally, the majority of the milk processed is of bovine (cow) origin; however, significant quantities of buffalo, goat and sheep milk are also manufactured into dairy products (Fox, 2003).

The behavior of the protein components to various treatments largely dictates how milk will behave during processing. In milk, the lactose, some of the mineral components and the native whey proteins are in true molecular solution. However, the casein and most of the calcium and phosphate are found in large macromolecular assemblies called casein micelles. The colloidal suspension of casein micelles in milk serum is a remarkably stable food protein system, which allows the milk to be processed under the rigorous conditions used in modern dairy factories.

Milk can be subjected to high temperatures and pressures, high shear and variations in concentration without appreciable damage to the casein micelle system. Even the extreme action of drying milk to a powder does not significantly alter the milk system as milk powders can be reconstituted to produce liquid milks which have many properties that are similar to those of the milk from which they were derived (Singh and Newstead, 1992; Kelly et al., 2003; Nieuwenhuijse and van Boekel, 2003; O’Connell and Fox, 2003).
The casein micelle

In order to understand and rationalize any changes to the properties and stability of milk, it is necessary to have some knowledge of the casein micelle structure and assembly. Despite extensive research efforts, the detailed structure and assembly of the casein micelle has not been unequivocally established, although several models have been proposed (Figure 8.1; Schmidt, 1982; Walstra and Jenness, 1984; Walstra, 1990; Holt and Horne, 1996; Horne, 1998; Walstra, 1999).

Evidence from electron microscopy and light scattering suggested that the casein micelle was assembled from smaller sub-units and, as a consequence, sub-micelle models of the casein micelle structure were developed. In the later iterations of these sub-micelle models, the casein proteins were hydrophobically aggregated to form the sub-micelle units, and these sub-micelle units were linked by colloidal calcium phosphate (CCP) to form the casein micelle. The distribution of $\kappa$-casein ($\kappa$-CN) between sub-micelles was heterogeneous, and the sub-micelles with high levels of $\kappa$-CN were located at the micelle surface whereas those with low levels of $\kappa$-CN were in the interior, thus giving a surface location to $\kappa$-CN that was consistent with experiment (Figure 8.1a; Schmidt, 1982; Walstra and Jenness, 1984; Walstra, 1990).

![Diagram of casein micelle](image)

In recent years, there has been growing speculation over the existence of sub-micelles and the validity of the sub-micelle models of the casein micelles (Holt and Horne, 1996; Horne, 1998; Walstra, 1999; Horne, 2002, 2006). In particular, there was evidence to show that the CCP was uniformly distributed through the casein micelle, which precluded sub-micelles being linked by CCP to form the micelle. In addition, it was considered to be unlikely that there would be heterogeneous populations of casein sub-micelles with different levels of $\kappa$-CN, or that assembly into casein micelles via calcium phosphate occurs only after the casein sub-micelles have been formed.

Recent electron micrograph images of casein micelles did not display the internal sub-structure expected for casein sub-micelles and it was considered that the appearance of sub-micelles in earlier electron micrographs may have been artifacts of the early preparation techniques for electron microscopy (McMahon and McManus, 1998; Horne, 2006). The sub-micelle model of the casein micelle was refined to change the role of CCP from that of linking the sub-micelles to a charge-neutralizing agent to allow for a uniform distribution of CCP, and the sub-micelles were now linked together via hydrophobic interactions (Walstra, 1999). However, new models for the casein micelle that do not rely on the formation of sub-micelles have been proposed (Holt and Horne, 1996; Horne, 1998), the most recent of which is commonly known as the dual-binding model, as shown in Figure 8.1b and described by David Horne in Chapter 5 of this volume.

Although various models of the casein micelles have been proposed, these have largely been derived from the same pool of research data and are therefore different depictions or interpretations of similar information. As a consequence, many of the salient features of the structure, assembly and stability of the different models are similar (Schmidt, 1982; Walstra, 1990; Holt and Horne, 1996; Horne, 1998; Walstra, 1999). Hydrophobic interactions and CCP are important in maintaining micelle integrity. Therefore, micelle integrity can be modified or destroyed by disruption to hydrophobic interactions or by the dissolution of the CCP. In all recent models, $\kappa$-CN has a preferential surface location with the C-terminal region protruding from the surface layer as a flexible hair.

These models of the casein micelles, with the surface layers of $\kappa$-CN and an internal structure maintained by hydrophobic interactions and CCP, have been used to explain micelle stability and the destabilization by the enzymes in rennet, by acidification or by the addition of alcohol (Walstra, 1990; Holt and Horne, 1996; Horne, 1998, 2003). However, less studied and less well understood are the mechanisms responsible for the changes occurring to the casein micelles during the heating of milk, in particular the interactions with the denatured whey proteins, the heat-induced, pH-dependent dissociation of the casein from the micelles and the eventual heat-induced coagulation of the casein micelles.

The heat treatment of milk

The effect of heat on the milk system is an important consideration in dairy chemistry as a heat treatment is involved in the manufacture of almost all milk products.
The heat treatment may range from thermization (about 65°C for 15 s) to sterilization (about 120°C for 10–20 min) or ultra-high temperature (UHT) treatment (typically 138–142°C for several seconds). As the thermal history of milk influences the behavior of the milk in subsequent applications, the effects of heat on milk have been the subject of intensive, if somewhat intermittent, research, and many reviews and books on the subject are available (Singh and Creamer, 1992; International Dairy Federation, 1995, 1996; O’Connell and Fox, 2003; Singh, 2004).

When milk is heated, a number of competitive and often interdependent reactions occur and the importance of each reaction is determined by the heating conditions as well as by factors such as milk composition or concentration. When considering the protein components of milk, reactions of particular importance are whey protein denaturation, the interactions of denatured whey proteins with other proteins (including those of the casein micelles) and casein micelle dissociation reactions. These three reaction processes can markedly modify the physico-chemical properties of milk and may play a major role in determining the stability of milk and the functional performance of heated milk products.

**Whey protein denaturation**

The whey proteins as found in milk are typical globular proteins with well-defined secondary and tertiary structures. In contrast to the highly stable caseins, the globular whey proteins (especially α-lactalbumin [α-LA] and β-lactoglobulin [β-LG]) retain their native conformations only within relatively limited temperature ranges. Exposing the whey proteins to extremes of temperature results in the denaturation and aggregation of the proteins; this process can be expressed using the simple reaction scheme as shown in Equation 8.1.

For protein species where the native protein is in the form of non-covalently linked oligomers (such as dimeric β-LG), the first step in the denaturation process is the reversible dissociation of the oligomer into monomeric species (Equation 8.1a). The monomeric protein can then undergo partial unfolding of the native structure (Equation 8.1b). In principle, this unfolding step is reversible; however, in reality, and in complex mixtures such as milk, the unfolding process is accompanied by the exposure of reactive amino side-chain groups, which allows irreversible aggregation reactions to occur. The unfolded whey protein can undergo aggregation reactions with other (unfolded) whey proteins, with aggregates or with the casein micelles (represented by A in Equation 8.1c).

At a fundamental level, protein denaturation is often defined as any non-covalent change to secondary or tertiary structure of the protein molecule (Equation 8.1b). From this denatured state, the protein can revert to its native state (refold) or interact with other components in the system (aggregate). Under this definition, α-LA is generally regarded as one of the most heat-labile whey proteins whereas β-LG is one of the most heat-stable whey proteins (Ruegg et al., 1977).

However, for the dairy industry, it is the irreversible aggregation processes that follow the unfolding of the native whey proteins that largely determine the functional properties of dairy products. Hence, it is common practice to define whey protein
denaturation as the formation of irreversibly denatured and aggregated whey proteins (Sanderson, 1970a; Singh and Newstead, 1992; Kelly et al., 2003), and therefore this encompasses only the irreversible process shown in Equation 8.1c. Unless otherwise stated, the irreversible denaturation process is the definition used in this chapter. Using this definition for the denaturation of whey proteins in milk, the immunoglobulins are the most heat labile and α-LA is the most heat stable of the whey proteins, with β-LG and bovine serum albumin being intermediate (Larson and Rolleri, 1955). In general, significant denaturation of the major whey proteins—α-LA and β-LG—occurs only on heating milk at temperatures above ≈70°C.

\[
\begin{align*}
\text{Equation 8.1a} & \quad (P_N)_n \rightleftharpoons nP_N \\
\text{Equation 8.1b} & \quad P_N \rightleftharpoons P_U \\
\text{Equation 8.1c} & \quad P_U + A \rightarrow (P - A)
\end{align*}
\]

**Assessment of the denaturation of whey proteins in milk**

A considerable amount of research has been directed towards determining and understanding the denaturation processes of the major whey proteins when milk is heated. Early studies precipitated the casein and denatured whey protein by adjustment of the pH to the isoelectric point of the casein (≈pH 4.6), and analyzed the supernatant and the original milk for protein nitrogen, which gave estimates of initial native whey protein levels and levels after heat treatment (Rowland, 1933). However, a rapid method for determining the native whey protein levels was required for assessing milk powders for suitability in applications in the bakery industry (Harland and Ashworth, 1947), and also for categorizing milk powders based on the heat treatments received during the powder manufacturing process (Sanderson, 1970b, 1970c). From these requirements, the whey protein nitrogen index (WPNI) method was developed.

In the WPNI method, the casein and the denatured whey protein were precipitated and separated from the native whey proteins by saturation of the milk with salt, and the supernatant containing the native whey proteins was analyzed for protein content. This was originally achieved by dilution and pH adjustment of the supernatant to produce a turbid solution, with the turbidity related to the level of native whey protein originally present (Harland and Ashworth, 1947; Kuramoto et al., 1959; Leighton, 1962). However, the WPNI method displayed considerable variability in the degree of turbidity developed for samples with similar levels of whey protein denaturation. To overcome this, Sanderson (1970a) combined a dye-binding method for determining the total protein content of milk with the original WPNI method, thus giving a more accurate and reliable method for determining the WPNI.

The original WPNI method or one of its variants is still the industry standard for determining the native whey protein levels of milk powder products, and is still widely used to classify milk powders according to the heat treatments received (Singh and Newstead, 1992; Kelly et al., 2003). However, a recent report indicates that Fourier transform near-infrared spectroscopy may have potential as a rapid
method for the determination of the WPNI of milk powders in the dry state, eliminating the necessity of reconstitution, precipitation and filtration (Patel et al., 2007).

Although the WPNI method can give an estimate of the level of whey protein denaturation, research into the denaturation and interactions of the individual whey proteins requires more accurate separation and analysis procedures. There are numerous quantitative methods for separating and determining the level of the individual whey proteins in milk and these can be used directly or adapted to determine the level of denaturation after defined heat treatments. Methods that have been used include polyacrylamide gel electrophoresis (PAGE; e.g. Hillier and Lyster, 1979; Dannenberg and Kessler, 1988c; Kessler and Beyer, 1991; Anema and McKenna, 1996), capillary electrophoresis (e.g. Fairise and Cayot, 1998; Butikofer et al., 2006), differential scanning calorimetry (e.g. Ruegg et al., 1977; Manji and Kakuda, 1987), high-performance liquid chromatography (HPLC; e.g. Kessler and Beyer, 1991) and various immuno-based assays (e.g. Lyster, 1970).

In general, good correlations have been observed when the various methods for determining whey protein denaturation have been compared (Manji and Kakuda, 1987; Kessler and Beyer, 1991; Anema and Lloyd, 1999; Patel et al., 2007). These methods are more time-consuming than the traditional WPNI methods and therefore cannot be used for routine analysis and classification of milk products. However, they have higher accuracy and reproducibility and can be used to determine the denaturation behavior of the individual whey proteins. In addition, variations of the techniques or coupling to additional detection devices can provide further information on the interactions of the denatured whey proteins with other components in the milk (e.g. Lowe et al., 2004; Patel et al., 2006, 2007).

**Kinetic evaluation and modelling of whey protein denaturation**

Early studies showed that the denaturation of whey proteins was a kinetic phenomenon, and therefore dependent on both the temperature and the time of the heat treatment (Rowland, 1933; Harland and Ashworth, 1945). Even though these early studies considered the whey protein components as a single entity, it was noted that the denaturation of the whey proteins did not follow a simple exponential law and was not a first-order (unimolecular) process; therefore, it was concluded that the effect of temperature on the rate constants was difficult to determine. In addition, there was a change in temperature dependence at temperatures above about 80°C, which was probably the first indication of the complex nature of the irreversible denaturation of the whey proteins in milk (Rowland, 1933).

Although early studies on the effect of temperature and heating time on the denaturation of the individual whey proteins had been performed (e.g. Harland and Ashworth, 1945; Gough and Jenness, 1962), it was the kinetic study of Lyster (1970) over a wide temperature range (68–155°C) that conclusively demonstrated the complexity of the denaturation process of the individual whey proteins. Lyster (1970) found that the denaturation of α-LA appeared to follow first-order kinetics and that the denaturation of β-LG was second order. Arrhenius plots for the denaturation of both α-LA and β-LG indicated that the irreversible denaturation reaction was not a simple process, as...
a change in temperature dependence was observed at about 80–90°C for both α-LA and β-LG (Figure 8.2). The rate constants increased more rapidly with an increase in temperature in the low temperature ranges than at higher temperatures.

Further studies on the kinetic evaluation of the denaturation of whey proteins in milk confirmed this complexity of the denaturation process and provided relationships between compositional aspects and the rate of denaturation (Lyster, 1970; Hillier and Lyster, 1979; Manji and Kakuda, 1986). However, it was the kinetic and thermodynamic studies of Dannenberg and Kessler (1988c) that provided insights into the possible mechanisms responsible for the complex temperature dependences of the denaturation of α-LA and β-LG. Dannenberg and Kessler (1988c) found that, in milk, the denaturation of β-LG had an order of ≈1.5, which is now generally accepted, and that the denaturation of α-LA was pseudo first order. On the basis of the thermodynamic evaluations of the denaturation reactions of β-LG and α-LA in the two temperature ranges (i.e. at temperatures above and below the marked change in temperature

![Figure 8.2](image-url)

**Figure 8.2** Effect of milk concentration on the Arrhenius plot for the thermal denaturation of β-LG (a) and α-LA (b) over a 75–100°C temperature range. ○: 9.6% total solids milk; ●: 19.2% total solids milk; □: 28.8% total solids milk; ■: 38.4% total solids milk. Part A was adapted with permission from Anema (2000). Copyright (2000) American Chemical Society. Part B was adapted with permission from Anema (2001). Copyright (2001) Blackwell Publishing.
dependence for the denaturation reactions; Figure 8.2), information on the possible rate-determining steps in the denaturation reactions was obtained.

At temperatures below about 90°C for β-LG and 80°C for α-LA, the high values for the activation energies and enthalpies indicated that a large number of bonds were disrupted and the positive activation entropies indicated a lower state of order of the reaction products. These kinetic and thermodynamic parameters were interpreted as indicating that the unfolding (reversible denaturation) of the whey proteins was the rate-determining step in the lower temperature ranges.

At higher temperatures, above 80°C for α-LA and above 90°C for β-LG, considerably lower activation energies and enthalpies were typical of chemical reactions and the negative activation entropies indicated a higher state of order. These parameters suggested that chemical (aggregation) reactions were the rate-determining step in the higher temperature ranges. Subsequent studies on the kinetic and thermodynamic evaluation of the denaturation reactions have supported these interpretations in skim and whole milk under industrial processing conditions (Anema and McKenna, 1996; Oldfield et al., 1998a).

The denaturation reactions of both β-LG and α-LA are enhanced when the pH of the milk is increased from the natural pH and are retarded when the pH is decreased (Law and Leaver, 2000). The denaturation of β-LG was retarded when all components in the milk were concentrated, although the effect was less pronounced as the temperature was increased (Figure 8.2a; Anema, 2000). In contrast, the denaturation of α-LA was hardly affected by milk concentration, with similar rates of denaturation at all milk concentrations regardless of the heating temperature (Figure 8.2b; Anema, 2001).

The seemingly contrasting effects of milk concentration on the denaturations of α-LA and β-LG have been explained by detailed studies on the effect of the concentrations of the individual components of milk on the denaturation reactions. Increasing the protein concentration of milk, while maintaining essentially constant non-protein soluble component concentrations, increased the rate of denaturation of both α-LA and β-LG (Figure 8.3; Law and Leaver, 1997; Anema et al., 2006), with a similar effect at all temperatures (Anema et al., 2006).

Increasing the non-protein soluble components while maintaining constant protein concentrations retarded the denaturation of both β-LG and α-LA; however, the effects on these two proteins were somewhat different (Figure 8.3; Anema et al., 2006). For β-LG, increasing the non-protein soluble components caused a substantial retardation of denaturation in the lower temperature range and this effect became less pronounced at higher temperatures.

In contrast, the effect of increasing the non-protein soluble components on α-LA denaturation was less pronounced than for β-LG and was similar at all temperatures investigated (Figure 8.3). The increasing lactose concentration, the major component of the non-protein soluble components, could explain much of the effect of increasing non-protein soluble components; however, clearly other composition factors such as pH and ionic components also have an effect (Figure 8.3; Anema et al., 2006).

From these results, it was possible to explain the effects of milk concentration on the denaturation of β-LG and α-LA. For α-LA, on increasing the total solids concentration of the milk (both protein and non-protein soluble components), the retardation
of the reaction rate by increasing the non-protein soluble components concentration was almost exactly offset by the increase in the denaturation rate for α-LA on increasing the protein concentration. As this effect was similar at all temperatures, increasing total solids appeared to have no effect on the rate of denaturation of α-LA (Figures 8.2b and 8.3b; Anema, 2001; Anema et al., 2006).

However, for β-LG, the retardation in the rate of denaturation on increasing the concentration of the non-protein soluble components was not completely offset by the increasing rate of denaturation on increasing the protein concentration; therefore β-LG denaturation was retarded by increasing the total solids concentration of the milk. However, the non-protein soluble components were less effective in retarding the denaturation of β-LG at higher temperatures and, as a consequence, the increase in total solids concentration appeared to have a smaller effect on the denaturation of β-LG at the higher temperatures and particularly above about 90°C (Figures 8.2a and 8.3a; Anema, 2000; Anema et al., 2006). The effects of the non-protein soluble components concentration or the lactose concentration on the denaturation of β-LG and α-LA have been discussed in terms of the preferential hydration theory (Anema, 2000; Anema et al., 2006).
Interactions between denatured whey proteins and κ-CN/casein micelles

An understanding of the denaturation reactions of the whey proteins provides information on the initial steps of a complex series of aggregation reactions that can occur when milk is heated. This aggregation process can involve other milk protein components and may involve numerous reaction pathways or interaction processes. Although the reactions of the denatured whey proteins with other milk protein components are important, these types of reactions are considerably more difficult to measure than the irreversible denaturation processes, particularly in a complex mixture of components such as is found in (skim) milk.

Interactions between denatured whey proteins and κ-CN in model systems

One of the major reactions of interest is the interaction between the denatured whey proteins and the casein micelles, particularly interactions of denatured β-LG with κ-CN at the micelle surface. Early studies on model systems indicated that there was an interaction between β-LG and κ-CN when these components were heated together (Zittle et al., 1962; Long et al., 1963; Sawyer et al., 1963). These conclusions were drawn from electrophoretic studies, which showed that the discrete bands assigned to κ-CN and β-LG observed in unheated solutions produced species of intermediate mobility when the solutions were heated together. Sedimentation velocity experiments also confirmed complex formation, as the β-LG–κ-CN complex formed on heating had markedly higher sedimentation coefficients than did the individual proteins when heated separately (Zittle et al., 1962).

Once interaction between κ-CN and denatured β-LG had been confirmed, subsequent investigations in heated model systems were aimed at determining the types of bonds involved in complex formation, the stoichiometry of the complexes formed and the involvement of other whey proteins (particularly α-LA). It was shown that reducing agents dissociated the heat-induced complexes and that thiol-blocking agents prevented the formation of the complexes (Sawyer et al., 1963). These results supported earlier suggestions that the free thiol group of β-LG was involved in the interactions (Trautman and Swanson, 1958; Zittle et al., 1962) and it was suggested that intermolecular disulfide bonds were formed between κ-CN and denatured β-LG (Sawyer et al., 1963). This has been corroborated by numerous subsequent studies (e.g. Grindrod and Nickerson, 1967; Purkayastha et al., 1967; Sawyer, 1969; Tessier et al., 1969).

Some studies indicated that the heat-induced self-aggregation of β-LG was limited when κ-CN was present, which suggested that κ-CN formed complexes with intermediate species of aggregated β-LG (Sawyer, 1969; McKenzie et al., 1971). In contrast, other studies indicated that the aggregation of β-LG was not a prerequisite for interaction with κ-CN (Euber and Brunner, 1982). The reason for these apparently conflicting observations may have been resolved through the detailed study of Cho et al. (2003), in which many of the possible pathways involved in the aggregation of β-LG with κ-CN in heated protein model systems were elucidated. Cho et al. (2003) proposed that, when mixtures of β-LG and κ-CN were heated, the free thiol of β-LG was
exposed and this initiated a series of thiol–disulfide exchange reactions of β-LG with other denatured β-LG molecules or with κ-CN. The products formed ranged from 1:1 β-LG–κ-CN complexes to large heterogeneous aggregates and the product mix was dependent on the ratio of κ-CN to β-LG. The aggregate species were held together by either or both disulfide bonds and hydrophobic interactions.

Although there have been some indications that there may be an interaction between α-LA and κ-CN on heating (Shalabi and Wheelock, 1976; Doi et al., 1983), other studies have reported that interaction between these proteins does not occur (Baer et al., 1976; Elfagm and Wheelock, 1978). It is now generally believed that interactions between α-LA and κ-CN will occur only if β-LG (or another whey protein with a free thiol) is present during heating, and this may require the initial formation of a β-LG–α-LA complex, which subsequently interacts with κ-CN (Baer et al., 1976; Elfagm and Wheelock, 1978).

There is considerable evidence to show that disulfide bonds are involved in the aggregated species formed between the denatured whey proteins and κ-CN; however, there are reports that suggest that non-covalent bonding may be important in these interactions, particularly in the early stages of heating and at lower heating temperatures (Sawyer, 1969; Haque et al., 1987; Haque and Kinsella, 1988; Hill, 1989). In addition, other studies have shown that, although a substantial part of the denatured whey proteins in heated milk are involved in disulfide-bonded aggregates, there is a significant proportion that can be recovered as monomeric protein under dissociating but non-reducing conditions, indicating that non-covalent interactions are also involved (Oldfield et al., 1998b; Anema, 2000). As Cho et al. (2003) have suggested, it is likely that both hydrophobic and disulfide interactions are important in the early stages of aggregate formation, with the interaction mechanism dependent on the composition of the system and the conditions of heating.

**Interactions between denatured whey proteins and κ-CN/casein micelles in milk systems**

Most of the early studies examining the heat-induced interactions between denatured whey proteins and κ-CN were in model systems using purified proteins in buffer systems. Milk is considerably more complex, with numerous protein species that could potentially interact on heating. A number of the milk proteins have free thiol groups and/or disulfide bonds. Although β-LG is the major whey protein component, denatured α-LA and bovine serum albumin can also be involved in thiol–disulfide exchange reactions and therefore can be incorporated in the aggregated products. For the caseins, both κ-CN and αs2-CN have disulfide bonds and therefore both could participate in thiol–disulfide exchange reactions with denatured β-LG or other denatured thiol-bearing whey proteins. As a consequence of this complexity, there are numerous potential thiol–disulfide interaction pathways, as well as non-covalent interactions, and therefore the separation and analysis of the reaction products can be a difficult process.

However, the studies on the interactions between the proteins in heated milk suggest that, despite the complexity of the system, the reactions between β-LG and κ-CN
may be similar to those occurring in the model system. In early electrophoretic studies on heated milk, it was noted that the bands corresponding to β-LG disappeared, along with a reduction in the intensity of the bands corresponding to casein. This was accompanied by the formation of bands corresponding to new (heterogeneous) components (Slatter and van Winkle, 1952; Tobias et al., 1952). When sulfhydryl-blocking agents were added, the band pattern was comparable with that of the original skim milk, indicating that sulfhydryl-disulfide exchange reactions were involved in the interaction mechanisms (Trautman and Swanson, 1958). Subsequent studies confirmed that an interaction between denatured β-LG and κ-CN on the casein micelles occurred on heating milk although, as expected, the other denatured whey proteins were also involved in the interactions (Snoeren and van der Spek, 1977; Elfagn and Wheelock, 1978; Smits and van Brouwershaven, 1980; Noh et al., 1989a, 1989b; Corredig and Dalgleish, 1996a, 1996b; Oldfield et al., 1998b; Corredig and Dalgleish, 1999).

Unlike κ-CN, αs2-CN does not readily interact with denatured whey proteins when milk is heated, although some interactions in UHT milks have been reported (Snoeren and van der Spek, 1977; Patel et al., 2006). This low reactivity may be due to the location of αs2-CN in the interior of the casein micelles, which makes it less accessible for interaction, whereas κ-CN is located at the casein micelle surface and therefore may be more accessible for interaction (Walstra, 1990; Horne, 1998). Interestingly, in pressure-treated skim milk, disulfide-bonded aggregates between αs2-CN and the denatured whey proteins are observed, suggesting that the disulfide bonds of αs2-CN may become accessible to thiol groups of the denatured whey proteins when the casein micelle structure is disrupted under pressure (Patel et al., 2006).

The degree of interaction of the denatured whey proteins with the casein micelles is dependent on many variables including the time, temperature and rate of heating, the milk and individual protein concentrations, the milk pH and the concentration of the milk salts (Smits and van Brouwershaven, 1980; Corredig and Dalgleish, 1996a, 1996b; Oldfield et al., 2000; Anema and Li, 2003b; Oldfield et al., 2005). For example, when the temperature of milk is gradually increased above 70°C, as in indirect heating systems, most of the denatured β-LG and α-LA associates with the casein micelles, presumably as disulfide-bonded complexes with κ-CN at the micelle surface (Smits and van Brouwershaven, 1980; Corredig and Dalgleish, 1996b).

In contrast, when milk is heated rapidly, as in direct heating systems, only about half of the denatured β-LG and α-LA associates with the casein micelles, with the rest remaining in the milk serum (Singh and Creamer, 1991a; Corredig and Dalgleish, 1996a; Oldfield et al., 1998b). Corredig and Dalgleish (1999) suggested that, on heating milk, α-LA and β-LG initially aggregate in the serum phase at a ratio dependent on the initial individual whey protein concentrations. These complexes subsequently associate with κ-CN at the casein micelle surface on prolonged heating. However, Oldfield et al. (1998b) proposed that, under rapid heating rates, β-LG forms aggregates in the serum before interacting with the casein micelles and this limits the level of association with the casein micelles, whereas, at slower heating rates, monomers or smaller aggregates of β-LG may interact with the micelles and this may allow higher association with the casein micelles.
The pH of the milk at heating is important in determining the level of interaction between the denatured whey proteins and the casein micelles. Many of the early studies on the effect of pH were attempts to explain the unusual heat stability characteristics of the milk at very high temperatures. When milk is heated at high temperatures (\( \approx 140^\circ \text{C} \)), the heat coagulation time/pH profiles (HCT/pH profiles) of most milks show increasing heat stability with increasing pH to a maximum at about pH 6.7, followed by decreasing stability to a minimum at about pH 6.9, and increasing stability again as the pH is increased further (Rose, 1961). Considerable research has been undertaken over decades in an attempt to explain this unusual pH-dependent heat stability of milk and numerous factors are known to influence the heat stability behavior. This has been covered in many review papers on the heat stability of milk (Singh and Creamer, 1992; International Dairy Federation, 1995; O’Connell and Fox, 2003; Singh, 2004).

The results from these studies on the heat stability of milk have influenced the direction of the future research on the effects of heat on milk and in particular the interactions between denatured whey proteins and \( \kappa \)-CN/casein micelles. Therefore, it is appropriate to briefly review aspects of the pH-dependence of heat stability that are relevant to understanding the interaction between denatured whey proteins and \( \kappa \)-CN/casein micelles. Electron microscopic studies showed that, when milk was heated at high temperatures (90–140°C) for long times (30 min) at pH below 6.7, the denatured whey proteins complexed on to the micelle surfaces as filamentous appendages. However, when the milk was heated at higher pH, the denatured whey proteins were found in the serum phase as aggregated complexes (Creamer et al., 1978; Creamer and Matheson, 1980). These were the first indications that the pH at heating may influence the interactions between the denatured whey proteins and the casein micelles when milk is heated at high temperatures.

Kudo (1980) showed that the amount of non-sedimentable protein in milk heated at pH 6.5 was lower than that in unheated milk; however, the level of non-sedimentable protein increased with the pH at heating so that, above pH 6.7, the level was markedly higher than in the unheated milk and increased with increasing pH. Kudo (1980) concluded that the denatured whey proteins co-sedimented with the casein micelles at low pH (\( \approx \text{pH 6.5} \)), whereas most of the denatured whey proteins along with some casein (particularly \( \kappa \)-CN) was released from the casein micelles at pH above 6.8. It was also proposed that the transition from whey-protein-coated casein micelles to protein-depleted forms with changing pH at heating could explain the pH-dependence of the heat stability of milk at high temperatures.

Singh and Fox (1985a, 1985b, 1986, 1987a, 1987b, 1987c), in a series of extensive studies, showed that the dissociation of \( \kappa \)-CN-rich protein on heating was dependent on the pH at heating. At pH below about 6.8, little dissociation of micellar \( \kappa \)-CN occurred whereas, at higher pH, particularly above pH 6.9, high levels of \( \kappa \)-CN dissociated from the micelles, with the level increasing proportionally with increased pH. The whey proteins, particularly \( \beta \)-LG, played an important role in the heat-induced pH-dependent dissociation of \( \kappa \)-CN (Singh and Fox, 1987b), as did mineral components such as calcium and phosphate (Singh and Fox, 1987c). The results from these extensive studies have been used to develop detailed mechanisms for the
pH-dependent heat stability of milk and concentrated milk systems (O’Connell and Fox, 2003; Singh, 2004).

Initially, it was reported that the dissociation of κ-CN from micelles occurred only when milk at high pH (above \( \approx \text{pH 6.8} \)) was heated at high temperatures, particularly 90°C or above (Singh and Fox, 1985b). However, subsequent studies demonstrated that, at these pH values, the dissociation of κ-CN occurred as soon as the temperature was raised above ambient, with the level of dissociated κ-CN increasing proportionally with temperature up to 90°C. In these studies, the dissociation of \( \alpha_s \)-CN (\( \alpha_{s1} \)-CN and \( \alpha_{s2} \)-CN combined) and β-CN showed an unusual temperature dependence. Increasing levels of these caseins dissociated as the temperature was increased up to about 70°C, with the levels then decreasing again at higher temperatures (Figure 8.4; Anema and Klostermeyer, 1997; Anema, 1998).

A subsequent study showed that the unusual temperature dependence of \( \alpha_s \)-CN and β-CN was a consequence of the whey proteins, particularly β-LG. When whey-protein-depleted milk was heated, the levels of \( \alpha_s \)-CN and β-CN dissociating from the casein micelles increased with increasing temperature up to 90°C. When compared with heating standard milk, this indicated that higher levels of \( \alpha_s \)-CN and β-CN dissociated from the micelles in the whey-protein-depleted milks at temperatures above about 70°C (Anema and Li, 2000).

It was postulated that all the caseins dissociated from the micelles on heating. On subsequent cooling, the dissociated κ-CN stabilized the dissociated \( \alpha_s \)-CN and β-CN as small serum phase aggregates if the heating temperature was below about 70°C. However, above about 70°C, κ-CN was associated with denatured whey proteins. It was already known that the complex formed between κ-CN and denatured β-LG was less effective at stabilizing \( \alpha_s \)-CN and β-CN in the presence of calcium ions than uncomplexed κ-CN (Zittle et al., 1962); therefore, this interaction may have prevented κ-CN from stabilizing the other caseins and they either reassociated with the casein micelles or formed larger aggregates on subsequent cooling (Anema and Li, 2000).

Early studies on the effect of the pH at heating on the interaction of denatured whey proteins with the casein micelles tended to use relatively large pH steps. In a model milk system containing casein micelles and β-LG, about 80% of the denatured β-LG associated with the casein micelles when the milk was heated at pH 5.8 or pH 6.3, whereas only about 20% associated with the casein micelles at pH 6.8 or pH 7.1 (Smits and van Brouwershaven, 1980).

The studies on the heat-induced, pH-dependent dissociation of κ-CN from the casein micelles showed that this dissociation of κ-CN was accompanied by increases in the levels of denatured whey proteins remaining in the serum (Singh and Creamer, 1991b), and this was confirmed by Anema and Klostermeyer (1997) and Oldfield et al. (2000), who reported that 80–90% of the denatured whey proteins associated with the casein micelles when milk was heated at pH below 6.7, whereas only about 20% of the denatured whey proteins associated with the casein micelles at pH above 6.8. Corredig and Dalgleish (1996b) measured the ratio of β-LG or α-LA to κ-CN in the colloidal phase obtained from heated milk adjusted to pH 5.8, 6.2 or 6.8. Although the denatured whey proteins interacted with the casein micelles at a faster rate at lower pH and at higher temperatures, the ratios of denatured whey proteins to
κ-CN on the casein micelles were not markedly different under the different heating conditions.

Recent studies demonstrated the extreme importance of pH on the association of denatured whey proteins (α-LA and β-LG) with the casein micelles when milk was heated above 70°C, particularly at pH 6.7 or below, where differences in association behavior could be measured at pH differences as small as 0.05 pH units (Anema and Li, 2003a, 2003b; Vasbinder and de Kruijf, 2003). From these studies, it was shown that about 80% of the denatured whey protein was associated with the casein

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**Figure 8.4** Effect of temperature and pH on the level of protein in the supernatants obtained from 10% total solids reconstituted skim milk samples heated for 30 min: κ-casein (a); αs-casein (b); β-casein (c). ○: pH 6.3; ●: pH 6.5; □: pH 6.7; ■: pH 6.9; △: pH 7.1. Adapted with permission from Anema and Klostermeyer (1997). Copyright (1997). American Chemical Society.
micelles at pH 6.5 and that this level of association decreased linearly as the pH at heating was increased, so that only about 30% was associated at pH 6.7. At higher pH (above pH 6.7), very low levels of denatured whey proteins associated with the casein micelles (Figure 8.5).

Although the heat-induced pH-dependent dissociation of κ-CN from the casein micelles could explain the low levels of denatured whey proteins interacting with the casein micelles at pH above 6.8, it had been reported that very little κ-CN dissociated from the casein micelles at pH below 6.8 (Singh and Fox, 1985b; Nieuwenhuijse et al., 1991; Singh, 2004). Therefore, it was initially unknown why small shifts in pH between pH 6.5 and pH 6.7 affected the association of denatured whey proteins with the casein micelles when milk was heated. The level of κ-CN in the serum phase was low; therefore, it was initially believed that κ-CN was not involved in this partition of the whey proteins between the serum and colloidal phases (Oldfield et al., 1998b; Anema and Li, 2003b; Vasbinder and de Kruif, 2003).

However, more recent studies showed that the heat-induced dissociation of κ-CN was pH dependent from pH 6.5 to pH 7.1, with a linear increase in serum phase κ-CN as the pH was increased throughout the pH range from 6.5 to 7.1 (Figure 8.6a), and that the level of serum phase κ-CN was correlated with the level of serum phase denatured whey protein (Figure 8.6b; Anema, 2007). The differences in the level of dissociated κ-CN between the earlier studies and the later studies may be related to the centrifuging conditions, which may have masked the effects at the lower pH, especially under conditions where the particles are less hydrated and more readily deposited.

Although the level of κ-CN in the serum phase at pH below 6.7 was relatively low (less than ≈30% of the total κ-CN), the ratio of denatured whey protein to κ-CN was high and relatively constant (at about 2.5 whey proteins to each monomeric κ-CN) for the serum phase proteins at all pH. In contrast, the ratio of denatured whey protein to

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**Figure 8.5** Level of whey proteins associated with the casein micelles/non-sedimentable whey proteins in skim milk samples that were heated at 90°C for various times. The pH values of the milk samples prior to heating were: ●: pH 6.5; ○: pH 6.55; ▼: pH 6.6; ▽: pH 6.65; ■: pH 6.7; △: pH 6.9; ●: pH 7.1. Adapted with permission from Anema et al. (2004a). Copyright (2004) American Chemical Society.
The whey proteins in milk

κ-CN was only about 1:1 for the whey protein associated with the casein micelles at pH 6.5 and this decreased to about 0.5:1 at pH 7.1 (Anema, 2007). Intensive studies on the soluble whey-protein–κ-CN complexes formed when milk was heated at the natural pH also showed that κ-CN was intimately involved in the serum phase aggregates and that a high ratio of denatured whey proteins to κ-CN was observed (Guyomarch et al., 2003). Electron micrographs of the serum phase whey protein–κ-CN aggregates indicated that these particles were roughly spherical with a relatively uniform size of about 20–50 nm (Parker et al., 2005).

There is still some debate over the sequence of events in the interaction reactions between the denatured whey proteins and κ-CN. Some reports suggest that κ-CN dissociates from the micelles early in the heating process and that the denatured whey proteins subsequently interact with the κ-CN either in the serum phase or on

Figure 8.6 (a) Effect of the pH at heating on the level of non-sedimentable κ-casein in milk: ●: serum phase κ-casein in unheated milk; ▽: serum phase κ-casein in milk heated at 90°C for 20 min; ▾: serum phase κ-casein in milk heated at 90°C for 25 min; ▼: serum phase κ-casein in milk heated at 90°C for 30 min. (b) Relationship between the serum phase denatured whey protein and the level of serum phase κ-casein for the heated milk samples: milk heated at 90°C for 20 min; ●: milk heated at 90°C for 25 min; ○: milk heated at 90°C for 30 min. Adapted with permission from Anema (2007). Copyright (2007) American Chemical Society.
the micelles, with a preferential serum phase reaction (Anema and Li, 2000; Anema, 2007). This proposal was supported by the observations that the dissociation of κ-CN is a rapid process and that significant dissociation of κ-CN can occur at temperatures below those where the denaturation of whey proteins occurs (Anema and Klostermeyer, 1997). In addition, significant dissociation of κ-CN occurs in systems that have been depleted of whey proteins (Anema and Li, 2000). The higher ratio of denatured whey protein to κ-CN for the serum phase, regardless of the pH at heating or the level of dissociated κ-CN, may also suggest a preferential serum phase reaction between the denatured whey proteins and κ-CN (Anema, 2007).

However, other reports suggest that, on heating milk, the denatured whey proteins first interact with the casein micelles and that the whey protein–κ-CN complex subsequently dissociates from the casein micelles (Parker et al., 2005; Donato and Dalgleish, 2006). This proposal was supported by the observation that the addition of sodium caseinate to milk did not increase the level of serum phase complexes between the denatured whey proteins and κ-CN, which was interpreted as indicating that the complexes between the denatured whey proteins and κ-CN were formed on the casein micelle surface regardless of the pH at heating (Parker et al., 2005).

In addition, it was suggested that two mechanisms occur depending on the pH at heating. This was based on observations that the protein composition of the serum phase appeared to vary markedly depending on whether the milk was heated at pH above or below the natural pH of the milk (Donato and Dalgleish, 2006), although other studies did not display a marked difference in composition (Anema, 2007). At this stage, further detailed investigations are required to clarify whether dissociation of κ-CN occurs before or after interaction with the denatured whey proteins.

The pH-dependent changes in the association of the denatured whey proteins with the casein micelles, and the dissociation of κ-CN from the micelles, can have a marked effect on some of the physical properties of the milk. A marked increase in casein micelle size was observed when high levels of denatured whey protein were associated with the colloidal phase, as is observed on heating milk at pH 6.5. This change in size was less pronounced as the pH at heating the milk was increased to pH 6.7 and a decrease in casein micelle size was observed when significant levels of κ-CN were dissociated from the colloidal phase, as is observed on heating milk at pH above 6.7 (Figure 8.7a; Anema and Li, 2003b). Similar changes in viscosity (Figure 8.7b) and turbidity (Figure 8.7c) with the pH at heating were also observed (Anema et al., 2004c).

The difficulty in interpreting these changes in size, viscosity and turbidity is determining whether the association of the denatured whey proteins with the casein micelles is directly responsible for the change in size/volume of the casein micelles by increasing the diameters of the individual particles as the proteins interact, or whether there is some associated phenomenon, such as aggregation of the casein micelles, that is related to the level of whey protein or κ-CN that is in the serum phase or associated with the casein micelles. The strong relationship between the level of whey protein associating with the colloidal phase and the size/volume of the casein micelles, the observation that the size change plateaus on prolonged heating, and the relationships between the protein composition of the micelles and size, viscosity and turbidity seem
The whey proteins in milk

Figure 8.7  Effects of the pH at heating on the changes in the size of casein micelles (a), the viscosity of the milk (b) and the turbidity of the milk (c). The milk samples were heated at 90°C for various times and the pH values of the milk samples prior to heating were: ●: pH 6.5; ○: pH 6.55; ▽: pH 6.6; ▽: pH 6.65; ■: pH 6.7; □: pH 6.9; ◆: pH 7.1. Some of the particle size and viscosity results were adapted with permission from Anema et al. (2004c). Copyright (2004) Elsevier.

to suggest that the size changes are a direct consequence of the distribution of protein between the colloidal and serum phases, rather than an associated aggregation reaction (Anema and Li, 2003b; Anema et al., 2004c).

Determination of the specific disulfide bonds formed between κ-CN and β-LG
Although many types of bonding may be involved in the early stages of interactions between the denatured whey proteins and κ-CN, there is clear evidence that disulfide
bonds are involved in complex formation when model systems and milk are heated. Recent studies have focused on determining the specific thiol groups of κ-CN and, in particular, β-LG that are involved in the disulfide bonding between these two protein species when they are heated in model systems or milk. Understanding the specific disulfide bonds involved in the interaction process may provide useful insights into the mechanisms for the denaturation and subsequent aggregation reactions of the whey proteins in milk.

Native β-LG has two disulfide bonds and one free thiol group at Cys$^{121}$ (Qin et al., 1999), whereas κ-CN is found as a heterogeneous polymeric protein cross-linked by disulfide bonding via the two Cys groups in the monomer protein (Rasmussen et al., 1999). It was believed that the formation of disulfide bonds between the denatured β-LG and other milk proteins, including κ-CN, during heating first involved the dissociation of the β-LG dimer to monomer species, followed by the unfolding of the native structure, exposing the buried side groups including the reactive free sulfydryl at Cys$^{121}$. The exposure of this free Cys$^{121}$ then initiated a series of intermolecular thiol–disulfide exchange reactions with other denatured whey proteins and with κ-CN on the casein micelle surface (Snoeren and van der Spek, 1977; Iametti et al., 1996; Hoffmann and van Mil, 1997; Verheul et al., 1998; Vásbinder and de Kruif, 2003; Creamer et al., 2004).

However, studies on pure β-LG indicated that, in the early stages of the denaturation process, non-native monomeric β-LG species are formed, which may be intermediates in the intermolecular aggregation processes. These non-native monomeric species are stable on subsequent cooling and can be separated by alkaline PAGE techniques (Manderson et al., 1998; Hong and Creamer, 2002) or by gel permeation chromatography (Iametti et al., 1996; Croguennec et al., 2003, 2004). It was hypothesized that the non-native monomers were formed from intramolecular thiol–disulfide exchange reactions between the free Cys$^{121}$ of β-LG and the Cys$^{106}$–Cys$^{119}$ and/or Cys$^{66}$–Cys$^{160}$ disulfide bonds within the same β-LG monomer (Iametti et al., 1996; Manderson et al., 1998; Hong and Creamer, 2002; Croguennec et al., 2003, 2004).

With the advent of sensitive mass spectrometric techniques, the identification of the Cys residues involved in disulfide-bonded protein species was possible. The strategy for the identification of specific disulfide bonds involves a number of steps (GilleceCastro and Stults, 1996; Gorman et al., 2002; Lowe et al., 2004). For the sample under analysis, the protein species are first hydrolyzed under conditions where no further thiol–disulfide bond exchange reactions are likely to occur. The peptides formed from this hydrolysis are separated, usually by reverse phase HPLC, and the mass of individual peptides is determined by mass spectrometry (MS). The identification of individual peptides can be achieved by comparing the measured masses with those of expected peptides for the hydrolysis of the protein under study. For the disulfide-bonded peptides, usually other criteria also need to be satisfied, such as the peptides being present in non-reduced hydrolysates but absent in the reduced system.

Further confirmation can be gained by the use of tandem MS, where single molecular ions are isolated and analyzed in the first mass analyzer and then passed into a collision cell where fragmentation of the peptide is induced by collision with an inert gas (collision-induced dissociation [CID]) and the fragments are characterized in the
second mass analyzer. From the mass of the fragments, the sequence of the amino acids in the peptides can be achieved, providing conclusive characterization of the peptides (Gorman et al., 2002; Lowe et al., 2004).

Using these types of mass spectrometric techniques, a stable non-native monomeric \(\beta\)-LG with a free sulfhydryl group at position Cys\(^{119}\) rather than the natural position of Cys\(^{121}\) was found in heated \(\beta\)-LG solutions (Croguennec et al., 2003), confirming that intramolecular thiol–disulfide exchange within monomeric \(\beta\)-LG could occur. It was suggested that this \(\beta\)-LG with the free thiol at Cys\(^{119}\) may be the activated monomer that was proposed as the starting point for intermolecular aggregation reactions leading to large polymers, although it was equally possible that unfolded protein with a free thiol at the natural position of Cys\(^{121}\) was what activated monomer (Croguennec et al., 2003, 2004; Creamer et al., 2004).

A more recent investigation on the disulfide bonding patterns in heated \(\beta\)-LG found that a significant proportion of Cys\(^{160}\) was in the reduced form after heating \(\beta\)-LG in solution, indicating that the Cys\(^{66}\)–Cys\(^{160}\) disulfide bond was broken during the early stages of heating, and that this may occur concurrently with the interchange of the free thiol from Cys\(^{121}\) to Cys\(^{119}\) (Creamer et al., 2004). It was suggested that a monomeric \(\beta\)-LG species with a free thiol at Cys\(^{160}\) may be (one of) the reactive species involved in the intermolecular thiol–disulfide bonding responsible for cross-linking in heat-induced whey protein aggregates because of its position near the C-terminal end of the protein.

Attempts have been made to identify the specific Cys residues involved in disulfide bonds formed between \(\kappa\)-CN and \(\beta\)-LG when these proteins are heated together. Livney and Dalgleish (2004) compared masses of peptides from tryptic digests of heated \(\kappa\)-CN/\(\beta\)-LG mixtures with theoretical values and concluded that Cys\(^{106/119/121}\) of \(\beta\)-LG were involved in disulfide bonds with both Cys\(^{11}\) and Cys\(^{88}\) of \(\kappa\)-CN (note that the hydrolysis pattern does not allow the separation of the three Cys\(^{106/119/121}\) residues of \(\beta\)-LG unless CID is used for sequencing). Although some peptides involving Cys\(^{66}\) and Cys\(^{160}\) of \(\beta\)-LG and the two Cys residues of \(\kappa\)-CN were also identified based on mass comparisons, the high abundance of disulfide-bonded peptides containing Cys\(^{106/119/121}\) led these authors to conclude that \(\beta\)-LG with a free thiol at Cys\(^{119/121}\) was the predominant species that was involved in intermolecular disulfide bonding. The potential disulfide-bonded species were characterized based on mass analysis alone; no confirmatory experiments, such as comparing reduced with non-reduced systems to ensure that the proposed intermolecular peptides were disulfide bonded, or confirming sequences by CID–MS to preclude mis-identification of similarly massed peptides, were performed.

In a novel study, Lowe et al. (2004) used an activated monomeric \(\kappa\)-CN where the reduced thiol groups were blocked with thionitrobenzoate (TNB). \(\beta\)-LG was added to the mixture and the system was heated under very mild conditions (60°C). The TNB groups on the thiols of \(\kappa\)-CN are good leaving groups and, when a reactive thiol from \(\beta\)-LG is exposed, it is capable of interacting with the activated TNB groups on \(\kappa\)-CN in a specific 1:1 oxidative reaction forming a disulfide-bonded complex and releasing the TNB as a brightly colored compound. This approach allowed the formation of specific disulfide bonds between \(\kappa\)-CN and \(\beta\)-LG under mild heating conditions.
Because of the chemical nature of the reaction, it limited further thiol–disulfide exchange reactions, which allowed specific interactions between β-LG and κ-CN to be monitored during the early stages of the denaturation of β-LG.

The interacted β-LG–κ-CN complexes were hydrolyzed with trypsin and separated by reverse phase HPLC followed by MS. In addition, disulfide bonding was confirmed by comparing HPLC traces of non-reduced systems with those of reduced systems, and the identities of some peptides were confirmed by sequencing using CID–MS. Although it was possible to identify disulfide bonds between Cys^{106/119/121} of β-LG and Cys^{88} of κ-CN, Cys^{160} of β-LG was found to have formed disulfide bonds with both Cys^{11} and Cys^{88} of κ-CN as major products (Figure 8.8). This supported the earlier findings on pure β-LG, that intramolecular thiol–disulfide exchange may precede the intermolecular reactions and that the non-native monomeric form of β-LG species with a free thiol at Cys^{160} is likely to be (one of) the reactive monomer species that initiates intermolecular thiol–disulfide exchange reactions (Creamer et al., 2004).

Lowe et al. (2004), using the techniques developed for the model system, expanded the study to examine the specific disulfide bonds involved in aggregation between β-LG and κ-CN in heated milk systems. Interestingly, no disulfide bonds between Cys^{106/119/121} of β-LG and the two Cys residues of κ-CN could be found, even though the disulfide bond between Cys^{88} of κ-CN and Cys^{106/119/121} of β-LG was readily identified in the model system. In the heated milk system, it was found that Cys^{160} of β-LG formed disulfide bonds with both Cys^{88} and Cys^{11} of κ-CN, as was found in the model system (Figure 8.8). In independent studies, a similar disulfide bond between Cys^{160} of β-LG and Cys^{88} of κ-CN was identified in a heated model goat milk system consisting of isolated casein micelles and β-LG in milk ultrafiltrate, although it appears that no attempts were made to isolate and characterize other intermolecular disulfide bonds between these protein species (Henry et al., 2002).

From these observations, it was concluded that, in the model system of β-LG and activated κ-CN, non-native monomeric β-LG species with a free thiol and either Cys^{119} or Cys^{121} (but probably not Cys^{106}) could be the reactive monomer that is involved in intramolecular thiol–disulfide exchange reactions, as Cys^{119/121} was involved in disulfide bonds with κ-CN. However, further intramolecular thiol–disulfide exchange reactions in heated β-LG must precede or occur concurrently with the intermolecular reactions, as disulfide bonds between Cys^{160} of β-LG and the two Cys residues of κ-CN were also observed as major products in the model system (Figure 8.8).

In the heated milk system, no peptides involving Cys^{106/119/121} and the two Cys residues of κ-CN were isolated and only peptides involving Cys^{160} and Cys^{66} with both Cys residues of κ-CN were found (Figure 8.8). As Cys^{160} and Cys^{66} are involved in a disulfide bond in native β-LG, this indicates that intramolecular thiol–disulfide exchange reactions in β-LG precede the intermolecular thiol–disulfide exchange reactions and that a β-LG (monomeric) species with a free thiol group at Cys^{160} may play a significant role in the inter-protein disulfide bonding that occurs in heat-induced milk or whey protein systems.

The differences in reaction products between the model systems and milk may be a consequence of factors such as the heating conditions, the nature of the reactions...
**Figure 8.8** Diagram indicating the identified peptides on the linear sequences of κ-CN and β-LG and the intermolecular disulfide bonds formed between κ-CN and β-LG on heating model systems and milk. The horizontal box lines represent the protein sequence, the lines over the boxes represent the peptides and S–S indicates the presence of a disulfide bond. Arrows indicate the major proteolytic sites, the chymosin site for κ-CN and the rapid tryptic sites for β-LG. Potential glycosylation and phosphorylation sites are indicated for κ-CN. Reproduced with permission from Lowe et al. (2004). Copyright (2004) American Chemical Society.
(oxidative interaction compared with thiol–disulfide interchange reactions) and the fact that the $\kappa$-CN in milk is found within the casein micelles whereas in the model system it is not (Lowe et al., 2004). Because of the C-terminal location of Cys$^{\beta-LG}$, when this Cys is in the free thiol form and not linked to Cys$^{66}$, it may be able to productively react with the disulfide bonds of $\kappa$-CN to give stable $\kappa$-CN–whey protein aggregates (Lowe et al., 2004).

Relationships between denaturation/interactions of the whey proteins in heated milk and the functional properties of milk products

When milk is heated, there are numerous changes to the milk system, including changes to the proteins, the milk salts (including mineral equilibria between the colloidal and serum phases) and lactose; many of the changes can involve more than one of the milk constituents (International Dairy Federation, 1995). The changes can be irreversible or reversible to various extents depending on the changes being monitored and the conditions of the heat treatment. Although the changes to the protein system are an important determinant of the functional properties of milk products, all other changes to the milk system should also be considered to obtain a full understanding of the relationship between heat treatments, interactions and functional performance. However, there are limited examples of changes to components other than the proteins and the functional behavior of milk products, and therefore this review is restricted to some examples of the relationships between the changes in the milk protein system and the functional performance of the milk.

Examples of the relationships between whey protein denaturation and the functional properties of milk

In the early days of milk powder manufacture, it was recognized that the level of whey protein denaturation could be used as an index for the extent of heat treatment the milk had received during the manufacture of the milk powders, and that the functional properties of the milk products were related to some extent to the heat treatment that the milk had received during processing and therefore the level of whey protein denaturation (Harland and Ashworth, 1947; Larson et al., 1951; Harland et al., 1952).

Even as early as 1952, the concept of “tailor-made” milk powders was discussed, where powders were processed to provide specific requirements, such as low-heat powders for beverage applications and cottage cheese manufacture, and high-heat powders for bakery applications (Harland et al., 1952). Although there were no standards of quality for processing at this time, it was recognized that the proper control of processing conditions, particularly preheating of the milk, was necessary to produce satisfactory products and that measurement of the level of whey protein denaturation could be used as an objective method for determining the suitability of milk (powder) products for particular commercial and functional applications.
The heat treatment of milk, whether in liquid milk applications or prior to drying for milk powder manufacture, remains one of the major processes for manipulating the functional properties of milk products, and products such as milk powders are still generally classified according to the heat treatments applied using one of the derivatives of the WPNI test (Singh and Newstead, 1992; Kelly et al., 2003). With the extensive research on the denaturation of the whey proteins, and the ability to predict the denaturation levels after defined heat treatments, it would be envisaged that the level of denaturation of the whey proteins could be used as an indicator of the functional properties of milk products. In a broad sense, this is true. For example, certain heat classifications of milk powders will give improved functionality for particular applications over other classes of milk powders. Some of the general applications of different heat-classified milks and their functional uses, in particular for milk powder products, have been summarized in numerous publications (Singh and Newstead, 1992; International Dairy Federation, 1995, 1996; Kelly et al., 2003).

However, a huge range of temperature and heating time combinations are available to denature the whey proteins when milk is heated. As a consequence, specific correlations between the level of whey protein denaturation and the functional properties of milk across all possible heating conditions and milk sources do not exist. For example, the WPNI method was developed for assessing the suitability of milk powders for use in bakery applications; however, it was noted that a powder with a low WPNI did not always correspond to good baking qualities (Harland and Ashworth, 1947). Some of these variations are due to factors such as natural variations in the initial whey protein levels in the milk (Harland et al., 1955; Sanderson, 1970b); however, others are due to the methods of heat treatment during milk processing. As such, the WPNI or level of whey protein denaturation is at best a guide for the suitability of powders for specific applications, or an in-factory guide on processing conditions. Many manufacturers impose additional specifications to the milk powders to ensure suitability in their specific applications (Sanderson, 1970c; Singh and Creamer, 1991a).

Some of the most detailed studies on the relationship between the functional performance of milk and the heat treatment conditions or whey protein denaturation levels have been reported for acid gel or yoghurt systems. Parnell-Clunies et al. (1986) showed correlations between the level of whey protein denaturation and the firmness and apparent viscosity of yoghurt, regardless of the method used to heat the milk (batch [85°C], high-temperature short-time [98°C] and UHT [140°C] heating systems for different holding times). However, other properties, such as water-holding capacity/syneresis, were more dependent on the heating system used and it was concluded that high levels of whey protein denaturation in milk were not necessarily associated with an improved water-holding capacity in yoghurt.

In extensive studies, Dannenberg and Kessler (1988a, 1988b) examined the relationship between the denaturation level of whey proteins in milk and the functional performance (firmness, flow properties and syneresis) of the milk in set yoghurt applications. There was a clear relationship between the level of whey protein denaturation in the milk and the firmness, with a higher firmness at higher levels of whey protein denaturation. Similar results were obtained for the flow properties of the yoghurt. However, very high levels of whey protein denaturation appeared to be detrimental,
with a decrease in the firmness and flow properties at denaturation levels above about 95% (Dannenberg and Kessler, 1988b).

For syneresis of the yoghurt, a negative relationship between the level of whey protein denaturation in the milk and the level of serum expelled from the yoghurt was observed (Dannenberg and Kessler, 1988a). Despite the apparent correlations between denaturation and firmness, flow properties and syneresis, there were significant variations at each denaturation level, indicating that the temperature of heating used to denature the whey proteins, rather than just the whey protein denaturation level, may be an important factor in determining the functional performance in acid gels.

In a study on reconstituted whole milk, McKenna and Anema (1993) also observed a positive correlation between the denaturation of the whey proteins in the milk and the firmness of the yoghurt made from the milk regardless of whether the heat treatment was performed before or after powder manufacture (Figure 8.9a). However, when individual heating conditions were examined, it was also noted that excessive heat treatment/denaturation of the milk could be detrimental to the firmness of the set yoghurt (McKenna and Anema, 1993). A less clear relationship between syneresis and the level

![Figure 8.9](http://www.fil-idf.org)

**Figure 8.9** Relationship between the level of whey protein denaturation in reconstituted whole milk and the firmness (a) and syneresis (b) of acid gels prepared from the heated milks. The milks were heated only before powder manufacture (●), heated only after reconstitution (▼) or heated both before powder manufacture and after reconstitution (○). Adapted with permission from the results of McKenna and Anema (1993). Copyright (1993) International Dairy Federation (http://www.fil-idf.org).
of whey protein denaturation was observed, with the level of syneresis appearing to have a greater dependence on the heating conditions (temperature, time and before/after reconstitution) than on the level of denaturation itself (Figure 8.9b; McKenna and Anema, 1993), which supports the findings of Parnell-Clunies et al. (1986).

Examples of the relationships between the level of interactions of whey proteins with κ-CN/casein micelles and the functional properties of milk

A major limitation in using whey protein denaturation as an index of the functional properties of milk is that it does not consider the subsequent interaction reactions of the denatured whey proteins. These interactions will be dependent on the conditions of denaturation such as temperature and time as well as on the properties of the milk such as pH, concentration and composition. It is considerably more complex to investigate the subsequent aggregation reactions, as there are potentially numerous pathways and there is great difficulty in isolating and characterizing the specific reaction products. However, in recent years, some effort has been made in identifying the interaction reactions of the denatured whey proteins with other components in milk and in some cases their effects on the functional properties of the milk.

Anema et al. (2004a) showed that small changes in pH from the natural pH of milk at the time of heating markedly affected the properties of acid gels prepared from these heated milks. During acidification of the heated pH-adjusted milks, the acid gelation curves were progressively shifted to higher firmness as the pH at heating was increased (Figure 8.10a) so that the final firmness of the acid gels (at pH 4.2) was almost doubled as the pH at heating was increased from pH 6.5 to pH 7.1 (Figure 8.10). The effect was particularly pronounced in the milks that were heated for times sufficient to fully denature the whey proteins (Figure 8.10b). This effect of small changes in the pH of the milk at the time of heating on the firmness of acid gels prepared from the heated milk has been independently confirmed (Lakemond and van Vliet, 2005; Rodriguez del Angel and Dalgleish, 2006).

In addition to influencing the final firmness of the acid gels, the pH at the heat treatment of the milk also influenced the pH at which the milk started gelling/aggregating during acidification (Vasbinder and de Kruif, 2003; Anema et al., 2004a, 2004b; Lakemond and van Vliet, 2005; Rodriguez del Angel and Dalgleish, 2006). On subsequent acidification of milk samples that were heated over a pH range from pH 6.5 to 7.1, those samples heated at higher pH (pH 7.1) started gelling at significantly higher pH on acidification than those samples heated at a lower pH. These effects were very dependent on the temperature at which the milks were acidified (Anema et al., 2004b).

The changes in acid gel firmness on changing the pH at heating of the milk could not be related solely to the level of whey protein denaturation (Figure 8.11a). Small changes in the pH of the milk before heating markedly affect the distribution of the denatured whey proteins and κ-CN between the colloidal and serum phases (Figure 8.5; Anema and Li, 2003a, 2003b; Vasbinder and de Kruif, 2003; Rodriguez del Angel and Dalgleish, 2006). Although heating milk prior to acidification markedly
increased the firmness of the acid gels (i.e. acid gels prepared from heated milks always had a considerably higher firmness than acid gels prepared from unheated milks [Dannenberg and Kessler, 1988b; Lucey et al., 1997; Lucey and Singh, 1998]), the distribution of the denatured whey proteins and κ-CN between the colloidal and serum phases also appeared to influence the firmness of the acid gels. When the final firmness of the acid gels was plotted against the level of non-sedimentable denatured whey proteins in the milk, the results for all pH values were close to a single line (Figure 8.11b).

Anema et al. (2004a) concluded that, although the denatured whey proteins associated with the micelles have a significant effect on the final firmness of the acid gels, those denatured whey proteins that remain in the serum appear to have a more dominant influence over the final firmness than those associated with the casein micelles. For samples where virtually all the whey proteins were denatured, the final gel

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**Figure 8.10** (a) Changes in firmness ($G'$) with time after the addition of glucono-δ-lactone (GDL) for heated (90°C/30 min) skim milk samples. (b) Changes in the final firmness (final $G'$) for acid gels prepared from milk samples heated for various times at 90°C. The pH values of the milk samples prior to heating were: ●: pH 6.5; ○: pH 6.55; ▽: pH 6.6; ▽: pH 6.65; ■: pH 6.7; □: pH 6.9; ●: pH 7.1. In all samples, the pH was readjusted back to pH 6.7 before addition of GDL and this pH reduced to ≈pH 4.2 after 5.5 h. Reproduced with permission from Anema et al. (2004a). Copyright (2004) American Chemical Society.
The whey proteins in milk

Strength for acid gels prepared from milks in which all the denatured whey proteins were in the serum phase was found to be essentially a factor of two higher than that for acid gels prepared from milks in which all the whey proteins were associated with the casein micelles (Figure 8.12; Anema et al., 2004a).

Rodriguez del Angel and Dalgleish (2006) separated the non-sedimentable whey protein–κ-CN aggregates from milks heated at different pH using size exclusion chromatography and related the peak area of these aggregates to the firmness of the acid gels. They also concluded that the gel firmness appeared to be strongly dependent on the formation of soluble complexes in the milks and that there appeared to be a linear relationship between the level of soluble aggregates in the heated milk and the final strength of the acid gels.

Based on these results, a hypothesis on the roles of the non-sedimentable and micelle-bound denatured whey protein–κ-CN has been developed (Anema et al.,...
The increased pH of gelation and the increased acid gel strength of heated milk when compared with unheated milk has been attributed to the incorporation of the whey proteins as well as casein (micelles) in the acid gel structure during the acidification of milk (Lucey et al., 1997; Lucey, 2002; Graveland-Bikker and Anema, 2003).

In milk, the casein is insoluble at its isoelectric point (pH 4.6), whereas the native whey proteins remain soluble at all pH. Therefore, unheated milk starts aggregating when the milk pH approaches the isoelectric point of casein and visible gelation is observed at about pH 4.9. However, for heated milk, the denatured whey proteins are insoluble at their isoelectric points (pH 5.3 for β-LG, the major whey protein). Therefore, on acidification of heated milk, the proteins will start aggregating at a much higher pH, closer to the isoelectric points of the whey proteins. As a consequence, the contribution of the denatured whey proteins to the acid gel structure and the firmness of the acid gels is markedly higher than that observed for unheated milk (Lucey et al., 1997; Graveland-Bikker and Anema, 2003).

The pH at heating the milk will produce casein micelle particles with markedly different compositions (Figures 8.5 and 8.6; Anema and Li, 2003b; Vasbinder and de Kruif, 2003). The isoelectric point of the casein micelles is ≈pH 4.6, whereas the isoelectric point of the whey proteins is about pH 5.3. Therefore, on the acidification of milks heated at different pH, different aggregation and gelation behavior is observed. For the milks heated at high pH, the serum phase denatured whey proteins/κ-CN can aggregate separately and at a higher pH than the casein micelles. As the isoelectric point of these serum phase protein components will be higher than that of the casein micelles, the pH at which aggregation occurs will be progressively shifted to higher pH as the heating pH and the concentration of the serum phase denatured whey proteins/κ-CN are increased (Anema et al., 2004a; Rodriguez del Angel and Dalgleish, 2006).

![Figure 8.12](image_url) Comparison between the final firmness (final $G'$) and the level of denatured whey protein (open symbols) and the level of soluble denatured whey protein (filled symbols) for acid gels prepared from heated (90°C/30 min) skim milk samples. The pH values at heating of the milks were: ●, ○: pH 6.5; ■, □: pH 6.55, ▲, △: pH 6.6; ▼, ▽: pH 6.65; ●, ◆: pH 6.7; dotted, ●, ○: pH 6.9; dotted, ■, □: pH 7.1. Adapted with permission from Anema et al. (2004a). Copyright (2004) American Chemical Society.
In addition, the dissociation of $\kappa$-CN from the casein micelles may also contribute to the higher aggregation pH as the pH at heating is increased, particularly above pH 6.7. Lower levels of $\kappa$-CN on the micelles will reduce the density of the surface hairy layer. This may cause the surface hairy layer to collapse at a higher pH, or this layer may have a reduced efficiency in stabilizing the casein micelles. Either effect will allow the $\kappa$-CN-depleted micelles to aggregate at a pH that is markedly higher than that observed for the native casein micelles or for casein micelles in milk heated at a lower pH.

The firmness of acid gels can be related to the number and properties of the contact points between the protein components in the acid gel (van Vliet and Keetals, 1995; Lucey et al., 1997). As the pH at heating of the milk is increased, the level of serum phase denatured whey protein–$\kappa$-CN complexes increases and therefore there are a greater number of particles to aggregate during the subsequent acidification to form the acid gels. There is also the potential for the formation of a more complex acid gel structure when the milk is heated at high pH, where there are high levels of serum phase denatured whey protein–$\kappa$-CN complexes, than when the milk is heated at low pH, where most of the denatured whey protein and $\kappa$-CN are associated with the casein micelles. In the latter case, the acid gel process will probably involve only entire whey protein–casein micelle complexes. Therefore, there may be fewer contact points in the acid gels formed from milk with the denatured whey proteins associated with the micelles than in those formed from milk with soluble denatured whey proteins and hence a gel with a lower firmness is observed (Anema et al., 2004a, 2004b).

The large strain deformation properties also gave some indication of the types of bonds involved in the acid gel network. In these experiments, the strain was increased at a constant rate and the stress was monitored until the gel structure yielded and the stress decreased. The maximum in the strain versus stress curves was considered to be the point at which the gel structure broke (Figure 8.13). As the pH at heating was increased, the breaking stress of the acid gels prepared from the heated milks was found to increase markedly; however, the breaking strain was virtually unchanged.

For a gel to break on increasing the strain, the strands within the gel network are first straightened and then stretched until rupture of the strands or the bonds within (van Vliet and Keetals, 1995; Mellema et al., 2002). Therefore, the breaking strain is dependent on factors such as the degree of curvature of the strands, with a higher breaking strain with higher strand curvature. As the breaking strain of the acid gels did not change with the pH at heating of the milk, despite the marked change in final firmness (Figures 8.10 and 8.12), this indicates that the relative curvature of the individual strands within the gel network was the same for all acid gel samples.

The types of bonds involved in the acid gel network will have an influence on the breaking stress (Mellema et al., 2002). The breaking of strands containing covalent bonds would require a greater force than the breaking of strands held together by non-covalent bonds, as covalent bonds have higher bond energies. Therefore, a change in the number or distribution of covalent bonds within the gel network may explain the differences in breaking stress as the pH of the milk at heating was changed (Figure 8.13). It seems unlikely that the difference in breaking stress can be due to a greater degree of disulfide bonding within the gelled sample; although there may be continuing thiol–disulfide exchange reactions occurring during acidification (Vasbinder...
et al., 2003), the physical number of disulfide bonds is unlikely to be markedly different between the samples.

The denatured whey proteins, along with some of the κ-CN, are progressively transferred to the serum phase when the pH of the milk is increased before heating (Figure 8.5). As these interactions involve disulfide bonding, this indicates that the interaction between the denatured whey proteins and κ-CN is transferred from the colloidal phase (casein micelle) to the serum phase as the pH of the milk at heating is increased. On subsequent acidification, both non-sedimentable and colloidal phase denatured whey proteins are incorporated in the acid gel structure. The non-sedimentable denatured whey protein–κ-CN complexes can form strands that may be involved in interconnecting the colloidal particles. As the non-sedimentable aggregates are disulfide bonded, those samples heated at high pH and with high levels of non-sedimentable whey protein–κ-CN aggregates will have a greater number of these strands interconnecting the residual casein micelles.

In contrast, the samples heated at lower pH will have the denatured whey proteins predominantly associated with the casein micelles and therefore fewer of the whey protein–κ-CN aggregates interconnecting the colloidal particles. Therefore, the samples heated at higher pH may have a greater number of disulfide bonds interconnecting the colloidal particles and therefore a higher breaking stress whereas, for the samples heated at lower pH, most of the disulfide bonds are on the colloidal particles and fewer disulfide bonds interconnect the colloidal particles, which may explain the lower breaking stress (Figure 8.13).

**Examples of the effect of denaturing whey proteins separately from casein micelles on the functional properties of milk**

Interestingly, different effects are observed when the whey proteins are denatured and aggregated separately from the casein micelles than when the milk is heated.
For example, Lucey et al. (1998) showed that acid gels prepared from milk samples where the whey proteins were denatured in the presence of casein micelles had a markedly higher firmness than acid gels prepared from milk samples where the whey proteins were pre-denatured and added back to the casein micelles. In fact, in many cases, the samples with denatured whey proteins added back to the casein micelles produced acid gels with firmness similar to or only slightly higher than those prepared from unheated milks.

In a separate study, Schorsch et al. (2001) prepared model milk systems in which the whey proteins were either heated in the presence of casein micelles or heated separately and added back to the casein micelles. Acid gels were prepared from these model milk systems. It was shown that the acid-induced gelation occurred at a higher pH and in a shorter time when the whey proteins were denatured separately from the casein micelles than when the whey proteins were heated in the presence of casein micelles. However, the gels formed were weaker and more heterogeneous because of the particulate nature of the denatured whey proteins.

It was suggested that the large denatured whey protein aggregates, as formed when the whey proteins were heated separately from the casein micelles, hinder the formation of a casein gel network when the milk is subsequently acidified and that a weak acid gel with a heterogeneous structure results. When the whey proteins are heated in the presence of the casein micelles, the denatured whey proteins interact with the $\kappa$-CN at the casein micelle surface and, on subsequent acidification, the denatured whey protein–casein micelle complexes aggregate to form a firmer acid gel with a more homogeneous structure (Schorsch et al., 2001). This proposal is supported by early studies, which showed that the aggregation of the denatured whey proteins, and in particular $\beta$-LG, formed large aggregate species when heated in the absence of $\kappa$-CN, whereas aggregation was limited when the whey proteins were heated in the presence of $\kappa$-CN (McKenzie et al., 1971).

### Conclusions

A considerable amount of work has gone into understanding the irreversible denaturation reactions of the whey proteins in heated milk systems. These detailed studies have produced models that allow reasonably accurate prediction of the level of whey protein denaturation in milks under a wide range of heating conditions, even in milk samples with markedly modified concentrations and compositions. However, with a few exceptions, monitoring of the whey protein denaturation levels provides only a crude indication of the functionality of the milk system. As a consequence, more recent research efforts have focused on trying to understand the specific interaction reactions of the denatured whey proteins with other proteins in the milk system. Early indications suggest that these types of studies on the interactions of denatured whey proteins may provide greater insights into the functional properties of heated milk products than can be obtained by monitoring just whey protein denaturation levels.

These initial studies on protein interactions have been conducted under relatively narrowly defined conditions (temperatures, heating times, pH, milk concentrations and milk compositions). It is likely that changes to these variables will markedly influence
the interaction behavior and will explain the changes in functional behavior when the heating conditions are changed (even though the whey protein denaturation levels may be similar). Although studies on understanding the specific interactions between milk proteins, particularly in complex systems such as milk, are extremely difficult, these types of studies should continue to give useful insights into the behavior of milk proteins during heating and the functional behavior of the heated milk products.

References


Effects of drying on milk proteins

Pierre Schuck

Abstract

Dehydration by spray drying is a valuable technique for water evaporation, using hot air to stabilize the majority of dairy ingredients. In view of the increased development of filtration processes, the dairy industry requires greater understanding of the effects of spray drying on the quality of protein powders. Several publications have reported that the proteins have an important role in the mechanisms of water transfer during drying and rehydration. The residence time of the droplet and then the powder is so short that it is very difficult to study the mechanism of the structural change in the protein without fundamental research into relationships with the process/product interactions. Following an introduction to spray drying, this chapter on the effects of drying on milk proteins covers five areas, i.e. the world dairy powder situation, properties of spray-dried milk products, principles of spray drying, drying of proteins and rehydration of dried protein powders.

Introduction

The purpose of the dehydration of milk and whey is to stabilize these products for their storage and later use. Milk and whey powders are used mostly in animal feeding. With changes in agricultural policies (such as the implementation of the quota system and the dissolution of the price support system in the European Union), the dairy industry was forced to look for better uses for the dairy surplus and for the
by-products of cheese (whey) produced from milk and buttermilk produced from cream. Studies on the reuse of protein fractions with nutritional qualities and functionality led us to believe that they could have multiple applications.

In the past 25 years, the dairy industry has developed new technological processes for extracting and purifying proteins (casein, caseinates, whey proteins, etc.) (Kjaergaard et al., 1987; Maubois, 1991) such as:

- dairy proteins and whey concentrates (Le Graët and Maubois, 1979; Goudédranche et al., 1980; Madsen and Bjerre, 1981; Maubois et al., 1987; Caron et al., 1997);
- micellar casein concentrates (Fauquant et al., 1988; Schuck et al., 1994a);
- micellar casein (MC) (Pierre et al., 1992; Schuck et al., 1994b);
- whey concentrates;
- selectively demineralized concentrates (Jeantet et al., 1996); and
- super-clean skim milk concentrates (Piot et al., 1987; Vincens and Tabard, 1988; Trouvé et al., 1991; Schuck et al., 1994a).

mainly because of the emergence of filtration technology (microfiltration, ultrafiltration, nanofiltration and reverse osmosis).

Most of these proteins, used as either nutritional or functional ingredients, are marketed in dehydrated form (Figure 9.1). The application of different processing steps allows the production of a wide range of different dried and stable intermediate dairy products. Many new uses for these constituents emerged with the manufacture of formula products, substitutes and adapted raw materials.

The most frequently used technique for the dehydration of dairy products is spray drying. It became popular in the dairy industry in the 1970s but, at that time, there were few scientific or technical studies on spray drying and in particular none on the effects of spray drying parameters or on the effects of the physico-chemical composition and microbiology of the concentrates on the powder quality. Manufacturers acquired expertise in milk drying and eventually in whey drying processes through trial and error. Because of the variety and complexity of the mixes to be dried, a more rigorous method based on physico-chemical and thermodynamic properties has become necessary. A better understanding of the biochemical properties of milk products before drying, water transfer during spray drying, the properties of powders and influencing factors is now essential in the production of milk powder. The lack of technical and economic information and of scientific methods prevents the manufacturer from optimizing dairy plants in terms of energy costs and powder quality.

The aim of this chapter is to give a brief summary of the process of the spray drying of dairy products and to review present knowledge on the properties of spray-dried milk products, modeling and simulation of water transfer processes (drying and rehydration), dairy powders and spray-drying equipment and energy consumption.

**World dairy powder situation**

There has been a change in the nature of dairy powders over the last 15–20 years (CNIEL, 1991, 2005). A decrease in production has occurred mainly for skim and
Figure 9.1 Fractionation of milk.
Effects of drying on milk proteins

fat-filled milk powders but the production of whole milk powder and whey powder increased between 1986 and 2004. This increase was reflected in the types of whey and derived powders (protein concentrates) produced. Cheese production from cow’s milk increased between 1986 and 2004, with a corresponding increase in whey production. Having fallen in 2004, the production of dry milk products did not recover significantly in the first months of 2005, the decline being due mainly to slower growth of milk supplies in many parts of the world.

Whole milk powder (WMP)

According to the International Dairy Federation (2005), the production of WMP was different in different parts of the world: WMP production increased, mainly in China, Latin America and New Zealand, but was almost unchanged in Europe and Australia. The most important producer is now China, with 0.9 million tonnes, accounting for more than one-quarter of recorded production and ahead of the EU25 (with 0.85 million tonnes). The next highest producers are New Zealand, Brazil, Argentina and Australia: all these countries together cover over 90% of world production. In the long term, the EU and Australia are likely to lose their share of world production.

In the medium and long term, the production of WMP is driven mainly by demand. WMP provides milk where raw milk is in very short supply compared with demand. In countries where reconstitution of dried milk is common, WMP is ideal for bridging regional and seasonal deficits and this is also the reason why WMP is the best product to meet this demand. WMP does not play a significant role in the international dairy trade, but the function of the WMP produced is just to balance the market over long distances and across seasons within a country.

Skim milk powder (SMP)

With SMP, the situation is somewhat different from WMP. As well as the function of balancing seasonal and regional markets, there is also the need to clear temporary surpluses when there is no other use for skim milk. This explains the decline in SMP production in 2004, because the overall availability of milk as a raw material fell short of the demand. As a result, world production of SMP fell by 0.3 million tonnes in 2004. Most of this reduction took place in the EU, with production falling by 250,000 tonnes in one year, to the extent that EU25 production in 2004 was less than EU15 production in 2003. The USA and New Zealand also reduced their output. A degree of recovery occurred in 2005, but it was only modest. This recovery is unlikely to replace the volumes made available in 2003 and 2004 by the clearing of stocks, in particular in Oceania, the USA and the EU.

Whey products, casein and other dairy ingredients

More liquid whey is now generated from cheese production, following the general growth in this sector. Although the increased application of new technologies, e.g. ultrafiltration, microfiltration (Maubois, 1991) and an increasing number of new processing techniques to exploit milk components, has changed the composition of the
liquid that remains, the major part is still ordinary sweet whey. Thus, the main product is still sweet whey powder, but other derivatives such as whey protein concentrate, lactose and demineralized and delactosed products are also being produced and are evolving even faster than ordinary whey powder. However, the statistics are difficult to obtain. For the EU15, whey powder production was estimated at 1.3 million tonnes.

Other important producers are the USA and Canada, with a combined production of 500,000 tonnes. In 2004, production stagnated or even declined, attributable in part to the growing use of liquid whey for other derivatives and a reduction in casein manufacture, as well as low prices.

Casein production increased in the EU15 in 2004, but fell again in 2005. However, in Eastern Europe and in Australia, recovery can be expected.

Properties of spray-dried milk products

A dairy powder is characterized not only by its composition (proteins, carbohydrates, fats, minerals and water) but also by its microbiological and physical properties (bulk and particle density, instant characteristics, flowability, floodability, hygroscopicity, degree of caking, whey protein nitrogen index, thermostability, insolubility index, dispersibility index, wettability index, sinkability index, free fat, occluded air, interstitial air and particle size), which form the basic elements of quality specifications; there are well-defined test methods for their determination according to international standards (Pisecky, 1986, 1990, 1997; American Dairy Products Institute, 1990; Masters, 1991). These characteristics depend on drying parameters (type of tower spray drier, nozzles/wheels, pressure, agglomeration and thermodynamic conditions of the air, such as temperature, relative humidity and velocity) and the characteristics of the concentrate before spraying (composition/physico-chemical characteristics, viscosity, thermo-sensitivity and availability of water). Several scientific papers on the effects of technological parameters on these properties have been published (Hall and Hedrick, 1966; De Vilder et al., 1979; Baldwin et al., 1980; Pisecky, 1980, 1981, 1986; Kessler, 1981; Bloore and Boag, 1982; De Vilder, 1986; Tuohy, 1989; Ilari and Loisel, 1991; Masters, 1991; Mahaut et al., 2000). Water content, water dynamics and water availability are among the most important properties (Figure 9.2).

The nutritional quality of dairy powders depends on the intensity of the thermal processing during the technological process. The thermal processing induces physico-chemical changes that tend to decrease the availability of the nutrients (loss of vitamins, reduction of available lysine content and whey protein denaturation) or to produce nutritional compounds such as lactulose (Straatsma et al., 1999a, 1999b).

Principles of spray drying

According to Pisecky (1997), spray drying is an industrial process for the dehydration of a liquid containing dissolved and/or dispersed solids (e.g. dairy products), by
transforming the liquid into a spray of small droplets and exposing these droplets to a
flow of hot air. The very large surface area of the spray droplets causes evaporation of
the water to take place very quickly, converting the droplets into dry powder particles.
Indeed, when a wet droplet is exposed to hot dry gas, variations in the temperature
and the partial pressure of water vapor are established spontaneously between this
droplet and the air:
- heat transfer from the air to the droplet occurs under the influence of the tem-
  perature variation;
- water transfer occurs in the opposite direction, explained by variation in the
  partial pressure of water vapor between the air and the droplet surface.

Air is thus used both for fluid heating and as a carrier gas for the removal of water. The air enters the spray drier hot and dry and leaves wet and cool. Spray drying is a
phenomenon of surface water evaporation maintained by the movement of capillary
water from the interior to the surface of the droplet. As long as the average moisture
is sufficient to feed the surface regularly, the evaporation rate is constant. If not, it
decreases.

The drying kinetics are related to three factors:

1. Evaporation surface created by the diameter of the particles. Spraying increases
   the exchange surface: 1L of liquid sprayed in particles of 100μm diameter
   develops a surface area of 60m², whereas the surface area is only approxi-
   mately 5dm² for one sphere of the same volume.
2. Difference in the partial pressure of water vapor between the particle and the
drying air: a decrease in the absolute humidity of the air and/or an increase
   in the air temperature tend to increase the difference in the partial pressure of
   water vapor between the particle and the drying air.
3. Rate of water migration from the center of the particle towards its surface: this
   parameter is essential for the quality of dairy powders. Indeed, it is important
that there is always water on the surface of the product so that the powder surface remains at the wet bulb temperature for as long as possible. The rate of water migration depends on the water diffusion coefficient, which varies according to the biochemical composition, water content and droplet temperature. This is why calculation of this coefficient is complex and the mathematical models suggested are not easily exploitable by the dairy industry.

To define the components of a spray-drying installation, according to Masters (1991) and Pisecky (1997), the main components of the spray drier shown in Figure 9.3 are as follows:

- A drying chamber (Figure 9.3, 7). The chamber can be horizontal (box drier) although, in the dairy industry, the chamber design is generally vertical with a conical or flat base.
- An air disperser with a hot air supply system such as a main air filter, supply fan, air heater and air disperser (Figure 9.3, 3). The air aspiration is performed through filters, the type depending on the local conditions and the nature of...
the product to be treated. The air can be heated in two different ways: by direct heating (gas) and/or by indirect heating (vapor, gas, oil or electricity). The air flow chamber can be in co-current, counter-current or mixed mode.

- An atomizing device with a feed supply system such as feed tank, feed pump, water tank, concentrate heater and atomizing device. There are three types of atomizing device: rotary atomizer (wheel or disk), nozzle atomizer (pressure, pneumatic or sonic) and combined (rotary and pneumatic) (Figure 9.3, 3).

- A powder recovery system. Separation of the dried product can be achieved by a primary discharge from the drying chamber followed by a secondary discharge from a particulate collector (using a cyclone, bag filter or electrostatic precipitation), followed by total discharge from the particulate collector and finishing with final exhaust air cleaning in a wet scrubber and dry filter (Figure 9.3, 8 and 9).

According to Sougnez (1983), Masters (1991) and Pisecky (1997), the simplest types of installation are single-stage systems with a very short residence time (20–60 s). Thus, there is no real balance between the relative humidity of the air and the moisture content of the powder. The outlet temperature of the air must therefore be higher and as a result the thermal efficiency of the single-stage spray drier is then reduced. This type of drying chamber was the standard equipment for drying milk in the 1960s. Space requirements were small and building costs were low. Generally, installations without any post-treatment system are suitable only for non-agglomerated powders not requiring cooling. If necessary, a pneumatic conveying system could be added to cool the powder while transporting the chamber fraction and the cyclone fraction to a single discharge point.

The two-stage drying system consists of limiting the spray-drying process to a process with a longer residence time (several minutes) to provide a better thermodynamic balance. This involves a considerable reduction in the outlet air temperature and also an increase in the inlet air temperature. A second final drying stage is necessary to optimize the moisture content by using an integrated fluid bed (static) or an external fluid bed (vibrating), the air temperatures of which are 15–25°C lower than with a single-stage system to improve and/or preserve the quality of the dairy powder (Figure 9.3, 11 and 14). Consequently, the surrounding air temperature at the critical drying stage and the particle temperature are also correspondingly lower, thus contributing to further economy improvement. The integrated fluid bed can be either circular (e.g. Multi Stage Drier [MSD™] chamber) or annular (e.g. Compact Drier [CD] chamber).

Two-stage drying has its limitations but it can be applied to products such as skim milk, whole milk, pre-crystallized whey, caseinates, whey proteins and derivatives. The moisture content of the powder leaving the first stage is limited by the thermoplasticity of the wet powder, i.e. by its stickiness in relation to the water activity and the glass transition temperature (Roos, 2002). The moisture content must be close to 7–8, 9–10 and 2–3% for skim/whole milk, caseinate/whey protein and pre-crystallized whey powders respectively. The two-stage drying techniques can be applied to the production of both non-agglomerated and agglomerated powders, but this technique is very suitable for the production of agglomerated powders, by separating the non-agglomerated particles from the agglomerates (i.e. collecting the cyclone fractions...
and reintroducing these fine fractions [called fines] into the wet zone around the atomizer of the chamber).

The three-stage drying systems, with an internal fluid bed as a second stage in combination with an external vibrating fluid bed as a third-stage drier, first appeared at the beginning of the 1980s and were called Compact Drier Instantization (CDI) or MSD™. Today, they dominate the dairy powder industry (Figure 9.3). Three-stage systems combine all the advantages of extended two-stage drying, using spray drying as the primary stage, fluid bed drying of a static fluid as the second drying stage and drying on an external vibrating fluid bed as the third drying stage. The final drying stage terminates with cooling to under the glass transition temperature. Evaporation performed at each stage can be optimized to achieve both gentle drying conditions and good thermal economy.

The CD is suitable for producing both non-agglomerated and agglomerated powders of practically any kind of dried dairy product. It can also cope successfully with whey powders, fat-filled milk and whey products as well as caseinates, both non-agglomerated and agglomerated. It has a fat content limit of about 50% fat in total solids. Powder quality and appearance are comparable with those of products from two-stage drying systems but they have considerably better flowability and the process is more economical. In comparison with the CD, the MSD™ can process an even wider range of products and can handle an even higher fat content. The main characteristic of MSD™ powder is related to very good agglomeration and mechanical stability, low particle size fractions (below 125 μm) and very good flowability.

Optimization of the process allows considerable improvement in the drying efficiency and the quality of the product obtained is generally better. The various advantages are:

- improved thermal efficiency: significant reduction in the outlet air temperature, permitting an increase in the inlet air temperature;
- reduction in material obstruction: the capacity in one volume is two or three times higher than for a traditional unit;
- considerable reduction in powder emission to the atmosphere: a reduction in the drying air flow and an increase in powder moisture content decrease the loss of fine particles in the outlet air;
- improved powder quality in relation to the agglomeration level, solubility, dispersibility, wettability, particle size, density, etc.

There are other examples of drying equipment such as the “tall form drier”, the “Filtermat® drier,” the “Paraflash® drier” and the “Tixotherm® drier.” All these towers have characteristics related to the specific properties of the product being dried (e.g. high fat content, starch, maltodextrin, egg and hygroscopic products).

## Drying of proteins

The native properties of milk components are substantially unaffected by moderate drying conditions. Depending on the preheating conditions, drier design and temperature
Effects of drying on milk proteins

The properties of spray-dried powder may vary significantly. An evaporating milk droplet in a spray drier in co-current air flow initially does not appreciably exceed the wet bulb temperature and can be held effectively at temperatures below 60°C. As the falling temperature period is approached in the course of further evaporation, the temperature rises to a final value determined by the final temperature of the drying gas and the residence time in the drier. Under properly controlled spray drying conditions, the changes in milk protein structure and solubility are minor. Spray drying does not denature the whey protein significantly and the levels of denatured whey protein in dairy powders are more or less equal to those of condensed milk and heated milk, which is substantially more denatured than during spray drying. The best example is in relation to the whey protein nitrogen index (WPNI).

According to Pisecky (1997), the WPNI expresses the amount of undenatured whey protein (milligrams of whey protein nitrogen per gram of powder). It is a measure of the sum of heat treatments to which the milk has been subjected prior to evaporation and spray drying. The heat treatment of a concentrate and, after, of a powder has only a negligible effect on the WPNI. The main operation to adjust the required value is the preheating process, i.e. time/temperature combination. However, there are many other factors that influence the WPNI, including the total amount of whey protein and the overall composition of the processed milk as influenced by animal breed and seasonal variations. The individual design of the processing equipment, i.e. the pasteurizer and holding tubes, also has great significance. Therefore, it is difficult to predict the conditions of achieving the required WPNI on a general basis. Obviously, the primary purpose of preheat treatment is to ensure the microbiological quality of dairy products.

In milk powder production, the influence of the heat treatment on the denaturation of whey proteins for achieving the desired properties of the final products is just as important. SMP for cheese manufacture should have as much undenatured protein as possible, i.e. it should be low heat (WPNI > 6), whereas for bakeries high heat powder with high denaturation is required (WPNI < 1.5). For ice cream, chocolate and confectionery, medium heat powder is required. According to Schuck et al. (1994a), the use of microfiltration (pore diameter, 1.4 μm), coupled with a low heat treatment during vacuum evaporation, allows the production of a “low low heat” SMP with a WPNI close to 9 mg of whey protein nitrogen/g of powder, a bacterial count <1000 CFU/g powder, a solubility index >99.5%, a dispersibility index >98.5% and a wetability index <15 s. Such a powder after water rehydration has the same renneting time as the original raw milk and can be used as a reference powder for either industrial or scientific purposes.

The stability of protein powders during storage is critically affected by the moisture content and the storage temperature. More precisely, such stability is governed by the water activity (a_w) and the glass transition (T_g) temperature. The a_w should be close to 0.2 at 25°C for optimal preservation, with an ideal moisture content determined by using the sorption isotherm of some dairy powders. For example, the corresponding moisture contents for skim milk, whey and protein powders must be close to 4, 2–3 and 6% respectively. The optimal storage temperature must be below the T_g temperature, which is close to 40–50°C at 0.2 a_w.
Example of dairy protein concentrates and powders

In this study, micellar casein concentrate (MCC) was prepared by microfiltration and diafiltration (pore diameter, 0.1 μm) on an MFS 19 (Tetra Laval, Århus, Denmark; 4.6 m²) at 50°C, according to Fauquant et al. (1988) and Pierre et al. (1992) at 200 g/kg total solids. The whey protein concentrate (WPC) was obtained by ultrafiltration and diafiltration of the microfiltrate (0.1 μm) obtained on a DDS module (GEA, Soeborg, Denmark) with the plane membrane (10000 g/mol molecular weight cut-off, 9 m², 50°C) at 200 g/kg total solids. The sodium, calcium and potassium caseinate concentrates were reconstituted from sodium, calcium and potassium caseinate powder at 190 g/kg total solids. The microfiltration retentate (R4 MF) was obtained by microfiltration (0.1 μm) on an MFS 19 (Tetra Laval, Århus, Denmark; 4.6 m²) at 50°C. The volume reduction ratio was 4.

The ultrafiltration retentate (R4 UF) was manufactured by ultrafiltration on a 2-S37 module with M₁ membranes (Tech Sep, Rhône Poulenc, St Maurice de Beynost, France; 100 000 g/mol nominal molecular weight cutoff, 6.8 m²) at 50°C. The volume reduction ratio was 4. NaCl solution, CaCl₂ solution, sodium phosphate solution at pH 7.1 (for MCC) or pH 6.6 (for WPC) and sodium citrate solution at pH 7.1 (for MCC) or pH 6.6 (for WPC) in 205 5 g/kg total solids were added to the MCC or the WPC to obtain a concentrate with 12% (w/w) (NaCl, CaCl₂ and sodium phosphate solution) and 30% (w/w) (sodium citrate) of mineral salts/total solids. After addition of salt, the pH was adjusted to 7.1 (for MCC) or 6.6 (for WPC) with 1 N KOH (NaCl, CaCl₂ and sodium phosphate solution) or with 1 N HCl (sodium citrate) at 20°C.

The spray drying of the concentrates was performed at Bionov (Rennes, France) in a three-stage pilot plant spray drier (GEA, Niro Atomizer, St Quentin en Yvelines, France) according to Schuck et al. (1998a) and Bimbenet et al. (2002) to obtain a micellar casein powder (MCP) or a whey protein powder (WPP). The temperature of the concentrate before drying was 40°C for MCC and 20°C for WPC. The atomizer was equipped with a pressure nozzle (0.73 mm diameter orifice) and a four-slot core (0.51 mm nominal width), providing a 60° spray angle. The evaporation capacity was 70–120 kg/h (depending on the inlet and outlet air temperatures and the air flow). The pressure in the nozzle was 16 MPa. The inlet temperature was 208°C for WPC and 215°C for MCC, the integrated fluid bed air temperature was 70°C for MCC and WPC and the outlet temperature was 80°C for WPC and 70°C for MCC. The inlet air humidity was controlled and adjusted by a dehumidifier (Munters, Sollentuna, Sweden). For each MCP or WPP, two granulations were obtained (i.e. non-granulated [NG] and granulated [G] powders) by reintroduction of the fine particles after the cyclones at the top of the spray drier.

Research approach using drying by desorption

Principles

The concentrates were dried in a water activity meter (Novasina RTD 200/0, Pfäffikon, Switzerland) at 20°C (constant temperature). The concentrate (100 mg) was placed in a plastic support (area, 95 mm²) with a zeolite WE 291 drier below (7 g) (Bayer, Puteaux, France). This method was used to simulate the conditions
of spray drying by establishing a difference in vapor pressure equilibrium between the dairy concentrate and the drying air, and to determine the water transfer from inside the dairy concentrate to the surface. The relative humidity (RH) was measured versus time following water transfer from the dairy concentrate to the zeolite. The final slope of the absolute value of the decrease in RH ($\beta$) represented the ability to remove bound water from the solute at the end of the drying phase (Schuck et al., 1998b, 1999); the lower the $\beta$ slope, the greater the difficulty to remove water at the end of drying and the higher the bound water content.

Desorption results
The drying slopes ($\beta$) of the various dairy products tested are shown in Table 9.1. The dairy products, ranging from the highest to the lowest absolute value of the slope, were skim milk ($\beta = 0.90\%/min$), R4 UF ($\beta = 0.75\%/min$), R4 MF ($\beta = 0.70\%/min$) and MCC ($\beta = 0.34\%/min$). These results could be explained by a decrease in water diffusion through the dried product, i.e. the final residue obtained at the end of drying, when the micellar casein concentration increased. Water transport was probably affected by the high micellar casein content of the sprayed droplet in the atomization tower, and, similarly, when the powder granule was dissolved in water. These results are in accordance with the results of Schuck et al. (1994a, 1994b).

Drying of the caseinates showed that sodium caseinate and potassium caseinate dried more easily ($\beta = 0.64$ and 0.65%/min respectively) than calcium caseinate ($\beta = 0.51\%/min$) (Table 9.1). The limitation of water diffusion through the calcium caseinate may be explained by the structure of this colloidal dispersion. Whereas, in calcium caseinate, the casein sub-units are more aggregated because of calcium binding, in sodium and potassium caseinates, the caseins are more soluble. These results

| Table 9.1 Drying by desorption of dairy concentrates |
|---------------------------------|----------------|
| Dairy concentrate               | $\beta$ Slope (%/min) |
| Skim milk                       | 0.90            |
| R4 UF                           | 0.75            |
| R4 MF                           | 0.70            |
| MCC                             | 0.34            |
| WPC                             | 0.68            |
| Sodium caseinate                | 0.64            |
| Potassium caseinate             | 0.65            |
| Calcium caseinate               | 0.51            |
| WPC + NaCl                      | 0.24            |
| WPC + CaCl$_2$                  | 0.46            |
| WPC + Phosphate                 | 0.41            |
| WPC + Citrate                   | 0.48            |
| MCC + NaCl                      | 0.19            |
| MCC + CaCl$_2$                  | 0.36            |
| MCC + Phosphate                 | 0.49            |
| MCC + Citrate                   | 0.45            |

R, Retentate; UF, Ultrafiltration; MF, Microfiltration; MCC, Micellar Casein Concentrate; WPC, Whey Protein Concentrate
showed that the water bound in a micellar structure was more strongly bound than that bound to the soluble caseins in sodium caseinate. The situation was intermediate for calcium caseinate. We assumed that these differences in water transfer during drying could be explained by the casein structure. The decrease in water inside the dairy concentrate led to a decrease in the water concentration on the surface of the concentrate in the water activity meter or on the surface of the droplet during spray drying and decreased the drying kinetics.

These results were confirmed by the desorption drying of two different classes of proteins, i.e. MCC (micellar structure), with a β value of 0.34%/min, and WPC (globular structure), with a β value of 0.68%/min (Table 9.1). These two different types of protein had the same protein content (89% of total solids) and the same water content before desorption drying, but not the same drying time or β value. All these results show that the drying rate is dependent on the nature and the structure of the casein. Water may be less available during the drying of a protein with a micellar structure than during the drying of a protein with a globular structure.

Mineral additions to WPC decreased the β value, the smallest decrease being with citrate (29% reduction) and the greatest decrease being with NaCl (65% reduction) (Table 9.1). A decrease in the β value means a lower rate of water transfer at the end of drying (Schuck et al., 1998b). The decrease in water transfer during the drying of modified WPC could be explained by the high hygroscopicity of the added mineral salts. This result suggested that the water is more closely bound to the mineral salts than to the whey proteins at the end of drying. The mineral addition to WPC under the test conditions had little effect on the whey protein structure but probably had some effect on the increase in bound water in the modified WPC.

Addition of NaCl to MCC decreased the β value (0.19%/min) (Table 9.1). Water in a NaCl-containing casein system is more rotationally mobile than water in a casein model system without NaCl, with the same water activity (Curme et al., 1990). For high electrolyte concentrations, the amount of bound water decreases because of the suppression of the electrical double layer surrounding the protein molecule; this is directly related to the hydration of ions (Na⁺, Cl⁻) and hence to the ability to separate water molecules from the protein molecules (Robin et al., 1993). Water is less closely bound to micellar casein in the presence of NaCl (Cayot and Lorient, 1998). The decrease in water transfer at the end of drying can be explained by the hygroscopicity of NaCl. The water is more closely bound to NaCl than to the micellar caseins.

Addition of CaCl₂ to MCC increased the β value (0.36%/min) (Table 9.1). Moreover, addition of CaCl₂ to milk decreases micellar solvation (Tarodo de la Fuente, 1985; Van Hooydonk et al., 1986; Jeurnink and de Kruif, 1995; Le Ray et al., 1998). Firstly, the water inside the micellar structure in the case of MCC without salt addition might be less available than the water inside the micellar structure in the case of MCC with CaCl₂. Secondly, the lower water content inside micellar structures with the addition of calcium might lead to an increase in water transfer during drying.

Addition of phosphate ions to MCC increased the β value (0.49%/min) (Table 9.1). The increase in water transfer was explained by the partial solubilization of caseins (Le Ray et al., 1998), although an increase in casein micelle solvation and an increase in viscosity were observed. These results may be discussed in terms of the
strength of water binding to caseins, either in a soluble form as in sodium caseinate (Schuck et al., 1998b) or in the micellar structure (MCC) as in the current experiments reported here. The water bound to micellar caseins was probably less easy to remove than the water bound to soluble caseins. Partial solubilization of caseins improved the water transfer during dehydration of phosphate solution + MCC.

Addition of citrate solution to MCC at 30% total solids increased the β value (0.45%/min) (Table 9.1). Addition of citrate induces the release of large amounts of soluble casein from the micellar phase because of solubilization of colloidal calcium phosphate (Le Ray et al., 1998). Similar results occurred with the addition of phosphate ion. Solubilization of the micellar casein improved the water transfer during dehydration of solution with added citrate.

**Industrial implications**

**Introduction**

Several studies (Masters, 1991; Pisecky, 1997) have reported that the moisture content can vary according to the product for the same outlet air temperature in the spray drier. For example, Pisecky (1997) has reported that the moisture content is close to 4% and 5% for WMP and SMP respectively for an outlet air temperature close to 85°C. On the other hand, to produce WMP and SMP at the same moisture content (4%, for example), the outlet air temperatures must be different (80 and 90°C respectively). All these differences can be explained by the effects of the chemical composition of the ingredients on the availability of the water that must be transferred from the droplet to the drying air.

The aims of this chapter were to evaluate water transfer during the spray drying of different dairy concentrates using thermodynamic and chemical approaches. Whey protein concentrates and isolates (WPC35, WPC50, WPC70, WPI90) with or without heat denaturation, MC, sodium caseinate (NaCas) and milk with and without whey protein enrichment were dried in a three-stage pilot plant spray drier. When the concentrate temperature, air flow rate, concentrate flow rate, total solids content of the concentrate, inlet air temperature absolute humidity, inlet air temperature before and after heating, and outlet air temperature after drying are known, it is possible to determine the specific energy consumption (SEC), which is the ratio of the energy consumed to the evaporation of 1 kg of water (measured in kJ/kg water) (Bimbenet et al., 2002).

Thus, if free water is spray dried the energy spent in terms of the SEC would be close to 2500 kJ/kg water. If the concentrate has greater and greater amounts of bound water to free water, the SEC increases up to 10,000 kJ/kg water. The significance of a very high SEC is related to the water, which is less and less available, limiting water transfer, and thus increasing the surface temperature of the droplet and hence increasing the risk of protein denaturation of the powder.

**Whey proteins**

The results presented in Table 9.2 show that water transfer during spray drying decreased when the whey proteins were native proteins. For the same moisture, the
SEC for drying was higher when (a) the native whey protein content increased in WPC and in milk and (b) the whey proteins were heat denatured in WPC35, but the SEC was lower when (c) the whey proteins were heat denatured in WPC50, WPC70 and WPI90. These results may be explained by the availability of the water (bound and unbound) in the concentrate in relation to the nature and the content of the whey proteins.

Caseins
The results presented in Table 9.2 show that water transfer during spray drying decreased when the micellar casein content increased. For the same moisture, the SEC for drying was higher when (a) the micellar casein content increased in MC compared with skim milk and (b) casein remained in a micellar state (as in MC) compared with a soluble state (e.g. in NaCas). These results may be explained by the availability of the water in the concentrate in relation to the content and the structure of the caseins. Water is more available when the caseins are soluble than when they are in a micellar state.

All these results also show that water transfer depends on the relationship between the water and the protein components and that these components should be taken into account when optimizing spray-drying parameters for proteins.

Rehydration of protein powders
Most food additives are prepared in powder form and need to be dissolved before use. Water interactions in dehydrated products and dissolution are thus important factors in food development and formulation (Hardy et al., 2002). Dissolution is an essential quality attribute of a dairy powder as a food ingredient (King, 1966). Many sensors and analytical methods such as the insolubility index (International Dairy Federation, 1988; American Dairy Products Institute, 1990), nuclear magnetic resonance (NMR) spectroscopy (Davenel et al., 1997), turbidity, viscosity and particle size distribution (Gaiani et al., 2006) can now be used to study water transfer in dairy protein concentrates during rehydration. Using combinations of these methods, it is very easy to determine the different stages of the rehydration process, i.e. wettability, swellability, sinkability, dispersibility and solubility.
The insolubility index (ISI, in %), described by the IDF standard (International Dairy Federation, 1988), for skim milk, is the volume of sediment (for 50 mL) after rehydration (10 g of powder in 100 mL of distilled water, at 25°C), mixing (for 90 s, at 4000 rev/min) and centrifugation (for 300 s, at 160 g). With this method, the quantity of insoluble material (true and false not differentiated) can be determined.

NMR spectroscopy is a technique for determining the rate of dissolution, the time required for complete reconstitution of powders and the transverse relaxation rate of reconstituted solutions. A 40 mm diameter glass tube filled with 20 mL of water at 40°C was put into the gap of the magnet of a Minispec Bruker PC 10 NMR spectrometer operating at a resonance frequency of 10 MHz. A suitably designed funnel and an electric stirrer (glass spatula) were inserted into the tube. The method was first described by Davenel et al. (1997). They showed that the solubilization rate was independent of the quantity of powder poured (up to 20 g powder/100 mL water) and increased with the stirring rate. In subsequent experiments, the rotation rate of the stirrer was adjusted after starting to 1150 rev/min for spray-dried powders and 1 g of powder was poured into the water.

The NMR measurements were generally continued until the solution was completely reconstituted, except if insoluble material was formed. Each decay curve was obtained by sampling a maximum of 845 spin echoes of a Carr-Purcell-Meiboom-Gill (CPMG) sequence every 20 s during the reconstitution period. Interpulse spacing between 180° pulses was fixed at 2 ms to limit the effect of diffusion caused by stirring. The NMR kinetic method was used in triplicate. The CPMG curves were well approximated by the sum of two exponential curves to determine the protons attributed to water protons in fast exchange with exchangeable protons of non-dissolved powder particles, and the protons attributed to water protons and exchangeable protons in the reconstituted phase (Davenel et al., 1997). With this method, it is possible to differentiate between the truly insoluble material and the falsely insoluble material. The falsely insoluble material can be explained by the low water transfer during rehydration and not by denatured protein, which is truly insoluble (Schuck et al., 1994b).

For viscosity measurement, a rheometer can be used to obtain viscosity profiles. In our study, the blades were placed at right angles to each other to provide good homogenization. Industrial dissolution processes usually include stirring at a constant speed and the experiments were therefore designed to provide a constant shear rate (100 s⁻¹). MCP was added to the rheometer cup manually. The aqueous phase used was distilled water at a volume of 18 mL. The powder was dispersed in the rheometer cup 50 s after starting the rheometer. Dissolution is highly dependent on temperature and concentration. The total nitrogen concentration employed to study these effects was about 5% (w/v) and the temperature was about 24°C (Gaiani et al., 2005, 2006).

The experiments to provide the turbidity profiles were carried out in a 2 L vessel equipped with a four-blade 45° impeller rotating at 400 rev/min. A double-walled jacket vessel maintained the temperature at 24°C. The turbidity sensor was placed 3 cm below the surface of the water and was positioned through the vessel wall to avoid disturbance during stirring. Turbidity changes accompanying powder rehydration.
were followed using a turbidity meter. The apparatus used light in the near-infrared region (860 nm), the incident beam being reflected back at 180° by any particle in suspension in the fluid to a sensitive electronic receptor (Gaiani et al., 2005).

A laser light diffraction apparatus with a 5 mW He–Ne laser operating at a wavelength of 632.8 nm can be used to record particle size distributions. The particle size distribution of dried particles was determined using a dry powder feeder attachment and the standard optical model presentation for particles dispersed in air was used. To measure the particle size distribution of micellar casein in concentrates, 0.5 mL of suspension was taken from the rheometer cup and introduced into 100 mL of pre-filtered distilled water (membrane diameter, 0.22 μm) to reach the correct obscuration. The results obtained corresponded to average diameters calculated according to the Mie theory. The criterion selected was d(50), meaning that 50% of the particles had diameters lower than this criterion (midpoint of cumulative volume distribution) (Gaiani et al., 2005, 2006).

Using this combination of three methods, it was possible to follow the water transfer during rehydration and to obtain the wetting time, determined using the first peak of increased viscosity and turbidity, and the swelling time, determined using the second peak of viscosity in relation to the increase in particle size. The rehydration time was then determined according to stabilization of the viscosity, turbidity and particle size values.

The results in Table 9.3 show that rehydration of MCP occurs in different stages: first there is wetting and swelling of the particles, followed by slow dispersion to reach a homogeneous fluid, in agreement with Gaiani et al. (2005, 2006). Using an NMR method, Davenel et al. (1997) also demonstrated two stages during MC rehydration, attributed to water absorption by powder and solubilization of particles (i.e. swelling and dispersion stages). They estimated the water uptake by the powder at around 5 g water/g powder during the first 20 min of rehydration but could not identify a wetting stage with this method.

MCPs with a high ISI (14.5 mL) are generally considered to be poorly soluble powders in which rehydration of the micelle remains incomplete (Jost, 1993).

<table>
<thead>
<tr>
<th>Powders</th>
<th>RP using NMR (min)</th>
<th>ISI using IDF Standard (mL)</th>
<th>WT (min)</th>
<th>ST (min)</th>
<th>DT + SolT (min)</th>
<th>RT (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCP G</td>
<td>22</td>
<td>14.5</td>
<td>1</td>
<td>2</td>
<td>804</td>
<td>807</td>
</tr>
<tr>
<td>MCP NG</td>
<td>8</td>
<td>3.5</td>
<td>3</td>
<td>17</td>
<td>551</td>
<td>571</td>
</tr>
<tr>
<td>MCP + Carbohydrate G</td>
<td>18</td>
<td>5.0</td>
<td>1</td>
<td>nm</td>
<td>nm</td>
<td>116</td>
</tr>
<tr>
<td>MCP + Carbohydrate NG</td>
<td>nm</td>
<td>nm</td>
<td>2</td>
<td>0</td>
<td>95</td>
<td>97</td>
</tr>
<tr>
<td>MCP + NaCl G</td>
<td>9.5</td>
<td>0.9</td>
<td>nm</td>
<td>nm</td>
<td>nm</td>
<td>nm</td>
</tr>
<tr>
<td>MCP + CaCl₂ G</td>
<td>∞</td>
<td>14.5</td>
<td>nm</td>
<td>nm</td>
<td>nm</td>
<td>nm</td>
</tr>
<tr>
<td>MCP + SCS/SPS G</td>
<td>6/5</td>
<td>&lt;0.5</td>
<td>nm</td>
<td>nm</td>
<td>nm</td>
<td>nm</td>
</tr>
<tr>
<td>WPP G</td>
<td>5</td>
<td>&lt;0.5</td>
<td>4</td>
<td>0</td>
<td>0</td>
<td>4</td>
</tr>
<tr>
<td>WPP NG</td>
<td>15</td>
<td>&lt;0.5</td>
<td>17</td>
<td>0</td>
<td>0</td>
<td>17</td>
</tr>
</tbody>
</table>

MCP, micellar casein powder; G, granulated; NG, non granulated; SCS, sodium citrate solution; SPS, sodium phosphate solution; WPP, whey protein powder; RP, reconstitution period; ISI, insolubility index; WT, wetting time; ST, swelling time; DT, dispersibility time; SolT, solubility time; RT, rehydration time = WT + ST + DT + SolT; ∞, infinite delay; nm, not measured
Addition of NaCl to the MC concentrate before spray drying considerably reduced the ISI and reconstitution period (RP) values (ISI 0.9 mL; RP 9.5 min) (Table 9.3). It has been hypothesized that the significant decrease in the RP value is more probably related to the hygroscopic strength of NaCl.

Addition of sodium citrate solution (SCS) or sodium phosphate solution (SPS) resulted in fast solubilization, as shown by the very low RP value and by the ISI value lower than 0.5 mL (Table 9.3). The resulting solution consisted of casein micelles in the form of sodium caseinate, associated with the occurrence of a single proton population, characterized by NMR relaxation rates that were much lower than the relaxation rate measured with reconstituted MC. This could be attributed to a decrease in the amount of hydration water induced by the change in micelle structure. The transparency of the solution indicated the formation of soluble caseins related to greater quantities of calcium complexes.

Reconstitution of MCP in the presence of CaCl$_2$ led to considerable changes in protein structure, associated with instability of the casein micelles, which began to precipitate just after mixing, as shown by the high ISI and the non-measurable RP value due to experimental delay. This precipitate probably resulted from aggregation of casein micelles or sub-micelles through a decreasingly negative charge on the protein by additional calcium binding, leading to a reduction in electrostatic repulsion (Dalgleish, 1982). In this case, the high ISI of these solutions was related to the presence of insoluble substances whereas, in the case of rehydration of MCP, the high ISI represented only the low water transfer rate in the casein (Table 9.3).

On the other hand, the rehydration of whey powders was totally different (Table 9.3). As the wettability of whey powders is poor, the instability at the beginning of the profile could be due to lump formation going past the sensor, as reported by Freudig et al. (1999). For non-granulated (NG) WPI powder, the very long signal instability could be explained by a tendency for the lumps to be stuck together in a thick layer of wet particles, due to the small size of the particles (Kinsella, 1984). Powder swelling was not reported for WPI powders, probably because globular protein powders bind less water than intact casein micelle powders (Kinsella, 1984; Robin et al., 1993). De Moor and Huyghebaert (1983) also reported that whey powders have a lower water holding capacity than casein powder.

As expected, granulation had a positive effect on wetting. The wetting time was systematically better for granulated particles. This phenomenon is well known, as fast wetting is enhanced, with large particles forming large pores, high porosity and small contact angles between the powder surface and the penetrating water (Pisecky, 1986; Freudig et al., 1999; Gaiani et al., 2005). A surprising influence of granulation on the rehydration time was observed. Depending on the nature of the protein, the granulation influence involved opposite effects. WPI rehydration was enhanced for granulated particles whereas the rehydration time was shorter for non-granulated particles of MCP. The controlling stage for whey proteins is wetting (Baldwin and Sanderson, 1973; Schubert, 1993). As granulation improves the wetting stage, the rehydration of whey powders is enhanced for granulated particles. In contrast, the controlling stage for casein proteins is dispersion. Indeed, even with a shorter wetting time, a granulated powder is slower to rehydrate than a non-granulated powder (Gaiani et al., 2005).
These results are not compatible with those of other studies, in which it was generally accepted that a single particle size around 200\(\mu\)m (Neff and Morris, 1968) or 400\(\mu\)m (Freudig et al., 1999) represented optimal dispersibility and sinkability. In fact, this optimal particle size depends on the composition of the dairy powder. As shown in Table 9.3, if the industry wishes to optimize powder rehydration, it seems to be better to rehydrate granulated powders if the protein is whey and to rehydrate non-granulated powders if the protein is casein.

**Conclusions**

The aim of this chapter was to explain the process of dehydration, i.e. spray drying, in order to understand the effects of spray drying on the quality of protein powders (micellar caseins [MC] and globular proteins [WPC]) during drying and rehydration. We then demonstrated that the quality of these powders depends on the chemical environment.

It is very important for the dairy industry to understand that enrichment of milk in micellar casein (by ultrafiltration or microfiltration) decreases water transfer during the drying and rehydration processes. Insolubility (International Dairy Federation, 1988) is related to the decrease in water transfer required for rehydration and not to thermal denaturation, and decrease in water transfer is related to the micellar structure. The destabilization of the micellar structure induced by the addition of phosphate or citrate solution to MC increases water transfer during drying and during rehydration. Water transfer in WPC or MC with added carbohydrates is improved during rehydration. Addition of NaCl to MC decreases water transfer during drying but increases water transfer during rehydration, and thus is related to the hygroscopicity of the carbohydrate and the NaCl.

The industrial requirement for protein powders with specific properties is expanding. As powder is the easiest way to carry and store milk derivatives, an understanding of the rehydration behavior of a dairy powder will become more and more important in the future.

Moreover, it is essential for both dairy powder producers and dairy powder users to have a method to evaluate the rehydration behavior of dairy powders. As demonstrated in several studies, the industry should take into account certain technological factors such as granulation and the incorporation mode, and also the nature of the protein being rehydrated, to optimize the rehydration of a dairy powder. In contrast to other studies, we found that improving the wetting stage by using granulated powders did not systematically improve the total rehydration. Depending on the nature of the protein, it seems to be better to work with granulated (for whey) or non-granulated (for micellar casein) powders to obtain more rapid rehydration (Gaiani et al., 2007).

In conclusion, water transfer in dairy protein concentrates during dehydration and during rehydration depends on the aqueous environment, the nature of the mineral salts and the structure of the dairy proteins (MC or WPC). The water–protein interaction requires further study, to understand the effects of preheat treatment and spray drying on the functional properties of protein powders.
References


Changes in milk proteins during storage of dry powders

Kerianne Higgs and Mike Boland

Abstract

Milk proteins undergo chemical changes, even in dried powders. This chapter reviews the changes undergone by caseins and whey proteins in milk powders and in purified protein products. Maillard compounds are of particular importance in milk powders, milk protein concentrates and whey protein concentrates, i.e. products where lactose is present. Caseins undergo isopeptide bond formation, as the result of dephosphorylation of phosphoserine and subsequent reactions of the dehydroalanine produced.

We discuss the nutritional significance of these changes. Both the Maillard reaction and the formation of isopeptides lead to loss of bioavailable lysine. This is not a problem in dairy proteins, which contain an excess of lysine, but it can affect the nutritional value of protein blends which may be limiting in lysine.

Introduction

Milk is an unstable foodstuff, prone in particular to microbiological degradation, but also to long-term chemical change. A large part of the world’s dairy production occurs in areas remote from the markets in which it is consumed and production is often seasonal, requiring storage to smooth out supply. The production of milk powders and other dried milk-protein-containing products has been the method of choice for over
100 years for the storage and shipping of milk over long distances and/or times, as it confers stability and massively reduces weight and bulk.

Milk powders were known to the Chinese and were described by Marco Polo. The production of milk powders was described by Nicolas Appert in the early nineteenth century and commercial processes for the spray drying of milk were patented in the USA in 1872 and 1905. This opened the way for large-scale industrial production of milk powders throughout the twentieth century.

Table 10.1 summarizes the biggest exporters and importers of milk powders in 2006. In the same period, more than 7 million tonnes of milk powder was produced globally.

In addition to milk powders, dried dairy protein products that are traded on the world market, largely as food ingredients, include casein and caseinate, whey powders, whey protein concentrates, whey protein isolates, milk protein concentrates, milk protein isolates and specialist nutritional powders and blends, which may also contain hydrolyzed dairy proteins.

Milk powders are used primarily for making reconstituted and recombined milks, usually sold to consumers in UHT format, although substantial amounts are used to make other dairy products such as yoghurt and ice cream as well as being minor ingredients in a wide range of non-dairy foods.

The main uses for dried milk protein products are nutritional and products include infant formulas, medical foods, specialist foods for weight management and foods for muscle building (where high protein and high levels of branched-chain amino acids are desirable). They are also used in non-nutritional applications including desserts, confectionery, toppings, imitation cheeses, sauces and dressings and coffee whiteners, where their functional properties are important.

These products, especially those from New Zealand and Australia, often have long storage times because of geographic distance to market and seasonal production. Research from our laboratories and that of others has shown that a range of reactions can occur in the dry powders and that they can affect powder functionality and nutritional value. The quality of dried milk protein products deteriorates on storage at ambient temperatures because of two main reactions: the Maillard reaction and isopeptide bond formation. There are also some minor reactions, which are covered later.

<table>
<thead>
<tr>
<th>Table 10.1</th>
<th>Main exporters and importers of milk powders in 2006 (000 tonnes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Export</td>
<td>New Zealand</td>
</tr>
<tr>
<td>SMP</td>
<td>245</td>
</tr>
<tr>
<td>WMP</td>
<td>630</td>
</tr>
<tr>
<td>Total</td>
<td>875</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Import</th>
<th>Philippines</th>
<th>Mexico</th>
<th>Algeria</th>
<th>Indonesia</th>
<th>Russia</th>
<th>China</th>
</tr>
</thead>
<tbody>
<tr>
<td>SMP</td>
<td>135</td>
<td>172</td>
<td>110</td>
<td>135</td>
<td>75</td>
<td>–</td>
</tr>
<tr>
<td>WMP</td>
<td>52</td>
<td>–</td>
<td>60</td>
<td>–</td>
<td>30</td>
<td>60</td>
</tr>
<tr>
<td>Total</td>
<td>187</td>
<td>172</td>
<td>170</td>
<td>135</td>
<td>105</td>
<td>60</td>
</tr>
</tbody>
</table>

Source: Barry Wilson’s Dairy Industry News, 18:8, August 2006
Standard abbreviations for many of the dried milk products are used throughout the world and are used in this chapter: SMP = skim milk powder; WMP = whole milk powder; WPC = whey protein concentrate; WPI = whey protein isolate (usually >85% protein); MPC = milk protein concentrate; MPI = milk protein isolate. Milk powders sold internationally must conform to standard protein levels and fat levels for WMP. A number following the abbreviation for the other products is the percentage of protein by weight in the dry powder. Caseins and caseinates are not usually abbreviated and are typically >90% protein by dry weight.

A range of chemical reactions that modify proteins can occur in dried milk products, particularly at elevated temperature. The most important of these both involve lysyl side chains and are the formation of Maillard and pre-Maillard compounds, in the presence of sugars, and the formation of isopeptide bonds, particularly in products containing phosphoseryl residues, such as casein.

The formation of Maillard and pre-Maillard compounds

The Maillard reaction occurs when lysine-containing proteins interact with reducing sugars. The first stable compound formed during the Maillard reaction is the Amadori product (so-called because it is the result of a class of reaction called the Amadori rearrangement), shown in Figure 10.1. These compounds block the ε-amino groups of lysine residues, reducing the bioavailability of that essential amino acid. This reaction is dependent on a reducing sugar, usually lactose, being present as the co-reactant (Erbersdobler, 1986).

Highly pure protein products such as casein and caseinate do not suffer significantly from this reaction, as not enough lactose is present. The reaction is particularly important in WMPs, SMPs, WPCs and MPCs and can occur to a limited extent in some MPIs and WPIs. The rate of reaction is critically dependent on the level of moisture (water activity) and the temperature as well as the lactose content.

Advanced Maillard reaction products are partly responsible for the development of aromas and color during food processing and preparation.

Measuring lactulosyl lysine levels

Lactulosyl lysine formation is conveniently monitored by measuring furosine concentrations and can be indirectly measured by monitoring the amount of available amine. The results from both these methods have been shown to correlate well with the lactulosyl lysine measured in WPCs directly by mass spectrometry (Figure 10.2).

Rates of formation of lactulosyl lysine

We studied the rate of lactosylation for WPC products as dry powders. The rates were found to be dependent on the lactose concentration (a consequence of processing to reach desired protein levels), water activity ($a_W$) and temperature ($T$). The trials lasted
only a few months and extrapolation beyond that timeframe cannot be done with confidence.

Detailed kinetics developed using WPC80 for a range of \( a_W \) and \( T \) values allowed the rates of lactosylation to be predicted for periods up to 4 months.

For the kinetic evaluation, Figure 10.1 can be simplified to

\[
\text{reactants} \quad \xrightarrow{k_1} \quad \text{lactulosyl lysine} \quad \xrightarrow{k_2} \quad \text{advanced Maillard products}
\]

where \( k_1 \) and \( k_2 \) are the rate constants for the formation and degradation of lactulosyl lysine.

The rate equation for the formation of lactulosyl lysine is

\[
\frac{dL}{dt} = k_1[\text{reactants}]^{n_1} - k_2[L]^{n_2}
\]  \hspace{1cm} (10.1)

where \( L \) represents lactulosyl lysine and \( n_1 \) and \( n_2 \) are the rate orders for the formation and degradation of lactulosyl lysine respectively.
Because the formation of lactulosyl lysine in dairy powders uses only a fraction of the available reactants, the first reaction can be considered to be zero order and the degradation of lactulosyl lysine is a first-order process. This simplifies the equation to

$$\frac{dL}{dt} = k_1 - k_2[L]$$  \hspace{1cm} (10.2)
Rearrangement of this equation and solving for \([L]_t\) gives

\[
[L]_t = \frac{k_1}{k_2} - \left( \frac{k_1}{k_2} - [L]_0 \right) \exp(-k_2 t)
\]  

(10.3)

Furosine is a hydrolysis product of lactulosyl lysine and gives a direct measure of its concentration (Figure 10.2). Therefore, furosine can be substituted for lactulosyl lysine in the rate equations.

The samples used were individual samples that were removed from controlled storage at seven time points (2, 6, 12, 24, 40, 78 and 116 days). They were analyzed for furosine and the values were plotted against time. The rates were determined using non-linear regression with Sigma Plot 8.0. At 40°C, the samples at time points after 40 days for water activities of 0.54 and 0.80 showed advanced Maillard browning and were not included in the regression analysis. \(R^2\) values for the regressions were between 0.95 and 0.99.

The lactosylation rate constants were greater at higher temperatures and at higher water activities. The rates were low at low water activity \((x_W = 0.33)\), with small increases with temperature; however, at higher water activities, the rates increased substantially with increasing temperature (Figure 10.3).

It was observed for a number of WPC80 specifications that lactosylation appeared to stop when only 2.5 (average) or 3 lactose molecules had been bound per \(\beta\)-lactoglobulin molecule. This corresponded to about 20% of the total lysine being blocked. This condition was specific to the 80% protein products, which contained

![Figure 10.3](image_url)  

*Figure. 10.3* Rate constants for the formation of furosine in a WPC80.
about 12% lactose. WPC56 products showed a much greater degree of lactosylation. However, it should be noted that much higher levels of lactosylation have reportedly been seen in overseas laboratories in WPC80 samples stored for long periods (W. J. Harper, Ohio State University, 2003, personal communication).

Formation of isopeptide bonds

Isopeptide bonds are formed largely by the breakdown of the phosphoseryl side chains that are present in products containing casein, to form dehydroalanyl side chains (Friedman, 1999). The latter are reactive and will form cross-links, mainly with adjacent lysyl (but also with histidinyl or cysteinyl) side chains to form lysinoalanyl, histidinoalanyl or lanthionyl isopeptides respectively. This reaction is not known to be significant in whey products, which do not contain significant amounts of phosphoseryl residue.

The main isopeptide product on digestion gives lysinoalanine, which renders lysine non-bioavailable. Additional minor reactions form histidinoalanine and lanthionine on digestion. The latter compound, although only a minor component, is also important because it renders cysteine partially non-bioavailable and the sulfur amino acids are often nutritionally limiting in milk proteins. (Note—lanthionine formation blocks the bioavailability of cysteine, which is not normally considered to be an essential amino acid, because it can be synthesized from methionine; however, methionine is itself a nutritionally limiting amino acid in casein.) Studies have indicated that, although lanthionine and histidinoalanine linkages are formed under the alkaline conditions encountered during processing, it is only lysinoalanine that is formed in the neutral conditions encountered in powders.

Lysinoalanine is usually measured directly in protein hydrolysates as part of an extended amino acid analysis. In casein products, measurement of available amine is a good alternative.

Rates of formation of lysinoalanine

The rates of formation of lysinoalanine in caseinates have been investigated by W. Thresher (1996, 1997, personal communications), for a range of temperatures and water activities. The rate constants for lysinoalanine formation as a function of temperature are shown in Figure 10.5.
Amino acids other than lysine

Cysteine, methionine and tryptophan are other essential amino acids that could be rendered non-bioavailable by reacting during processing and/or storage. Cysteine undergoes β-elimination to give dehydroalanine when treated with alkali. This can then react with a lysine residue to give lysinoalanine. Cysteine can also react with dehydroalanine to give lanthionine. Chemically determined values for cysteine and lysine availability have been found to correlate well with rat protein efficiency ratios for heat- and alkali-treated caseinates (W. Thresher, 1996, personal communication).

Tryptophan residues are relatively stable during processing and storage. They are not easily oxidized and have been found to be relatively resistant to oxidizing lipids, alkali, quinones and reducing sugars (Nielsen et al., 1985). Any losses are small and not significant when compared with losses of other amino acids such as methionine and lysine. However, we have seen small amounts of oxidized tryptophan residues in digests of trim milk purchased from the supermarket (Figure 10.6).

Methionine is relatively easily oxidized to the sulfoxide, but methionine in the sulf oxide form is still bioavailable (Nielsen et al., 1985).

Table 10.2 indicates levels of key essential amino acids following either alkali treatment of casein or extensive lactosylation of WPC. Note that the losses of amino acids other than lysine are considerably lower than the losses of lysine. The conditions used for the casein are well beyond any normal exposure during processing or storage.
Figure. 10.6 Time-of-flight mass spectrometry of the $M^{+2}$ ion from the tryptophan-containing $\kappa$-casein peptide SPAQILQWQVLSNTVPAK. The chemical structures in the figure show the various levels of oxidation found.

Table 10.2 Amino acid concentrations of a control and an alkali-treated casein, and a control and a lactosylated WPC56 (Sources: Nielsen et al., 1985; K Higgs, unpublished results)

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>Concentration (mg/g crude protein)</th>
<th>Concentration (mg/g sample)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control casein</td>
<td>Alkali-treated casein$^a$</td>
</tr>
<tr>
<td>Tryptophan</td>
<td>13.4</td>
<td>12.1 (90%)</td>
</tr>
<tr>
<td>Lysine</td>
<td>91.0</td>
<td>60.8 (67%)</td>
</tr>
<tr>
<td>Methionine</td>
<td>31.4</td>
<td>25.3 (80%)</td>
</tr>
<tr>
<td>Cysteine</td>
<td>4.4</td>
<td>3.3 (75%)</td>
</tr>
</tbody>
</table>

$^a$Casein was heated for 4 h at 80°C in 0.15 M NaOH (Nielsen et al., 1985)

$^b$WPC was heated at 40°C, $\delta_w$ 0.75 for 100 h. The median number of lactosyl groups on $\beta$-lactoglobulin was 5 (K Higgs, unpublished results)

Implications for nutritional value of milk proteins

Milk protein is rich in essential amino acids, with many, including lysine, well exceeding recommended requirements (Table 10.3). Use of these proteins as the predominant nutritional source thus poses few problems if some of the lysine is non-bioavailable.
Changes in milk proteins during storage of dry powders

Lysine

The main concern during the storage of most nutritional proteins is the loss of lysine as a result of Maillard reactions or isopeptide bond formation.

Lactulosyl lysine renders lysine non-bioavailable. The protein efficiency ratio (PER) was found to be decreased in a WPC56, with an average of five lactulosyl lysine residues per protein molecule (Figure 10.7). A more detailed study using skim milk diets with pigs confirmed that lactulosyl lysine was non-bioavailable (Rerat et al., 2002). That study also found a decrease in the digestibility of lysine, phenylalanine, valine, cystine, aspartic acid, glycine and methionine residues. This decrease

<table>
<thead>
<tr>
<th>Essential amino acid (AA)</th>
<th>Recommended requirement** (mg/g protein)</th>
<th>Caseinate</th>
<th>MPI</th>
<th>WPC</th>
<th>Soy</th>
<th>Wheat</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isoleucine</td>
<td>28</td>
<td>46</td>
<td>44</td>
<td>54</td>
<td>47</td>
<td>33</td>
</tr>
<tr>
<td>Leucine</td>
<td>66</td>
<td>91</td>
<td>103</td>
<td>119</td>
<td>85</td>
<td>68</td>
</tr>
<tr>
<td>Lysine</td>
<td>58</td>
<td>77</td>
<td>81</td>
<td>94</td>
<td>63</td>
<td>27</td>
</tr>
<tr>
<td>Sulfur AA**</td>
<td>25</td>
<td>33</td>
<td>39</td>
<td>52</td>
<td>24</td>
<td>39</td>
</tr>
<tr>
<td>Aromatic AA</td>
<td>63</td>
<td>106</td>
<td>102</td>
<td>68</td>
<td>97</td>
<td>78</td>
</tr>
<tr>
<td>Threonine</td>
<td>34</td>
<td>43</td>
<td>45</td>
<td>66</td>
<td>8</td>
<td>29</td>
</tr>
<tr>
<td>Tryptophan</td>
<td>11</td>
<td>12</td>
<td>14</td>
<td>20</td>
<td>11</td>
<td>11</td>
</tr>
<tr>
<td>Valine</td>
<td>35</td>
<td>57</td>
<td>57</td>
<td>51</td>
<td>49</td>
<td>43</td>
</tr>
<tr>
<td>Histidine</td>
<td>19</td>
<td>29</td>
<td>27</td>
<td>21</td>
<td>29</td>
<td>—</td>
</tr>
</tbody>
</table>

*For 2–5 year olds
**Includes cysteine, cystine and methionine

Figure. 10.7 Plot of lactosylation level against PER. The PER value for ANRC casein was used for zero lactosylation.

Lysine

The main concern during the storage of most nutritional proteins is the loss of lysine as a result of Maillard reactions or isopeptide bond formation.

Lactulosyl lysine renders lysine non-bioavailable. The protein efficiency ratio (PER) was found to be decreased in a WPC56, with an average of five lactulosyl lysine residues per protein molecule (Figure 10.7). A more detailed study using skim milk diets with pigs confirmed that lactulosyl lysine was non-bioavailable (Rerat et al., 2002). That study also found a decrease in the digestibility of lysine, phenylalanine, valine, cystine, aspartic acid, glycine and methionine residues. This decrease
suggests that lactulosyl lysine residues hinder the release and therefore utilization of adjacent amino acids.

Lysinoalanine is not a bioavailable source of lysine (Robbins et al., 1980; Friedman, 1999). Alkali treatment of casein with 0.2 N NaOH at 80°C for 1 h reduced the PER of casein from 3.09 to 0.02 for a diet containing 10% casein (Possompes et al., 1989; cited in Friedman, 1999).

Milk proteins are unusually rich in lysine (Table 10.3) and can stand to lose a significant proportion of the lysine before it becomes limiting (around 50% in the case of whey proteins and 25% in the case of casein). The real concern arises when milk proteins, and particularly whey proteins, are being added to a mixture to provide a source of lysine supplementation. When this is the case, it is particularly important to ensure that the lysine content has not been compromised after long storage. Steps to ensure this include keeping the product at temperatures less than 25°C for most of the time and ensuring that the water activity in the product is kept low, preferably at or below 0.3. Experiments carried out in our laboratories to determine bioavailable lysine showed that loss of lysine on storage was negligible at 30°C, whereas losses were severe at 40°C (Figure 10.8).

**Sulfur amino acids**

Modification of sulfur amino acids is possible through loss of cysteine via lanthionine formation and through oxidation of methionine to methionine sulfoxide. Whey proteins are relatively rich in sulfur amino acids and casein has more than adequate quantities. It should be noted that these scores are relative to World Health Organization (WHO) requirements for children aged 2–5 years. The requirement for rat diets is higher and this is thought to be because of the requirement of the rat for hair production. (Hair is rich in sulfur amino acids.)

Lanthionine is a problem only for caseinates and lanthionine formation is known to occur during caseinating. The rates of lanthionine formation in dry powder have not been studied. This should be done, if only to briefly investigate lanthionine levels in old stocks or library samples.

Oxidation of methionine to the sulfoxide does not alter its bioavailability *per se*; however, it has been claimed that the presence of the sulfoxide side chain in intact proteins may hamper digestion of the protein, thus affecting its overall bioavailability (Anon, 1973).

**Other amino acids**

One essential amino acid known to be destroyed during some processes is tryptophan. Extensive investigations by Nielsen et al. (1985) on whey proteins, casein and WMP showed that tryptophan remained relatively intact even when substantial lysine modification had occurred. Tryptophan analysis gave a result of 100% (within experimental error) and a bioavailability of >90% in milk powder that had been stored at 60°C for 5 weeks, which resulted in a loss of 80% of available amine. This powder was considered to show “advanced” Maillard browning.
Samples of WPC80 powders that had earlier been shipped from New Zealand to the USA or Europe were obtained and analyzed for lactosylation levels. The powders were at the time less than 2 years old and thus were considered to be current stock. The powders had average bound lactose levels of between 0.7 and 1.2. This correlates to between 87–95% of the lysine in the products being bioavailable (estimated by extrapolation from the remaining lysine in β-lactoglobulin). This compares well with freshly produced WPC80 powders, which had an average number of lactose bound of 0.6, or 96% of available lysine remaining (Figure 10.9).

**Figure. 10.8** Loss of bioavailable lysine with time for SMP, WMP and WPC with a water activity ($a_w$) of 0.3 during storage at (a) 30°C and (b) 40°C. The solid and dashed lines on the graphs show the levels at which lysine becomes limiting in WPC, and in SMP and WMP respectively. (K. Higgs, unpublished results.)

**Product-specific storage trials**

Samples of WPC80 powders that had earlier been shipped from New Zealand to the USA or Europe were obtained and analyzed for lactosylation levels. The powders were at the time less than 2 years old and thus were considered to be current stock. The powders had average bound lactose levels of between 0.7 and 1.2. This correlates to between 87–95% of the lysine in the products being bioavailable (estimated by extrapolation from the remaining lysine in β-lactoglobulin). This compares well with freshly produced WPC80 powders, which had an average number of lactose bound of 0.6, or 96% of available lysine remaining (Figure 10.9).
We did a 2-year storage study on >80% protein powders to determine changes, if any, in nutritional properties, but these powders were kept at constant temperature and were not exposed to any of the temporal variations possible during shipping and storage in overseas warehouses.

A decrease in available amine of 5% was seen in MPC85 when stored at 20°C for 2 years; this increased to 10% when the storage temperature was increased to 30°C. A single MPI stored for 2 years gave consistent available amine results over the storage period.

The caseinates and caseins in the 2-year study showed no definitive trend in available amine values. Most values remained consistent over the 2 years. This was expected as a previous 3-month study showed that storage temperatures in excess of 30°C were required for significant levels of lysinoalanine formation.

**Conclusions**

Milk proteins do undergo change on the storage of dry powders. Powders containing appreciable amounts of lactose (milk powders and WPCs) form pre-Maillard reaction products, rendering lysine non-bioavailable, whereas those containing casein undergo formation of isopeptide bonds, reducing the availability of lysine and sulfur amino acids. Because milk proteins are rich in lysine, loss of some lysine will not be
a significant problem; however, when milk proteins are used as “balancers” in formulations with other proteins that are poor in lysine, attention should be given to the storage history of the protein.

Both the Maillard reaction and the formation of isopeptide bonds are undesirable and are best avoided by ensuring that shipping and storage temperatures do not exceed 30°C for significant periods. This should not be a problem in temperate climates; however, in climates where high temperatures routinely occur, consideration should be given to storage in a cool store.

References


Abstract

Because of their high nutritional quality and versatile functional properties, milk proteins are widely used as ingredients in many manufactured food colloids, e.g. dairy desserts, nutritional beverages, ice cream, yoghurt, spreads, confectionery and baked goods. Milk proteins perform a wide range of key functions in prepared foods, including emulsification, thickening, gelling and foaming. The functionality of milk proteins is a consequence of their molecular structures and interactions. An important functionality of milk proteins in food colloids is their ability to facilitate the formation and stabilization of oil droplets in emulsions. The ability of milk protein products to adsorb at the oil–water interface and to stabilize emulsions is influenced by the structures, flexibility and aggregation state of the constituent proteins.

This chapter deals mainly with the properties and functionalities of food emulsions formed with a range of milk protein products, and how they are influenced by different environmental and processing conditions. Of particular importance are the effects of pH, calcium ions and protein content and the influences of thermal and high-pressure processing. The chapter...
focuses on the structure and composition of adsorbed protein layers, competition between proteins and the creaming and flocculation behaviors of emulsion droplets.

Introduction

Milk proteins possess functional properties that provide desirable textural and other attributes to the final product and, for this reason, have found numerous applications in traditional dairy products and other foods. The functional properties of milk proteins, such as emulsification, thickening, gelling, flavor binding and foaming, contribute to the sensory characteristics and the stability of the manufactured foods. Several types of milk protein products, e.g. caseins and caseinates, whey protein concentrates (WPCs) and whey protein isolates (WPIs), milk protein concentrate (MPC) powders and hydrolyzed proteins, are manufactured from milk by the dairy industry.

Caseinates are produced from skim milk by adding acid (hydrochloric acid or lactic acid) or microbial cultures to precipitate the casein from the whey at pH 4.6. The acid-precipitated casein can then be resolubilized with alkali or an alkaline salt (using calcium, sodium, potassium or magnesium hydroxide) to about pH 6.7 and spray dried to form caseinate. Caseinates have exceptional water-binding capacity, fat emulsification properties and whipping ability, and a bland flavor. Emulsion-type products, e.g. coffee whiteners, whipped toppings, cream liqueurs and low-fat spreads, are an important application of caseinates in the food industry. In recent years, the use of casein and caseinate in dietary preparations, nutritional products and medical applications has increased; many of these preparations are also oil-in-water emulsions containing relatively small amounts of fat.

WPC and WPI are concentrated forms of whey protein components. Ultrafiltration, diafiltration and ion-exchange technology are used to concentrate and separate the protein from other components. The whey protein is then dried to obtain WPC or WPI, both of which are highly soluble, with protein levels ranging from 80–95%. Both WPC and WPI have a wide range of food applications and, because of their high protein content, can function as water-binding, gelling, emulsifying and foaming agents. Processing treatments used in the manufacture of WPC and WPI may sometimes cause some protein denaturation, which tends to affect their functionality.

MPCs are processed directly from skim milk by ultrafiltration/diafiltration and can have a range of protein contents from 56–82%. MPCs are used as functional ingredients in a wide variety of foods, including beverages, processed cheese and confectionery.

The functionality of milk proteins in processed foods is determined by their molecular structures and interactions with other food components, such as fats, sugars, polysaccharides, salts, flavors and aroma compounds. The type and the strength of various interactions determine the structure, texture, rheology, sensory properties and shelf life of manufactured food products. Much knowledge on the structure and properties of individual milk protein components has been gained, but less is known about interactions between different components that occur in a food system as a result of processing and formulation. Controlling these interactions is of key importance for achieving the desired functional properties in food products.
significance for the development of novel products and processes as well as for the improvement of conventional products and processes (Table 11.1).

This chapter focuses on the emulsifying properties of milk proteins, as this functional property is very important in all the food applications of milk protein products. The adsorption behavior of different milk protein products at oil-in-water interfaces and the stability of the resulting emulsions are considered, focusing on the work carried out in our laboratory at Massey University.

**Adsorption of milk proteins during the formation of emulsions**

Emulsions are composed of oil droplets (average range 0.5–5 μm diameter) enveloped by a continuous film of surfactant material that stabilizes the droplets. In the food industry, homogenization is widely used for finely dispersing oils in food products and proteins are most commonly used as emulsifying agents. The state of the droplet size distribution after homogenization reflects the emulsifying capacity of the proteins, the energy input during formation as well as the effects of various factors, such as pH, temperature, ionic strength and ratio of the two phases, on the surface activity of the proteins (Walstra, 1993; Dickinson, 1998a). In addition, the droplet size distribution influences markedly the properties of food emulsions, such as stability, viscosity, texture and mouthfeel.

During homogenization, the milk protein, in the form of individual molecules or protein aggregates, becomes rapidly adsorbed at the surface of the newly formed oil droplets. The amount of protein present at the interface per unit surface of dispersed phase is defined as the protein load, which is usually expressed as milligrams of protein per unit area of the dispersed phase (mg/m²). The protein load determines the amount of protein required to make an emulsion with a desired oil volume and droplet size and is dependent on the concentration and the type of protein as well as on the conditions used for emulsion formation. The factors that affect the protein load include protein concentration, volume of oil, energy input, state of protein aggregation, pH, ionic strength, temperature and calcium ions (Dickinson and Stainsby, 1988; Walstra, 1993).

### Table 11.1 Functional properties of milk proteins in food systems

<table>
<thead>
<tr>
<th>Functional property</th>
<th>Food system</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solubility</td>
<td>Beverages</td>
</tr>
<tr>
<td>Emulsification</td>
<td>Coffee whiteners, cream liqueurs, salad dressings, desserts</td>
</tr>
<tr>
<td>Foaming</td>
<td>Whipped toppings, shakes, mousses, cakes, meringues</td>
</tr>
<tr>
<td>Water binding</td>
<td>Bread, meats, bars, custard, soups, sauces, cultured foods</td>
</tr>
<tr>
<td>Heat stability</td>
<td>UHT- and retort-processed beverages, soups and sauces, custard</td>
</tr>
<tr>
<td>Gelation</td>
<td>Meats, curds, cheese, surimi, yoghurt</td>
</tr>
<tr>
<td>Acid stability</td>
<td>Acid beverages, fermented drinks</td>
</tr>
</tbody>
</table>
The properties of the adsorbed layers depend on the amounts and structures of the proteins present during homogenization. Proteins are amphipathic molecules containing both polar and non-polar parts and orientate at the interface in such a way that a substantial proportion of the non-polar amino acids remains in contact with the oil phase and the polar groups are in contact with the aqueous phase (Dickinson, 1992, 1998a). The main thermodynamic driving force for the adsorption of proteins is the removal of hydrophobic residues from the unfavorable environment of the bulk aqueous phase by displacement of structured water molecules from the close vicinity of the interface. An additional important driving force is the unfolding and reorganization of the native protein structure, which is due to interaction with the interface. By adsorbing at the interface, the protein reduces the free energy of the system and hence the interfacial tension.

Once a protein is adsorbed at an interface, it undergoes unfolding and rearrangement to form a stabilizing adsorbed layer (Dickinson, 1992; Dalgleish, 1996) and the extent of unfolding depends on the flexibility of the protein molecule, i.e. on the strength of the forces maintaining the secondary and tertiary structures. Because the caseins have rather flexible structures, they unfold rapidly at the interface and may form extended layers up to about 10 nm thick (Dalgleish, 1990). Dalgleish (1999) suggested that casein molecules are stretched to their maximum extent when their overall surface coverage is less than about 1 mg/m². Conversely, the presence of excess casein increases the monolayer coverage to a maximum value of 3 mg/m², the parts of the molecules in contact with the interface adopt a more compact conformation and the hydrophilic moieties protrude further from the interface.

Whey proteins (such as β-lactoglobulin), which give adsorbed layers that are only about 2 nm thick, change conformation and unfold their structure to some extent at the surface (Dalgleish and Leaver, 1993; Mackie et al., 1993; Dalgleish, 1995; Dickinson and McClements, 1995; Dalgleish, 1996; Fang and Dalgleish, 1998). The adsorbed whey protein structure lies somewhere intermediate between the native structure and the fully denatured state, which may have a native-like secondary structure and an unfolded tertiary structure (Dickinson, 1998a). Additionally, the partial unfolding of the globular whey protein structure following adsorption causes exposure of the reactive sulfydryl group, leading to slow polymerization of the adsorbed protein in the aged layer via sulfydryl–disulfide interchange (Dickinson and Matsumura, 1991; McClements et al., 1993).
The amount of protein adsorbed on the interface of an emulsion droplet suggests the state of the protein adsorbed at the interface. If the protein load is \(<1\) mg/m\(^2\), it suggests that the protein molecules are fully unfolded. If the protein load is 1–3 mg/m\(^2\), a monolayer of globular proteins may be present or unfolded molecules may be adsorbed in the conformation of trains, loops and tails. Protein load values above 5 mg/m\(^2\) suggest the adsorption of aggregates of proteins or multilayers of proteins. Some proteins of higher molecular weight may also give higher protein loads (Phillips, 1981; Hunt and Dalgleish, 1994; Dam \textit{et al}., 1995; Srinivasan \textit{et al}., 1996).

Extensive studies on purified milk protein systems show that a disordered casein monomer may be regarded as a complex linear copolymer that adsorbs to give an entangled monolayer of flexible chains, having some sequences of segments in direct contact with the surface (trains) and other sequences of segments protruding into the aqueous phase (loops and tails) (Dickinson, 1998a). Based on various experimental studies and molecular modeling, the \(\beta\)-casein molecule has been shown to adsorb with an extensive hydrophobic region anchored directly at the surface and a hydrophilic region (40–50 residues at the N-terminus) protruding extensively into the aqueous phase. This is probably also the portion of the molecule that forms the hydrodynamically thick layer.

In contrast to the dangling tail predicted for \(\beta\)-casein, a loop-like conformation has been predicted for \(\alpha_{s1}\)-casein and it does not have such a pronounced inequality in the distribution of hydrophobic and hydrophilic residues in its primary structure. It has been suggested that \(\alpha_{s1}\)-casein adsorbs to the oil–water interface via peptides towards the middle of its sequence, rather than the end as in \(\beta\)-casein, and it may be this that causes the protein to form a thinner adsorbed layer than does \(\beta\)-casein (Dalgleish, 1996). The simple train–loop–tail model is not adequate to describe the molecular configuration of adsorbed \(\beta\)-lactoglobulin. A closely packed, dense and rather thin (2–3 nm at neutral pH) layer of \(\beta\)-lactoglobulin is formed, which can be modeled as a dense two-dimensional assembly of highly interacting deformable particles.

Milk proteins used by the food industry contain complex mixtures of proteins in various states of aggregation. The structures of the adsorbed layers formed with these complex mixtures are not understood in molecular detail. The most commonly used milk proteins, i.e. sodium caseinate and whey proteins, show excellent emulsifying ability and it is possible to make stable emulsions at a relatively low protein-to-oil ratio (about 1:60). In emulsions formed with sodium caseinate or whey proteins, the protein load increases with an increase in protein concentration until it reaches a plateau value of about 2.0–3.0 mg/m\(^2\) (Singh, 2005) (Figure 11.1). The emulsifying ability of “aggregated” milk protein products, such as MPC and calcium caseinate, is much lower than that of whey protein or sodium caseinate, i.e. much higher concentrations of protein are required to make stable emulsions and larger droplets are formed under similar homogenization conditions.

The surface protein concentration of emulsions formed with MPC is in the range 5–20 mg/m\(^2\) depending on the protein concentration used in making the emulsions (Euston and Hirst, 1999). At low protein-to-oil ratios, protein aggregates are shared by adjacent droplets, resulting in bridging flocculation and consequently a marked increase in droplet size. In addition, the spreading of protein at the interface is
Interactions and functionality of milk proteins in food emulsions

limited in these emulsions, because the aggregates are held together by calcium bonds and/or colloidal calcium phosphate and these bonds are unlikely to be affected during the emulsification process. The higher conformational stability of these aggregates will also contribute to their reduced emulsifying ability (Euston and Hirst, 1999; Srinivasan et al., 1999).

The composition of the interfacial layer is determined by the quantities and structures of the proteins present at the moment the emulsion is formed. If proteins are the only emulsifiers present, they will adsorb to the oil–water interface, generally in proportion to their concentration in the aqueous phase (Dalgleish, 1997). However, certain mixtures of caseins show competition during adsorption at oil–water interfaces and rapid exchanges between adsorbed and unadsorbed caseins after emulsion formation. Studies have demonstrated that β-casein, because of its greater surface activity, adsorbs in preference to αs1-casein in emulsions stabilized by mixtures of these proteins and that β-casein displaces αs1-casein rapidly from the droplet surface (Dickinson et al., 1988b). In binary mixtures containing β-lactoglobulin and α-lactalbumin, some limited competitive adsorption does occur, but little exchange between the adsorbed and unadsorbed protein occurs. The protein that arrives at the interface first during homogenization is the protein that predominates there afterwards.

In contrast to model systems, no competitive adsorption has been observed in emulsions stabilized by more complex casein mixtures, such as sodium caseinate (Hunt and Dalgleish, 1994). Interestingly, this behavior appears to be related to the ratio of protein to oil in the emulsions (Euston et al., 1995; Srinivasan et al., 1999). Srinivasan et al. (1999) has shown that, in sodium caseinate emulsions, when the ratio of protein to oil is very low (about 1:60), β-casein is preferentially adsorbed at the droplet surface but, when the total amount of protein is greatly in excess of the amount needed for full surface coverage, αs1-casein is adsorbed in preference to the other caseins. At all concentrations, κ-casein from sodium caseinate appears to be less readily adsorbed (Figure 11.2). The concentration dependence of the competitive

Figure 11.1 Influence of protein concentration on average droplet size \(d_{43}\) (left) and surface protein coverage (right) in emulsions (30% soya oil) made with sodium caseinate (●), calcium caseinate (▲), WPC (■) or MPC (◆) (from Singh, 2005, reproduced with the permission of The Royal Society of Chemistry).
adsorption of $\alpha_{s1}$-casein and $\beta$-casein in sodium caseinate emulsions may be a consequence of the different complexes that can be formed by caseins in solution (Rollema, 1992).

The preferential adsorption of $\beta$-casein, because of its high surface activity, appears to exist only at low concentrations where caseins may exist as monomers. With increasing protein concentration, caseins aggregate to form various complexes (Lucey et al., 2000) and it is likely that $\beta$-casein loses its competitive ability because of its self-aggregation to form micelles or through the formation of complexes with other caseins. Therefore, the surface composition of emulsions formed using a relatively high sodium caseinate concentration is likely to be determined by the surface activities and flexibilities of the casein aggregates and complexes. Although extensive information on the surface activity and hydrophobicity of individual caseins is available, little is known about how these characteristics are modified when casein molecules undergo self-association under different environmental conditions.

When the casein is in the highly aggregated form of casein particles, as in calcium caseinate or MPC, there is very little competitive adsorption and protein exchange (Euston and Hirst, 1999; Srinivasan et al., 1999). In these systems, the average surface composition is probably determined by the adsorption of protein aggregates of fixed composition. For instance, calcium caseinate solution consists of large $\alpha_{s1}$-casein-rich aggregates, which appear to dominate the droplet surface after emulsification (Srinivasan et al., 1999). When WPCs or WPIs are used to make emulsions, there is no preferential adsorption between $\beta$-lactoglobulin and $\alpha$-lactalbumin regardless of the protein-to-oil ratio in the emulsion (Euston et al., 1996; Ye and Singh, 2000, 2006a).

The aggregation state and the flexibility of protein molecules can be altered by changes in pH, addition of divalent cations and various processing treatments prior to emulsification. These changes will inevitably influence the adsorption behavior of

![Figure 11.2](image-url)
milk proteins at the oil–water interface. For example, addition of CaCl$_2$ at above a
certain critical concentration to a sodium caseinate or whey protein solution before
homogenization increases the droplet size, increases the surface protein coverage and,
in sodium caseinate emulsions, also affects the competition between different proteins
(Ye and Singh, 2000, 2001). The proportions of $\beta$-lactoglobulin and $\alpha$-lactalbumin
at the droplet surface remain unaffected by the addition of CaCl$_2$ to a whey protein
solution prior to emulsification. In contrast, addition of CaCl$_2$ to a sodium caseinate
solution markedly enhances the adsorption of $\alpha_s$-casein at the droplet surface, with a
much lesser effect on $\beta$-casein adsorption.

The effects of calcium on surface coverage and composition can be explained by
the binding of the ions to the negatively charged amino acid residues on the protein.
This reduces electrostatic repulsions between the protein molecules and increases the
potential for intermolecular associations. Because of the presence of clusters of phos-
phoserine residues, the caseins have stronger affinity than the whey proteins to bind
calcium. Consequently, the caseins (except $\kappa$-casein) are precipitated by calcium,
with $\alpha_s$-casein being the most sensitive to aggregation and precipitation by calcium.
In sodium caseinate emulsions, the increased surface coverage upon addition of cal-
cium prior to emulsification is probably due to adsorption of casein aggregates on to
the droplet surface (Ye and Singh, 2001). Greater $\alpha_s$-casein adsorption reflects its
stronger tendency to be aggregated by calcium ions in solution or at the interface.

The native whey proteins do not bind much calcium and are not precipitated in the
presence of calcium (Baumy and Brule, 1988), although heat-denatured whey proteins
are able to bind considerable amounts of calcium and undergo aggregation (Pappas
and Rothwell, 1991). The increase in surface protein coverage suggests the formation
of aggregates of whey proteins in the presence of calcium, which subsequently become
adsorbed during emulsification (Ye and Singh, 2000). This has been attributed to a
decrease in the denaturation temperature of the whey proteins in the presence of calcium.

All these results confirm that, under a given set of homogenization conditions, the
surface composition is largely dependent on the protein-to-oil ratio and the aggrega-
tion state of the proteins in solution. It appears that the structure of the interfacial
layer in emulsions can be manipulated by controlling the protein concentration, the
protein type and the ionic environment. Because of their different interfacial struc-
tures, these droplets would be expected to exhibit different reactivities, which could
be exploited to develop new food textures. Further studies are required for an under-
standing of the relationship between droplet surface structures and the sensitivity of
the droplets to different environments and processing conditions.

**Stability of milk-protein-based emulsions**

The term “emulsion stability” refers to the ability of an emulsion to resist any altera-
tion in its properties over the timescale of observation (McClements, 1999; Dickinson,
2003; McClements, 2005). An emulsion is thermodynamically unstable as the free
energy of mixing is positive because of the large interfacial area between the oil and
the aqueous phase. Therefore, the kinetic stability, i.e. the time period for which the
Emulsion is stable, is important (Damodaran, 1997; McClements, 1999; Dickinson, 2003; McClements, 2005). For instance, an emulsion can be considered to be “stable” if the inevitable process of separation has been slowed to an extent that it is not of practical importance during the shelf life of the product. An emulsion may become unstable because of a number of different types of physical and chemical processes. Physical instability refers to the change in spatial arrangement or size distribution of emulsion droplets, such as creaming, flocculation or coalescence, whereas chemical instability includes change in the composition of the emulsion droplet itself, such as oxidation, hydrolysis, etc. (McClements and Decker, 2000; McClements, 2005).

Creaming is the movement of oil droplets, under gravity or in a centrifuge, to form a concentrated layer at the top of an oil-in-water emulsion sample, with no accompanying change in the droplet size distribution. Creaming is reversible and the original uniform distribution of droplets can usually be obtained by gentle mixing. The creaming process can be explained by Stokes’ Law (Hunter, 1986; McClements, 2005):

\[
\nu_{\text{stokes}} = \frac{2r^2(\rho_1 - \rho_2)}{9\eta}
\]  

(11.1)

where \(\nu_{\text{stokes}}\) = velocity of creaming, \(r\) = emulsion droplet radius, \(\rho_1\) and \(\rho_2\) = density of the continuous phase and the dispersed phase respectively and \(\eta\) = shear viscosity of the continuous phase. The creaming rate can be reduced by lowering the radius, increasing the continuous phase viscosity or decreasing the difference in density between the two phases. However, this law often fails to define the rate of creaming due to flocculation or coalescence.

Coalescence, i.e. an increase in droplet size by accretion, gradually results in separation of the oil and the aqueous phase and is always irreversible. Coalescence requires rupture of the stabilizing film at the oil–water interface, but this occurs only when the layer of continuous phase between the droplets has thinned to a certain critical thickness (Dickinson and Stainsby, 1988; Britten and Giroux, 1991; Das and Kinsella, 1993; Walstra, 1993).

Flocculation has been defined as the reversible aggregation mechanism that arises when droplets associate as a result of unbalanced attractive and repulsive forces (Dalgleish, 1997). Generally, two types of flocculation are distinguished, i.e. depletion flocculation and bridging flocculation (Dickinson, 2003). The type of mechanism prevailing depends upon the interaction between the interfacial layer and the emulsion droplets.

Bridging flocculation normally occurs when a high-molecular-weight biopolymer at a significantly low concentration adsorbs to two or more emulsion droplets, forming bridges (Dickinson and Pawlowsky, 1998; Dickinson, 1998b; McClements, 1999; Dickinson, 2003; McClements, 2005; Fellows and Doherty, 2006). Depletion flocculation occurs as a result of the presence of unadsorbing biopolymer in the continuous phase, which can promote association of oil droplets by inducing an osmotic pressure gradient within the continuous phase surrounding the droplets (de Hek and Vrij, 1981; Dickinson, 1999; Tuinier and de Kruij, 1999; McClements, 2005).
Essentially, if the added biopolymer is either unadsorbed or poorly adsorbed, the biopolymer is squeezed out of the area between two approaching emulsion droplets. The concentration of biopolymer between the emulsion droplets becomes less than its overall solution concentration, resulting in osmotic imbalance. The net effect is that the particles are attracted towards each other, resulting in flocculation. The attraction energy is determined by the concentration of the polymer and the range of interaction depends on the radius of gyration of the polymer molecule. The bonds formed through the depletion flocculation mechanism are generally weak, flexible and reversible.

The ability of proteins to stabilize emulsions is the most important criterion besides the emulsion formation in most food applications. The forces involved in stabilizing and destabilizing emulsions include van der Waals’ attractive forces, electrostatic interactions and steric factors. At pH values away from their isoelectric point, as proteins are electrically charged, there is an electrostatic repulsion, which prevents dispersed droplets from closely approaching one another. With the possible exception of highly charged proteins, a predominant contribution to emulsion stabilization by protein comes from the steric stabilization mechanism. Interactions between droplets stabilized by proteins may be influenced by the presence of certain ions, particularly calcium, as proteins are capable of binding ions.

As long as sufficient protein is present during homogenization to cover the oil droplets, emulsions stabilized by milk proteins are generally very stable to coalescence over prolonged storage. However, these emulsions are susceptible to different types of flocculation, which in turn leads to enhanced creaming or serum separation. At low protein-to-oil ratios, there is insufficient protein to fully cover the oil–water interface during homogenization and this results in bridging flocculation. Another consequence of insufficient protein is coalescence of droplets during or immediately after emulsion formation. Bridging flocculation is commonly observed in emulsions formed with aggregated milk protein products, such as calcium caseinate or MPC, in which the droplets are bridged by casein aggregates or micelles. Optimum stability can generally be attained at protein concentrations high enough to allow full saturation coverage at the oil–water interface. However, at very high protein-to-oil ratios, the presence of excess, unadsorbed protein may lead to depletion flocculation in some emulsions. Both depletion flocculation and bridging flocculation cause an emulsion to cream more rapidly.

Depletion flocculation has been observed in sodium-caseinate-based emulsions but not in emulsions formed with calcium caseinate, MPC or whey proteins (Dickinson and Golding, 1997; Euston and Hirst, 1999; Srinivasan et al., 2001; Singh, 2005) (Figure 11.3). In sodium-caseinate-based emulsions, it was shown that, at a protein content of nearly 2.0 wt%, the emulsion droplets were protected from flocculation by a thick steric-stabilizing layer of sodium caseinate. The emulsion was stable against flocculation, coalescence and creaming for several weeks. However, when the protein content was increased to above 3.0 wt%, unadsorbed protein gave rise to depletion flocculation. Because of this depletion flocculation, the effective diameter of the droplets increased, resulting in a marked decrease in creaming stability with an increase in the caseinate concentration from 3 to 5 wt%. Further increasing the protein content to 6.0 wt% and above resulted in very high depletion flocculation, leading to a strong emulsion droplet network that was stable to creaming.
The differences in the creaming stabilities of emulsions made with different kinds of milk protein products are largely related to depletion flocculation effects (Singh, 2005). The depletion interaction free energy ($\Delta G_{\text{DEP}}$), of the order of a few $kT$, can be estimated using Equation (11.2) (Walstra, 1993):

$$\Delta G_{\text{DEP}} = -2\pi \gamma^2 \Pi (\gamma_d - 2\gamma_m/3)$$  \hspace{1cm} (11.2)

where $\Pi$ is the osmotic pressure of the polymer solution, represented as a fluid of hard spheres of radius $\gamma_m$, and $\gamma_d$ is the mean droplet radius. The osmotic pressure under ideal conditions is given by the following equation:

$$\Pi = cRT/M$$  \hspace{1cm} (11.3)

where $R$ is the molar gas constant, $T$ is the temperature, $M$ is the molecular mass of the polymer and $c$ is the number concentration of the polymer.

For depletion flocculation to occur, the polymer has to have a fairly high $M$, so that the $\gamma_m$ is relatively large. However, at a given $c$, $M$ is inversely proportional to $\Pi$. Therefore, an increase in the polymer molecular mass will reduce the osmotic pressure driving the depletion interaction. Hence, at a given concentration, the depletion interaction free energy is low for a polymer of low molecular mass, increases with an increase in molecular mass until it reaches a maximum and then decreases with a further increase in molecular mass. Similarly, a reduction in the polymer number concentration will reduce the osmotic pressure.

Although the exact state of the casein molecules in concentrated sodium caseinate solutions is unknown, a sodium caseinate solution has been reported to have a radius of gyration of about 20–30 nm, as determined by static light scattering (Lucey et al., 2000). It is likely that depletion flocculation in sodium caseinate emulsions

![Figure 11.3 Creaming stability and microstructure of emulsions made with sodium caseinate (●) or WPC (■) (30% oil). Scale bar represents 10 μm (from Singh, 2005, reproduced with the permission of The Royal Society of Chemistry).](image)
is caused by the presence of these casein aggregates formed from self-assembly of sodium caseinate in the aqueous phase of the emulsion at concentrations above 2 wt% (Dickinson and Golding, 1998).

Emulsions formed with whey proteins, MPC and calcium caseinate do not show depletion flocculation, probably because there are no suitably sized protein particles at the required concentrations in the aqueous phase. The molecular size of whey proteins is less than the optimum, whereas the casein micelles in MPC are too large to induce depletion flocculation. Calcium caseinate consists of mixtures of casein aggregates of different sizes, but the concentration of aggregates capable of inducing depletion flocculation is probably too low. The extent of creaming in these emulsions is largely determined by the particle size of the droplets. Generally, in these emulsion systems, the creaming stability increases with increasing protein concentration up to a certain concentration and then remains almost constant (Euston and Hirst, 1999; Srinivasan et al., 2001). However, the creaming stability of emulsions formed with calcium caseinate or MPC at relatively high protein concentration tends to be higher than that of whey-protein-stabilized emulsions. This can be attributed to an increase in the droplet density as a result of the presence of a much thicker and denser adsorbed protein layer at the droplet surface.

The addition of moderate amounts of CaCl₂ to emulsions containing excess sodium caseinate has been shown to eliminate depletion flocculation and to improve the creaming stability (Ye and Singh, 2001). This effect appears to be due to an increase in the average size of the casein aggregates in the aqueous phase, resulting in a large increase in the molecular mass of the caseins (Dickinson et al., 2001). In addition, there is a reduction in the concentration of unadsorbed caseinate. Both these effects are expected to cause a substantial reduction in the concentration of small particles, which are assumed to be the main depleting species responsible for inducing reversible flocculation in the calcium-free systems.

Presumably, the substantial reduction in osmotic pressure makes the magnitude of $\Delta G_{DEP}$ predicted from Equation (11.2) too small to cause depletion flocculation. Similarly, addition of NaCl at above a certain concentration reduces the extent of depletion flocculation of sodium caseinate emulsions and improves the creaming stability (Srinivasan et al., 2000). This effect is due to increased adsorption of protein at the droplet surface and hence a lower concentration of unadsorbed protein remaining in the solution. Decreasing the pH of emulsions formed with excess sodium caseinate also gradually eliminates depletion flocculation, through aggregation of adsorbed protein and a transfer of more protein to the droplet surface (Singh, 2005). Therefore, it seems to be possible to switch depletion flocculation off and on by controlling the concentration and the aggregation state of the casein molecules in the aqueous phase.

**Heat-induced changes in milk-protein-based emulsions**

Food emulsions are often heat treated at relatively high temperatures to provide a long shelf life to the product via microbial sterility. These heat treatments can cause
denaturation and aggregation of adsorbed and unadsorbed proteins, resulting in aggregation or coalescence of droplets and gel formation. Emulsions formed with whey proteins at neutral pH are stable against heating when the ionic strength and/or the concentration of protein in the emulsions are low. Addition of KCl at 100 mM or above has been shown to cause destabilization of whey protein emulsions, leading to gel formation (Hunt and Dalgleish, 1995).

Both the unadsorbed protein and the adsorbed protein are necessary for the heat-induced aggregation of whey-protein-stabilized emulsions. Aggregation of emulsion droplets is more extensive and proceeds more rapidly as the concentration of protein in the emulsion is increased, whereas removal of unadsorbed protein from the emulsion decreases the rate of droplet aggregation (Euston et al., 2000). During heat treatment, the protein-covered droplet appears to interact more readily with the unadsorbed protein than with another emulsion droplet. This has been explained by assuming that the relative surface hydrophobicities of the emulsion droplet and the unadsorbed denatured whey proteins are different. Interaction of two emulsion droplets through their respective adsorbed protein layers will have a relatively low probability because the surface hydrophobicity is likely to be relatively low. When an emulsion droplet and an unadsorbed protein molecule aggregate, at least one of them (the denatured protein molecule) has a relatively high surface hydrophobicity and this will increase the probability of interaction and aggregation (Euston et al., 2000).

In emulsions made with 3.0% WPI and 25% soya oil, the amount of adsorbed protein was shown to increase from 2.9 to 3.7 mg/m² within the first 10 min of heating at 75°C, but further heating had no effect (Sliwinski et al., 2003). At 90°C, the plateau value of about 4 mg/m² was reached within 5 min of heating. Studies on the effects of heating temperature in the range 50–90°C on WPI emulsions (pH 7.0) (Monohan et al., 1996; Demetriades and McClements, 1998) show that droplet aggregation occurs on heating in the range 75–80°C, which causes an increase in viscosity and a loss of creaming stability, but the degree of aggregation and the susceptibility to creaming decrease on heating at temperatures above 80°C.

It has been suggested that, in the temperature range 75–80°C, the whey protein molecules at the droplet surface are only partly unfolded and that not all of the hydrophobic amino acid residues are directed towards the oil phase. Consequently, the surface of the droplet is more hydrophobic, making it susceptible to droplet aggregation. At higher temperatures, the proteins become fully unfolded with all of the hydrophobic residues being directed into the oil phase, which makes the droplets less prone to aggregation. The role of sulfydryl–disulfide interchange reactions in droplet aggregation is not clear. It has been suggested that disulfide-mediated interactions during heat treatment are not critical during the initial stages of aggregation but they tend to strengthen the aggregates (Demetriades and McClements, 1998).

Recently, Dickinson and Parkinson (2004) and Parkinson and Dickinson (2004) reported that addition of a very small proportion of caseinate (0.03–0.15% of the total emulsion) can stabilize a whey-protein-based emulsion against heat treatment. The magnitude of the effect is dependent on the type of casein, with the order of effectiveness being β-casein > sodium caseinate > αs1-casein. The stabilizing effect of the casein in these mixed milk protein systems is strongly synergistic. The casein
polymer appears to be acting in a colloidal stabilizing capacity at a surface concentration very much lower than that at which it could be used as an emulsifying or stabilizing agent simply on its own. It has been suggested that adsorbed casein molecules keep the emulsion droplet surfaces sufficiently far apart to prevent the “normal” cross-linking processes that occur between whey-protein-coated droplets during heat-induced aggregation and gelation, because of the steric hindrance from the loops and tails of the disordered casein polymers (Parkinson and Dickinson, 2004).

In contrast to whey proteins, emulsions formed with sodium caseinate (2 wt% protein, 20% soya oil) are stable to heating at 90 °C for 30 min or 121 °C for 15 min, as determined by droplet size analysis (Hunt and Dalgleish, 1995; Srinivasan et al., 2002). However, the protein coverage and the adsorbed casein composition change upon heat treatment, indicating that interactions between unadsorbed caseinate molecules and caseinate at the droplet surface may occur during heating (Srinivasan et al., 2002).

Analysis of adsorbed caseins isolated from emulsions heated at 121 °C for 15 min has shown that a substantial proportion of the adsorbed caseinate is polymerized to form high-molecular-weight aggregates (Srinivasan et al., 2002), held together through covalent bonds other than disulfide bonds. These covalent bonds appear to form mainly between caseinate molecules at the surface of the same droplet because of the higher local concentrations of casein molecules at the droplet surface.

Interestingly, the adsorbed caseins also appear to undergo thermal degradation, resulting in the formation of low-molecular-weight species. Relatively high proportions of casein degradation products present at the droplet surface indicate that the adsorbed caseinate molecules are more susceptible to fragmentation during heating than those in solution and that these peptides remain adsorbed. This is probably due to different structures and conformations of the caseins at the droplet surface than of those in the solution.

The creaming stability of sodium caseinate emulsions has been found to improve upon heating, with the onset of depletion flocculation occurring at higher protein concentration than in unheated emulsions (Srinivasan et al., 2002). This can be attributed to a reduction in the number of unadsorbed caseinate molecules/aggregates in the aqueous phase as a result of increased surface coverage and heat-induced polymerization and degradation of the casein molecules. The improvement in the creaming stability in heated emulsions at low protein concentrations may be attributed to an increase in droplet density because of the presence of greater amounts of polymerized protein at the droplet surface.

The surface protein composition of emulsion droplets may also change during heat treatment in emulsions formed with whey proteins. For WPI-stabilized emulsions, the amount of β-lactoglobulin at the droplet surface was found to increase during heat treatment, whereas the amount of adsorbed α-lactalbumin decreased markedly (Ye and Singh, 2006a). It seems that β-lactoglobulin displaces α-lactalbumin from the interface on heating at temperatures up to 90 °C. The reason for this is not clear.

Similar phenomena were observed in studies of exchanges of caseins and whey proteins at the interfaces of oil-in-water emulsion droplets (Dalgleish et al., 2002). It was found that, at temperatures above 40 °C, addition of WPI to the aqueous phase of caseinate-stabilized emulsions caused a displacement of adsorbed caseins. As the
β-lactoglobulin and α-lactalbumin adsorbed, αs1- and β-caseins were desorbed, principally the former, whereas the αs2- and κ-caseins were not displaced. The rate of the displacement or exchange reaction was temperature dependent, being almost undetectable at room temperature, but complete within 2 min at 80°C. The displacement reaction was not affected by ionic strength; neither were any of the reactions apparently dependent on sulfydryl exchange reactions (Dalgleish et al., 2002). However, no exchange of proteins occurred when an emulsion prepared with WPI was treated with caseinate and heat treated at 80°C for 2 min (Brun and Dalgleish, 1999). This was surprising in view of the known interactions of whey proteins with αs2- and κ-caseins, involving sulfydryl–disulfide interchange reactions.

Pressure-induced changes in milk-protein-based emulsions

The effect of ultra-high pressure (100–1000 MPa) on the structures of milk proteins in aqueous solution has received considerable attention over the last few years (see Chapter 7). High pressure can disrupt the quaternary and tertiary structures of globular proteins with relatively little influence on their secondary structure. In addition, proteinaceous colloidal aggregates (e.g. casein micelles), which are held together by ionic and hydrophobic interactions, can be dissociated by high-pressure treatment (Gaucheron et al., 1997; Huppertz et al., 2004).

Whey proteins are sensitive to high-pressure treatments (López-Fandinó et al., 1996; Anema et al., 2005). Solution studies (Patel et al., 2005) of native β-lactoglobulin and whey protein products have shown that high-pressure treatment has a marked effect on the protein’s conformation and consequently its aggregation behavior; the aggregation is more extensive at high protein concentrations (Patel et al., 2005). The formation of aggregates is most probably due to the generation of intermolecular disulfide bridges through sulfydryl–disulfide interchange reactions (Patel et al., 2006).

In model oil-in-water emulsions, high-pressure treatment has been shown to have no effect on the droplet size distribution or the emulsion viscosity of sodium-caseinate-based emulsions at pH 7 (Dumay et al., 1996). However, high-pressure treatment significantly induced flocculation of emulsion droplets and increased the emulsion viscosity of oil-in-water emulsions stabilized by β-lactoglobulin or WPC at neutral pH (Dumay et al., 1996; Dickinson and James, 1998). The unfolded unadsorbed whey proteins in the emulsion treated by high pressure appear to be the major contributor to the cross-linking or flocculation of emulsion droplets because greater emulsion flocculation was observed in emulsions with higher proportions of unadsorbed protein in the aqueous phase.

As in the case of emulsions treated by heat processing, whey-protein-stabilized emulsions are more sensitive to pressure and temperature at pH values closer to the isoelectric point and at high ionic strength. In terms of the change in emulsion rheology, severe high-pressure treatment (800 MPa for 60 min) is equivalent to
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relatively mild thermal treatment (65°C for 5 min) (Dickinson and James, 1998). In a concentrated emulsion formed with β-lactoglobulin (1% protein and 40% vol% n-tetradecane), an emulsion gel was produced following high-pressure treatment. When β-lactoglobulin or WPC solution was treated by high pressure before emulsion formation, the emulsions had larger droplet sizes than emulsions made with the native protein (Galazka et al., 1995).

The results indicated a modification of protein structure, leading to the loss of emulsifying efficiency as a result of protein aggregation, despite an increase in surface hydrophobicity. After adsorption on the surface, the protein probably became partially unfolded at the interface and subsequent pressure treatment caused no further conformational change. No studies on the behavior of emulsions formed with aggregated milk proteins, such as micellar casein, upon high-pressure treatment have been reported.

Milk protein hydrolysates and oil-in-water emulsions

Milk protein hydrolysates have been used extensively in infant and specialized adult nutritional formulations. Extensively hydrolyzed proteins are more easily digested and have substantially reduced immunological reactivities. These formulations are essentially multicomponent emulsion systems and therefore the emulsifying properties of protein hydrolysates are important.

The flexibility and thus the availability of hydrophobic and hydrophilic segments within the protein chain can be improved by moderate enzymatic hydrolysis of globular proteins (e.g. whey proteins), thus improving the emulsifying properties of the protein. However, extensive hydrolysis (above 20% degree of hydrolysis), because of the production of many short peptides, has been found to be detrimental to the emulsifying and stabilizing properties of whey proteins (Singh and Dalgleish, 1998). The main form of instability in emulsions formed with highly hydrolyzed whey proteins is the coalescence that arises because of the inability of the predominantly short peptides to adequately stabilize the large oil surface generated during homogenization (Singh and Dalgleish, 1998; Agboola et al., 1998a, 1998b). Nevertheless, it seems to be possible to make a fairly stable emulsion using highly hydrolyzed whey proteins at high peptide concentrations (peptide-to-oil ratio about 1:1w/w), and at low homogenization pressures, as the sole emulsifier (Agboola et al., 1998a, 1998b). Under these conditions, there is a sufficient amount of high-molecular-weight peptides (>5000 Da) in the emulsion to cover and stabilize the emulsion droplets.

Addition of calcium at above 20 mM has been shown to reduce the emulsion stability of emulsions formed with whey protein hydrolysates (Ramkumar et al., 2000). This instability arises mainly from the binding of calcium to the adsorbed peptides, leading to a reduction in the charge density at the droplet surface, which would reduce the inter-droplet repulsion and enhance the likelihood of droplet flocculation. The formation of calcium bridges between peptides present on two different emulsion droplets would also enhance flocculation.
In these emulsions, it is observed that some very large droplets, apparently formed by coalescence, are also formed in the presence of calcium. This is likely to be due to the binding of calcium ions to the negatively charged peptides, causing aggregation of larger, more surface-active peptides. This situation would reduce the effective concentrations of emulsifying peptides available during emulsion formation.

Heat treatment of emulsions stabilized by highly hydrolyzed whey proteins at 121°C for 16 min results in destabilization of the emulsions, which appears to occur mainly via a coalescence mechanism (Agboola et al., 1998b). As the adsorbed peptide layers in these emulsions lack the cohesiveness of the parent proteins and have poor ability to provide steric or charge stabilization, increased collisions between the droplets during heating would cause droplet aggregation, leading to coalescence. It is also possible that desorption of some loosely adsorbed peptides occurs during heating, as indicated by the decrease in the amount of peptides associated with the oil surface after heating, which would also enhance coalescence.

It is well known that the addition of polysaccharide such as xanthan gum or guar gum (in the concentration range 0.01–0.4%) to protein-stabilized emulsions promotes droplet flocculation, through a depletion mechanism, which enhances the creaming rate (Singh et al., 2003). Similar effects are observed in emulsions stabilized with hydrolyzed whey proteins but, interestingly, this depletion flocculation also promotes the coalescence of droplets during the storage of emulsions (Ye et al., 2004a, 2004b).

The rate of coalescence is enhanced considerably with increasing concentration of polysaccharide in the emulsion up to a certain critical concentration. At a given concentration, the rate of coalescence was highest in the emulsions containing guar gum, whereas it was lowest in the emulsions containing κ-carrageenan (Figure 11.4), which
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could be explained on the basis of the relationship between the strength of the depletion potential and the molecular weight and radius of gyration of the polysaccharide.

Whey protein peptides adsorbed at the droplet surface in these emulsions would almost certainly have a reduced surface viscosity compared with intact proteins and this could lead to reduced stability to drainage and film rupture. It can be concluded that flocculation of droplets, through various mechanisms (e.g. depletion flocculation, calcium-induced aggregation, heat treatment), in these types of emulsions, where the interface is rather weak, would lead to coalescence during storage.

**Lactoferrin-based oil-in-water emulsions**

Bovine milk contains low levels of lactoferrin, an iron-binding glycoprotein with about 700 amino acid residues and a molecular weight of about 80,000 Da (Baker and Baker, 2005). The polypeptide is folded into two globular lobes, representing its N- and C-terminal halves, commonly referred to as the N-lobe and the C-lobe. The surface of the lactoferrin molecule has several regions with high concentrations of positive charge, giving it a high isoelectric point (pI ≈ 9). This positive charge is one of the features that distinguishes lactoferrin from other milk proteins, such as β-lactoglobulin, which have isoelectric points in the range 4.5–5.5 and are negatively charged at neutral pH. This unique difference could allow the formation of oil-in-water emulsions containing cationic emulsion droplets, through adsorption of lactoferrin, over a wider pH range.

Recent work (Ye and Singh, 2006b) has shown that, similar to other milk proteins (e.g. caseinate and β-lactoglobulin), lactoferrin adsorbs on to the interface of oil-in-water emulsion droplets and forms stable emulsions, but emulsion droplets with an overall positive surface charge are produced. In contrast to caseinate- and whey-protein-stabilized emulsions, the cationic emulsion droplets formed by lactoferrin are stable against a change in the pH from 7.0 to 3.0. For emulsions prepared under the same conditions (concentrations of oil and protein, pH, homogenization pressure), the droplet sizes in the lactoferrin emulsions are similar to those in β-lactoglobulin-stabilized emulsions, but the surface protein coverage (mg/m²) of the emulsions made at pH 7.0 is higher in lactoferrin emulsions, possibly because of its higher molecular weight.

The formation of a positively charged adsorbed layer in lactoferrin-stabilized emulsions over a wide pH range provides an opportunity for electrostatic interactions with other milk proteins that are mostly negatively charged around neutral pH. In aqueous solutions, lactoferrin tends to form a complex with β-lactoglobulin via electrostatic interactions (Wahlgren et al., 1993). Adsorption of such a complex on to the droplet surface during emulsion formation results in greater amounts of protein at the droplet surface and the formation of thick interfacial layers. It is interesting to note that oil-in-water emulsions formed using a binary mixture of β-lactoglobulin and lactoferrin are very stable, even though the overall charge (ζ-potential) of the emulsion droplets is close to zero. This suggests that steric repulsion plays an important role in this binary protein-stabilized emulsion.
Multilayered emulsions can be produced by interactions of oppositely charged milk proteins, i.e. lactoferrin and \( \beta \)-lactoglobulin or caseinate at neutral pH (Ye and Singh, 2007). A primary emulsion, containing either anionic droplets coated with \( \beta \)-lactoglobulin or cationic droplets coated with lactoferrin, can be produced. A secondary emulsion can then be made by mixing either \( \beta \)-lactoglobulin solution or lactoferrin solution with the primary emulsion (Ye and Singh, 2007).

For example, when the emulsions formed with lactoferrin (1 wt%, pH 7.0) were diluted with aqueous phase containing a range of \( \beta \)-lactoglobulin concentrations, the adsorption of \( \beta \)-lactoglobulin increased considerably with an increase in the \( \beta \)-lactoglobulin concentration up to 0.42 wt%, with very little change above this concentration. This increase in \( \beta \)-lactoglobulin on the surface of emulsions formed with lactoferrin was further confirmed by the change in the \( \zeta \)-potential. In the absence of \( \beta \)-lactoglobulin, the \( \zeta \)-potential of the emulsion droplets was around +50 mV, because the lactoferrin used to stabilize the droplets has a net positive charge at pH 7.0. The \( \zeta \)-potential became less positive, and eventually changed from positive to negative, as the \( \beta \)-lactoglobulin concentration in the emulsion was increased (Figure 11.5).

**Figure 11.5** Influence of addition of \( \beta \)-lactoglobulin into emulsions formed with 1 wt% lactoferrin (30 wt% soya oil, pH 7.0) on the \( \zeta \)-potential of the emulsion droplets (from Ye and Singh, 2007, reproduced with the permission of Springer).

Multilayered emulsions can be produced by interactions of oppositely charged milk proteins, i.e. lactoferrin and \( \beta \)-lactoglobulin or caseinate at neutral pH (Ye and Singh, 2007). A primary emulsion, containing either anionic droplets coated with \( \beta \)-lactoglobulin or cationic droplets coated with lactoferrin, can be produced. A secondary emulsion can then be made by mixing either \( \beta \)-lactoglobulin solution or lactoferrin solution with the primary emulsion (Ye and Singh, 2007).

For example, when the emulsions formed with lactoferrin (1 wt%, pH 7.0) were diluted with aqueous phase containing a range of \( \beta \)-lactoglobulin concentrations, the adsorption of \( \beta \)-lactoglobulin increased considerably with an increase in the \( \beta \)-lactoglobulin concentration up to 0.42 wt%, with very little change above this concentration. This increase in \( \beta \)-lactoglobulin on the surface of emulsions formed with lactoferrin was further confirmed by the change in the \( \zeta \)-potential. In the absence of \( \beta \)-lactoglobulin, the \( \zeta \)-potential of the emulsion droplets was around +50 mV, because the lactoferrin used to stabilize the droplets has a net positive charge at pH 7.0. The \( \zeta \)-potential became less positive, and eventually changed from positive to negative, as the \( \beta \)-lactoglobulin concentration in the emulsion was increased (Figure 11.5).

**Conclusions**

Milk proteins in soluble and dispersed forms have excellent surface-active and emulsion-stabilizing properties. Differences in the emulsifying abilities of milk proteins arise largely from the differences in structure, flexibility, state of aggregation and composition of the proteins. These attributes of milk proteins (and hence their emulsifying abilities) are modified through various interactions occurring during the processing of milk that is required to isolate the protein components as well as
Interactions and functionality of milk proteins in food emulsions during the manufacture of prepared foods. Emulsions with different surface compositions and structures can be made using different kinds of milk proteins and these emulsions exhibit different sensitivities to solution conditions, such as pH and ionic strength, and processing conditions, such as heat and high-pressure treatments. This could offer possibilities for the formation of emulsions with a range of functionalities for different food applications.

Most of the research during the last 20 years has been performed on oil-in-water emulsions using purified or simple mixtures of caseins and whey proteins, with a great deal of information now being available on the conformation of proteins at oil-water interfaces, competitive exchange reactions between adsorbed and unadsorbed proteins, protein–polysaccharide interactions and factors controlling the rheology and stability of emulsions. In addition, some understanding of how processing conditions (heat treatments, high-pressure treatments) influence interfacial structures and emulsion properties has been achieved.

There is much less understanding of the behavior of more complex mixtures of proteins in emulsions and the stability behavior of emulsions under processing environments commonly encountered in the food industry. In addition, there is an almost complete lack of understanding of the behavior of emulsions during oral processing in the mouth as well as during digestion processes. It is critical to understand the oral behavior of emulsions, as common sensorial attributes (e.g. creaminess, smoothness), and that the release of fat-soluble flavors are based on interfacial structures and rheological parameters. There is some evidence to show that the behavior of emulsions in the gastrointestinal tract is affected by their physico-chemical properties and that the properties of the interface modulate fat digestion and consequently influence the bioavailability of lipid nutrients. This area of research needs to be further developed before the knowledge can be used to develop novel products with health and sensory attributes.

References


Milk protein–polysaccharide interactions

Kelvin K. T. Goh, Anwesha Sarkar and Harjinder Singh

Abstract

Proteins and polysaccharides are common ingredients in many food formulations. They are generally responsible for imparting key sensory attributes (e.g. textural attributes, controlled flavor release) and are capable of modifying phase stability in colloidal food systems. Their physicochemical properties depend not only on the molecular parameters of the individual biopolymers but also on the nature of interactions between the protein and polysaccharide molecules.

This chapter provides a basic overview of the possible types and natures of the interactions that can occur between protein and polysaccharide molecules in aqueous solutions. A description of the phase diagram that is commonly used to estimate phase stability is given. Extensive research carried out in this field over the last few decades, outlining different milk protein–polysaccharide interactions, is summarized in tables. The last sections attempt to categorize the different types of interactions and their impact on the microstructures and the rheological properties of the systems. The chapter concludes by stressing the importance of understanding these interactions, which potentially provide food scientists with the opportunity to modify or create novel food structures and functionalities.

Introduction

Proteins and polysaccharides are broadly classified as biopolymers because of their large molecular structures. These macromolecules are known to play important...
physico-chemical roles, such as imparting thickening, stabilizing, gelling, emulsifying properties etc., in food products (Hemar et al., 2001a, 2001b; Dickinson, 2003; Dickinson et al., 2003). The physico-chemical properties of proteins and polysaccharides have individually been studied extensively over the last several decades. It is well established that the factors influencing the physico-chemical properties of these macromolecules in solution include molar mass, molecular conformation, polydispersity, charge density, concentration, pH, ionic strength, temperature, solvent quality and nature of molecular (intra-/inter-) interactions (Tolstoguzov, 1997a; Doublier et al., 2000; de Kruif and Tuinier, 2001).

In many food systems, their physical properties become more complex as both proteins and polysaccharides are present (either naturally or added as ingredients) in the complex multi-component mixtures. The overall stability and the microstructure of the food systems depend not only on the physico-chemical properties of proteins or polysaccharides alone, but also on the nature and strength of the interactions between proteins and polysaccharides (Dickinson, 1998). This chapter reviews a number of recent studies on protein–polysaccharide interactions, with a particular focus on milk proteins and a diverse range of polysaccharides in aqueous systems.

### Mixing behavior of biopolymers

When aqueous solutions of proteins and polysaccharides are mixed, one of four phenomena can arise: (a) co-solubility; (b) thermodynamic incompatibility; (c) depletion interaction (or flocculation); (d) complex coacervation (Figure 12.1) (Tolstoguzov, 1991, 1997a; Schmitt et al., 1998; Syrbe et al., 1998; de Kruif and Tuinier, 2001; Benichou et al., 2002; Dickinson, 2003; Tolstoguzov, 2003; de Kruif et al., 2004; Martinez et al., 2005). These phenomena can be explained as follows.

**Co-solubility** refers to the creation of a stable homogeneous solution, i.e. the generation of one phase in which the two macromolecular species either do not interact or exist as soluble complexes in the aqueous medium. When intermolecular attraction is absent, macromolecules are co-soluble only in dilute solution, where the entropy of mixing favors more randomness in the system (Tolstoguzov, 2003).

To achieve co-solubility from a thermodynamic viewpoint, the Gibbs’ free energy of mixing ($\Delta G_{\text{mixing}}$), given in Equation (12.1), must be negative. This means that the entropy of mixing should favorably exceed the enthalpy term (note: the highest level of entropy is achieved when the different kinds of molecules are randomly distributed throughout the system) (McClements, 2005). The expression for the Gibbs’ free energy accompanying mixing under standard conditions is given by

$$
\Delta G_{\text{mixing}} = \Delta H_{\text{mixing}} - T \Delta S_{\text{mixing}}
$$

where $\Delta G_{\text{mixing}}$, $\Delta H_{\text{mixing}}$ and $T \Delta S_{\text{mixing}}$ are the free energy, enthalpy (interaction energy) and entropy changes between the mixed and unmixed states respectively.

When the size of the molecules is small, as it is in the case of monomer sugars and hydrophilic amino acids, mixing the two species results in a co-soluble system.
However, with increasing molecular weight and concentration of the polymers, the system tends to become less co-soluble as a result of thermodynamic incompatibility (Tolstoguzov, 1991, 1997a), because the entropy of mixing of biopolymers is significantly lower than that of monomers. The bulky size and the rigid structure of biopolymer molecules decrease the entropy of mixing, resulting in a higher free energy. For a mixed biopolymer solution, the enthalpy–entropy balance generally results in mutual exclusion of one biopolymer from the local vicinity of the other biopolymer. This means that biopolymers in mixed solution show a preference to be surrounded by their own type; otherwise, their mixtures separate into liquid phases (Grinberg and Tolstoguzov, 1972; Tolstoguzov et al., 1985; Tolstoguzov, 1988, 1991; Grinberg and Tolstoguzov, 1997; Polyakov et al., 1997).

Thermodynamic incompatibility occurs when the two non-interacting macromolecular species separate into two different phases, as the enthalpy of mixing exceeds the entropy difference (Grinberg and Tolstoguzov, 1997; Schmitt et al., 1998; Benichou et al., 2002; Tolstoguzov, 2002). Each of the two distinct immiscible aqueous phases formed is loaded mainly with only one biopolymer species, i.e. a protein-rich phase and a polysaccharide-rich phase. Phase separation as a result of incompatibility can also occur if each biopolymer shows varying affinity towards the solvent (Tolstoguzov, 1991; Piculell and Lindman, 1992). In this case, solvent–protein (or solvent–polysaccharide) interactions are favored over protein–polysaccharide...
interactions and solvent–solvent interactions, leading to two phases—one enriched in protein and the other enriched in polysaccharide (Doublier et al., 2000).

Thermodynamic incompatibility can also arise within a mixture of polysaccharides or proteins. Some examples include: polysaccharides with different structures; proteins of different classes, such as water-soluble albumins with salt-soluble globulins; native and denatured forms of the same protein as well as aggregated and non-aggregated forms of the same protein (Tolstoguzov, 2002).

Thermodynamic incompatibility is highly dependent on pH and ionic strength and is prevalent when protein and neutral polysaccharide are present or when both protein and polysaccharide carry the same negative charge at neutral pH (Doublier et al., 2000). Although thermodynamic incompatibility is prevalent in mixed polymer systems, some of these systems do not achieve thermodynamic equilibrium within a limited timescale because of the presence of kinetic energy barriers. When the kinetic energy exceeds the thermal energy of the system, the molecules become “trapped” in a metastable state (McClements, 2005). Some examples of kinetic energy barriers include the formation of a gel network within an incompatible system or a highly viscous continuous phase that slows down the phase separation process. The choice of which phase to gel and the component used to promote gelation depends on the type of biopolymers used in the system (Bryant and McClements, 2000a, 2000b; Norton and Frith, 2001; Kim et al., 2006).

Depletion interaction (or flocculation) usually involves spherical particles in the presence of macromolecules (Asakura and Oosawa, 1954, 1958; Bourriot et al., 1999a). Phase separation of particulate suspensions is enhanced by the addition of a polymer. This phenomenon usually occurs in a colloidal dispersion in the presence of non-interacting polymers (e.g. polysaccharides in an emulsion, polysaccharides and colloidal casein micelles). The higher osmotic pressure of the polymer molecules surrounding the colloidal particles (as compared with the inter-particle region) causes an additional attractive force between the particles, leading to their flocculation. The attractive force depends on the size, shape and concentration of the polymer molecules and the colloidal particles (Hemar et al., 2001b).

When colloidal particles approach each other, the excluded (or depleted) layer starts to overlap, allowing more space for the polymer molecules. The increase in volume causes the total entropy of the system to increase (i.e. the free energy to decrease), which in turn encourages attraction interaction between the colloidal particles (de Bont et al., 2002). In a mixed protein–polysaccharide system where the protein species is casein micelles, phase separation is often attributed to a depletion flocculation phenomenon (Bourriot et al., 1999a; Tuinier and De Kruif, 1999; Tuinier et al., 2000), because of the large colloidal particle size of the casein micelles and because increasing the concentration of polysaccharides results in greater attraction between the casein micelles (Doublier et al., 2000).

Complex coacervation is the formation of electrostatic complexes between protein and polysaccharide molecules, leading to a two-phase system. One phase has both biopolymers in a complex matrix and the other phase contains mainly the solvent water and is depleted in both biopolymers. Complex coacervation commonly occurs between oppositely charged biopolymers. Complex coacervation between oppositely
charged proteins and polysaccharides was first reported with the mixing of gelatin and gum arabic in acetic acid solution (Tiebackx, 1911).

The term “coacervation” was first introduced in 1929 to describe a process in which aqueous colloidal solutions separate into two liquid phases: one rich in colloid, i.e. the coacervate, and the other containing little colloid (Bungenberg de Jong and Kruyt, 1929). If the two biopolymers are present in equal proportions by weight at a pH such that they carry net equal opposite charges, the yield of coacervate will be at its maximum (Schmitt et al., 1998). The size and the morphology of these structures may be exploited to bring about new functionalities and textural changes in processed foods.

Phase diagram

Mixing two aqueous solutions of protein and polysaccharide will give rise to a one-phase system or a two-phase system, depending on the solution composition and the environmental conditions, as depicted in Figure 12.1 (Tolstoguzov, 1991, 1997a; Schmitt et al., 1998; Syrbe et al., 1998; de Kruif and Tuinier, 2001; Benichou et al., 2002; Tolstoguzov, 2002; Dickinson, 2003; de Kruif et al., 2004; Martinez et al., 2005).

In a one-phase system, protein and polysaccharide can exist either as individual molecules or as soluble complexes that are uniformly dispersed throughout the entire system. However, with increasing molecular weight and concentration of the biopolymers, the system tends to become less co-soluble and gives rise to a two-phase system, i.e. the system separates into two distinct phases that have different biopolymer concentrations.

For a system with relatively strong net repulsion between protein and polysaccharide in aqueous solution, the two biopolymers move into two different phases as a result of thermodynamic incompatibility. Two distinct immiscible aqueous phases are formed and each is loaded mainly with only one biopolymer species, i.e. one phase is protein rich and the other phase is polysaccharide rich. A typical phase diagram for segregating a biopolymer system is shown in Figure 12.2 and has been explained by many researchers (Grinberg and Tolstoguzov, 1972; Polyakov et al., 1980; Antonov et al., 1982; Tolstoguzov et al., 1985; Grinberg and Tolstoguzov, 1997; Bourriot et al., 1999a; Closs et al., 1999; Clark, 2000; Lundin et al., 2003; Thaïudom and Goff, 2003; Tolstoguzov, 2003; Ercelebi and Ibanoglu, 2007).

The phase diagram consists of a typical binodal curve (the solid line curve), which divides the single-phase miscible region (below the curve) from the two-phase immiscible region (the shaded region). The binodal branches show the points of limited co-solubility. The points of the binodal curve connected by tie lines represent the compositions of the co-existing equilibrium phases. From the phase diagram, it is possible to determine the effective concentrations of biopolymers in the two phases and the concentrations at which maximal co-solubility of the biopolymers is achieved. In addition, it helps to establish which of the two biopolymers forms the continuous phase.
The interactions responsible for complex formation between biopolymers can be classified as: weak or strong, specific or non-specific, attractive or repulsive (Dickinson, 1998). The overall interaction between protein and polysaccharide is the sum of the following different intermolecular forces arising between the various segments and chains of the two biopolymers (Dickinson, 1998; Schmitt et al., 1998).

**Figure 12.2** A typical phase diagram showing a protein–polysaccharide solution with water as the solvent at a particular pH, temperature and ionic strength. A sample of composition O (which was initially made with A% protein and B% polysaccharide) separates out into two bulk polymer-rich phases. The protein-enriched phase will have composition C% protein whereas the polysaccharide-enriched phase will have composition D% polysaccharide. The binodal (solid) curve separates the single-phase region from the two-phase domain (obtained by direct observation of the phase separation in test tubes). The % protein in the polysaccharide phase will be negligible and vice versa. The tie line is obtained by joining C and D. The ratio DO/OC represents the volume ratio of the protein-rich phase C and the polysaccharide-rich phase D by the inverse-lever rule. If O is shifted along the tie line to $O_1$, the new phase volume ratio will be $DO_1/O_1C$. Although any composition lying on the same tie line results in the same effective concentration in the enriched phases, the phase volume varies. The line obtained by joining the mid points (+) of two or more tie lines gives the rectilinear diameter. The co-ordinates of the critical point E (obtained from the intersection of the binodal curve to the rectilinear diameter) show the composition of a system separating into two phases of the same volume and composition, which means that the separated-phase systems will have 50% protein and 50% polysaccharide in the same phase volume ratio. Point F is the separation threshold, which is the minimum critical concentration required for the biopolymers to separate into two phases.

**Nature of interactions in protein–polysaccharide systems**

The interactions responsible for complex formation between biopolymers can be classified as: weak or strong, specific or non-specific, attractive or repulsive (Dickinson, 1998). The overall interaction between protein and polysaccharide is the sum of the following different intermolecular forces arising between the various segments and chains of the two biopolymers (Dickinson, 1998; Schmitt et al., 1998).

**Repulsive interactions**

Repulsive interactions are always non-specific and of transient duration. They usually arise from excluded volume effects and/or electrostatic interactions and tend to be weak, except at very close range or at very low ionic strength.

The excluded volume or steric exclusion effects are the non-specific and transient interactions that can be found in non-ionic, non-penetrable polysaccharides
and polypeptides that cannot occupy the same solution volume (Tolstoguzov, 1991; Polyakov et al., 1997; Schmitt et al., 1998; Tolstoguzov, 2002, 2003). Excluded volume effects exhibit mutual spatial restrictions and competition between the biopolymers for solution space, i.e. there is a reduction in the mixing entropy of the system because of the reduction in the volume available for the biopolymer molecules to occupy.

Net repulsive interactions, because of electrostatic effects, depend largely on the pH and the ionic strength of the background electrolyte concentration. Electrostatic repulsive interactions are commonly found in mixtures of proteins and anionic polysaccharides under conditions where both biopolymers carry the same net charge, e.g. pH above the isoelectric point (pI) of the protein.

**Attractive interactions**

Attractive interactions between proteins and polysaccharides may be weak or strong and either specific or non-specific. Non-specific attractive interactions arise as a result of a multitude of weak interactions between groups on the biopolymers, such as electrostatic, van der Waals’, hydrogen bonding and hydrophobic interactions. Hydrogen bonding and hydrophobic interactions are actually collective interactions (e.g. electrostatic, van der Waals’ and steric overlap) including some entropy effects (McClements, 2005).

Electrostatic interactions are the most important forces involved in complex formation between proteins and ionic polysaccharides. These interactions between charged biopolymers lead to a decrease in the electrostatic free energy of the system. Moreover, the enthalpy contribution, because of interactions of oppositely charged biopolymers and liberation of counter ions along with water molecules, often compensates for the loss of configurational entropy of mixing rigid biopolymers (Piculell and Lindman, 1992; Tolstoguzov, 1997a).

Strong electrostatic attractive interactions between positively charged proteins (pH < pI) and anionic polysaccharides occur, especially at low ionic strength. Generally, two types of complexes are formed by electrostatic interactions (Tolstoguzov, 1997b; Schmitt et al., 1998; Tolstoguzov, 2002, 2003). Soluble complexes are obtained when the opposite charges carried by the two biopolymers are not equal in number, whereas insoluble complexes result when the net charge on the complex is close to zero.

van der Waals’ forces are extremely weak electrical attractions that arise because of temporary dipole interactions (Sherony and Kintner, 1971; Stainsby, 1980; Dickinson, 1998). Basically, every atom has an electron cloud that can yield a temporary electric dipole. The dipole in one atom can induce a corresponding dipole in another atom. This is possible only if the atoms are close. However, if they are too close, repulsive forces between the adjacent negatively charged electron clouds may not allow these van der Waals’ attractions. Although these transient electrical attractive forces are very weak, they influence macromolecular interactions together with other non-covalent forces described above (Damodaran, 1997).

Hydrophobic bonding is an entropy-driven long-range interaction between non-polar groups and is promoted by conformational and structural modifications of
Milk protein–polysaccharide interactions

biopolymers, mostly by unfolding of polymeric chains exposing hydrophobic groups. These kinds of interactions are promoted by an increase in temperature (Stainsby, 1980; Piculell and Lindman, 1992; Samant et al., 1993; Antonov et al., 1996; Tolstoguzov, 1997a).

Hydrogen bonding is a moderately strong bond —O—H⁺…δ⁻—O—, which becomes relatively insignificant at high temperature. These bonds are ionic in nature and refer to the interaction between hydrogen atoms attached to an electronegative atom (oxygen, sulfur) and another electronegative atom (e.g. the sulfur of a sulfate group). A classical example of hydrogen bonding has been shown in the complex coacervation of gelatin and pectin (Braudo and Antonov, 1993), which is obtained over a wide range of pH including the isoelectric pH (4.8) of gelatin. Protein–polysaccharide hydrogen bonding in gelatin–pectin, gelatin–alginate and chitosan–collagen over a wide range of pH has been well established by various studies (Taravel and Domard, 1995; Antonov et al., 1996).

Covalent bonds

Covalent bonds are very strong, specific, non-electrostatic and permanent linkages. There are two principal methods that can be used to generate a covalent linkage between proteins and polysaccharides. The most commonly used method utilizes the chemical reaction between amino groups of proteins and carboxylic groups of polysaccharides (the Maillard reaction) to give an amide covalent bond (Stainsby, 1980). Covalent bonds can also be generated enzymatically using the oxidoreductase family of enzymes (E.C. 1.XXX), which catalyze the oxidation of the phenolic group of tyrosine residues with carbohydrate groups containing phenolic residues, such as cereal arabinoxylans (Boeriu et al., 2004). Tyrosine-containing peptides have also been conjugated with ferulic acid (Oudgenoeg et al., 2001) and with whey proteins through the use of three different oxidoreductases (Faergemand et al., 1998).

Recently, cross-linking of proteins and polysaccharides using transglutaminase (E.C. 2.3.2.13) has been suggested (Flanagan and Singh, 2006). Many polysaccharides contain residual protein; for example, gum arabic, guar gum and locust bean gum all contain low levels of protein. Gum arabic (approximately 2% protein, depending on the source) consists of, among other sub-units, a glycoprotein and an arabino-galactan protein. Provided the residual protein in these polysaccharides contains lysine and/or glutamine residues, theoretically, the treatment of protein and polysaccharide mixtures with transglutaminase could lead to the formation of heteropolymers (i.e. protein–polysaccharide conjugates) in addition to homopolymers (cross-linked protein or cross-linked polysaccharide). Flanagan and Singh (2006) demonstrated that sodium caseinate–gum arabic conjugates catalyzed by transglutaminase can be produced.

These kinds of interactions are generally very stable to pH and ionic strength. Because of their stability properties, this type of bonding has been used intentionally to produce conjugated emulsifiers (Shepherd et al., 2000; Song et al., 2002; Akhtar and Dickinson, 2003, 2007; Neirynck et al., 2004; Dunlap and Côté, 2005; Benichou et al., 2007). In most of these studies, the covalent conjugation between protein and polysaccharides has been studied through use of the Maillard reaction.
Apart from these major interactions, ion bridging involving binding of cations such as Ca\(^{2+}\) may also contribute to some extent in protein–polysaccharide interactions although they do not have the predominant influence (Stainsby, 1980; Antonov et al., 1996; Dickinson, 1998). For example, firm sodium caseinate gels (\(G' > 100\) Pa) were formed using pectin concentrations \(\geq 0.6\%\) at one particular degree of methylation (=31\%) and amidation (=17\%) in the presence of Ca\(^{2+}\) ions (1.8 mM) at pH \(\approx 3.6\) (Matia-Merino et al., 2004).

**Milk protein–polysaccharide interactions**

Milk proteins together with polysaccharides dissolved in an aqueous phase form a pseudo-ternary system of milk protein–polysaccharide–water. Various interactions in these systems could lead to complex formation or bulk phase separation. Extensive studies on protein–polysaccharide interactions, particularly using well-studied milk proteins and commercially available polysaccharides, have been carried out (Dickinson, 1998). Table 12.1 and Table 12.2 show a compilation (non-exhaustive) of various milk protein (casein and/or whey protein) and polysaccharide mixtures in aqueous systems and the conditions under which different kinds of interactions occur.

**Rheological properties and microstructures of protein–polysaccharide systems**

The rheological properties of a solution containing only protein are expected to be different from those of a pure polysaccharide solution. Polysaccharide molecules generally have a greater effect than proteins in causing a significant increase in solution viscosity, because polysaccharide molecules are usually much larger and more extended (\(\approx 5.0 \times 10^5\) to \(2.0 \times 10^6\) Da) than globular proteins (\(\approx 1.0 \times 10^4\) to \(1.0 \times 10^5\) Da). Hence, polysaccharide molecules generally occupy larger hydrodynamic volumes, which give rise to higher solution viscosity. The above assumes that intermolecular interactions are absent or negligible (e.g. in dilute solution). When intermolecular interactions among neighboring polymer molecules (i.e. polysaccharide–polysaccharide or protein–protein interactions) are present, the rheological properties of the systems can be expected to change significantly.

Changes in rheological properties may arise as a result of an increase in the size of the particles (e.g. protein–polysaccharide complexes), or when depletion interactions occur in the mixed system or if one or more polymer species form continuous network structures. The overall effect results in the formation of different microstructures. Schematic illustrations of some possible microstructures formed from mixtures of proteins and polysaccharides under some specific conditions (e.g. pH, ionic strength, heat treatment, etc.) are shown in Figures 12.3a and 12.3b.

Various rheological techniques have been employed to characterize the physicochemical properties of protein–polysaccharide systems. Generally, if the mixtures are liquid like, viscosity measurements using rotational viscometers are commonly
### Table 12.1 Milk protein–polysaccharide interactions in aqueous systems

<table>
<thead>
<tr>
<th>No.</th>
<th>Casein–polysaccharide aqueous systems</th>
<th>Conditions</th>
<th>Interactions</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td><strong>Milk proteins</strong> (Casein micelles + Whey proteins) + <strong>Pectin</strong> (High methoxyl – 62.7% methylated)</td>
<td>20°C, pH 6.0–10.5, 0–0.5 M NaCl</td>
<td>Thermodynamic incompatibility</td>
<td>(Antonov et al., 1982)</td>
</tr>
<tr>
<td>2.</td>
<td><strong>Casein micelles + Alginate</strong></td>
<td>pH 7.2, 25°C</td>
<td>Thermodynamic incompatibility</td>
<td>(Suchkov et al., 1981, 1988)</td>
</tr>
<tr>
<td>3.</td>
<td><strong>Casein micelles</strong> (2.5%) + <strong>Pectin</strong> (Low methoxyl – 35%, High methoxyl – 73%, Low methoxyl amidated – 35% methylated and 20% amidated) (0.1–0.2%)</td>
<td>pH 6.7/5.3, 60°C</td>
<td>pH 6.7: Depletion interaction. Methylation affects interaction</td>
<td>(Maroziene and de Kruif, 2000)</td>
</tr>
<tr>
<td>4.</td>
<td><strong>Casein micelles</strong> (0.8–4%) + <strong>Galactomannans</strong> (Guar gum, Locust bean gum) (0.09–0.3%)</td>
<td>5/20°C, pH 6.8/7.0, 0.08/0.25 M NaCl, sucrose (10–40 w/w%)</td>
<td>Depletion interaction. Sucrose affects interaction</td>
<td>(Bourriot et al., 1999a; Schorsch et al., 1999)</td>
</tr>
<tr>
<td>5.</td>
<td><strong>Casein micelles</strong> (1.0%) + <strong>Carrageenan</strong> (ι, κ, λ-forms) (0.12%)</td>
<td>pH 6.7/7.0, 60/50/20°C, 0.25 M NaCl/0.05 M NaCl–0.01 M KCl</td>
<td>Depletion interaction</td>
<td>(Dalgleish and Morris, 1988; Langendorff et al., 1997, 1999, 2000; Bourriot et al., 1999b)</td>
</tr>
<tr>
<td>6.</td>
<td><strong>Sodium caseinate</strong> (0.1–0.5%) + <strong>Gum arabic</strong> (0.01–5%)</td>
<td>0.5 M NaCl, pH 2.0–7.0, slow acidification with glucono-δ-lactone</td>
<td>Soluble electrostatic complexation</td>
<td>(Ye et al., 2006)</td>
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<td>7.</td>
<td><strong>Casein micelles</strong> (0.1%) + <strong>Exopolysaccharide</strong> (5.0%) (Lactococcus lactis subsp. cremoris B40)</td>
<td>pH 6.6, 25°C</td>
<td>Depletion interaction</td>
<td>(Tuinier &amp; De Kruif, 1999; Tuinier et al., 1999)</td>
</tr>
<tr>
<td>8.</td>
<td><strong>Sodium caseinate + Maltodextrin</strong> (2:1, 1:1 and 1:4)</td>
<td>60°C, 2–4 days</td>
<td>Covalent conjugate via Maillard reaction. No phase separation</td>
<td>(Shepherd et al., 2000; Morris et al., 2004)</td>
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<td>9.</td>
<td><strong>Casein</strong> (β-Casein, αs-Casein) + <strong>Polysaccharide</strong> (Dextran, Galactomannan) (1:1)</td>
<td>60°C, 24 h</td>
<td>Covalent conjugate via Maillard reaction. No phase separation</td>
<td>(Dickinson and Semenova, 1992; Kato et al., 1992)</td>
</tr>
<tr>
<td>10.</td>
<td><strong>Sodium caseinate</strong> (6.0%) + <strong>Sodium alginate</strong> (1%)</td>
<td>pH 7.0, 23°C</td>
<td>Thermodynamic incompatibility</td>
<td>(Guido et al., 2002; Simeone et al., 2002)</td>
</tr>
<tr>
<td>No.</td>
<td>Whey protein–polysaccharide aqueous systems</td>
<td>Conditions</td>
<td>Interactions</td>
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<tr>
<td>1.</td>
<td>β-Lactoglobulin (β-Lg) (0.5%) + Chitosan</td>
<td>pH 3.0–7.0, 5 mM phosphate buffer</td>
<td>pH-dependent β-Lg–chitosan Soluble/insoluble complex coacervation</td>
<td>(Guzey and McClements, 2006)</td>
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<td></td>
<td>(Degree of deacetylation: 85%) (0–0.1%)</td>
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<td>2.</td>
<td>Heat-denatured whey protein isolate (HD-WPI) (8.0%) + Pectin (28, 35, 40, 47 and 65% methylation) (0.1–1.5%)</td>
<td>pH 6.0/7.0, 80°C/85°C, 5.0/10.0 mM CaCl₂</td>
<td>Thermodynamic incompatibility</td>
<td>(Beaulieu et al., 2001; Kim et al., 2006)</td>
</tr>
<tr>
<td>3.</td>
<td>β-Lg (12.0%) + Alginate (0.1–1.0%)</td>
<td>pH 7.0/(3.0–7.0), 87°C/30°C, high pressure</td>
<td>pH-dependent β-Lg–chitosan Soluble/insoluble complex coacervation</td>
<td>(Dumay et al., 1999; Harnsilawat et al., 2006)</td>
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<tr>
<td>4.</td>
<td>β-Lg (0.05%) + Pectin (Low methoxyl – 28.3/42.6%, High methoxyl – 71.3/73.4%) (0.0125%)</td>
<td>pH 4–40°C/25°C, pH 4.0–7.5/6.5, 0.11/0.1–1.0 M NaCl/87°C/high pressure</td>
<td>pH, ionic strength and temp.: Complex coacervation. Precipitation for modified pectin. Methylation affects complexation</td>
<td>(Dumay et al., 1999; Wang and Qvist, 2000; Girard et al., 2002, 2003a, 2003b, 2004; Kazmierski et al., 2003)</td>
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<tr>
<td>5.</td>
<td>Whey protein isolate (WPI) (5.0%) + Galactomannans (Locust bean gum) (0–0.4%)</td>
<td>pH 5–7</td>
<td>pH and concentration: Biphasic gel</td>
<td>(Tavares and Lopes da Silva, 2003)</td>
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<td>6.</td>
<td>HD-WPI (8.5%) + Xanthan gum (0–0.2%)</td>
<td>pH 7.0/5.4, high pressure treatment, 0.2 M NaCl</td>
<td>Native WPI: Co-solubility. HD-WPI: Thermodynamic incompatibility</td>
<td>(Bryant and McClements, 2000b; Li et al., 2006)</td>
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<td>7.</td>
<td>WPI (4–12.5%) + Xanthan gum (0.01–1.0%)</td>
<td>pH 5.5/6.0/6.5/7.0, 0.1/0.5 M NaCl, high-pressure treatment</td>
<td>Depletion interaction, pH-dependent electrostatic complexation</td>
<td>(Zasypkin et al., 1996; Laneuville et al., 2000; Hemar et al., 2001b; Benichou et al., 2007; Bertrand and Turgeon, 2007)</td>
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<td>8.</td>
<td>Bovine serum albumin (BSA) + Sulfated polysaccharides (κ-, κ-carrageenan, Dextran sulfate) (2.5:1 and 5:1)</td>
<td>pH 6.5–8, high-pressure treatment</td>
<td>Complex coacervation</td>
<td>(Galazka et al., 1996, 1997, 1999)</td>
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<td>9.</td>
<td>HD-WPI (10.0%) + κ-Carrageenan (0.5%)</td>
<td>pH 7.0, pH 1–12</td>
<td>Complex coacervation</td>
<td>(Mleko et al., 1997)</td>
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<td>10.</td>
<td>β-Lg (0.5–10.0%) + κ-Carrageenan (1.0%) (1.2, 5:1 and 10:1)</td>
<td>pH 7, 45–80°C, 0.1 M NaCl/0.01 M CaCl₂</td>
<td>Temp., pH and concentration dependent. Phase-separated bicontinuous gel formation</td>
<td>(Capron et al., 1999; Ould Eleya and Turgeon, 2000)</td>
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<th>No.</th>
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<tr>
<td>11.</td>
<td>β-Lg + Gum arabic (2:1)</td>
<td>pH 3.6-5.0, 0.005-10.7 mM NaCl</td>
<td>Complex coacervation</td>
<td>(Schmitt et al., 1998, 1999, 2000, 2001; Sanchez &amp; Renard, 2002; Sanchez et al., 2002, 2006)</td>
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<td>12.</td>
<td>WPI + λ-Carrageenan (1:1 to 150:1)</td>
<td>pH: Wide range, 0–0.1 M (NaCl/CaCl₂)</td>
<td>Electrostatic complexation. Precipitation</td>
<td>(Weinbreck et al., 2004a)</td>
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<td>13.</td>
<td>WPI + Gum arabic (2:1)</td>
<td>pH 4.0-7.0, 0–0.1 M NaCl</td>
<td>Complex coacervation. Glassy state</td>
<td>(Weinbreck et al., 2003a, 2004b, 2004c)</td>
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<td>14.</td>
<td>β-Lg + Carboxymethyl dextran (1:1 and 7:2)</td>
<td>pH 5.5/4.75, 4°C/25°C</td>
<td>β-Lg–carboxymethyl dextran covalent conjugate. No phase separation</td>
<td>(Hattori et al., 1994)</td>
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<td>15.</td>
<td>WPI + Carboxymethyl potato starch (2:1)</td>
<td>pH 7.0, 24°C</td>
<td>WPI–carboxymethyl starch covalent conjugate</td>
<td>(Hattori et al., 1995)</td>
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<td>16.</td>
<td>WPI + Exopolysaccharide (Lactococcus lactis subsp. cremoris B40) (2:1)</td>
<td>pH: Wide range, 25°C, 0–0.1 M (NaCl/CaCl₂), heat treatment of WPI</td>
<td>Electrostatic complexation. Precipitation HD-WPI: Depletion interaction</td>
<td>(de Kruif and Tuinier, 1999; Tuinier &amp; de Kruif, 1999; Weinbreck et al., 2003b)</td>
</tr>
<tr>
<td>17.</td>
<td>β-Lg + Pullulan</td>
<td>0.01 M NaCl, 4°C</td>
<td>Depletion interaction</td>
<td>(Wang et al., 2001)</td>
</tr>
<tr>
<td>18.</td>
<td>β-Lg + Carboxymethyl cellulose</td>
<td>60°C, 0.05–0.2 M, pH 2.5–7.0</td>
<td>Insoluble electrostatic complex, sedimentation</td>
<td>(Hidalgo and Hansen, 1969; Hansen et al., 1974)</td>
</tr>
<tr>
<td>19.</td>
<td>WPI + Maltodextrin (1:2 and 1:3)</td>
<td>80°C, 2 h, 79% RH</td>
<td>Covalent conjugation. No phase separation</td>
<td>(Akhtar and Dickinson, 2007)</td>
</tr>
<tr>
<td>20.</td>
<td>WPI/Whey Protein Concentrate (WPC) + Pectin (4:1, 2:1, 1:1 and 1:2)</td>
<td>60°C, 14 days, pH 7.0</td>
<td>Covalent conjugation. No phase separation</td>
<td>(Mishra et al., 2001; Neirynck et al., 2004)</td>
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Figure 12.3a  Schematic diagrams of some possible microstructures formed between non-interacting protein–polysaccharide mixtures. Circle (●) represents protein; coil structure represents polysaccharide molecules. (a) Flocculated protein network formed with polysaccharide filling the space in the network; (b) polysaccharide molecules overlap and form continuous “network” with protein filling the space; (c) particulate protein gel network formed with polysaccharide filling the space; (d) polysaccharide gel network formed with protein filling the space; (e) bicontinuous network formed from protein and polysaccharide; (f) polysaccharide gels dispersed among a weakly flocculated protein network; (g) protein gels dispersed among entangled polysaccharide molecules.

Figure 12.3b  Schematic diagrams of some possible microstructures formed between interacting protein–polysaccharide mixtures. Circle (●) represents protein; coil structure represents polysaccharide molecules. (a) Protein–polysaccharide complexes formed; (b) protein interacting with gelling polysaccharide helices; (c) polysaccharide interacting with protein particulate gel network; (d) polysaccharide gel helices interacting with protein particulate gel network.
used to obtain steady-state viscosity curves, yield stress, etc. Other simpler methods include the use of a kinematic viscometer (e.g. an Ubbelodhe capillary viscometer) to obtain a single point relative viscosity measurement. If the samples are viscoelastic (e.g. gels), rheometers are widely used to obtain rheological data (e.g. loss and storage moduli obtained within the linear viscoelastic region), by performing small deformation oscillatory measurements.

The rheological data yield information on the viscosity and viscoelastic properties of the mixed systems. Knowledge of the rheological properties of mixed protein–polysaccharide systems is essential to gain insights into the nature of the interactions and the resulting microstructure of the system. Fundamental understanding of the interactions at the molecular and colloidal levels will provide a strong foundation to exploit the physical functionality of such complex systems in different applications (e.g. microencapsulation technology, imparting specific sensory characteristics, time/temperature/pH/ionic control-release, emulsion stability, etc.).

In the following sections, various examples of mixed systems involving different milk proteins and polysaccharides are provided. An attempt is made to classify these mixed systems into two broad categories (i.e. interacting and non-interacting). Under each of these headings, the systems are further grouped according to whether they form or do not form gels (i.e. gelling or non-gelling). The discussion focuses mainly on the techniques used and the rheological properties of the systems.

**Non-interacting protein–polysaccharide mixtures**

Non-interacting protein–polysaccharide mixtures existing as one phase are rare but may occur when the two different molecular species have good chemical resemblance of their hydrophilic surfaces (Tolstoguzov, 1991, 2006). However, many polymer mixtures are thermodynamically incompatible and segregative interactions often occur in the absence of electrostatic interaction or in the presence of electrostatic repulsion (Neiser et al., 1998). Protein–polysaccharide mixtures that commonly exist as two separate phases are the result of either thermodynamic incompatibility or depletion flocculation (Doublier et al., 2000).

**Non-gelling phase-separated systems**

The following are examples of non-interacting, non-gelling protein and polysaccharide mixtures. The proteins and polysaccharides were mixed under conditions where the mixtures did not form gels. The rheological properties of these systems are discussed in relation to their interactions and the microstructures formed.

**Casein micelles and galactomannans**

A non-interacting protein–polysaccharide mixture where phase separation occurred was reported for a mixed system consisting of micellar casein (3%) and guar gum (0.2%) at pH 7 (Bourriot et al., 1999c). There was a significant change in the flow and viscoelastic properties compared with the individual biopolymer systems. There was an increase in the apparent viscosity in the mixed system. Furthermore, the mechanical spectra (elastic modulus $G'$, viscous modulus $G''$) of the frequency
sweeps showed slightly higher values of the moduli, which were less frequency dependent. The results suggested the formation of a weak network structure within the system because of flocculation of the casein micelles as the polysaccharide molecules were excluded from the protein phase.

The appearance of a slightly thixotropic behavior indicated that the network could be easily broken under shear because the network formed by the micellar casein was weakly flocculated and reversible, presumably attributable to a depletion flocculation mechanism. Also the lower the intrinsic viscosity of the polysaccharide, the higher was the concentration of polysaccharide required before phase separation occurred (Bourriot et al., 1999c). An increase in the concentration of the polysaccharide resulted in stronger flocculation of the casein micelles because the volume occupied by and the osmotic pressure from the surrounding polysaccharides increased.

Similar thixotropic behavior was reported for a ternary solution consisting of micellar casein, locust bean gum (LBG) and sucrose (Schorsch et al., 1999). At pH = 6.8, the casein micelles and the LBG were thermodynamically incompatible, behaving as a water-in-water emulsion. The presence of sucrose, even at high concentration (40%), did not significantly improve the compatibility of the biopolymers (Schorsch et al., 1999).

**Milk proteins and xanthan gum**

Another study investigated the interaction between xanthan gum (0–1% w/w, a polysaccharide with known “weak gel” properties) and different types of milk proteins (5% w/w, sodium caseinate [Na-CN], skim milk powder [SMP], whey protein isolate [WPI] and milk protein concentrate [MPC]) in aqueous solution at neutral pH (Hemar et al., 2001b). The microstructures of the mixtures were different depending on the xanthan gum concentration and the protein type. In the case of xanthan gum mixtures with either MPC or SMP, depletion flocculation of the casein micelles took place. The size of the depleted protein aggregates decreased with increasing xanthan gum concentration (the microstructure resembled a particulate network).

In the case of xanthan gum mixtures with either Na-CN or ultracentrifuged WPI, no phase separation occurred within the timescale of the experiment. This was attributed to the larger size of the casein micelles (average diameter ≈ 0.2 μm) compared with the nanometre size scale of WPI and Na-CN (0.05 μm) (Lucey et al., 2000). However, the rheological behavior of the mixtures was very similar to the rheological behavior of xanthan gum. The differences in the microstructures of the mixtures that were observed by confocal laser scanning microscope (CLSM) were not detected by viscosity measurements, probably because the weakly flocculated proteins were easily redispersed by the shearing action of the viscometer during measurement.

**Gelling phase-separated systems**

In a system where two biopolymer species (e.g. proteins and polysaccharides) do not interact, gelation of one or more of the components in a thermodynamically incompatible system will cause competition between phase separation and gelation (Neiser et al., 1998). Gelation basically means the formation of a three-dimensional aggregated
network structure, which is generally induced by heating, cooling, acidification, enzymatic treatments, high-pressure processing, etc.

Generally, heating enhances hydrophobic and covalent interactions. Unfolded proteins interact to give rise to aggregates in the case of whey protein (Kinsella, 1984; Boye et al., 1997). In mixed systems, the microstructure will depend on the rates of phase separation and gel formation (Tavares et al., 2005). The gel may appear to be homogeneous at a macroscopic level, but heterogeneous at a microscopic level. However, the rheological properties of such gels depend on the concentration and arrangement of each species in the different phases. If the gelling species is in the continuous phase, the gel strength is higher than if the gelling species is in the dispersed phase where the network is disrupted (Neiser et al., 1998).

**Whey protein and galactomannans**

One study was based on a mixture of LBG (a non-gelling neutral polysaccharide) and whey protein at neutral pH and pH 5 (close to the pI of whey proteins) (Tavares and Lopes da Silva, 2003). At neutral pH, it is known that whey protein forms clear fine-stranded gels (protein aggregation is hindered by electrostatic repulsion), whereas, at lower pH (e.g. pH 5), an opaque coarse particulate gel is formed (Langton and Hermansson, 1992; Aguilera, 1995). Rheological measurements showed that a WPI gel (13% w/w) had a stronger and more elastic character at pH 5 than at pH 7 because of the thick particulate network formed (Stading et al., 1993; Bertrand and Turgeon, 2007). For the protein gels at pH 7, increasing LBG concentration (>0.25%) decreased the onset temperature for gelation and decreased the gelation time. The presence of LBG was also found to increase the gel rigidity.

The authors attributed this to a decrease in macromolecular mobility within the network in the presence of LBG because of segregative interactions and the “local” concentration of each polymer species. The LBG molecules acted as fillers in the continuous protein network. At pH 5, the elastic character of the particulate gel network was shown to decrease in the presence of LBG, especially at low protein concentration (5%). It was suggested that LBG chains hampered protein–protein interactions and were detrimental to the development of a protein gel. However, at a higher protein concentration (13%), where sufficient particulate gel network was formed, LBG acted as fillers within the network, hence improving the gel strength.

In a subsequent study carried out using WPI and guar gum at pH 7, an increase in protein gel strength with decreasing degree of branching of the galactomannans was found (Tavares et al., 2005). Like LBG, guar gum was dispersed as droplets among the whey protein network at low concentration (0.2%). However, at higher gum concentration (0.6%), the dispersed droplets joined to form a continuous polysaccharide-rich phase. Despite the different microstructures observed, the linear viscoelastic profiles were rather similar, indicating that viscoelasticity was fairly insensitive to microstructural changes of this nature.

**WPI and xanthan gum**

A very similar trend was observed for whey protein and xanthan gum mixtures after heat treatment (Bertrand and Turgeon, 2007). The microstructures and rheological
properties of the gels were highly dependent on pH and salt. At pH 6.5, the presence of xanthan gum improved the elastic modulus of the WPI gel. This was attributed to segregative phase separation, where xanthan gum was dispersed among the protein gel network. However, lowering the pH decreased the elastic character of the gel. At pH 5.5 (close to the pI of WPI), the addition of xanthan gum decreased the elastic modulus of the gel. It was suggested that possible WPI–xanthan gum complexes formed decreased protein–protein interactions, producing a weaker gel network.

**β-Lactoglobulin and pectin**

A different type of network was formed in mixtures of β-lactoglobulin (8% w/w) and low methoxyl pectin (0.85% w/w) after thermal treatment at pH 6.8. The storage modulus of the mixed gel system was significantly lower than that of the protein gel alone. The microstructure observed by CLSM revealed phase separation, with β-lactoglobulin appearing as spherical colloidal particles distributed in a continuous pectin network (Donato et al., 2005). A similar type of protein-depletion-induced phase separation was reported for a mixed system containing aggregated whey protein and an exopolysaccharide (EPS) from lactic acid bacteria (Tuinier et al., 2000).

**β-Lactoglobulin and κ-carrageenan**

If two gelling species are present in a binary system, the mixed gels may form interpenetrating, coupled or phase-separated networks (Morris, 1986). Interpenetrating networks are the result of two independent continuous networks formed throughout the gel and only topological interactions exist between the networks. Coupled networks (ordered into junction zones, like those of a polysaccharide gel) are formed when favorable interactions between the two molecular species exist. However, such systems involving protein–polysaccharide interactions are uncommon (Rao, 1999).

Phase-separated networks are formed when one polymer species is incompatible with the other polymer species, forming phase-separated regions within the gel network (Piculell and Lindman, 1992; Turgeon and Beaulieu, 2001). An example of a phase-separated gel is that for κ-carrageenan and β-lactoglobulin (Capron et al., 1999). The mixed polymer formed a gel that was weaker than the carrageenan gel alone when the protein was in its native state. On heating the mixture to 90°C, holding for 30 min and then cooling to 20°C, the gel rheology indicated the melting of κ-carrageenan and the gelation of β-lactoglobulin above 65°C. There was no aggregation of κ-carrageenan with β-lactoglobulin on heating. The gelation time of β-lactoglobulin was reduced in the presence of κ-carrageenan, which was attributed to microphase separation, which caused an increase in the local concentration of β-lactoglobulin (Capron et al., 1999). On cooling, the mixed gel system formed a phase-separated bicontinuous network (Ould Eleya and Turgeon, 2000).

**Interacting protein–polysaccharide mixtures**

Another phase separation phenomenon is associative phase separation, where associative interactions are present. Associative interactions between protein and
polysaccharide can occur as a result of electrostatic interactions, hydrogen bonding, hydrophobic interactions or poor solvent conditions (Antonov et al., 1996; Gao and Dubin, 1999; Doublier et al., 2000; de Kruif et al., 2004). In some cases, complexes are formed via electrostatic interactions (known as coacervates). Coacervates of protein–polysaccharide can occur when the pH of the mixture is lower than the isoelectric point of the protein. At this pH, the protein possesses a net positive charge whereas the polysaccharide still possesses a negative charge. The result of the complexation is the formation of a solvent-rich phase and a coacervate-rich phase (Doublier et al., 2000; Ould Eleya and Turgeon, 2000).

The rheological properties of milk protein–polysaccharide complexes are related to the interaction between the complexes and the water molecules, which forms soluble (or liquid coacervate phase) or insoluble (or precipitate) complexes. The solubility of the complexes is based on the energetic difference between biopolymer–biopolymer and biopolymer–solvent interactions (Damodaran, 1997). The main parameters affecting the solubility of biopolymer complexes are charge density, pH, ionic strength and protein:polysaccharide (PP:PS) ratio (Schmitt et al., 1998). It has been suggested that a complex involving a strong polyelectrolyte will form a precipitate rather than a liquid coacervate. A number of examples of protein–polysaccharide systems with complex coacervations have been reviewed (Schmitt et al., 1998; Turgeon et al., 2003; de Kruif et al., 2004). Some examples of interacting polymers in mixed systems and the effect on their rheological properties are given below.

**Non-gelling phase-separated systems**

**β-Lactoglobulin and chitosan**

It has been reported that the solubility of a protein increases below its isoelectric pH when it complexes with an anionic polysaccharide (Tolstoguzov et al., 1985; Tolstogusov, 1986). A recent study of a β-lactoglobulin–chitosan complex showed that the complex was either soluble or insoluble, depending on the pH (Guzey and McClements, 2006). The interaction of soluble chitosan ($M_w = 15000\text{Da}$, degree of deacetylation = 85%, 0–0.1 wt%, 5 mM phosphate buffer) with β-lactoglobulin (0.5 wt% β-lactoglobulin, 5 mM phosphate buffer) in aqueous solutions studied at pH 3–7 showed that, at pH 3, 4 and 5, the majority of the β-lactoglobulin–chitosan complex in the solutions was soluble, but that at pH 6 and 7, a significant fraction of the two biopolymers was insoluble.

**Whey proteins and EPS**

“Soluble complexes” formed via electrostatic interactions were reported for EPS B40 (an EPS from *Lactococcus lactis* subsp. cremoris NIZO B40) and whey protein (PP:PS = 2:1) under specific pH and ionic conditions (with no macroscopic phase separation) (Weinbreck et al., 2003b). Decreasing the pH of the mixtures increased further aggregation of the complexes, which led to phase separation. In addition, increasing the ionic strength of the solution caused a shift to a lower pH value for the onset of complexation. In this study, complexation in this system led to a decrease in solution viscosity, as intramolecular repulsion of the EPS was reduced in the presence
of whey proteins. The decrease in viscosity was attributed to a reduction in the quantity of dispersed phase, i.e. water present within the complexes. Consequently, it was suggested that dilute solution viscosity measurement (which is related to the size of complexes) could be used to determine the optimum conditions for complexation (Weinbreck et al., 2003b). A potential benefit of this complexation is that it protects the protein from loss of solubility as a result of aggregation during thermal or high-pressure treatments (Imeson, 1977; Galazka et al., 1997).

**Whey proteins and gum arabic**

Viscosity curves were obtained to evaluate the “strength” of electrostatic interactions of whey protein–gum arabic coacervates (Weinbreck and Wientjes, 2004). This study showed that the stronger the interaction, the greater was the shear-thinning behavior and the slower was the reformation of the complexes after shearing. The highly viscous coacervate (at pH 4) was attributed to electrostatic interactions. At pH above the isoelectric point (without electrostatic interactions), the mixtures appeared to be more elastic than viscous.

**Sodium caseinate and gum arabic**

In contrast to whey proteins, sodium caseinate and gum arabic mixtures showed some peculiar behavior (Ye et al., 2006) as no coacervation was observed in these systems. Below a certain pH (pH 5.4), electrostatic interactions between sodium caseinate and gum arabic led to the formation of stable composite nanoparticles in the size range 100–200 nm. These complexes remained constant in particle size and were stable and soluble over a defined pH range (pH 3.2–5.4). This pH range was dependent on the ratio of sodium caseinate to gum arabic in the mixtures and also on the ionic strength.

The sodium caseinate–gum arabic particles associated to form large particles, which resulted in phase separation when the pH was lower than 3.0. A mechanism for the formation of these nanoparticles, based around self-aggregation of the casein and electrostatic interaction between the aggregated particles of casein and gum arabic, was proposed. As the pH of the mixture decreased below pH 5.4, the caseinate molecules tended towards small-scale aggregation prior to large-scale aggregation and precipitation at pH values closer to their pI (pH 4.6).

In this case, the gum arabic molecules may have attached to the outside of these small-scale aggregates in the early stages of aggregation through electrostatic interactions between negatively charged gum arabic and exposed positive patches on the surface of the caseinate aggregates. The presence of hydrophilic gum arabic molecules on the outside of the caseinate aggregates may have been enough to sterically stabilize these nanoparticles and consequently prevent self-aggregation. As the charge on the nanoparticles was quite low, e.g. ≈15 mV at pH 4.0, steric stabilization was probably important.

**Casein micelles and pectin**

Protein–polysaccharide interactions were shown to be pH dependent in the case of pectin and casein micelles (Ambjerg and Jørgensen, 1991; Maroziene and de Kruif, 2000).
At pH 6.7, pectin did not adsorb on to the casein micelles. With sufficient pectin present (0.1–0.2%), phase separation occurred because of depletion interactions of the casein micelles (≈0.1%). However, adsorption of pectin on to the casein micelles occurred at pH 5.3. Viscosity measurements were employed to study the changes that occurred at different polymer concentrations. At low pectin concentrations (≈0.1%) and at pH 5.3, bridging flocculation occurred. A maximum viscosity at this pectin concentration was attributed to bridging flocculation, as bridging among the casein particles was interpreted as having a larger effective volume. As the pectin concentration increased (>0.1%), the casein micelles became fully covered and interactions between the casein particles were reduced, as, typically, in acidified milk.

When the protein was fully covered by the pectin, the viscosity decreased to a certain extent but remained higher than for the pure milk samples (without pectin). The amount of pectin required for full coverage of the casein micelles differed depending on the type of pectin: high methoxyl (HM) < low methoxyl amidated (LMA) < low methoxyl (LM) pectin. Adding more pectin beyond the concentration for full coverage led to phase separation because of depletion interactions. A further increase in pectin reduced the thickness of the casein-depleted layer as the viscosity of the continuous phase became very high and formed gelled polymer networks (Maroziene and de Kruif, 2000). When the pH of the mixture was increased from 5.3 back to 6.7, desorption of pectin from the casein occurred, but over a much longer time scale (≈10–15 min) than for the adsorption process (Maroziene and de Kruif, 2000).

Gelling phase-separated systems

Sodium caseinate and pectin
The dynamic rheological properties of glucono-δ-lactone (GDL)-acidified protein gels (2% w/v Na-CN) were studied in the presence of LMA pectin (0.01–1% w/v) at pH 4 (Matia-Merino et al., 2004). The presence of pectin (0.01–0.05% w/v) was found to decrease the storage modulus and to increase the gelation time, because pectin adsorbed on to the casein particles. At pectin concentration >0.08% w/v, acid-induced gelation appeared to be completely inhibited over a time period of ≈9 h at 25°C.

Casein micelles and ι-carrageenan
For casein–carrageenan mixed systems, the attractive interactions involved the negatively charged sulfated groups of the polysaccharides and the positive “patches” between residues 97 and 112 of κ-caseins (Snoeren, 1975), despite a pH above the isoelectric point and an overall net negative charge of the casein micelles. The interaction between ι-carrageenan (0.5%) and skim milk (based on 3.3% protein) mixtures was studied above and below the coil–helix transition temperature of carrageenan (Langendorff et al., 1999). At temperatures above the coil–helix transition temperature, carrageenan did not adsorb to the casein micelles, resulting in depletion flocculation.

In contrast, at temperatures below the coil–helix transition temperature, attractive interactions between carrageenan and casein micelles occurred. The higher charge density of the double-helix form, as compared with the coil conformation of carrageenan,
probably explained the stronger attractive interaction between casein micelles and carrageenan. The presence of casein micelles increased the gel strength (indicated by higher $G'$ and $G''$) and the gelation temperature (from 39–47°C) when the mixtures were heated to 65°C and cooled to 25°C.

Depending on the concentration of carrageenan, different types of gel network were deduced from the frequency sweep. At low carrageenan concentrations (<0.2%), one type of network was formed on cooling. This was probably due to the bridging of casein micelles by the adsorbed carrageenan helical chains. The network was much more thermally stable than the pure carrageenan gels. At above 0.2% concentration, as well as the formation of a network as described above, a second network was formed, similar to that of a carrageenan gel in the absence of proteins. This was attributed to interactions between carrageenan chains (Langendorff et al., 1999). Among the different types of carrageenans, the amount required for full coverage increased from $\kappa < \iota < \lambda$ (Langendorff et al., 1997) because the charge density of the polymer determined the strength of adsorption (Pereyra et al., 1997; Maroziene and de Kruif, 2000).

It is clear from the above examples that mixed protein and polysaccharide systems can produce very different rheological properties. The rheological properties of these systems are the results of cumulative effects from the molecular parameters (e.g. size, conformation, charge density, concentration, PP:PS ratio) of the macromolecules, the conditions (e.g. pH, temperature, ionic strength) to which the mixed systems are subjected and the resulting interactions (e.g. type of interaction, strength of interactions, gels or aggregates) among the macromolecules. Understanding the rheological properties of these systems may help in the development of novel food structures with unique sensory properties and of functionalities such as in micro-encapsulation and controlled-release applications.

**Conclusions**

Protein and polysaccharide are the two main structural entities in foods and a great deal of work on the interactions between proteins and polysaccharides has been published over the last few decades. In recent years, excellent progress has been made on understanding the key variables and interactions that control the physical stability, rheology and microstructure of protein–polysaccharide mixtures. Although milk proteins, particularly whey proteins, have been most widely used in studies of protein–polysaccharide systems, most deal with the relatively simple binary combination of one protein and one polysaccharide. More complex systems, including ternary mixtures, still remain to be investigated in detail. At a practical level, it seems to be possible to manipulate these interactions and produce different microstructures by controlling internal (pH, ionic strength, biopolymer ratio, molecular weight and charge of the biopolymer) and external (temperature, pressure and shear rate) factors. However, there is a considerable challenge in understanding how different microstructures relate to the sensory properties of food products, such as mouthfeel and flavor release.
The formation of complexes through electrostatic and covalent interactions has been the subject of intensive studies, mainly because of potentially better functionality, e.g. rheology, gelation and interfacial properties, of composites compared with the protein or polysaccharide alone. Protein–polysaccharide complexes can serve as texturizing agents, encapsulating agents, fat replacers and stabilizers of emulsions and other dispersed systems. However, information on the detailed molecular structures of protein–polysaccharide complexes is still lacking and describing the experimental observations within the known theoretical frameworks remains a challenge.

It is now becoming apparent that the modification of food structure through modulation of macromolecular interactions can also be used to control the release of nutrients and bioactive components during digestion and to target where and how such components are released. The basic science underpinning these functions is largely unknown. New knowledge in this area will enable the development of composite food systems and ingredients that are superior in nutritional value and textural characteristics.

References


Interaction between milk proteins and micronutrients

T. Considine and J. Flanagan

Abstract

Milk proteins can interact with micronutrients via a variety of mechanisms, with hydrophobic interactions being of particular importance. This chapter focuses on the interaction of individual milk proteins as well as mixtures of milk proteins with a range of micronutrients including vitamins, fatty acids, sugars and minerals. Thus, milk proteins can be used as a carrier of micronutrients and thereby increase the nutritional benefit of milk and milk-based products.

It is widely known that the processing of milk proteins via heat or high pressure can result in modification to protein structure, resulting in altered interactions between the proteins and the micronutrient. Interestingly, the presence of some micronutrients can retard the denaturation of some milk proteins. The addition of specific micronutrients may be used as a processing tool to prevent denaturation of milk proteins under physical conditions which normally result in denaturation.

Introduction

The existence of a three-dimensional, folded protein structure is dependent on several forces. These include hydrogen bonding, hydrophobic interactions, van der Waals’ forces and electrostatic interactions. Some amino acid residues may exhibit a hydrophobic character, while electrostatic forces are based on interactions between charged residues. Some proteins contain two or more polypeptide chains. The result of interactions between these components is the quaternary structure, which under
normal physiological conditions is known as the “native state”. Thus, the conformation of a protein is extremely dependent upon the presence of amino acids and the variation of residues within the primary structure. Although proteins may be in the native state, interactions, through hydrophobic, electrostatic, van der Waals’ and other forces, are possible through exposed regions on the surface of the protein. It is through these mechanisms that interactions between milk proteins and various micronutrients such as retinol, fatty acids, minerals and surfactants, for example, can occur.

Protein structures can be readily destabilized from their native state by relatively minor changes in the environmental conditions. Variations in pH, temperature and pressure, for example, can all induce structural transitions in proteins. In some cases, the objective of processing is to induce changes in protein structure, e.g. the heating of whey proteins to form gels. In other cases, however, changes in the environmental conditions can elicit changes in protein structure that result in undesirable functional properties, e.g. loss of solubility or biological activity.

In addition to pH-, temperature- and pressure-induced changes in protein structure, the presence of micronutrients can affect how the protein structure reacts to variations in pH, temperature or pressure. By interacting with specific sites within the protein’s three-dimensional structure, micronutrients can render a protein more, or less, susceptible to denaturation.

Interaction between milk proteins and micronutrients

Micronutrients, such as retinol, sugars, vitamins, fatty acids and minerals, among others, may interact with milk proteins through a variety of mechanisms. The main mechanism is through hydrophobic interaction. In this respect, the majority of studies involving interactions between milk proteins and micronutrients focus on globular whey proteins with their hydrophobic cavities and extensive secondary and tertiary structure. Interactions between caseins and micronutrients are mostly based on electrostatic interactions.

Retinol

Most lipocalin molecules such as β-LG have clear biological roles as ligand carriers. Different researchers have used a variety of methods to determine binding constants, thus making comparison between studies difficult. For example, Muresan et al. (2001) compared fluorometry with equilibrium dialysis. The former yielded higher binding affinities than the latter. The pH, the genetic variant and the source of the protein all contribute to the discrepancies in the literature.

Papiz et al. (1986) identified that the structure of β-LG was remarkably similar to the structure of retinol-binding protein (RBP). Nonetheless, no definite biological function has been attributed to β-LG. In spite of this, various ligand-binding sites have been defined (Qin et al., 1998; Wu et al., 1999) and the structural changes
induced in different environments have been determined by X-ray crystallography (Qin et al., 1998).

However, little endogenous retinol is found bound to \( \beta \)-LG when it is first purified and the ligand most closely associated with the protein is palmitate (Pérez et al., 1989). Vitamin A in the unesterified form is readily oxidized by atmospheric oxygen. The acetate and palmitate esters of vitamin A are somewhat more stable towards oxidation than the free alcohol. \( \text{Cis–trans} \) isomerization is directly promoted by light containing wavelengths of less than 500 nm.

In foods, the retinyl esters and carotenoids are dissolved in the fat matrix, where they are protected from the oxidizing action of atmospheric oxygen by vitamin E and other antioxidants (Ball, 1988). Free retinol is a rather unstable compound, especially in an aqueous environment, but its stability is greatly improved when bound to a RBP (Futterman and Heller, 1972). With the addition of the double bond, free all-trans-3-dehydroretinol is much more sensitive to degradation by oxygen than free retinol (Schwieter et al., 1962). When all-trans-3-dehydroretinol is bound to RBP, however, it is found to be very stable.

In 1972, Futterman and Heller, using fluorescence measurements, first reported that bovine \( \beta \)-LG, like RBP, formed water-soluble complexes with retinol. It has been suggested by Fugate and Song (1980) that the \( \beta \)-LG binding site makes the retinol more rigid than in free solution. Using fluorescence titration and circular dichroism (CD), \( \beta \)-LG was found to display two high-affinity binding sites for retinol per protein dimer, each with an association constant of \( 2 \times 10^{-8} \text{ M} \) (Fugate and Song, 1980). Fluorescence studies exclude the possibility that both pockets accommodate retinol simultaneously (Futterman and Heller, 1972; Fugate and Song, 1980; Dufour and Haertle, 1990a).

There has been some debate regarding the binding site of retinol. Papiz et al. (1986) suggested that it bound inside the main hydrophobic binding pocket, whereas Monaco et al. (1987) proposed that a retinol-binding site for \( \beta \)-LG was an external, solvent-accessible hydrophobic cleft located between the three-turn \( \alpha \)-helix that is packed against the outer surface of the \( \beta \)-barrel and the \( \beta \)-barrel itself. Site mutation experiments, F136A and K141M, did not support this idea and suggested that retinol binds to an evolutionary conserved interior cavity rather than the surface pocket (Cho et al., 1994).

Several reports have indicated that Trp19 of \( \beta \)-LG is essential for the binding of retinol (Papiz et al., 1986). Katakura et al. (1994) used site-directed mutagenesis to investigate whether this completely conserved residue would be indispensable for forming the characteristic structure. Substituting Trp19 with tyrosine was shown not to be critical for the binding of retinol but was important for maintaining stability.

Dufour and Haertle (1991) monitored the binding of retinol, retinyl acetate, retinoic acid and \( \beta \)-carotene to native, esterified and alkylated \( \beta \)-LG by quenching of tryptophan fluorescence. The retinoids bound to native or modified \( \beta \)-LG in a 1:1 molar ratio with apparent dissociation constants in the range of \( 10^{-8} \text{ M} \), whereas the molar ratio was 1:2 for \( \beta \)-carotene–protein. Chemical modification of \( \beta \)-LG by methods such as methylation, ethylation (Dufour and Haertle, 1990a, 1990b) or alkylation (Dufour and Haertle, 1991) have been shown to enhance the binding affinity
for retinol, by opening up a second binding site. It may therefore be assumed that the partial change of β-LG secondary structure produced by these treatments does not destroy the structure of the retinol-binding pocket.

Very few studies have been carried out with ligand binding to α-LA, in comparison with the vast range of studies with β-LG. However, there is the potential of ligands binding to α-LA. Puyol et al. (1991) studied the binding of retinol and palmitic acid in a whey protein mixture. From this study, α-LA was shown to bind retinol more strongly than β-LG, but a much lower percentage of palmitic acid bound to α-LA in comparison with β-LG.

Futterman and Heller (1972) showed that, as with β-LG, BSA formed a strong fluorescent water-soluble complex with retinol. They also postulated that, although no detectable retinol is bound to serum albumin in vivo, the possibility exists that this protein could serve as an auxiliary carrier if excess free retinol were introduced into the circulation.

Raica et al. (1959) reported that a liposoluble substance such as retinol can also be bound to casein. Modification of casein micelle from its natural state (e.g. through acidification or rennet treatment of milk) affects the nutritional activity of retinol (Adrian et al., 1984).

Vitamin D

The affinity of β-LG for vitamin D₂ is about 10-fold greater than that for vitamin A and other retinoids (Dufour and Haertle, 1991; Cho et al., 1994; Wang et al., 1997a). Further work on the binding of vitamin D and cholesterol to β-LG has been explored by Wang et al. (1997a) and Kontopidis et al. (2004).

The binding of retinal, vitamin D₂ and retinyl palmitate by β-LG was studied by Wang et al. (1999). Analysis of competitive binding experiments with palmitate indicated that retinal and palmitate did not compete for the same site; however, vitamin D₂ appeared to displace palmitate at higher concentrations. Retinoids and vitamin D₂ were bound more tightly than palmitate.

Recently, Forrest et al. (2005) reported on the interactions of vitamin D₃ with β-LG A under a range of environmental conditions (i.e. pH and ionic strength). At pH 4.6, β-LG A occurs as octamers (Verheul et al., 1999), whereas dimers predominate at pH 6.6 and pH 8.0. Fogolari et al. (2000) demonstrated the importance of pH when binding ligands to β-LG. The results of Forrest et al. (2005) indicated that binding depended greatly on the solution conditions, e.g. at low pH, 2.5 (I = 0.15M), the EF loop (gate) is closed and thus vitamin D₃ was probably weakly bound in the external hydrophobic surface. Upon lowering the ionic strength to 0.08 M, binding increased. It was suggested (Arymard et al., 1996) that lowering the salt concentration allowed more surface binding. A dissociation constant of 0.02–0.29 μM was reported for β-LG A, with apparent mole ratios of vitamin D₃ bound per mole of β-LG A ranging from 0.51 to 2.04 (Forrest et al., 2005).

Two studies have discussed the stability of vitamin D₃ in cheese (Banville et al., 2000; Upreti et al., 2002) and there is one recent study of the binding of vitamin D₃ to β-casein. Forrest et al. (2005) reported on the interactions of vitamin D₃ with
β-casein under a range of environmental conditions. The binding constants of vitamin D$_3$ to β-casein were dependent on pH and ionic strength. In agreement with the study of Lietaer et al. (1991), an increase in binding as a function of ionic strength was apparent at pH 6.6. This was attributed to reduced solubility of the protein and enhanced hydrophobic interactions, creating more surface area for binding (Lietaer et al., 1991). Increased binding was associated with a weaker affinity, compared with lower ionic strength where binding was stronger. Although stronger interactions at low ionic strength were attributed to fewer protein interactions, the authors could not identify a reason for decreased binding at pH 8.

A dissociation constant of 0.06–0.26 μM was reported for β-casein, with apparent mole ratios of vitamin D$_3$ bound per mole of β-casein ranging from 1.16 to 2.05. It was suggested by Forrest et al. (2005) that the rheomorphic nature of β-casein allowed the hydrophobic area to bind strongly with vitamin D$_3$, in the most thermodynamically stable conformation. The hydrophobic interactions were aligned with the perturbation of phenylalanine and the quenching of tryptophan, both of which are located in the hydrophobic core.

**Vitamin C**

Few studies have explored the interactions between vitamin C and milk proteins. Binding of ascorbic acid to BSA was recorded by Tukamoto et al. (1974). Oelrichs et al. (1984) investigated the interactions between ascorbate and BSA. They suggested an intrinsic association constant of 2600 M$^{-1}$ at 20ºC. Dai-Dong et al. (1990) observed an increased stability of ascorbic acid in the presence of β-LG compared with in pure water, but also found that vitamin C was more thermostable when heated in the presence of β-LG.

In contrast to these studies, Puyol et al. (1994) reported the lack of interaction of ascorbic acid with β-LG or indeed any of the other whey proteins. Puyol et al. (1994) suggested that the discrepancy between their work and that of Dai-Dong et al. (1990) may have been related to the methods used. Monitoring the reducing ability of ascorbic acid may not reflect sufficient allowance for the effects of ascorbate losses through autoxidation. Puyol et al. (1994) also suggested that the antioxidant effect of reductive thiols in β-LG and serum albumin may have a protective effect.

**Other vitamins**

Milk also contains an array of vitamin-binding proteins, including vitamin-B$_{12}$-binding protein, folate-binding protein, vitamin-D-binding protein and riboflavin-binding protein. These proteins occur at low concentrations, but may play a significant role in the uptake of vital vitamins from the diet (Anderson and von der Lippe, 1979; Salter and Mowlem, 1983). Folate-binding proteins (FBPs) are specifically involved in the uptake of folate from the intestine. *In-vivo* studies on rats have shown that protein-bound folate is absorbed at a lower rate than free folate, resulting in increased retention time of folate, allowing it to reach its target tissues. FBPs also reduce the availability of folate to bacteria in the gut and hence may have antibacterial properties (Ford, 1974).
Raw bovine milk contains a riboflavin-binding protein (Kanno and Kanehara, 1985) and riboflavin bound to this milk protein has been shown to have similar antioxidant activities to riboflavin bound to egg white riboflavin-binding protein (Toyosaki and Mineshita, 1988). More recently, Nixon et al. (2004) investigated the source of the co-operativity between FBP and folate and their results suggested stoichiometric interactions. This area, including the binding of trace elements, has been reviewed in detail by Vegarud et al. (2000).

**Fatty acids**

Most of the fatty acids present in milk are found as triglycerides, which form the fat globule (Walstra and Jenness, 1984). The presence of β-LG increases the activity of ruminant pharyngeal lipase, which is deemed to be important during the neonatal period because levels of pancreatic lipase and bile salts are low at that age (Hamosh et al., 1981). The ability of β-LG to remove the released fatty acids, which would otherwise inhibit lipase activity (Calvo et al., 1990), is thus of great importance. Under these conditions, it could be possible that β-LG would bind large amounts of fatty acids, thus displacing retinol (Puyol et al., 1991). Thus, Pérez et al. (1992) proposed that ruminant β-LG, because of its activity to bind fatty acids, might play a role in the activity of pregastric lipases.

Diaz de Villegas et al. (1987) observed that bovine milk has several long-chain fatty acids that bind to β-LG. The predominant fatty acids bound to β-LG were myristic, palmitic and oleic acids, which together accounted for approximately 83% of the fatty acids from total lipids and 70% of the free fatty acids. These fatty acids also predominate in milk lipids.

Pérez et al. (1989) demonstrated that two types of lipids, namely free fatty acids and triglycerides, bound to β-LG. The total amount of fatty acids extracted from β-LG was 0.71 mol per mol of monomer protein. The predominant fatty acids were palmitic (31–35%), oleic (22–23%) and myristic (14–17%) acids, which combined account for 66–75% of the total fatty acids bound to β-LG. The unsaturated fatty acids extracted from β-LG were less than 31% of the total fatty acids and mainly oleic (22–23%) and palmitoleic (4–5%) acids.

As with retinol, there also seems to be controversy regarding the binding location of fatty acids. Narayan and Berliner (1998) suggested that fatty acids bind at the “external site” of β-LG. However, this conflicts with earlier studies by Puyol et al. (1991), which suggested competitive binding, and by Creamer (1995), which suggested an internal location as the primary binding site for fatty acids. Since then, several studies have shown ligands to bind in the internal cavity. Qin et al. (1998), using X-ray crystallography, showed 12-bromododecanoic acid binding inside the calyx and Wu et al. (1999) revealed that palmitate binds in the central cavity (Figure 13.1) in a manner similar to the binding of retinol to the related lipocalin, serum RBP. Ragono et al. (2000) provided further evidence for cavity binding of β-LG and palmitic acid, as did Zsila et al. (2002) using CD, electronic absorption spectroscopy and electrospray ionization mass spectrometry (ESI-MS) with cis-parinaric acid.
Konuma et al. (2007) examined palmitic acid binding to a dimeric β-LG mutant A34C using heteronuclear nuclear magnetic resonance (NMR) spectroscopy. Their results suggested a 1:1 binding stoichiometry. They indicated that the protein conformation should be complementary, at least in part, to the ligand’s structure, if tight binding (dissociation constant of \( \leq 10^{-7} \text{M} \)) is to occur. They further highlighted the role of the highly flexible loops above the barrel in ligand binding, which supports the work of Zidek et al. (1999) and Stone (2001). Konuma et al. (2007) hypothesized...
that the barrel’s entrance accommodates a variety of ligands, because of its plasticity, whereas the bottom of the cavity shows rigid and somewhat selective binding.

Thus, it has been established that β-LG strongly binds one mole of long-chain fatty acids (myristic, palmitic, stearic acid, etc.) per mole of monomeric protein (Spector and Fletcher, 1970; Frapin et al., 1993; Dufour et al., 1994). Frapin et al. (1993) explored the binding of a variety of fatty acids to β-LG and strength of binding was associated with chain length. As β-LG has fatty acids physiologically bound, the amount present probably depends on the isolation technique or whether the protein is delipidated. Thus, this consideration is necessary when determining apparent association constants.

Fatty acid binding to β-LG is sensitive to changes in pH. Changes in binding constants are observed over the pH range 5.5–8.5 (Pérez and Calvo, 1995). This may be due to the electrostatic interactions; for example, as the pH increases, β-LG becomes negatively charged, thus making it less electrostatically inviting for a negatively charged fatty acid. The two lysine residues at the opening of β-LG’s ligand-binding cavity—Lys60 and Lys69—are likely to play a significant role in ligand affinity. The inability of porcine β-LG to bind fatty acids may be due to the substitution of Lys69 by glutamate, as suggested by Frapin et al. (1993) and Pérez et al. (1993). Creamer (1995) also hypothesized that lysine was involved in the binding process, whereby, at neutral pH, the carboxylate group of the fatty acid salt bridged to the positively charged ε-amino group.

Puyol et al. (1991) studied the competition between the binding of retinol and free fatty acids to β-LG. They observed that, when the ratio between the concentrations of the total fatty acids (as palmitic acid) and retinol is similar to that found in milk, the fatty acids compete with retinol for binding to β-LG. Using intrinsic fluorescence studies, Frapin et al. (1993) and Dufour et al. (1994) suggested that an external, independent fatty-acid-binding site on the β-LG−retinol complex was in the groove between the α-helix and the β-sheets of the protein. Narayan and Berliner (1997) supported simultaneous binding of retinoids and fatty acids to β-LG. However, binding is more difficult to determine when several ligands are present.

The organic-anion-binding sites of albumin are composed of two parts: a pocket lined with non-polar amino acid chains and a cationic group located at or near the surface of the pocket (Swaney and Klotz, 1970). Most of the information available on the mechanism of binding has been obtained using organic dyes, anionic detergents and fluorinated or spin-labeled derivatives. Free fatty acid binding involves hydrophobic interactions with the hydrocarbon chain and electrostatic interactions with the carboxylate anion of BSA (Spector et al., 1969).

Andersson et al. (1971) suggested that the fatty-acid-binding sites are located in clefts between the globular regions of the albumin polypeptide. One tryptophan is located deep inside the globular structure whereas the other is superficially located fairly accessible to solvent. Several of the strong fatty-acid-binding sites are located within 10 Å of the buried tryptophan residue (Spector, 1975). Spector et al. (1969) reported that palmitate and palmitoleate were bound more tightly than oleate, linoleate, stearate or myristate and much more tightly than laurate. When a long-chain hydrocarbon did not contain a free carboxyl group (methyl palmitate, cetyl alcohol and hexane), they were bound to a limited extent.
The amount of fatty acids found bound to albumin was 4.8 mol per mol and the predominant acids were oleic, palmitic and stearic acids (Pérez et al., 1989). They suggested that the amount of fatty acids bound to milk albumin could be attributed to the equilibrium between the albumin that carries the fatty acids from lipolysis and the albumin that has been delivered to the tissues. However, milk albumin is not subject to this effect and the fatty acids bound are not taken up by the tissues. Although the number of high-affinity binding sites and the values of apparent association constants for fatty acids to β-LG are lower than those for albumin (Anel et al., 1989), the molar concentration of β-LG in milk is much higher than that of albumin, and therefore β-LG is considered to be the main fatty-acid-binding protein in ruminant whey (Pérez et al., 1990).

Recently, Barbana et al. (2006) reported that bovine holo-α-LA neither contains bound fatty acids in vivo nor has the ability to bind them in vitro. Cawthern et al. (1997) observed the lack of binding of stearic acid with bovine holo-α-LA, the fluorescent indicator acrylodated intestinal fatty-acid-binding (ADIFAB) protein. However, these results are in contrast to their spin resonance and intrinsic protein fluorescence results (Cawthern et al., 1997), which showed that stearic acid was bound to holo-α-LA with a dissociation constant of 10–100 × 10⁻⁶ M. On the other hand, interactions of apo-α-LA with fatty acids have been reported by Barbana et al. (2006). Bovine apo-α-LA displayed apparent affinity binding constants of 4.6 × 10⁶ and 5.4 × 10⁵ M⁻¹ for oleic acid and palmitic acid respectively using partition equilibrium and fluorescence spectroscopy showed a binding constant of 3.3 × 10⁶ M⁻¹ for oleic acid. The small fluorescence changes observed for palmitic acid made it difficult to obtain a binding constant.

**Sugars and polyols**

The effect of sugars on the unfolding and denaturation of proteins has often been described by the steric exclusion effect (McClements, 2002; Baier and McClements, 2005). This effect occurs only if solvent and cosolvent molecules have different sizes. As micronutrients are larger than water molecules, there is a region surrounding the protein molecule from which sugars are excluded and this preferential exclusion effect has been verified in a number of studies (Hammou et al., 1998).

As water is able to get into the layer surrounding the protein, a concentration gradient of the sugar molecules between the inner layer and the outer solution arises (Figure 13.2). This is a thermodynamically unfavorable situation because of the free energy that is required to maintain this concentration gradient. Subsequent movement of water molecules from the area surrounding the protein to outer parts leads to a dehydration of the protein molecule. This dehydration can result in tighter folding of the protein molecules. If the transfer free energy of the protein is greater for the native state than for the denatured state, such as in solutions with sugars, the protein is stabilized in its native state. Furthermore, McClements (2002) reported different molecular mechanisms of stabilization. Sugars may be more preferentially excluded from the denatured state than from the native state or they may be more preferentially accumulated by the native state than by the denatured state.
Interaction between milk proteins and Micronutrients

An alternative explanation of the effect of non-interacting species on the unfolding and denaturation of proteins has been put forward by Semenova et al. (2002), who proposed a direct hydrogen bonding between sugars and proteins, which results in additional hydration; however, the exclusion of sugars from the protein domain is not fully explained by this hypothesis.

The research group of Timasheff has been dominant in research into the interactions of proteins and sugars, or cosolvents as they describe them (Timasheff, 1993). Xie and Timasheff (1997) reported on the exclusion of trehalose from the domain of ribonuclease A at low temperatures. However, at 52°C, where ribonuclease is in the unfolded state, stabilization was brought about by preferential binding of the trehalose to the protein. Sugars showed similar binding affinities to the native and unfolded proteins, but had a lower affinity than water for the exposed peptide groups in the denatured protein. The authors concluded that, in the unfolded state, sugars are more preferentially excluded from the protein domain. The same group conducted a lot of earlier research showing the exclusion of water from the domains of a range of globular proteins, in the presence of sucrose (Lee and Timasheff, 1981), lactose and glucose (Arakawa and Timasheff, 1982). In all cases, they argued that the exclusion of sugars from the protein domain made unfolding of the protein less thermodynamically favorable.

The exclusion of non-interacting species from the protein domain was further shown by Lehmann and Zaccai (1984) who observed, with the aid of neutron small-angle scattering, that glycerol was excluded from the exterior of ribonuclease A at room temperature. Ebel et al. (2000), in a hydration study of rabbit muscle aldolase using a variety of sugars, claimed that the hydration parameter increased as the sugar size increased, indicating moderate exclusion volume effects contributions of the sugars. The effect of sugars on the structure of water may also be linked to the ability of the sugars to bind water.

A recent study comparing the effects of trehalose, maltose and sucrose on the structure of water found that trehalose binds to a larger number of water molecules.

Figure 13.2 Protein–cosolvent–solvent interactions as a result of (a) steric interaction or (b) differential interaction effects. In (b), the exclusion of the cosolvent from the region surrounding the protein is clearly shown. Taken from McClements (2002). Critical Reviews in Food Science and Nutrition, Taylor and Francis Publishers (www.informaworld.com).
than do maltose and sucrose, thus affecting the structure of water to a greater extent (Lerbret et al., 2005). In a rare study involving sugars and casein, Mora-Gutierrez and Farrell (2000) also proposed preferential exclusion of sugar molecules from the casein domain, resulting in preferential hydration of the caseins. The ability of sugars to alter the heat- and pressure-induced denaturation of milk proteins is discussed further in this chapter.

**Flavors**

The interaction of milk proteins and volatile flavor has been reviewed in detail by Kühn et al. (2006) and the reader should refer to this recent review for more in-depth discussion of protein–flavor interactions. However, this section covers the area briefly. A number of flavor compounds are known to bind to milk proteins. Despite this wide knowledge, there are large discrepancies in the binding data because of the use of different methodologies, which appears to be a common feature of determining binding constants.

β-LG is known to interact with a variety of flavor compounds including ionones (Dufour and Haertle, 1990b; Jouenne and Crouzet, 2000; Jung and Ebeler, 2003), lactones (Sostmann and Guichard, 1998; Guth and Fritzler, 2004), alkanes (Mohammadzadeh et al., 1967, 1969a, 1969b), aldehydes (van Ruth et al., 2002), esters and ketones (Guichard and Langourieux, 2000; Jouenne and Crouzet, 2000). In contrast to β-LG, very few studies have explored flavor binding to α-LA. The binding of aldehydes and methyl ketones (Franzen and Kinsella, 1974) and 2-nonanone and 2-nonanal (Jasinski and Kilara, 1985) to α-LA has been explored. BSA has been shown to bind alkanes (Mohammadzadeh et al., 1967, 1969a, 1969b) and Damodaran and Kinsella (1980a, 1980b, 1981) studied interactions of 2-nonanone and BSA. Jasinski and Kilara (1985) compared the binding of 2-nonanone and nonanal to BSA.

The binding of flavors to caseins or sodium caseinate has also received some attention, including the binding of diacetyl (Reineccius and Coulter, 1969), vanillin (McNeill and Schmidt, 1993), β-ionone, n-hexanol, ethylhexanoate and isoamyl acetate (Voilley et al., 1991) to sodium caseinate.

The most extensively studied flavor compound is 2-nonanone. Thus the binding strengths of this flavor to the whey proteins can be compared. Although different authors have reported different affinity constants, the trend BSA > β-LG > α-LA occurs. Table 13.1 illustrates the interactions between 2-nonanone and various milk proteins (Kühn et al., 2006).

**Minerals**

The abilities of certain milk proteins, in particular the caseins, to bind calcium are extremely well known. The extent of binding of the caseins is directly related to the number of phosphoserine residues and thus follows the order $\alpha_{s2}^\gamma > \alpha_{s1} > \beta > \kappa$-casein (Rollema, 1992). Increased binding of calcium to the caseins results in reduced negative charges on the casein molecule, resulting in diminished electrostatic repulsion and consequently inducing precipitation.
Table 13.1 Binding data for the interactions between 2-nonanone and milk proteins (25°C): 
n, number of binding sites per monomer; \( K \), intrinsic binding constant; reproduced with the permission of Kühn et al. (2006); copyright 2006 Journal of Food Science, Institute of Food Technologists Wiley-Blackwell Publishing Ltd

<table>
<thead>
<tr>
<th>Method</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Equilibrium dialysis</td>
<td>Jasinski and Kilara (1985)</td>
</tr>
<tr>
<td>Fluorescence spectroscopy</td>
<td>Liu et al. (2005)</td>
</tr>
<tr>
<td>Headspace SPME</td>
<td>Zhu (2003)</td>
</tr>
<tr>
<td>Headspace SPME</td>
<td>Zhu (2003)</td>
</tr>
<tr>
<td>Static headspace analysis</td>
<td>Charles et al. (1996)</td>
</tr>
<tr>
<td>Equilibrium dialysis</td>
<td>Jasinski and Kilara (1985)</td>
</tr>
<tr>
<td>Equilibrium dialysis</td>
<td>Jasinski and Kilara (1985)</td>
</tr>
<tr>
<td>Liquid–liquid partitioning</td>
<td>Damodaran and Kinsella (1980a)</td>
</tr>
<tr>
<td>Equilibrium dialysis</td>
<td>Jasinski and Kilara (1985)</td>
</tr>
<tr>
<td>PFG-NMR spectroscopy</td>
<td>Jung et al. (2002)</td>
</tr>
</tbody>
</table>

*WPC, whey protein concentrate

Caseins with low numbers of phosphoserine residues, such as \( \alpha_{s1} \)-casein B, \( \alpha_{s1} \)-casein C and the \( \alpha_{s2} \)-caseins, are insoluble in \( Ca^{2+} \) concentrations above about 4 mM. However, \( \beta \)-casein is soluble at high concentrations of \( Ca^{2+} \) (0.4 M) at temperatures below 18°C, but \( \beta \)-casein is very insoluble above 18°C, even in the presence of low concentrations of \( Ca^{2+} \) (4 mM). \( \kappa \)-Casein, with only one phosphoserine, binds little calcium and remains soluble in \( Ca^{2+} \) at all concentrations. Although \( \kappa \)-casein does not bind calcium to any great extent, its ability to stabilize \( \alpha_{s1} \)-, \( \alpha_{s2} \)- and \( \beta \)-caseins against precipitation by \( Ca^{2+} \) is well known and plays a large part in the stabilization of the casein micelle. This is discussed in more detail in Chapter 5 in this volume.

Whereas intact casein has been shown to bind zinc and calcium, tryptic hydrolysates of \( \alpha_{s1} \)-, \( \alpha_{s2} \)-, \( \beta \)- and \( \kappa \)-caseins also display mineral-binding properties. Termed caseinophosphopeptides (CPPs), these peptides can bind and solubilize high concentrations of calcium because of their highly polar acidic domain. Consumption of high concentrations of calcium in early life contributes to the development of maximal bone density, which in turn can prevent osteoporosis in later life (FitzGerald and Meisel, 2003). In addition, calcium-binding CPPs can have an anti-cariogenic effect in that they inhibit caries lesions through recalcification of the dental enamel (FitzGerald, 1998). CPPs have also been reported to improve the intestinal absorption of zinc, as studied using an isolated perfused rat intestinal loop system (Peres et al., 1998).

Lactoferrin has the ability to bind iron very strongly. In vivo, the ferric III form of iron is bound to lactoferrin (Anderson et al., 1989). Considerable interest has been expressed in supplementing bovine-milk-based infant formulas with lactoferrin, as
bovine milk contains much lower levels of lactoferrin than human milk and lactoferrin, isolated from human milk, can bind two moles of iron per mole of protein (Bezwoda and Mansoor, 1986). The biological importance of lactoferrin has been reviewed recently by Lönnerdal (2003). Nagasko et al. (1993) reported that lactoferrin can bind iron at sites other than its chelate-binding sites, probably on the surface of the molecule.

Wieczorek et al. (1992) studied the interaction of milk proteins with fluoride ions over a range of pH values. They found that fluoride does not interact with α-, β- or κ-casein. However, interaction between α-LA and fluoride ions was observed at pH 3.9. Wieczorek et al. (1994) subsequently showed that both α-LA and apo-α-LA failed to bind fluoride ions at pH 4.6, but did at pH 3.7. Other studies involving interactions of minerals/ions and milk proteins are listed in Table 13.2.

**Surfactants**

Some interesting non-food-grade ligands, e.g. 1-anilino-8-naphthalensulfonate (ANS), sodium dodecyl sulfate (SDS) and alkysulfonate (AL) ligands have been widely used to study ligand binding to β-LG. These ligands provide useful information regarding the structure of the molecule. SDS is an amphiphilic ligand that binds strongly to a small number of sites on β-LG at low SDS concentration (Ray and Chatterjee, 1967; Jones and Wilkinson, 1976; Lamiot et al., 1994).

Seibles (1969) studied the interaction of dodecyl sulfate with three of the variants of β-LG. Two moles of dodecyl sulfate were bound to all of the genetic variants of β-LG studied. Creamer (1995) demonstrated that SDS had a profound effect on the equilibrium unfolding of bovine β-LG by maintaining β-LG in the native confirmation despite high concentrations of urea.

Busti et al. (1999) examined the interaction of AL with β-LG and demonstrated one binding site per molecule. Their results suggested that the protective action of AL was exerted by the fraction of the AL bound to the monomer. Also, the efficiency of AL stabilizing native β-LG was related to the length of the hydrocarbon tail. Protein–surfactant interactions are greatly influenced by the surfactant’s hydrophilic group and aliphatic chain length (Ananthapadmanabhan, 1993).

Waninge et al. (1998) showed a substantial increase in unfolding temperature of β-LG–SDS complex at a molar ratio of 1:1. In contrast, a decrease in the unfolding temperature was observed with the addition of an anionic surfactant dodecyl trimethyl ammonium chloride (DOTAC) at a similar ratio. Whereas DOTAC was easily removed via dialysis, it was impossible to remove SDS by dialysis.

Lu et al. (2006) showed that the anionic surfactant sodium perfluorooctanoate was a strong denaturant of β-LG. However, the denaturing ability of sodium

<table>
<thead>
<tr>
<th>Milk protein</th>
<th>Mineral</th>
<th>Reference</th>
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<tbody>
<tr>
<td>β-LG A</td>
<td>Chromium</td>
<td>Divsalar et al. (2006a)</td>
</tr>
<tr>
<td>β-LG A and B</td>
<td>Lead</td>
<td>Divsalar et al. (2005)</td>
</tr>
<tr>
<td>Caseins and β-LG</td>
<td>Mercury</td>
<td>Mata et al. (1997)</td>
</tr>
<tr>
<td>α-LA</td>
<td>Copper</td>
<td>Permyakov et al. (1988)</td>
</tr>
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perfluorooctanoate could be tempered with cationic surfactants, such as alkyl trimethyl ammonium bromide. Sodium n-alkyl sulfates (sodium octyl sulfate, decyl sulfate and dodecyl sulfate) were shown to induce non-native α-helical intermediates of thiol-modified β-LG (Chamani, 2006). Maulik et al. (1996) observed the binding of cetyl trimethyl ammonium bromide with β-LG and reported a two-stage interaction by first-order kinetics. Tetradecyl trimethyl ammonium bromide (TTAB) was also shown to interact with α-LA and cause protein unfolding below TTAB concentrations of 2 mM (Housaindokht et al., 2001).

The interactions of sucrose esters with the casein micelle (Fontecha and Swaisgood, 1995) and β-casein (Clark et al., 1992) have also been studied. Creamer (1980) examined the effect of SDS on the β-casein self-association. The results indicated that SDS binds on an external site of β-casein, such that the hydrophobic tail of SDS becomes involved in the casein self-association. This is supported by the lack of displacement of ANS by SDS. It was also postulated that SDS binds to sites in or on the protein such that the amino acid residues involved in the self-association reaction can interact more favorably with one another. At low concentrations, SDS is thought to bind to a limited number of sites. Despite the increase in the negative charge of the protein, the normal monomer–polymer equilibrium moves predominantly to the polymer in solution whereas, at high concentrations, only protein monomers are present.

**Phospholipids**

Most of the work involving interactions of phospholipids and milk proteins has arisen because of interest in molecular assembly to replicate cell walls. Bos and Nylander (1996) concluded from their studies that both electrostatic and hydrophobic interactions are important for the incorporation of β-LG into phospholipid monolayers. Contrasting effects of β-LG on two zwitterionic phospholipids was reported by Lefevre and Subirade (1999). Whereas dipalmitoyl phosphatidylcholine was unaffected by the presence of β-LG, the conformational disorder of milk sphingomyelin was increased due to hydrophobic interactions with β-LG. Lefevre and Subirade (2001) also studied the effect of phospholipids on the structure of β-LG.

Montich and Marsh (1995) showed, using electron spin resonance, that binding of α-LA to dimyristoyl phosphatidylglycerol bilayers at pH 4.0 caused a motional restriction throughout the full length of the lipid acyl chain. The authors concluded that this restriction in motion would be of direct relevance to the insertion and translocation of protein in the molten globule state across lipid membranes. Earlier work had shown interaction between the milk proteins α-LA and β-LG and phospholipids in monolayer or vesicle formation (Cornell and Patterson, 1989; Kim and Kim, 1989; Cornell et al., 1990).

**Other micronutrients**

Some of the milk proteins, most particularly the whey proteins, have been used as model proteins in studies involving a range of other micronutrients. The interaction
of small heat-shock proteins, such as alpha-crystallin, prevents the precipitation of α-LA when in the molten globule state (Lindner et al., 1997). This finding was confirmed by Sreelakshmi and Sharma (2001) who found that the active site of alpha-crystallin by itself can maintain a significantly denatured and unfolded protein in soluble form. Zhang et al. (2005) reported on the chaperone-like activity of β- and α-caseins. β-Casein was found to be able to suppress the thermal and chemical aggregation of insulin, lysozyme and catalase.

The use of milk proteins as chaperones for drugs has also been studied. The interaction of chlorpromazine with β-LG and αs-casein affected the proteins in different ways. Far UV CD studies revealed that chlorpromazine increased the secondary structure of β-LG, whereas the structure of casein became further disordered (Bhattacharyya and Das, 2001). Divsalar et al. (2006b) also reported on the interaction between genetic variants of β-LG and an anti-cancer component.

Effect of processing on milk protein structure

Heat has been used extensively in food processing for centuries and is a widely applied treatment in food production, primarily for the control of microbial populations. Fields of application are pasteurization under mild temperatures and sterilization under higher temperatures. However, heating may also affect texture as well as taste development and may result in flavor and color changes. The latter effects are often described as disadvantages of heat treatment. Changes in the organoleptic properties are generally as a result of structural changes occurring within the constituents of the food, namely the proteins, polysaccharides or fats.

Another technology that is similar in its control of the microbial population of food products is high-pressure treatment. Foods are preserved with minor changes in texture, flavor or color, in contrast to heat processing, and high pressure is usually called cold preservation technology. High pressure is a long-used technique in Japan and has also become increasingly popular worldwide. However, high-pressure treatment may cause some conformational and structural changes to the individual constituents of the food, possibly resulting in altered functional and organoleptic properties.

Heat and high-pressure treatment may cause the denaturation of globular whey proteins such as β-LG; although there may be differences in the mechanisms behind the denaturation process, the general process appears to be similar. These processes have been examined in detail in Chapter 7.

The denaturation of whey proteins during the heat treatment of milk, the interactions of the denatured whey proteins with other milk components and the effect of these reactions on the physical and functional properties of milk products have been extensively reported and reviewed in great detail (Singh and Creamer, 1992; O’Connell and Fox, 2003; Singh and Havea, 2003). Recent studies have shown that heat-induced aggregation and gelation occur along detailed pathways and are influenced by the types of proteins and forces (disulfide bonding and hydrophobic interactions) present (Havea et al., 1998, 2000, 2001; Schokker et al., 1999, 2000; Abbasi and Dickinson, 2002).
There is an increasing number of studies investigating the effect of high pressure on whole milk and individual constituents; they have been reviewed by Huppertz et al. (2002) and Trujillo (2002). Much of the research interest has focused on the effect of pressure treatment on the physical and functional properties of milk products; however, some studies have dealt with the changes to the individual proteins during the denaturation process. There have been numerous articles citing the mechanisms of heat and pressure denaturation of milk proteins and interested readers are referred to the recent reviews of Huppertz et al. (2006) and Considine et al. (2007a), which cite most of these articles.

**Protein denaturation by thermal and pressure treatments and effect of micronutrients**

As a consequence of their lack of defined secondary and tertiary structure, the caseins have not been suitable candidates for observing changes in protein denaturation. In contrast, the whey proteins have been studied widely as a model globular protein because of their well-defined secondary and tertiary structures, as outlined above.

Interactions between whey proteins and other species induced by either heat treatment or pressure treatment may be divided into two separate classes: covalent interactions and non-covalent interactions. The most important covalent interaction involving whey proteins upon storage is their reaction with reducing sugars via the Maillard reaction to form discolored protein powders, which also have reduced solubilities and nutritional properties. Non-covalent interactions can also occur; these, too, may lead to a loss of protein solubility after association of the proteins with polysaccharides and these non-covalent interactions are primarily driven by reversible electrostatic interactions.

In the present work, the effects of non-interacting species on the unfolding and structural transitions of whey proteins are of specific interest. The marked increase in the thermal and conformational stability of globular proteins in aqueous media in the presence of sugars is well known and has been extensively studied.

**Processing treatments involving ligands**

Whereas the majority of studies have explored ligand binding after heating or pressure treatment of β-LG, only a few studies have looked at the effect of various ligands during the processing treatment (Dufour et al., 1994; Stapelfeldt and Skibsted, 1999; Considine et al., 2005a, 2005b, 2007b). The studies of Considine et al. (2005a, 2005b, 2007b) showed that ligands can retard the heat or pressure denaturation of β-LG, with the type of ligand having an impact on this process. For example, during the heat denaturation of β-LG, both SDS and palmitate stabilized the native structure of β-LG against heat-induced structural flexibility, subsequent unfolding and denaturation up to approximately 70°C, whereas both retinol and ANS provided very little stabilization. When a similar range of ligands was used during pressure denaturation, a similar effect was noted, i.e. higher pressures were required to cause unfolding of β-LG when a ligand was present (Figure 13.3).
It was noted in these studies, and in the comparison study of heat and pressure using myristate and conjugated linoleic acid as ligands, that $\beta$-LG unfolds slightly differently with respect to the type of treatment (Figure 13.3). This mechanism is discussed in detail elsewhere (Considine et al., 2005a, 2005b, 2007b).

Celej et al. (2005) compared the effects of the binding of two ANS derivatives, namely 1,8-ANS and 6-anilinonaphthalene-2-sulfonic acid (2,6-ANS), on BSA thermostability. 1,8-ANS had a stronger effect on BSA thermal stability and they also indicated that the binding parameters are quite different and suggested that stereochemistry is also an important factor in determining protein–ligand interactions. Thus electrostatic interactions should also be considered along with hydrophobic interactions. The authors emphasized the importance of free ligand concentration rather than ligand to protein mole ratio when determining protein stability.

As discussed earlier, the binding of retinol to casein is through hydrophobic interactions (Poiffait and Adrian, 1991). $\beta$-Casein is the most hydrophobic casein and has a highly charged N-terminal domain, containing an anionic phosphoserine cluster, that is clearly distinct from a very hydrophobic C-terminal domain (Swaisgood, 2003). There has been little work on the ability of the caseins to bind retinol. However, Poiffait and Adrian (1991) reported that casein plays an important role in stabilizing retinol over time or during heat treatment. However, information in this area is limited.

**Processing treatments involving sugars**

In an extensive study, Garrett et al. (1988) examined the thermal denaturation of the individual whey proteins $\beta$-LG and $\alpha$-LA and unfractionated whey protein in the presence and absence of sucrose using UV absorption and light scattering methods. The authors found that, in the presence of sucrose, there was increased exposure of the tryptophan residues of both $\beta$-LG and whey protein after heating to 90°C. It must be noted that the whey protein results were based on soluble protein only—insoluble protein that formed after heating had been removed centrifugally. The authors postulated that the hydrophobic groups in the protein’s interior had paired up with sucrose after the protein had undergone a transitionary conformational change following heat treatment. In this way, aggregation of the protein was inhibited, although the sugars were reported to have promoted conformational changes.
The conformational-promoting properties of sugars as reported by Garrett et al. (1988) were contrary to the conformational-retaining properties of sugars as found by Boye and Alli (2000), who reported on the thermal denaturation (by differential scanning calorimetry [DSC]) of 1:1 mixtures of α-LA and β-LG in the presence of a range of sugars. They found that sugars protected against heat-induced denaturation and the protection offered was in the order galactose = glucose > fructose = lactose > sucrose > absence of sugars for increases in the thermal transition temperature of β-LG. No significant effects of sugar were observed with apo-α-LA. Interestingly, an earlier study by the same authors solely on α-LA found an increase in the thermal transition temperature of both the apo and holo forms of α-LA when either 50% sucrose or 50% glucose was added; this increase was fully reversible in the holo form, but only partly reversible in the apo form (Boye et al., 1997). The thermal transition temperature of β-LG was found to be increased in the presence of sucrose, lactose and glucose at 10–50% (Boye et al., 1996b).

Jou and Harper (1996), using the DSC technique, found an increase in the thermal transition temperature of whey protein concentrates following the addition of sugars, with the protection offered by the sugars in the order maltose > trehalose > sucrose. Lactose was also found to provide some protection against heat-induced denaturation. A similar increased thermal stability effect on the heat denaturation temperature of β-LG was observed for sorbitol (Harwalkar and Ma, 1992).

Dierckx and Huyghebaert (2002) followed the heat-induced gelation of a whey protein isolate solution using DSC and dynamic rheology. They found that, by adding increasing concentrations of sucrose or sorbitol, both the thermal transition temperature of the protein denaturation process and the gelation temperature were increased, with a linear relationship existing between the transition and gelation temperatures. They suggested that, because of the differences in the gelation mechanisms observed at different pH values, sucrose and sorbitol affected protein–protein interactions in gels through enhancement of hydrophobic interactions.

Kulmyrzaev et al. (2000) had conducted an earlier study on the effect of sucrose on the thermal denaturation, gelation and emulsion stabilization of whey protein isolate and, although they also observed increases in the thermal transition temperatures on the addition of increasing concentrations of sucrose and improved gel formation and enhanced emulsification flocculation, they postulated that sucrose played different roles in a pre-denatured (improved heat stability) and a post-denatured (enhanced protein–protein interactions) whey protein solution system.

In a study on the effects of different lactose concentrations (within a naturally occurring range) on the formation of whey protein microparticulates, Spiegel (1999) put forward a two-stage process in the aggregation of whey proteins. Up to approximately 85°C, the aggregation of whey proteins is limited by the slow unfolding of the individual proteins; above 100°C, however, aggregation is the rate-limiting step as the rate of unfolding is high. Lactose (at 500 mM) was also found to increase the temperature of the denaturation of whey protein isolate at pH 9.0 by approximately 3°C. However, the authors realized the effect that the Maillard reaction was having in these systems, a factor that some reports seem to ignore.
Baier and McClements (2001) found that increased concentrations of sucrose (up to 40%) could increase the thermal stability of BSA; gels formed from these systems had a higher gelation temperature and a lower complex shear modulus. Similar effects were found in a more recent study (Baier and McClements, 2003). A further study by the same group (Baier et al., 2004) showed that 40% glycerol increased the temperature of gelation of BSA, but no change in the temperature of denaturation of BSA with increasing concentration of glycerol was detected.

Only some studies have proposed that heat-induced protein unfolding is promoted in the presence of sugars, with the denatured state formed being less susceptible to aggregation than the denatured state formed under normal heating conditions. It appears that this issue has yet to be fully resolved.

Some early DSC work (Dumay et al., 1994) showed that the presence of 5% sucrose was enough to reduce the extent of β-LG unfolding by 22% following high-pressure treatment at 450 MPa for 15 min. In a subsequent study, Dumay et al. (1998) found that adding sucrose to β-LG solutions prior to pressure-induced gelation resulted in gels with decreased pore size and strand thickness. They attributed this to a reduction in the number of protein–protein interactions occurring under the influence of pressure.

Keenan et al. (2001) found that low concentrations of sucrose aided in the pressure-induced gel formation of a range of milk-protein-containing systems, but that gel formation was reduced at higher sucrose concentrations. In another group of studies, the pressure-induced gelation properties of skim milk powder were found to be improved by adding low concentrations of sucrose, glucose or fructose, whereas high (45–50%) sugar concentrations inhibited gel formation (Abbasi and Dickinson, 2001).

Boye et al. (1996a) found that lactose, sucrose and glucose increased the temperature of denaturation of BSA, with 50% glucose having a greater stabilizing effect than 50% sucrose. Wendorf et al. (2004) studied the ability of different proteins (ribonuclease A, BSA and egg white lysozyme) to adsorb to a liquid–solid interface in the presence of a range of sugars. They found that the ability of sugars to reduce protein adsorption followed the trend trisaccharides > disaccharides > 6-carbon polyols > monosaccharides and this was explained by the stabilization of the protein in the native state in solution.

Other studies have also shown the beneficial effects of sugars in protecting against denaturation induced by freeze drying, spray drying and chemicals. At low temperatures, high concentrations of sugars cause a substantial increase in solution viscosities and can thus affect protein denaturation. Tang and Pikal (2005) showed that, by negating the thermal stabilizing effects of sucrose by adding denaturants, the increased stability of β-LG in the freeze drying process could be directly attributed to a viscosity effect. Murray and Liang (1999) explored the addition of sucrose, trehalose, lactose and lactitol to whey protein concentrate solutions prior to spray drying and found that the foaming properties of the spray-dried powders were dramatically decreased when sugars were absent. Trehalose was particularly successful in retaining the original foaming properties of both whey protein concentrate and β-LG, but did not perform as well in spray-dried BSA powders (Murray and Liang, 1999). Trehalose has also been found to be effective in stabilization against chemical denaturants.
Conclusions

The interaction of milk proteins with various micronutrients is primarily governed by the physico-chemical properties of the protein. The whey proteins, with extensive secondary and tertiary structure and significant hydrophobicity (albeit largely shielded in the native form), tend towards hydrophobic interactions with ligands and fatty acids. Steric exclusion effects govern the interaction of sugars and polyols with proteins, thus affecting their denaturing properties in the presence of pressure or heat. Electrostatic interactions drive the association of minerals and proteins.

In the food industry, an increasing emphasis is being placed on foods that will have a physiologically functional benefit, in addition to the nutritional benefit of the food. This is being driven by consumers who are becoming increasingly more health aware and health responsible. The challenge for the food scientist is now to deliver the required physiologically functional activities into the final food product, while retaining product quality and shelf life. Knowledge of the interactions of these micronutrients with milk proteins, a major component in many food products, is necessary to achieve this aim. A relevant example of this is a recent patent by Swaisgood et al. (2001) who described the potential of using β-LG as a protein ingredient for carrying lipophilic nutrients such as a range of vitamins, cholesterol and CLA. The binding constants for linoleic acid, CLA and CLA's methyl ester (CLAME) were determined by fluorescence. The use of these β-LG complexes in foods, especially low-fat foods and low-fat dairy foods, is discussed.

A further use of a β-LG–nutrient complex may be in the personal care/hygiene sector. Topical formulations would contain complexes of β-LG with vitamins A or E or CLA and other essential fatty acids. This carrier could be used in a variety of forms including sprays, emulsions, mousses, liquids, creams, oils, ointments and gels. In addition, protein structure can be tailored, by using processing treatments to induce structural changes that may lead to increased interactions with micronutrients.

References


Model food systems and protein functionality

W. James Harper

Abstract

Fabricated foods generally comprise a mixture of components made up of lipids, proteins, simple and complex carbohydrates, emulsifiers and salts, which are capable of interacting with each other and modifying the final characteristics of the food. Often processing utilized in the manufacture of the food also modifies these interactions. Model food systems were first developed because of the disparity between laboratory functional tests for proteins and the functionality in the food. Harper (1984) stated that “disparity between (laboratory) test results and actual functionality in final food formulations necessitates closer scrutiny by researchers of traditional experimental methods”. This disparity results from alteration of functionality (and structure of the protein) through component interactions as well as changes brought about by heat and shear during processing.

Model food systems today find utility for investigating the functionality of many other food components, including starches, gums and emulsifiers, as well as factors affecting areas of continued interest (lipid oxidation, Maillard reaction, etc.). Therefore, model food systems provide a means of determining how the ingredients and the process alter the characteristics of the final product, as well as evaluating the sensitivity of the characteristics of the food to the different ingredients and processing steps.

Model food systems are based on the formulation and processing of real foods, using laboratory and pilot plant facilities. Generally, ingredients that do not have a main effect on
the final characteristics of the product are eliminated. One potential limitation is the use of processing equipment that does not scale up to commercial equipment.

The utilization of carefully selected statistical designs is essential to unravel the multiple interactions that do occur and to optimize food formulation and processing.

A major limitation of model food systems is that they do not provide any information as to the mechanisms by which the ingredients and the process control the final characteristics of the product. Thus they have application to only the food under investigation. They do have a major role in food product development.

Introduction

The utilization of proteins in food for nutritional and functional purposes goes back many centuries, but understanding of the relationship of structure and function has been given close attention only during the past 30–40 years (Owusu-Apenten, 2004). Numerous studies and many reviews have contributed to gaining an understanding of just how proteins act in a complex food system and how the structure and the function are altered by the other ingredients in the food, its intrinsic properties and its processing. These include: Anfinsen (1972), Kinsella (1982), Nakai (1983), Mulvihill and Fox (1987), Mangino et al. (1994), Zayas (1996), Li Chan (2004), Luyten et al. (2004) and Owusu-Apenten (2004).

There are two broad ways of gaining knowledge of the structure and function of protein systems: (a) study of pure proteins in simple systems and (b) study of commercial proteins in the food systems in which they are used. These are entirely different (Luyten et al., 2004; Owusu-Apenten, 2004) and provide quite different information. Functionality tests can be very useful in obtaining reproducible functional properties, even though such tests cannot be used to predict the final characteristics in a real food system (de Wit, 1984, 1989; Harper, 1984; Owusu-Apenten, 2004). Some differences include the following:

- In pure structure/function studies, pure proteins are generally used and are used at concentrations much lower than those used in food systems (Owusu-Apenten, 2004).
- In food systems, proteins are seldom pure and may actually involve complex mixtures of proteins from a given food source (such as milk proteins, egg proteins, soy proteins, etc.) or proteins from multiple food sources (i.e. meat and soy and milk and gluten) or proteins that have been selectively denatured to provide the desired functionality (Mangino et al., 1994).
- In structure/function studies care is taken to avoid interactions with other components and to avoid modifying the secondary and tertiary structure during the experiments. The proteins are fully hydrated (Kinsella, 1982; Owusu-Apenten, 2004).
- In food systems the proteins are constantly exposed to other ingredients, which can modify the structure and hence function, as well as being modified by processes that often include pH, heat and shear (Lee et al., 1992; Kilara, 1994).
Competition for water can also modify functionality, as can changes in intrinsic properties (Zayas, 1996).

- In structure/function studies outcome is generally measured for a specific and single response (Owusu-Apenten, 2004).
- In food systems the ingredients can influence product functionality at different points in the process or functionality can be expressed in more than one outcome with respect to the characteristics of the food (de Wit, 1984, 1989; Harper, 1984).

There is no question that proteins and other hydrocolloids are important and are required to give the food desirable characteristics. However, our knowledge remains incomplete today, because we still cannot fully predict the characteristics of a formulated food on the basis of our knowledge of the structure and function of pure proteins or hydrocolloids under strictly controlled conditions (Kinsella, 1982; de Wit, 1984, 1989; Harper, 1984; Owusu-Apenten, 2004; Zayas, 1996).


**Protein functionality in foods**

Proteins used in foods include plant proteins (soy, wheat, rice, corn and other plant sources), milk proteins (caseins, caseinates, whey proteins and milk protein concentrates [both caseins and whey proteins]), egg proteins (egg white and egg yolk proteins), wheat proteins, meat proteins and fish proteins. Each type of protein exhibits different functional properties and has application in different types of food products (Inglett and Inglett, 1992; Kinsella, 1982; Lee et al., 1992; Kilara, 1994; Mangino et al., 1994; Owusu-Apenten, 2004).

Major functionalities of food proteins include solubility, emulsification, gelation and foaming, water binding and heat stability. As shown in Table 14.1, different types of foods have different functional requirements and may require multiple functionalities.

Factors that may modify the protein, during processing, and hence its effect on the product characteristics include heat, shear, salts and other hydrocolloids (de Wit, 1984; Mangino et al., 1987; Yada, 2004).

**Role of interactions in determining food characteristics**

Interactions between ingredients and modifications caused by processing are the primary reasons why the functionality of proteins and other colloids cannot be predicted...

The following diagram provides an overview of the potential interactions that can occur in a food product (adapted from Harper, 1984):

![Diagram of food system interactions]

Essentially, almost everything can modify the functionality of everything else. Salts are somewhat unique in that they do not in themselves affect product characteristics, but can act on proteins, surfactants, polysaccharides and, to some extent, polar lipids to modify the functionality of each in the food system. The nature and the extent of these interactions will be modified by pH, ionic strength, ingredient concentrations and process-induced modifications. Some examples include the following:

- Surfactants and proteins can interact competitively at the surface of an oil to modify the characteristics of the emulsion, such as stability, size distribution and light scattering. The extent to which a given component will dominate the characteristics will depend upon the relative concentrations (Figure 14.1) and the chemical natures of the surfactant and the protein.
- Starch is frequently used to provide texture to food products. However, the viscosity during processing and the final viscosity can be greatly altered by interactions with other components. The data in Figure 14.2 show that interactions that are observed with two-component systems do not always predict the effect of three- and four-ingredient interactions. In addition, “who sees who first” can further modify other interactions and change product characteristics.

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**Table 14.1 Multiple functionality in selected food products**

<table>
<thead>
<tr>
<th>Food type</th>
<th>Multiple functionalities</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beverages</td>
<td>Solubility, Heat stability, pH stability, Color</td>
</tr>
<tr>
<td>Baked goods</td>
<td>Emulsification, Foaming, Gelation</td>
</tr>
<tr>
<td>Dairy analogs</td>
<td>Gelation, Foaming, Emulsification</td>
</tr>
<tr>
<td>Egg substitutes</td>
<td>Foaming, Gelation</td>
</tr>
<tr>
<td>Meat emulsions</td>
<td>Emulsification, Foaming, Gelation, Adhesion/Cohesion</td>
</tr>
<tr>
<td>Soups and sauces</td>
<td>Viscosity, Emulsification, Water adsorption</td>
</tr>
<tr>
<td>Infant formulas</td>
<td>Emulsification, Heat stability</td>
</tr>
<tr>
<td>Toppings</td>
<td>Foaming, Emulsification</td>
</tr>
<tr>
<td>Frozen desserts</td>
<td>Foaming, Gelation, Emulsification</td>
</tr>
</tbody>
</table>

Adapted from Kinsella (1982), de Wit (1984), Kilara (1994) and Owusu-Apenten (2004)
Figure 14.1 Effect of emulsifier (mono- and diglycerides) and protein on the stability of an oil-in-water emulsion (coffee whitener). Adapted from Harper (1984).

Figure 14.2 Effect of components singly and in combination on the peak viscosity of potato starch (starch (S)): S + X = starch + xanthan gum; S + M = starch + mono- and diglycerides; S + F = starch + fructose; S + X + M = starch + xanthan gum + mono- and diglycerides; S + X + F = starch + xanthan gum + fructose; S + M + F = starch + mono- and diglycerides + fructose; S + X + M + F = starch + xanthan gum + mono- and diglycerides + fructose. Adapted from WJ Harper, unpublished.

Processing effects

The functionality of commercial food proteins, and other hydrocolloids, can be modified both during their production and during the processing of the food product itself. An overview of the conversion of a raw protein source to a functional food ingredient and the subsequent further processing during food manufacture is outlined in Figure 14.3.

During the production of commercial food proteins for use as food ingredients, the proteins may be exposed to a wide range of processing steps that can include thermal processes (pasteurization, sterilization), shear (pumping, mixing, homogenization), pressure (high pressure processing, retorting), concentration (membrane processing, evaporation, drying) and precipitation (heat, acid, salts, solvents). Each of these steps will modify the functional properties of the protein and thus will affect the final characteristics of the food (Kinsella, 1982; de Wit, 1984, 1989; Harper, 1984; Dybing and Smith, 1991; Kilara, 1994; Zayas, 1996; Owusu-Apenten, 2004). Such processes can alter functionality in food through a number of different modifications of the
protein, including changes in sulphydryl interactions, modification of secondary and quaternary structure and shifts in the hydrophilic/lipophilic balance (Kinsella, 1982).

Subsequent processing during use of the protein as a functional ingredient in food will bring further changes in the system, especially those occurring in the presence of other interacting ingredients. Such changes in the characteristics of the food generally cannot be predicted; thus, there is a need for the use of model food systems as an intermediate step in product development (Owusu-Apenten, 2004).

**Uses of model food systems**

Model food systems can be used in a variety of ways (de Wit, 1984; Harper, 1984; Owusu-Apenten, 2004), including the following:

- determining the relative significance of the main effects of ingredients;
- studying factors in food that affect chemical and physical changes (Maillard reaction, lipid oxidation, etc.);
evaluating the sensitivity of the food to alterations in formulation and processing;

- defining ingredient interactions;
- optimizing the formulation for robustness;
- determining critical steps in the processing of the product;
- determining interrelationships between ingredients and the process;
- as a means of tailor-making ingredients for a specific food application;
- evaluating and minimizing the sensitivity of product attributes to the formulation and the process.

Owusu-Apenten (2004) stated that the advantages of the use of model food systems over standard functionality tests included: (a) their ease of use, (b) the lack of a need for specialized equipment and methodologies, (c) the ability to aid in product optimization, and (d) the ability to test for multiple factors and interactions with respect to formulation and processing.

**Initial steps to developing model food systems**

The approach to the development of a model food system will be the same, whether the ingredient being investigated is a protein, lipid, emulsifier, starch or gum.

The development of a model food system begins by reviewing as many formulations as can be found and selecting those that are common to all formulations at a concentration that is at the central point of the various formulas (Harper, 1984). Next, a small-scale process for making the products is developed using processing steps and conditions as close to the commercial process as possible. When more than four or five ingredients are involved, it is often necessary to do a screening experiment to eliminate ingredients that do not have a main effect on important characteristics.

Each different food will have different characteristics, which may include taste, color and texture, that can be modified by the formulation and the process. Key attributes and methods for their evaluation need to be selected. Generally, the methods for evaluation are different from those that are used in research (Owusu-Apenten, 2004).

**Statistical design**

Statistical design is an essential component in the use of model food systems because of its ability to provide information of ingredient and processing interactions (Dziezak, 1990; Earle et al., 2001; Hanrahan and Lu, 2006). Most fabricated food products have from 5 to 25 variables when both the ingredients and the processing steps are taken into consideration. This makes full factorial designs, which would exceed several hundred experiments, an impractical choice. Thus fractional factorial screening designs are generally required.

For most food products, the experimental design is a step-wise process, starting with screening experiments to minimize the variables that do not have main effects on the characteristics of the products. One of the most common screening designs is the Plackett-Burman, which can be used with up to 36 variables (Mullen and Ennis, 1985; Hanrahan and Lu, 2006). The screening experiments allow determination of the main effects that can be used in further fractional factorial designs to get a better understanding.
of ingredient and process interactions and to generate response surfaces that give an understanding of the nature of the interactions (Hanrahan and Lu, 2006).

In developing a fractional factorial experimental design in model food systems, it is necessary to know: (a) the critical factors associated with the ingredients and the process, (b) the region of interest where the factor levels influencing the product characteristics are known, (c) that the factors vary continuously throughout the experimental range tested, (d) that a mathematical function relates the variable factors to the measured response, and (e) that the response defined by the function is a smooth curve.

Numerous studies have used statistical design and response surface methodology to determine the effect of interactions on product characteristics and to optimize specific characteristics in a food (Dziezak, 1990).

In developing an experimental design, consultation with a statistician familiar with the factors that affect the outcomes of the specific design is needed to avoid common pitfalls, which include: (a) critical factors may not be correctly defined or specified, (b) the range of factors selected is too narrow or too broad so that the optimum cannot be defined, (c) lack of the use of good statistical practices, (d) too large a variation in the range of the factors utilized, introducing bias and error, (e) over-reliance on computer-generated results and (f) a need to make sure that the results make good sense.

Applications of model food systems

Initially, model food systems were applied to milk proteins to gain a better understanding of what was required to get desired characteristics in complex food products that could not be predicted from standard functionality tests. de Wit (1998) stated “Information obtained from functional characterization tests in model systems is more suitable to explain retroactively protein behaviours in complex food systems than to predict functionality”. What has been learned using milk proteins in model food systems has been shown to be equally applicable to other food proteins. In addition to understanding the protein being used, there is a need to know the functionality of other ingredients in the food, the probability of how they will interact and modify the function of the food protein, and the use of statistical design to gain the full potential of the model system approach.

The effect of model food systems to assess their performance in foods has been applied to a large number of different types of foods, as shown in Table 14.2. These include bakery products, dairy products, dairy analogs, meat products, sauces and dressings, fermented foods, wine and infant formulas.

The examples of the model food systems used to illustrate applications in this chapter are primarily from the first generation category. They include bakery products, dairy analogs, meat products, salad dressings and sauces.

Bakery products

Bread represents a system where the methods of evaluation of ingredients have been standardized and covered by AACC-approved methods (AACC methods 10–9, 10–10 and 10–11). Details of the procedures and evaluation techniques have been given
by various investigators (Lindbloom, 1997; Pomerance, et al., 1984; Ranhortra et al., 1992; Fenn et al., 1994; Cauvain and Young, 2006). In general, the substitution or addition of other proteins (milk, whey proteins, etc.) leads to a loss of loaf volume (Harper, et al., 1980; de Wit, 1984). Harper and Zadow (1984) found that heat treatments that prevented loss of loaf volume in bread made with milk powder were ineffective in preventing loss of loaf volume in bread made with whey protein concentrates.

Model food systems have been used widely in cake systems: pound cake (Lee, 1999), Madeira cake (de Wit, 1984), white cake (Harper et al., 1980) and angel food cake (Kissell and Bean, 1978). Of these, angel food cake has received the most attention (Lowe et al., 1969; DeVilbiss et al., 1974; Cunningham, 1976; Regenstein et al., 1978; Johnson and Zabik, 1981a, 1981b; Ball and Winn, 1982; Froning et al., 1987; Froning, 1988; Martinez et al., 1995).

The primary protein evaluated has been egg white, for which the cake height and the cake texture can be related to the individual egg white proteins (Johnson and Zabik, 1981a, 1981b; Ball and Winn, 1982). Attempts to replace egg white with whey proteins have never been completely successful (DeVilbiss et al., 1974; Harper et al., 1980). Arunepanlop et al. (1996) were able to replace 25–50% of the egg white with whey protein and could achieve greater replacement by the addition of xanthan gum. Cake volume is essentially the same as with egg white, but the cakes collapse upon baking. This emphasizes the requirement for both foaming and gelation (Owusu-Apenten, 2004). This is due in part to the lower gelation temperature for foams made with egg white (Pernell et al., 2002) and in part to the shear-induced denaturation of egg white with mixing (DeVilbiss et al., 1974).

Other proteins evaluated for angel food cake include blood plasma protein (Kahn et al., 1979; Raeker and Johnson, 1995) and dried beef plasma (Duxbury, 1988).

The factors to consider in developing a model food system for bakery products are outlined briefly in Table 14.3.

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Table 14.2 Model food systems used to assess functionality in foods

<table>
<thead>
<tr>
<th>First generation model foods</th>
<th>Additional examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cakes (angel food, chocolate, yellow, pound)</td>
<td>Low fat spreads</td>
</tr>
<tr>
<td>Meringues</td>
<td>Beer batters</td>
</tr>
<tr>
<td>Bread</td>
<td>Beef patties</td>
</tr>
<tr>
<td>Coffee whitener</td>
<td>Gravies</td>
</tr>
<tr>
<td>Ham, restructured meats</td>
<td>Meat emulsions</td>
</tr>
<tr>
<td>Infant formula</td>
<td>Cream</td>
</tr>
<tr>
<td>Salad dressing</td>
<td>Milk</td>
</tr>
<tr>
<td>Sausage</td>
<td>Cheese</td>
</tr>
<tr>
<td>Starch pudding</td>
<td>Processed cheese</td>
</tr>
<tr>
<td>Whipped topping</td>
<td>Soups and sauces</td>
</tr>
<tr>
<td>Ice cream</td>
<td>Surimi</td>
</tr>
<tr>
<td></td>
<td>Wine</td>
</tr>
<tr>
<td></td>
<td>Yoghurt</td>
</tr>
</tbody>
</table>

*aAdapted from Harper (1984) and de Wit (1984)*

*bAdapted from Owusu-Apenten (2004).*
Model food systems and protein functionality

Dairy analogs

Dairy analogs include coffee whiteners, whipped toppings and processed cheese products.

Coffee whiteners

Coffee whiteners, first developed in the 1950s, generally are protein-stabilized oil-in-water emulsions with vegetable oil as the dispersed phase. A model system, developed by Harper and Raman (1979) and Harper et al. (1980), utilized caseinate, soy bean oil, carbohydrate, phosphate, emulsifier and a gum (xanthan gum or carrageenan). The role of the ingredients has been reviewed by Knightly (1969) and Patel et al. (1992) and the process has been reviewed by Owusu-Apenten (2004). Factors affecting the properties of coffee whiteners are presented in Table 14.4.

Patented processes include using milk protein retentate (Kosikowski and Jimenez-Florez 1987), reformed casein micelles (McKenna et al., 1992), phosphate-modified milk protein (Melachouris et al., 1994) and soy proteins (Melmychyn, 1973).

Alternative proteins that have been suggested to replace caseinate include milk protein concentrate (Euston and Hirst, 2000), whey protein (Hlavacek et al., 1970; Gruetzmacher and Bradley, 1991; Euston and Hirst, 2000), wheat protein (Golde and Schmidt, 2005; Patil et al., 2006), soy protein (Hlavacek et al., 1970; Golde and Schmidt, 2005), peanut protein (Malundo et al., 1992) and cottonseed protein (Choi et al., 1982).

Coffee whiteners are evaluated to ensure that they do provide an emulsion with a small particle size to maximize whiteness, minimize astringency of the coffee by binding with the coffee tannins, maintain stability in hot coffee under acidic conditions, minimize feathering in the presence of hard water salts and readily disperse in

<table>
<thead>
<tr>
<th>Table 14.3 Factors affecting functionality of protein in bakery products</th>
</tr>
</thead>
<tbody>
<tr>
<td>Product type</td>
</tr>
<tr>
<td>Bread</td>
</tr>
<tr>
<td>Cakes</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 14.4 Factors affecting functionality of protein in coffee whiteners</th>
</tr>
</thead>
<tbody>
<tr>
<td>Product type</td>
</tr>
<tr>
<td>Coffee whiteners</td>
</tr>
</tbody>
</table>

Golde and Schmidt (2005) compared coffee whiteners made from sodium caseinate, soy protein isolate and wheat protein isolate and found that they gave similar whiteness (L*) to the coffee. However, the liquid coffee whiteners made with wheat protein tended to separate upon storage and the whiteners made with soy protein isolate tended to show feathering.

**Whipped toppings**

Most commercial whipped toppings contain sodium caseinate as the protein of choice (Knightly, 1968). Other proteins used for whipped toppings include whey protein concentrate (Peltonen-Shalaby and Mangino, 1986; Liao and Mangino, 1987) and soy protein isolates (Kolar et al., 1979; Lah et al., 1980; Chow et al., 1988; Abdullah et al., 1993; Shurtleff and Aoyagi, 1994).

Whipped toppings are high-fat, foamed emulsions with about 40% total solids—model food systems generally also contain sugars, gums and small molecular weight emulsifiers (Knightly, 1968; Harper et al., 1980). The model system differs from whipping or foaming tests with respect to both compositions and much lower fat contents (Owusu-Apenten, 2004). Min and Thomas (1977) found that calcium addition to a 15%-fat-containing whipped topping stabilized with sodium caseinate gave improved stability to the system. Peltonen-Shalaby and Mangino (1986) showed that pasteurization also improved the overrun of the topping. Liao and Mangino (1987) used whey proteins to make a model whipped topping and found a positive correlation between exposed hydrophobicity and overrun. Factors affecting the properties of whipped toppings are presented in Table 14.5. Other factors that affect overrun and stability include the hardness of the fat, the type and percentage of emulsifier and the equipment used for mixing (Harper, 1984).

<table>
<thead>
<tr>
<th>Table 14.5 Factors affecting functionality of protein in whipped toppings</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Product type</strong></td>
</tr>
<tr>
<td>Whipped toppings</td>
</tr>
</tbody>
</table>

**Salad dressings**

Salad dressings are high-fat emulsions, frequently stabilized by high shear in the presence of egg yolk as the primary emulsifier (Parker et al., 1995). Mayonnaise—a spoonable dressing—contains 75% oil by definition. Subsequently, starch pastes were used to make a spoonable dressing with about 40% oil. Today, the most common dressings are pourable, with a wide range of oil contents and are stabilized primarily by xanthan gum (Franco et al., 1995).
Model food systems have been used to gain better understanding of both ingredients (Smith, 1977; Paredes et al., 1988) and processing (Parker et al., 1995).

Smith (1977), using a central composite statistical design, found that the coefficients of the regression analysis were larger for the interaction terms than for the main effect terms in pourable salad dressing with 40% oil and containing egg, vinegar, xanthan gum and mustard powder. The order of addition was also found to be important to the viscous properties of the pourable salad dressing.

**Meat products**

Model meat products, including beef, pork, lamb, poultry or fish, have been utilized for recombined meats (ham, steaks, etc.) and meat macro-emulsions that include bologna, sausages, liver sausages, frankfurters and meat loaves.

Non-meat proteins have been injected into beef and ham, together with water, followed by tumbling to maintain nutritionally equivalent protein levels, increase yield and improve texture (Zayas, 1996; Yada, 2004; Szerman et al., 2007). Szerman et al. (2007) found whey protein isolates to be superior to vegetable proteins on the basis of flavor.

Meat emulsions generally have size distributions between 0.1 and 50 μm and many investigators suggest that they are three-dimensional gel networks with entrapped fat (Regenstein, 1978; Krisman and Sharma, 1990; Xiong et al., 1992; Correia and Mittal, 1993; Barbut, 1995). However, most reviewers continue to classify them as meat emulsions (Gordon, 1969; Webb, 1974; Owusu-Apenten, 2004).

The factors that affect the functionality of proteins in these products include: meat extraction temperature, emulsification temperature, shear during emulsification, fat melting point, pH, ionic strength, ratios of ingredients, salt, soluble protein concentration and type of salt (salt, phosphates, citrates, etc.).

Achievement of functionality has been determined by a number of different methods, including:

- emulsification capacity (EC) (Swift et al., 1961; Swift and Sulzbacher, 1963);
- emulsion activity (EA) (Acton and Saffle, 1972);

Although the tests for EC and ES for comminuted meat products are widely used, there does not appear to be much collaborative testing of the different methods (Owusu-Apenten, 2004).

The type of protein affects the EC of meat emulsions, with isolated muscle proteins giving different EC values. In general, the EC was in the order of myosin > actomyosin > actin for different types of meat (Tsai et al., 1972; Galluzzo and Regenstein, 1978; Li Chan et al., 1984).

Substitution of meat protein by other protein in meat emulsions, as measured by large deformation rheological testing, showed:

- gluten, soy protein isolate or egg white increased the yield after the cooking of meat emulsions (Randall et al., 1976);
- corn germ protein at 2% substitution reduced the shear force and reduced cooking losses (Mittal and Usborne, 1985);
- partial substitution of meat with sodium caseinate, soy protein isolate, whey protein concentrate or wheat germ protein all increased cook yield, increased protein level and decreased fat in frankfurters, without affecting quality (Atughonu et al., 1988);
- addition of bovine blood plasma to meat emulsion products improved emulsion stability and yield, and contents of protein, phenylalanine and valine (Marquez et al., 1997).

Use of model food systems for other food components

In addition to evaluating the performance of proteins in food systems, a wide range of other applications has been utilized. During the past several years, more than 200 papers have been published on other uses of model food systems. A full review of such uses is outside the scope of this chapter. However, selected applications from studies over the past several years are cited both to provide a basis for understanding the scope of the use of model food systems in the food industry and to provide a starting point for obtaining more detailed information.

Applications include:
- factors affecting flavor release in foods (Bylaite et al., 2005; Heineman et al., 2005; Conde-Petit et al., 2006; Nongonierma et al., 2006; Seuvre et al., 2007);
- factors affecting D values in food (Rodriguez et al., 2006);
- lipid oxidation (Jaswir et al., 2004; Sakanaka and Tachibana, 2007; Wijeratne et al., 2006);
- water migration in foods (Guignon et al., 2005; Doona and Moo, 2007);
- Maillard reaction investigations (Severini et al., 2003; Song and Roos, 2006; Acevedo, 2006; Casal, 2006);
- effects of high pressure processing of food (Severini et al., 2003; Sila et al., 2007).

Limitations of model food systems

Model food systems can tell you “what”, but cannot tell you the mechanism(s) by which the effects occur. Frequently, the results with a model system cannot be scaled up to full commercial practice because of differences in equipment and processes. However, they do provide insight into directions to take to overcome scale-up problems. Generally, the results are valid only within the parameters that have been established. Optimization of a food system can sometimes be outside the limits of either the processing equipment or the functionality of a specific ingredient.

Conclusions

Historically, model food systems were used first to improve the functionality of milk proteins in food systems. Currently, there are very few publications on the use of
milk proteins for these purposes, although it is known that a number of dairy food companies use model food systems in their product development programs. Today, the publications concerning model food systems have a much broader usage, with attention being given to a better understanding of how complex food systems affect such things as oxidation, Maillard reactions, shelf life, etc.

Model food systems can be a valuable tool in product development with respect to developments of both formulations and manufacturing processes and have a role in the development of ingredients for new foods.

Model food systems do not provide information on why interactions occur, but can provide insights as to which interactions need basic study to provide a more robust product.

In the future, it can be expected that model food systems will continue to provide a better understanding of how interactions modify the functionality of proteins in complex food systems and continue to give insight on how to use this information to interface with studies on the basis of protein structure/function.

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Sensory properties of dairy proteins

M. A. Drake, R. E. Miracle and J. M. Wright

Abstract

Production and applications for dairy proteins are increasing globally. The dairy protein category is a wide one, encompassing milk proteins to whey proteins and many subcategories of these products such as caseins, hydrolysates and serum or native whey proteins. While functionality and nutrition continue to be key aspects of these products, flavor is a critical parameter that should not be overlooked. An array of sensory analysis techniques can be applied to measure flavor intensities and flavor variability and to determine flavor sources when applied in conjunction with analytical chemistry. This chapter addresses the current status and ongoing research on the sensory properties of dairy proteins.

Introduction

Dairy proteins are valuable dried ingredients with a host of functional and nutritional properties (Foegeding et al., 2002; Miller, 2005; O’Connell and Flynn, 2007). Within the category of dairy proteins, there are a variety of ingredients including whey proteins and milk proteins of various protein contents. Dried caseins and caseinates as well as serum proteins (“native” whey proteins or whey proteins separated from milk prior to the cheesemaking process) are also contained within this category. Dairy
Sensory properties of dairy proteins

proteins (primarily dried) are used in an increasingly wide array of ingredient applications for functionality but, with the current consumer focus on health and nutrition, these ingredients are also used widely to enhance nutrition.

Milk proteins play a crucial role in the flavor of all dairy foods. As part of the sensory experience, proteins provide mouthfeel, viscosity and structure to dairy foods. Amino acids and peptides can elicit basic tastes but can also serve as the starting substrates for numerous volatile aroma-active compounds. Proteolysis and the subsequently released amino acids and peptides are the sources as well as the substrates for many desirable and undesirable flavors in cheeses and other fermented dairy products (Singh et al., 2003, 2005; Carunchia Whetstine et al., 2005a; Drake et al., 2007). Heat processing influences the flavor potential of proteins via denaturation and the release of sulfurous compounds and the typical eggy aroma of scalded milk. Denaturation can also make proteins more susceptible to breakdown and thus influences flavor and flavor development in this manner as well. Theoretically, pure undegraded protein should be flavorless. However, dairy proteins as food ingredients are not 100% protein. Fat, ash, carbohydrate and other components are present in various amounts and also clearly influence the final flavor and flavor stability of dairy proteins.

As with all foods, flavor plays a large role in acceptance and product success. Dried ingredients certainly affect the quality of the final product (Caudle et al., 2005) and the sensory properties of these valuable ingredients should not be overlooked. Dried dairy proteins should ideally be bland or mild and dairy-like in flavor. Recent research has demonstrated that dairy proteins are not flavorless and display a wide array of flavor variability. Understanding and documenting the flavor of these proteins is the key to strategic research and marketing. This chapter addresses and reviews current research on the sensory properties of dairy proteins.

Sensory analysis

Sensory analysis is a scientific discipline that encompasses the depth and breadth of all properties of a food that are perceived by the human senses (Drake, 2007). As such, sensory properties are crucial for product success. Dairy foods continue to enjoy a positive flavor perception by the consumer (Drake and Gerard, 2003; Russell et al., 2006) and this competitive edge should ideally be maximized. This means that a complete understanding of flavor, flavor variability, sources of flavors and consumer perception is mandatory.

A wide array of sensory tests is available to objectively or subjectively measure the sensory properties of foods. These tests and their specific application to dairy products are covered in several recent textbooks and review articles (Lawless and Heymann, 1999; Singh et al., 2003; Drake, 2004, 2007; Meilgaard et al., 2007). Two basic categories exist: analytical tests and affective tests. Several types of tests exist within each category and selection of the specific and appropriate test is dependent on the specific objective in mind. Analytical sensory tests are a group of sensory tests that are objective in nature and use trained or screened panelists. Some examples include descriptive analysis, discrimination tests and threshold tests. Affective tests
are subjective tests and comprise tests that use consumers in qualitative or quantitative measurements.

**Whey proteins**

Whey proteins are recovered from membrane processing and concentration of the liquid whey stream resulting from cheesemaking. Thus, one source of flavor and flavor variability of whey proteins is the flavor of the liquid whey source. The flavor of liquid whey varies, not surprisingly, with the type of cheese (Tomaino *et al*., 2004; Gallardo Escamilla *et al*., 2005). The flavor (sensory perception and volatile components) of fresh liquid whey from thermophilic starters (pasta filata cheeses) will differ from that from mesophilic starter cultures (Cheddar cheese). The flavor of whey from direct acid-set curd will deviate further (Table 15.1). The addition of enzymes such as lipases will increase the free fatty acid content of the whey and this will also influence flavor in the form of rancid, waxy and/or animal flavors depending on the lipase and the milk source.

Within a single type of cheese, the flavor of the whey will vary depending on starter culture rotation and/or other variables in the cheesemaking process. Carunchia Whetstine *et al*., (2003) documented tremendous variability in flavor and volatile compound profiles within and between two Cheddar cheese facilities with starter culture rotation. These results were further confirmed by Karagul-Yuceer *et al*., (2003a). Free fatty acid profiles and proteolysis were also distinct. Tomaino *et al*., (2004) documented that cold storage of liquid pasteurized whey increased lipid oxidation products and resulted in cardboard flavors. They speculated that lactic starter culture enzymes accelerated these storage-induced changes because the concentrations of lipid oxidation products were higher in fermented whey than in whey from direct

### Table 15.1 Flavor profiles of fluid whey obtained from Mozzarella cheese, Cheddar cheese or acid casein manufacture

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Cheddar</th>
<th>Mozzarella</th>
<th>Acid casein</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heated milk</td>
<td>41a</td>
<td>20c</td>
<td>27b</td>
</tr>
<tr>
<td>Caramelized milk</td>
<td>21a</td>
<td>4b</td>
<td>4b</td>
</tr>
<tr>
<td>Natural yoghurt</td>
<td>4c</td>
<td>29a</td>
<td>5c</td>
</tr>
<tr>
<td>Stale</td>
<td>9b</td>
<td>9b</td>
<td>22a</td>
</tr>
<tr>
<td>Rancid</td>
<td>16c</td>
<td>24b</td>
<td>38a</td>
</tr>
<tr>
<td>Oaty</td>
<td>8a</td>
<td>1b</td>
<td>7a</td>
</tr>
<tr>
<td>Dirty</td>
<td>2b</td>
<td>7b</td>
<td>35a</td>
</tr>
<tr>
<td>Acid</td>
<td>5b</td>
<td>39a</td>
<td>36a</td>
</tr>
<tr>
<td>Sweet</td>
<td>37a</td>
<td>8b</td>
<td>12b</td>
</tr>
<tr>
<td>Bitter</td>
<td>2b</td>
<td>14a</td>
<td>7ab</td>
</tr>
<tr>
<td>Salty</td>
<td>6c</td>
<td>12b</td>
<td>18a</td>
</tr>
</tbody>
</table>

*a* Adapted from Gallardo Escamilla *et al*., (2005)

*b* Attributes were scored on a scale from 0 to 100

*c* Means in a row followed by different letters are different (p < 0.05)
acid coagulation. Further, differences in lipid oxidation products were observed in
fresh whey from three different single mesophilic starter strains. Clearly, the initial
raw product stream in whey protein processing displays tremendous flavor, flavor
variability and flavor precursors.

It is not unexpected then that finished dried protein concentrates and isolates also
display flavor variability. Liquid whey is subjected to a host of processing techniques
to concentrate and separate the whey protein. Pasteurization, membrane filtration, con-
centration and spray drying are all steps that can induce the formation of flavor com-
 pounds. Although there is a general process of whey protein production, each facility
is distinct, with facility-specific storage parameters and/or time/temperature profiles
that further contribute variability to the finished product. In the case of whey protein
from colored Cheddar cheese, a bleaching process with hydrogen or benzoyl perox-
ide is also involved. The process of oxidizing the whey stream to decolorize it will
result in a host of possible flavors and flavor precursors. There is currently no recent
published work on the impact of these specific processing steps (other than storage of
liquid whey, mentioned in the previous paragraph) on final whey protein flavor.

Application of defined sensory analysis in combination with instrumental volatile
analysis has shed light recently on the sources of many dairy flavors and will ultimately
aid in the identification of methods to control flavor. This approach recently led to the
identification of a method to enhance the nutty flavor in Cheddar cheese (Avsar et al.,
2004; Carunchia Whetstine et al., 2006) and the specific identification of a cabbage
off-flavor in whey protein isolate (Wright et al., 2006). The reader is referred to three
recent reviews on the application of these techniques to control the flavor in dairy prod-
ucts (Singh et al., 2003; Drake, 2004; Drake et al., 2006). A host of defined flavors in
whey proteins have been documented (Drake et al., 2003; Carunchia Whetstine et al.,
2005b; Drake, 2006; Russell et al., 2006; Wright et al., 2006) (Table 15.2) and vola-
tile compound flavor variability has also been documented (Morr and Ha, 1991; Mills,
1993; Quach et al., 1999; Carunchia Whetstine et al., 2005b; Wright et al., 2006).

The many processing variables listed above undoubtedly contribute to these differ-
ences in flavor among fresh products. Figure 15.1 demonstrates the flavor variability
in fresh (<1 month old) product collected from different manufacturers. Products 4,
5, 6, 8, 9 and 10 were manufactured from Mozzarella or white Cheddar cheese whey
whereas products 1, 2, 3 and 7 were manufactured from colored Cheddar cheese
whey. These differences in flavor and volatile compounds also suggest that there are
some flavors and volatile components that are specifically formed from whey pro-
tein manufacturing bleaching processes and this is certainly an area of research that
should be investigated in ongoing efforts to minimize whey protein flavor. The flavor
intensities of many whey proteins are comparable with those of soy proteins (Russell
et al., 2006) and this is a crucial issue for competitive global marketing.

Storage of whey proteins is another source of flavor variability. The purported shelf
life of whey protein concentrate (WPC80) and whey protein isolate (WPI) varies
from 12 to 24 months depending on the supplier. There is no published work that
demonstrates the flavor stability of these products with storage although recent vola-
tile compound work on WPC80 subjected to accelerated storage conditions has dem-
onstrated key volatile component changes with storage (Javidipour and Qian, 2008).
Furthermore, these products are often agglomerated to enhance their functional properties. The agglomeration process (rewet or single pass) can include addition of lecithin to further increase wettability.

Both of these processes (agglomeration and agglomeration with lecithin) may impact flavor and decrease shelf life. Figure 15.2 demonstrates the flavor changes with storage time of non-agglomerated, agglomerated and instantized (agglomerated with lecithin) WPC80 from a single supplier, as documented by a trained sensory panel. The results suggest that agglomeration, especially agglomeration with lecithin,
Figure 15.1  Principal component biplot of descriptive sensory analysis of WPC80. The WPC80s are represented by numbers.

Figure 15.2  Flavor changes during storage for 12 months at 21°C of non-agglomerated, agglomerated and instantized (agglomerated with lecithin) WPC80s. C, control non-agglomerated product; A, steam-agglomerated product; AL, product agglomerated with lecithin.

affects the storage stability of WPC80, with more rapid development of fatty, cucumber and lipid oxidation types of off-flavors.

Samples of agglomerated and non-agglomerated WPI and WPC80 were collected from suppliers whose products were previously noted to develop a cucumber off-flavor.
The samples were stored at 21°C and were monitored by descriptive sensory analysis (rehydrated to 10% solids w/w) and by instrumental volatile analysis (headspace solid phase microextraction gas chromatography–olfactometry (HS-SPME GC–O) with gas chromatography–mass spectrometry (GC–MS)).

Samples for volatile analysis were prepared as previously described (Wright et al., 2006). Briefly, 20 g of each reconstituted whey protein, a stirring bar and 1 g of NaCl were placed in a 40 mL amber glass SPME vial and sealed air tight with a Teflon™-sided silicon septum (PTFE/silicon) and a plastic cap (Supelco, Bellefonet, Pennsylvania, USA). Samples were heated to 40°C and stirred for 30 min before the SPME fiber (three phase: 2 cm – 50/30 μm DVB/Carboxed™/PDMS Stable Flex, Supelco) was exposed in the headspace at a depth of 3.8 cm for an additional 30 min prior to injection on to the gas chromatograph. The fiber was desorbed at 250°C for 5 min in the injection port fitted with an SPME inlet at a depth of 7.6 cm.

GC–O analysis was performed using an HP 5890 series II gas chromatograph equipped with a flame ionization detector (FID), a sniffing port and a splitless injector. For GC–MS, samples were prepared analogously prior to injection on to the GC–MS system by a CTC Analytics CombiPal Autosampler (Zwingen, Switzerland). The fiber was desorbed at 250°C for 5 min in the injection port fitted with an SPME inlet at a depth of 50 mm. GC–MS analysis was performed using an Agilent 6890N gas chromatograph with a 5973 inert MSD with a DB-5 ms (20 m × 0.25 mm internal diameter × 0.25 μm film thickness) column. Each sample was analyzed in triplicate.

The sensory profiles confirm that agglomerated samples, particularly those with lecithin, are more prone to cucumber off-flavor (Figures 15.3 and 15.4). A number of

![Biplot (axes F1 and F2: 58.72%)](image)

**Figure 15.3** Flavor changes during storage for 18 months at 21°C of non-agglomerated, agglomerated and instantized (agglomerated with lecithin) WPC80s. L, lecithinated; C, control; S, steam agglomerated. Number indicates months of storage.
compounds with cucumber or fatty aromas were recorded by GC–O in samples with and without cucumber flavor (Table 15.3). These compounds were unfortunately at or below MS detection limits by the extraction technique used. The aroma of a compound when it is isolated is not necessarily the aroma or flavor that compound elicits when it is in a food matrix, which means that sensory analysis of the compound in the food matrix is recommended (Drake and Civille, 2003). The character of an aroma can also change with compound concentration (Drake and Civille, 2003).

However, when suspect compounds were placed into WPC80 without cucumber flavor within their reported threshold range and were presented to trained panelists ($n = 8$), many of them elicited cucumber flavors (Table 15.4), suggesting that one or a combination of these compounds is responsible for this off-flavor that develops during the storage of whey proteins. In agreement with the previous example (Figure 15.2), these compounds are also lipid oxidation compounds, again indicating that lipid oxidation is a major source of off-flavor development in these protein products.

Recent work has suggested that native whey proteins might provide a product with the functional and nutritional benefits of whey proteins with superior flavor properties. Native whey proteins are simply whey proteins that are removed from fluid milk prior to the initiation of cheesemaking. In fact, serum or whey proteins can be removed from fluid milk and the cheesemaking procedure can subsequently be initiated as normal with few or no effects on cheese yield. As the

Figure 15.4 Flavor changes during storage for 18 months at 21°C of non-agglomerated, agglomerated and instantized (agglomerated with lecithin) WPIs. L, lecithinated agglomerated; C, control. Number indicates months of storage.
Table 15.3 Aroma-active “green” compounds identified by HS-SPME GC–O from stored agglomerated and non-agglomerated WPC80 and WPI. C refers to non-agglomerated product, S to steam agglomerated product and L to product agglomerated with added lecithin. The number following the treatment letter designation indicates storage time at 21°C in months

<table>
<thead>
<tr>
<th>WPC80</th>
<th>Post Peak Intensity</th>
<th>RI</th>
<th>Method of ID</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound</td>
<td>Odor</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>C8</td>
<td>C10</td>
<td>C12</td>
</tr>
<tr>
<td>Hexanal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E-2-nonenal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E,Z-2,6-nonadienal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Isobutyl-methoxy-pyrazine</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E,Z-2,4-nonadienal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6-Decenal</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>WPI</th>
<th>Post Peak Intensity</th>
<th>RI</th>
<th>Method of ID</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C8</td>
<td>C10</td>
<td>C12</td>
</tr>
<tr>
<td>Methyl-2-butenol</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hexanal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E-2-nonenal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E,Z-2,6-nonadienal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Isobutyl-methoxy-pyrazine</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E,Z-2,4-nonadienal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6-Decenal</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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**Notes:**

- Odor description at the gas chromatograph sniffing port
- Mean post peak intensities as determined by two experienced sniffers at the gas chromatograph sniffing port ([37]van Ruth, 2001)
- Retention index was calculated from GC-O data on a DB-5 column
- Compounds were identified by comparison with authentic standards on the following criteria: retention index (RI) on a DB-5 column, odor property at the gas chromatograph sniffing port and mass spectra in the electron impact mode. Positive identifications indicate that mass spectral data were compared with authentic standards.
- MS = mass spectra
- ND = not detected
native whey proteins have not been subjected to the normal cheesemaking and whey protein processing procedures, their flavor profiles are remarkably bland and nearly free from flavor (Figure 15.5). Significant economic challenges face the industrial scale-up of these products but, at a minimum, research with these products may reveal key information to minimizing the flavor of traditional whey proteins.

Dairy products and, by association, dairy ingredients continue to enjoy a positive flavor image by consumers (Drake and Gerard, 2003; Russell et al., 2006). However, many consumers are unaware that whey proteins are dairy proteins and this is a
sensory issue because consumer perception influences consumer liking. Drake (2006) documented that consumers were generally less sure of their responses when asked to comment on the properties of specific protein types. The US consumer was generally less informed about whey proteins and was more confident and aware of soy protein than New Zealand consumers (Jones et al., 2008). In a follow-up study (Childs et al., 2007), focus groups with US consumers confirmed that most US consumers were unaware that whey proteins were dairy or milk proteins. Consumer education is a current challenge to the dairy protein industry.

One other issue pertinent to whey proteins and consumer acceptance is whether the flavor variability documented by trained panelists is detected by consumers and/or whether it affects the quality of the finished product. Intuitively, the freshest and highest quality ingredients make the best finished product. However, research also indicates that consumers can discern differences in whey protein flavors and that these flavors carry through into ingredient applications (Drake, 2006). Figure 15.6 demonstrates this concept with protein beverages manufactured from different fresh WPC80s. The nature of the off-flavor and the ingredient application will also influence flavor carry-through (Drake, 2006).

Some ingredient applications will be more tolerant than others of variability in the flavor of the ingredients. Childs et al. (2007) recently demonstrated flavor and texture/mouthfeel differences between meal replacement beverages and bars made with whey proteins or soy proteins or mixtures of whey and soy proteins. The ingredient applications were made using standard formulas to allow direct comparison of the influence of the different proteins. Trained panelists documented discernible flavor carry-through of whey and soy proteins in vanilla meal replacement beverages (Figure 15.7). In contrast, no differences in flavor between bars made with whey or soy proteins were noted although several differences in bar texture were impacted by the protein type (Figures. 15.8 and 15.9).
Figure 15.7 Trained panel flavor and mouthfeel profiles of vanilla meal replacement shakes made with whey protein, soy protein or a mixture of whey protein and soy protein. * Indicates significant attributes ($p < 0.05$). Adapted from Childs et al. (2007).

Figure 15.8 Trained panel flavor profiles of peanut-butter-flavored meal replacement bars made with whey protein, soy protein or a mixture of whey protein and soy protein. No attribute differences were noted ($p > 0.05$). Adapted from Childs et al. (2007).
Milk proteins

Milk protein concentrates (MPCs) and isolates (MPIs) represent a newer category of dried dairy ingredients that are rapidly gaining in popularity. These products are manufactured by concentrating milk proteins (whey proteins and caseins) from fluid milk by membrane processing followed by spray drying. Recent work in the primary author’s laboratory has addressed the sensory properties of milk proteins across increasing protein concentration. MPCs with lower protein contents (56, 70% protein dry weight) are characterized by fluid milk types of flavors: cooked/milky, sweet aromatic, sweet taste and cereal (Figure 15.10). As the protein content is increased, the flavor profiles change and MPC77, MPC80 and MPI are characterized by tortilla, brothy, cardboard and animal flavors and higher astringency.

Changes in flavor with increasing protein content were also observed when the sensory properties of lower protein WPCs were compared with those of WPC80 and WPI. Increases in whey protein content likewise resulted in decreases in sweet aromatic and milky flavors. These changes in flavor are probably directly linked to changes in composition and different concentrations of resulting volatile components. A comparison of aroma-active volatile components isolated from WPC80/WPI and whey powder revealed few differences (Mahajan et al., 2004; Carunchia Whetstine et al., 2005b). Differences in flavor are probably due to differences in the relative abundance of specific compounds. Similarly, volatile compound changes are evident in MPCs and MPIs as the protein content is increased. MPCs with higher final protein content have lower sulfur compound response as well as lower aldehyde...
levels when analyzed by HS-SPME techniques (Table 15.5). Changes appear to be due to changes in relative abundance rather than the evolution of new compounds.

**Caseins and hydrolysates**

Caseins represent the primary protein constituent of milk; whey or serum proteins are the other fraction. Just as whey proteins comprise a large group of functional ingredients, so do caseins and caseinates. Caseins are traditionally produced by acid or rennet precipitation of casein followed by spray or roller drying. Caseinate or soluble casein is produced when casein curd (usually acid precipitated) is treated with alkali at pH 6–7 and fully dissolved prior to spray drying (O’Connell and Flynn, 2007). Potassium, sodium and calcium are commonly used counter-ions.

Caseins display a unique set of functional properties including solubility and heat stability and are thus used for a host of ingredient applications. Caseins have relied on functionality for their success because a host of relatively intense and unpleasant flavors, including sulfur, animal, tortilla, musty, cardboard, burnt feathers, glue and bitter taste, have been associated with them (Ramshaw and Dunstone, 1969; Walker and Manning, 1976; Drake et al., 2003; Karagul-Yuceer et al., 2003b) (Figure 15.11). Micellar casein can be manufactured by membrane fractionation and spray drying of fluid milk and may represent a blander option. The net result is a more mildly flavored product that still displays some of the previously reported flavors.

Dairy protein hydrolysates are another promising category of protein-derived ingredients with valuable functional and nutritional properties (Nnanna and Wu, 2007).
Table 15.5 Mean relative concentration (ppb) of selected volatile components extracted from the headspace of rehydrated (10% solids w/w) domestic and international MPCs with various protein contents; all concentrations given as mean relative concentration and (standard deviation)

<table>
<thead>
<tr>
<th>MPC Sample</th>
<th>Protein (%)</th>
<th>Dimethyl sulfide</th>
<th>Propional, 2-methyl-</th>
<th>Furan, 2-methyl-</th>
<th>Butanal, 3-methyl-</th>
<th>Butanoic acid, methyl ester</th>
<th>Hexanal</th>
<th>2-Heptanone</th>
<th>Heptanal</th>
<th>Hexanoic acid, methyl ester</th>
<th>Pentanoic acid, 1-methyl ethyl ester</th>
<th>Benzaldehyde</th>
<th>Furan, 2-pentyl-</th>
<th>Octanal</th>
<th>2-Nonanal</th>
<th>Nonanal</th>
<th>Octanoic acid, methyl ester</th>
<th>Decanal</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 56</td>
<td>0.66 (0.18)</td>
<td>2.33 0.22</td>
<td>0.07</td>
<td>0.22</td>
<td>2.40</td>
<td>0.33</td>
<td>0.44</td>
<td>0.15</td>
<td>0.50</td>
<td>0.32</td>
<td>0.07</td>
<td>0.21</td>
<td>0.07</td>
<td>0.07</td>
<td>0.38</td>
<td>0.71</td>
<td>0.37</td>
<td>0.44</td>
</tr>
<tr>
<td>2 56</td>
<td>0.45 (0.04)</td>
<td>2.73 0.21</td>
<td>0.35</td>
<td>0.43</td>
<td>7.44</td>
<td>0.70</td>
<td>1.72</td>
<td>1.11</td>
<td>0.46</td>
<td>1.49</td>
<td>0.10</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
<td>0.07</td>
<td>0.61</td>
<td>0.71</td>
<td>0.81</td>
</tr>
<tr>
<td>3 56</td>
<td>0.53 (0.00)</td>
<td>0.15 0.69</td>
<td>0.02</td>
<td>0.01</td>
<td>3.12</td>
<td>0.74</td>
<td>0.49</td>
<td>0.06</td>
<td>0.49</td>
<td>0.61</td>
<td>0.10</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
<td>0.02</td>
<td>0.80</td>
<td>0.30</td>
<td>0.81</td>
</tr>
<tr>
<td>4 56</td>
<td>0.26 (0.02)</td>
<td>0.11 0.09</td>
<td>0.01</td>
<td>0.18</td>
<td>3.96</td>
<td>0.59</td>
<td>0.39</td>
<td>0.07</td>
<td>0.53</td>
<td>0.77</td>
<td>0.01</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.80</td>
<td>0.26</td>
<td>0.10</td>
</tr>
<tr>
<td>5 70</td>
<td>0.23 (0.01)</td>
<td>0.11 0.44</td>
<td>0.00</td>
<td>0.02</td>
<td>2.25</td>
<td>0.29</td>
<td>0.40</td>
<td>0.04</td>
<td>0.45</td>
<td>0.95</td>
<td>0.01</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.80</td>
<td>0.17</td>
<td>0.10</td>
</tr>
<tr>
<td>6 70</td>
<td>0.39 (0.05)</td>
<td>0.08 0.16</td>
<td>0.00</td>
<td>0.31</td>
<td>0.27</td>
<td>0.36</td>
<td>0.05</td>
<td>0.12</td>
<td>0.52</td>
<td>0.26</td>
<td>0.01</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.01</td>
<td>0.30</td>
<td>0.18</td>
<td>0.16</td>
</tr>
<tr>
<td>7 70</td>
<td>0.18 (0.02)</td>
<td>0.09 0.42</td>
<td>0.07</td>
<td>0.42</td>
<td>3.27</td>
<td>0.39</td>
<td>0.44</td>
<td>0.30</td>
<td>0.53</td>
<td>1.03</td>
<td>0.07</td>
<td>0.12</td>
<td>0.06</td>
<td>0.02</td>
<td>0.02</td>
<td>0.22</td>
<td>1.37</td>
<td>1.04</td>
</tr>
<tr>
<td>8 70</td>
<td>0.34 (0.06)</td>
<td>0.07 0.43</td>
<td>0.07</td>
<td>0.30</td>
<td>2.34</td>
<td>0.20</td>
<td>0.58</td>
<td>0.15</td>
<td>0.52</td>
<td>0.23</td>
<td>0.02</td>
<td>0.02</td>
<td>0.03</td>
<td>0.02</td>
<td>0.02</td>
<td>0.20</td>
<td>1.28</td>
<td>0.20</td>
</tr>
<tr>
<td>9 70</td>
<td>0.00 (0.03)</td>
<td>0.29 0.07</td>
<td>0.14</td>
<td>0.28</td>
<td>0.94</td>
<td>0.15</td>
<td>0.94</td>
<td>0.03</td>
<td>0.58</td>
<td>1.61</td>
<td>0.00</td>
<td>0.03</td>
<td>0.03</td>
<td>0.03</td>
<td>0.01</td>
<td>0.10</td>
<td>0.80</td>
<td>0.01</td>
</tr>
<tr>
<td>10 70</td>
<td>0.11 (0.02)</td>
<td>0.04 0.30</td>
<td>0.08</td>
<td>0.84</td>
<td>0.82</td>
<td>0.39</td>
<td>0.21</td>
<td>1.21</td>
<td>0.52</td>
<td>0.19</td>
<td>0.02</td>
<td>0.02</td>
<td>0.03</td>
<td>0.02</td>
<td>0.02</td>
<td>0.16</td>
<td>0.17</td>
<td>1.41</td>
</tr>
<tr>
<td>11 80</td>
<td>0.01 (0.02)</td>
<td>0.01 0.24</td>
<td>0.01</td>
<td>0.24</td>
<td>0.21</td>
<td>0.01</td>
<td>0.04</td>
<td>0.02</td>
<td>0.01</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
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<td>0.75</td>
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<tr>
<td>12 80</td>
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<td>0.03</td>
<td>0.19</td>
<td>1.92</td>
<td>0.30</td>
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<td>0.00</td>
</tr>
<tr>
<td>13 80</td>
<td>0.00 (0.00)</td>
<td>0.03 0.14</td>
<td>0.04</td>
<td>0.14</td>
<td>0.49</td>
<td>0.82</td>
<td>0.09</td>
<td>0.19</td>
<td>0.26</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.26</td>
<td>1.69</td>
<td>2.39</td>
</tr>
<tr>
<td>14 80</td>
<td>0.06 (0.00)</td>
<td>0.02 0.69</td>
<td>0.06</td>
<td>0.20</td>
<td>0.47</td>
<td>0.21</td>
<td>0.32</td>
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<td>0.49</td>
<td>0.46</td>
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<td>0.04</td>
<td>0.04</td>
<td>0.32</td>
<td>0.08</td>
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</tr>
</tbody>
</table>

Table 15.5 (Continued) Mean relative abundance of volatile compounds averaged across protein levels; MPC volatile compound mean relative concentration (ppb); all concentrations given as mean relative concentration

<table>
<thead>
<tr>
<th>Protein (%)</th>
<th>Dimethyl sulfide</th>
<th>Propional, 2-methyl-</th>
<th>Furan, 2-methyl-</th>
<th>Butanal, 3-methyl-</th>
<th>Butanoic acid, methyl ester</th>
<th>Hexanal</th>
<th>2-Heptanone</th>
<th>Heptanal</th>
<th>Hexanoic acid, methyl ester</th>
<th>Pentanoic acid, 1-methyl ethyl ester</th>
<th>Benzaldehyde</th>
<th>Furan, 2-pentyl-</th>
<th>Octanal</th>
<th>2-Nonanal</th>
<th>Nonanal</th>
<th>Octanoic acid, methyl ester</th>
<th>Decanal</th>
</tr>
</thead>
<tbody>
<tr>
<td>56 0.48</td>
<td>2.87 0.17</td>
<td>0.18 0.27</td>
<td>0.59 0.35</td>
<td>0.50 0.80</td>
<td>0.98 0.24</td>
<td>0.26</td>
<td>1.16</td>
<td>0.37</td>
<td>0.15</td>
<td>0.05</td>
<td>0.22</td>
<td>0.02</td>
<td>0.02</td>
<td>0.22</td>
<td>0.85</td>
<td>0.54</td>
<td>0.12</td>
</tr>
<tr>
<td>70 0.21</td>
<td>2.93 0.11</td>
<td>0.07 0.38</td>
<td>1.65 0.31</td>
<td>0.52 0.71</td>
<td>0.37 0.14</td>
<td>0.19</td>
<td>0.85</td>
<td>0.54</td>
<td>0.12</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.54</td>
<td>0.12</td>
<td>0.02</td>
</tr>
<tr>
<td>80 0.02</td>
<td>2.04 0.02</td>
<td>0.06 0.17</td>
<td>1.46 0.07</td>
<td>0.41 1.09</td>
<td>0.53 0.08</td>
<td>0.24</td>
<td>0.39</td>
<td>0.47</td>
<td>0.71</td>
<td>0.05</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.47</td>
<td>0.71</td>
<td>0.02</td>
</tr>
</tbody>
</table>
Hydrolysis improves digestibility and hydrolysates are used widely in infant formulas. Recent research has demonstrated that peptides with specific bioactive properties can also be generated and certainly an array of functional properties such as solubility and heat stability can be altered via hydrolysis.

Whey protein hydrolysates are commonly added to meal replacement bars to inhibit bar hardening with storage time (Childs et al., 2007). These products can be manufactured from casein, whey protein or milk protein by enzymatic hydrolysis and are classified based on their degree of protein hydrolysis (molecular weight). Flavor is a significant challenge to increased usage of these products, particularly in beverages. Brothy and free fatty acid flavors and bitter taste are distinct (Figure 15.12) and

![Figure 15.11](image1)  
*Figure 15.11*  Trained panel flavor profiles of rehydrated caseins (10% solids [w/w]). Caseins 1 and 2 are rennet caseins, caseins 3 and 5 are acid caseins and casein 4 is a sodium caseinate.

![Figure 15.12](image2)  
*Figure 15.12*  Trained panel flavor profiles of commercial rehydrated whey protein hydrolysates (5% solids [w/w]). PH1 has a higher degree of hydrolysis than PH2–PH6. PH2–PH6 represent different enzymatic digestions.
intensities can vary with the degree of hydrolysis, the specific processing steps, the enzyme used and the protein source. Because of their intense aromas and flavors, these products should be rehydrated to a lower solids concentration prior to sensory analysis (e.g. 5% w/w compared with 10% w/w for all other dairy proteins).

**Flavor binding**

Although somewhat beyond the scope of this chapter, it is important to note that, in addition to displaying and contributing flavors, dairy proteins can interact and bind with desirable flavors in foods and influence flavor in this manner as well. An excellent review on this subject has been published recently (Kuhn et al., 2006). Most of the research in this arena has been conducted with instrumental analysis (e.g. headspace analysis and calculation of binding constants) and very little research to relate these results directly back to sensory perception has been attempted. Future research should address this issue.

**Conclusions**

Applications for dairy proteins continue to increase and flavor will remain a crucial aspect. An abundance of research on the functional properties of dairy proteins exists, but there is still a relative dearth of information on the flavor of dairy proteins. Flavor sources, flavor formation during processing and flavor carry-through and stability in ingredient applications are key areas for future research.

The flavor of dairy proteins and their flavor performance in ingredient applications will ultimately influence their widespread usage and competitiveness with other protein sources. Published research has only recently begun to reflect and emphasize the importance of this issue. The positive flavor image of dairy foods, combined with the numerous functional and nutritional benefits of dairy proteins, provides a powerful marketing juggernaut for these products, but specific flavor properties and the flavor variability of these proteins should not be overlooked in ongoing research.

**Acknowledgments**

The authors gratefully acknowledge Dairy Management, Inc. and the California Dairy Research Foundation for providing financial support.

**References**


Milk protein gels

John A. Lucey

Abstract

The formation and the properties of the main types of milk protein gels are described, i.e. casein gels made with rennet or acid, heat-induced whey protein gels and gels made by a combination of approaches. The impact of various factors on these gelation properties is discussed. Recent key advances are highlighted, including the use of high pressure and transglutaminase cross-linking of proteins and new insights into the ubiquitous use of thermal processing to alter the texture of these gels.

Introduction

Gelation of the proteins in milk is the basis for the manufacture of cheese and fermented milk products. Various different approaches can be used to destabilize the milk proteins including heat (whey proteins), use of rennet enzyme (caseins) and acidification (caseins and denatured whey proteins). Combinations of these approaches can also be used to form dairy products, e.g. the use of a low concentration of rennet in cottage cheese (or quarg), which is primarily a cultured product. Yoghurt is a cultured product in which caseins and denatured whey proteins are responsible for the gelation properties.

Milk protein gels are irreversible, in contrast to many polysaccharide gels which are thermoreversible. Milk gels are often classified as particle gels although it is now recognized that they are not simple particle gels, as the internal structure of...
the casein particle plays an important role in their rheological properties (Horne, 2001, 2003). The properties of milk protein gels have been reviewed (Green, 1980; de Kruif et al., 1995; Lucey, 2002; van Vliet et al., 2004). The casein particles in rennet gels undergo rearrangement, fusion and syneresis in the process of forming cheese curd; thus they are inherently dynamic in nature and the rearrangement processes involved have been studied (Dejmek and Walstra, 2004).

**Rennet-induced gels**

**Introduction**

Coagulation of milk by rennet probably occurred initially by accident, as warm milk was stored in sacks made from the stomachs of ruminant animals that contained some residual proteinase enzymes. Crude extracts, prepared from the fourth stomach of young calves (called rennets, which are a type of aspartic proteinase), have been used for cheesemaking for thousands of years. Pepsin is the predominant proteinase in adult mammals. Naturally produced calf chymosin (EC 3.4.23.4) may contain up to six molecular species, which have slight differences in their amino acid residues (Crabbe, 2004). Chymosin has been cloned into several genetically modified organisms to produce fermentation-derived chymosin, which is widely used in many countries around the world (Crabbe, 2004). The rennet coagulation of milk has been reviewed (Dalgleish, 1987, 1993; Hyslop, 2003; Horne and Banks, 2004).

**Primary phase of rennet coagulation**

The basic building blocks of rennet-induced gels are the casein micelles. Both $\alpha_s$- and $\beta$-caseins are sensitive to precipitation by the Ca$^{2+}$ in milk and are protected by association with $\kappa$-casein, which is one reason for the formation of micelles. $\kappa$-Casein molecules are thought to have a predominantly surface position on micelles (although some $\kappa$-casein is also present in the interior of the micelle), where the hydrophilic C-terminal apparently acts as a “hairy” layer providing steric stabilization and a barrier against association with other micelles (Walstra, 1990; Chapter 5 in this volume).

The two stages of the rennet coagulation of milk are shown in Figure 16.1. In the primary phase of rennet coagulation, the C-terminal part (residues 106–169) of the $\kappa$-casein molecule is hydrolyzed and this hydrophilic peptide diffuses away from the micelle (called para-casein) into the serum phase. This macropeptide is called caseinomacropeptide (CMP) or, if it is highly glycosylated, glycomacropeptide (GMP). Most microbial coagulants, including those derived from *Rhizomucor miehei*, hydrolyze the same Phe$_{105}$–Met$_{106}$ bond as chymosin; however, *Cryphonectria parasitica* hydrolyses the Ser$_{104}$–Phe$_{105}$ bond (Drohse and Foltmann, 1989). The proteolysis of other proteins in milk by chymosin occurs at a much slower rate (Crabbe, 2004).

The enzymatic reaction in milk appears to obey first-order kinetics. The proteolysis of $\kappa$-casein is usually described by standard Michaelis-Menten kinetics, although Hyslop (2003) questioned whether this was truly appropriate. It should be noted that
the primary phase and the secondary phase of clotting overlap as the aggregation begins before the enzymatic reaction is complete.

**Secondary phase of rennet coagulation**

The stability of the casein micelles of milk is attributed to their net negative charge and to steric repulsion by the flexible macropeptide region of \( \kappa \)-casein (the so-called hairs that extend out into the solution), calcium-induced interactions between protein molecules, hydrogen bonding and electrostatic and hydrophobic interactions. The release of the CMP (or GMP), which diffuses away from the micelles, leads to a decrease in the zeta potential, by \( \approx 5–7 \text{ mV} (\approx 50\%) \), which reduces electrostatic repulsion between rennet-altered micelles. Removal of the “hairs” results in a decrease in the hydrodynamic diameter by \( \approx 5 \text{ nm} \) and a loss of steric stabilization, and causes a slight minimum in the viscosity during the initial lag phase of renneting.

Various attempts have been made to model the aggregation reaction (Horne and Banks, 2004). The nature of the attractive forces during the aggregation of casein micelles is still not completely clear, although calcium bridges, van der Waals’ forces and hydrophobic interactions may be involved. Destabilized micelles will aggregate only in the presence of free \( \text{Ca}^{2+} \). Rennet acts on casein at temperatures as low as 0°C, but milk does not clot at temperatures below 18°C whereas aggregation is very rapid at high temperature (e.g. 55°C).

When milk is clotted under normal conditions of pH and protein content, the viscosity does not increase until the enzymatic phase is mostly complete, i.e. at \( >60\% \) of the (visual) rennet coagulation time. Coagulation does not occur until the enzymatic phase is at least \( \approx 87\% \) complete. Sandra et al. (2007) studied the rennet gelation process using diffusing wave spectroscopy, which allowed gelation to be monitored without the need for dilution. Sandra et al. (2007) suggested that partially renneted casein micelles do not begin to approach one another until the extent of breakdown of the \( \kappa \)-casein hairs has reached about 70%; above this point, they interact increasingly strongly with an increase in the extent of proteolysis. This interaction

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**Figure 16.1** The two stages of the rennet coagulation of milk.
initially restricts the diffusive motion of the particles rather than causing true aggregation. Only after more extensive removal of the protective κ-casein hairs does true aggregation occur, with the appearance of a space-filling gel (as defined by rheology terms, such as having a loss tangent value <1). A micrograph of a rennet-induced skim milk gel is shown in Figure 16.2.

Srinivasan and Lucey (2002) studied the impact of plasmin enzyme on the rennet coagulation of skim milk. They found that hydrolysis of αs- and β-caseins (as plasmin hardly degrades κ-casein) accelerated the rennet coagulation time. Srinivasan and Lucey (2002) hypothesized that plasmin could have degraded non-κ-casein “hairs” present on the surface of micelles and that this could have reduced the repulsive barrier to aggregation of rennet-altered micelles such that aggregation could take place at a lower degree of κ-casein hydrolysis.

Completely hydrolyzed micelles initially form small linear chains and these continue to aggregate to form clumps, clusters and eventually a system-spanning network that has a fractal-like appearance.

Little aggregation occurs at low temperatures (e.g. <15°C), which is usually taken as an indication of the importance of hydrophobic interactions. It is more likely that, with decreasing temperature, the activation free energy for flocculation increases, presumably because of the presence of β-casein chains on the outside of the micelle (Walstra, 1993). There is an increase in the strength of rennet gels at lower temperatures (where hydrophobic interactions are weak), reflecting swelling of casein particles, which results in an increase in the contact area between aggregated particles and strands.

Figure 16.2 A confocal laser scanning micrograph of a rennet gel made from skim milk. Protein is white; dark areas are water. Scale bar = 10μm.
Monitoring gelation

There have been several recent reviews of techniques to monitor milk gelation (Lucey, 2002; O’Callaghan et al., 2002; Klandar et al., 2007). The interest in monitoring gelation comes from the desire by cheesemakers to know when it is the “optimum” time to initiate cutting as well as from the desire by researchers to better understand this complex process.

Two promising techniques for the study of milk gels are diffusing wave spectroscopy and ultrasonic spectroscopy (Alexander and Dalgleish, 2004; Dalgleish et al., 2006). These techniques could be used to complement existing approaches. For example, Wang et al. (2007) used both ultrasonic and (traditional) rheological methods to investigate the effects of milk pretreatment at ultra-high temperatures on the rennet gelation of a whey-protein-free casein solution. Wang et al. (2007) found that the ultrasonic velocity was able to measure the enzymatic hydrolysis and aggregation process, but was not as sensitive in detecting gel formation. In contrast, the oscillatory rheological method was not able to detect the enzymatic hydrolysis reaction, but was very suitable for characterizing the formation of a gel network.

Rheological properties of rennet-induced milk gels

Rennet-induced gels are viscoelastic and their rheological properties can be characterized using dynamic low-amplitude oscillatory rheology, which determines both the viscous component and the elastic component. These measurements should be performed in the linear viscoelastic range, where the deformation (strain) is proportional to the applied stress. Often, for rennet and acid gels, that means trying to operate at \( \leq 3\% \) strain, which can be difficult during the early stage of gel formation for many (of the popular) controlled stress rheometers (because of the very low torque resulting on the measuring geometry of the rheometer from such a weak gel). Some new techniques/software can be used to reduce this problem in commercial rheometers (e.g. Lauger et al., 2002).

Parameters that can be determined include the elastic or storage modulus \((G')\), which is a measure of the energy stored per oscillation cycle, the viscous or loss modulus \((G'')\), which is a measure of the energy dissipated as heat per cycle, and the loss tangent (tan \(\delta\)), which is the ratio of the viscous properties to the elastic properties \((\text{loss tangent} = G''/G')\). The loss tangent is related to the relaxation of bonds in the gel during deformation and is a useful parameter.

During gelation, there is a lag period before a measurable storage modulus value is obtained (this depends on the sensitivity of the rheometer to measure events close to the gelation point). The loss tangent decreases from \( \gg 1 \) to \(<1\) at the gelation point and then attains a relatively constant value (about 0.35 for rennet gels). The dynamic moduli initially increase relatively rapidly and then, after a period of several hours, tend to plateau. In commercial practice, rennet-induced gels are cut once they have attained a certain firmness (usually assessed subjectively by the cheesemaker) or, more commonly, at a fixed time after rennet addition. The increase in the moduli after gelation probably reflects ongoing fusion of micelles, which results in
an increase in the contact area between aggregated particles, and possibly the incorporation of additional particles into the gel network.

Some micelles that have incomplete hydrolysis of their $\kappa$-casein hairs could be trapped within the space-filling network at the point of network formation and they might later become attached to the matrix once their $\kappa$-casein hairs get completely hydrolyzed. Mellema et al. (2002) reported that their analysis suggested that nearly all casein was incorporated in the rennet gel, at least very soon after network formation. Mellema et al. (2002) also considered that changes in the storage modulus and microstructure during aging could be explained in terms of (various types of) rearrangements of the gel network at various length scales.

Typical plateau values for the storage modulus of rennet-induced gels (made from unconcentrated milk) range from 100 to 200 Pa. Both moduli have lower values at low frequencies, reflecting relaxation of more bonds when the timescale of the applied stress is longer. The loss tangent at low frequencies is an important indicator of rearrangements as this is approximately the same timescale as that over which rearrangement processes related to syneresis in rennet gels are estimated to occur (van Vliet et al., 1991).

The development of the complex or shear modulus as a function of time after rennet addition can be replotted against a reduced time $t/t_g$, where $t_g$ is the gelation time. Various individual renneted milk curves can be normalized against their complex or shear modulus value at a low multiple (two or three) of the $t_g$. These various curves then collapse into a single or master curve because of the scaling behavior of the dynamics of the gel formation process (Horne, 1995, 1996). Various mathematical, empirical and kinetic models have been applied to predict the development of gel firmness or shear moduli; their effectiveness in performing this function has been reviewed recently by Horne and Banks (2004).

**Syneresis of rennet-induced milk gels**

The syneresis of rennet-induced gels has been reviewed (Walstra et al., 1985; Pearse and MacKinlay, 1989; Walstra 1993; van Vliet and Walstra, 1994; Dejmek and Walstra, 2004). Rennet-induced gels remain stable for several hours if left undisturbed. They rapidly synerese if disturbed by cutting or by wetting the gel surface. A rennet-induced gel may lose up to two-thirds of its volume (as whey) under quiescent conditions and more than 90% if external pressure is applied (Dejmek and Walstra, 2004).

Cheesemaking can be viewed as a dehydration process and syneresis is the crucial method by which most of the moisture is lost from curd particles. As syneresis is the main method that cheesemakers have to control cheese moisture, it is also the process that is most often manipulated; it also helps to facilitate differentiation between cheese varieties. Most of the water in milk gels is not chemically “bound” to proteins but rather is physically entrapped in the network structure (van Vliet and Walstra, 1994).

Because of the complexity of the syneresis process, researchers have often used thin gel slabs to monitor one-dimensional shrinkage (e.g. van Dijk and Walstra, 1986). The one-dimensional syneresis of rennet gels is related to the flow of liquid
(whey) through the network (because liquid flows out of the gel concomitantly with gel shrinkage) and is governed by the equation of Darcy:

\[ v = \frac{B \cdot p}{\eta \cdot x} \]

where \( v \) is the superficial flow velocity of the syneresing liquid, \( B \) is the permeability coefficient, \( \eta \) is the viscosity of the liquid, \( p \) is the pressure acting on the liquid and \( x \) is the distance over which the liquid must flow.

It is believed (Walstra, 1993) that there is an internal (endogenous) pressure or driving force within rennet gels that is responsible for the shrinkage of the gel once the initial gel is disturbed (presumably this overcomes the yield stress of the system). It has not been possible to measure this small endogenous pressure experimentally. Endogenous syneresis pressure (i.e., the pressure within the rennet gel causing the syneresis) is not constant. It increases initially as a function of time after renneting but decreases at longer times, presumably because of fusion of para-casein micelles and a reduction in permeability of the contracting network. In practice, syneresis in curd particles occurs in three dimensions simultaneously and is much harder to study than the one-dimensional model.

In rennet-induced milk gels, the mechanism responsible for the strong tendency of the gels to exhibit syneresis is related to the (extensive) rearrangements of the casein network that occur after gel formation. As acid-induced gels undergo much less rearrangement, they synerese less. The rearrangement process is accelerated, and is more extensive, at high temperatures and lower pH values (<6.5 but >5.1) (the loss tangent is also higher under these conditions). Aging of rennet-induced gels results in a coarsening (sometimes called microsyneresis) of the gel (i.e., as a result of more rearrangements) and there is an increase in the permeability and the fractal dimensionality.

Rearrangements of casein particles into a more compact structure would increase the number of bonds and hence decrease the total free energy of the system (Walstra, 1993). However, the casein particles are already part of the gel network, which must be deformed or broken locally to form new junctions. Breakage of the bonds in the strands requires a sufficiently low yield stress if it is to be exceeded. In cheesemaking, conditions such as cutting, stirring, acid production and the increase in the cooking temperature all encourage syneresis and the rearrangement processes that facilitate syneresis of the gel network. If the strands become too thick (e.g., because of a very high casein concentration), syneresis hardly occurs.

One-dimensional syneresis of rennet-induced skim milk gels was studied in gels with different thicknesses and at pH values of 6.4 and 6.0 using a laser displacement sensor (Lodaite et al., 2000). Syneresis was much faster at the lower pH and the initial syneresis rate increased linearly with slab thickness.

Several (mostly empirical) techniques have been used to estimate the syneresis of rennet gels, including shrinkage of gel slabs, determining the volume of whey expelled as a function of time, the dry matter content or density of curd particles and low-resolution nuclear magnetic resonance (NMR) (Dejmek and Walstra, 2004). A recent development has been a light backscatter sensor, with a large field of view...
relative to curd size, for continuous on-line monitoring of coagulation and syneresis to help cheesemakers improve their control over the moisture content of the curd (Fagan et al., 2006).

**Some factors influencing the texture of rennet-induced gels**

Many factors influence the milk clotting process and the consistency of rennet gels including pH, temperature, casein content, ionic strength, enzyme concentration, calcium content, presence of homogenized fat globules, concentration of denatured whey proteins and casein hydrolysis by proteinases such as plasmin. These factors have been reviewed many times (Dalgleish, 1987, 1993; Green and Grandison, 1993; Lomholt and Qvist, 1999; Hyslop, 2003; Horne and Banks, 2004) because of the importance of rennet gels for the cheese industry.

The effects of pH (5.19–6.21) and NaCl concentration (0, 1.75 and 3.50%) on the rheological and microstructural properties of rennet-induced casein gels made from ultrafiltered skim milk (19.8%, w/w casein) were recently investigated (Karlsson et al., 2007a). Low pH and high NaCl concentration reduced the rate of development of the gel elasticity after coagulation. Strain at fracture and stress at fracture 48 h after coagulation showed maximum and minimum values at pH 5.8 and 5.29 respectively. The microstructure examined with confocal laser scanning microscopy was unaffected by the changes in pH and the concentrations of NaCl, probably because of the very high volume fraction of caseins in this type of gel (Karlsson et al., 2007a).

Rennet-induced coagulation of ultrafiltered skim milk (19.8%, w/w casein) at pH 5.8 was studied and compared with coagulation of unconcentrated skim milk of the same pH (Karlsson et al., 2007b). At the same rennet concentration, coagulation occurred at a slower rate in ultrafiltered skim milk but started at a lower degree of κ-casein hydrolysis, compared with the unconcentrated skim milk. Confocal laser scanning micrographs revealed that, during storage for up to 60 days (at 13°C), the microstructure and the size of the protein strands of the ultrafiltered gel hardly changed, probably because of the high zero shear viscosity of the concentrated system (Karlsson et al., 2007b).

Plant coagulants obtained from the flowers of Cynara sp. have been used to make rennet gels and cheeses (Esteves et al., 2001, 2002, 2003). These coagulants are less sensitive to changes in gelation temperature, they cause more casein rearrangements during gelation and they have higher values for the storage modulus (at least during the initial stages of gelation), compared with gels made with chymosin (probably because of greater proteolysis of the caseins).

Choi et al. (2007) demonstrated that the concentration of insoluble calcium phosphate (CCP) associated with the casein micelles had an important influence on the properties of rennet gels. Removal of some CCP from milk prior to gelation using calcium-chelating agents lowered the storage modulus of rennet gels because of the reduction in the amount of CCP cross-linking in the casein micelles. Reduction in the CCP content prior to rennet gelation resulted in gels with higher loss tangent values, indicating greater bond mobility.

Choi et al. (2007) also studied the impact of preacidification of milk prior to gelation. They found that gels made at pH 6.4 had higher storage modulus values than gels
made at pH 6.7 probably because of the reduction in electrostatic repulsion, whereas the CCP content only slightly decreased at this pH value. The storage modulus values were highest at pH 6.4 and decreased with decreasing pH from 6.4 to 5.4 because of the reduction in CCP cross-linking within the casein micelles (Choi et al., 2007).

### Milk heat treatment

It is well known that severe heat treatment of milk at temperatures sufficiently high to denature the whey proteins results in an increased rennet coagulation time as well as weaker gels (Lucey, 1995). There are some reports that the interaction of denatured whey proteins with the $\kappa$-casein inhibits the primary phase of rennet action on $\kappa$-casein (to some extent). For example, Reddy and Kinsella (1990) reported that very high heat treatments decreased the initial velocity ($V_i$) and GMP release. However, most studies have concluded that the secondary phase of the coagulation process is the step that is mainly inhibited by the presence of denatured whey proteins on the micelle surface. These denatured whey proteins probably sterically interfere with the (normal) aggregation of rennet-altered micelles (Lucey, 1995).

Vasbinder et al. (2003) concluded that whey protein denaturation had only a small effect on rennet activity and that the release of GMP (or the formation of para-$\kappa$-casein) was similar in heated and unheated milks. Anema et al. (2007) adjusted the pH of the milk prior to heat treatment, which allowed them to manipulate the distribution of denatured whey proteins and $\kappa$-casein between the serum and micellar phases; they reported that the retardation in rennet gelation as a result of heat treatment was observed regardless of whether the denatured whey proteins were associated with the casein micelles or in the serum phase.

### Enzymatic cross-linking of caseins

Transglutaminase (TGase; EC 2.3.2.13) catalyzes covalent intermolecular protein cross-linking through an acyl-transfer reaction, between the $\gamma$-carboxyamide group of a peptide-bound glutamine residue (acyl donor) and the primary amino group of an amine (acyl acceptor). The application of TGase in various types of dairy products has been reviewed (Jaros et al., 2006). In a system where caseins and whey proteins are available as substrates for TGase, such as milk, the caseins are preferentially cross-linked over native whey proteins (Han and Damodaran, 1996).

Lorenzen (2000) incubated preheated milk with TGase for various incubation times prior to rennet addition and found that increasing TGase incubation times, as well as an increasing intensity of preheat treatment of the milk, resulted in increasing coagulation times up to the point of a complete absence of coagulation. Lorenzen (2000) attributed the reduced rennetability of preheated milk to a “surface sealing” of the casein micelles with cross-linked $\beta$-lactoglobulin, leading to a steric inhibition of the release of the macropetide from the surface of the casein micelle.

O’Sullivan et al. (2002b) also attributed the loss of rennetability to the impact of TGase cross-linking on the primary enzymatic phase, i.e. reduced rate of hydrolysis of $\kappa$-casein. Huppertz and de Kruif (2007) criticized the analytical method used by O’Sullivan et al. (2002b) to study the hydrolysis reaction because they suggested that
this method detects only the products of hydrolysis of non-cross-linked milk; hydrolysis products of cross-linked κ-casein would not be adequately detected because the macropeptide remains attached to the micelle. Huppertz and de Kruijf (2007) suggested instead that TGase treatment affects mainly the secondary stage of rennet-induced coagulation. They suggested that this inhibition was due to the progressive cross-linking of the κ-casein located on the surface of the casein micelles, which provided additional steric hindrance to the aggregation of renneted micelles.

High hydrostatic pressure

High hydrostatic pressure influences various properties of milk including a reduction in the size of the casein micelles, denaturation of β-lactoglobulin and a reduction in the CCP content. Huppertz et al. (2005) studied the impact of milk heat treatment (90°C for 10 min) and subsequent application of high-pressure treatment at pressures from 0 to 600 MPa. Heated unpressurized milk or heated milk treated for 0 min at 100 MPa was not coagulable by rennet; however, heated milk treated at 250–600 MPa for 0–30 min had a rennet coagulation time equal to, or lower than, that of unheated unpressurized milk; the coagulation time decreased with increasing pressure and treatment time. The strength of the rennet-induced coagulum from heated milk treated at 250–600 MPa for 30 min, or 400 or 600 MPa for 0 min, was considerably greater than that of the rennet-induced coagulum from unheated unpressurized milk. There was also an increase in the yield of cheese curd by ≈15%.

Acid-induced milk gels

Impact of acid on casein micelles

In cultured products, such as yoghurt, as the pH of milk is reduced, CCP is dissolved and the internal casein micelle structure is altered because of the loss of CCP. Little casein dissociation occurs at the high temperatures (>40°C) commonly used for yoghurt manufacture. Aggregation of casein occurs as the isoelectric point (pH ≈ 4.6) is approached (Horne, 1999). Native casein micelles (in milk of normal pH) are stabilized by their negative charge and steric repulsion (Lucey and Singh, 2003). Lucey (2003) distinguished three (arbitrary) pH regions in the acidification of milk from pH 6.7 to 4.6:

(a) pH from 6.7 to ≈6.0. The decrease in pH causes a reduction in the net negative charge on the casein micelles, thereby reducing electrostatic repulsion. As only a relatively small amount of CCP is dissolved above pH 6.0, the structural features of the micelles are relatively unchanged.

(b) pH from ≈6.0 to ≈5.0. The decrease in pH causes a reduction in the net negative charge on the casein micelles, thereby reducing electrostatic repulsion. As the κ-casein “hairs” on the micelle surface are charged, these charged “hairs” may shrink/collapse as the pH decreases. The net result is a decrease in both electrostatic repulsion and steric stabilization. The CCP within the casein micelles is dissolved completely by pH ≈ 5.0.
(c) $pH \leq 5.0$. The net negative charge on the casein micelles declines with the approach of the isoelectric point and there are increased $+/-$ charge interactions (and van der Waals' forces). The reduction in electrostatic repulsion allows increased hydrophobic interactions (Horne, 1998, 2001). In unheated milk gels where acidification is the only coagulation method, gelation occurs at around pH 4.9; if acidification is performed at very high temperatures, a higher gelation pH is observed.

On acidification, casein particles aggregate as a result of (mainly) charge neutralization. Acidification eventually leads to the formation of chains and clusters that are linked together to form a three-dimensional network (Kaláb et al., 1983). Acid casein gels can be formed from sodium caseinate (this ingredient is sometimes used as a yoghurt stabilizer). Direct acidification of milk at a low temperature and subsequent warming is another approach to acid gel formation. Glucono-δ-lactone (GDL) is also used to acidify milk but these acid-induced gels have different rheological and structural properties from gels produced by bacterial cultures (Lucey et al., 1998a).

Hydrophobic interactions are unlikely to play a direct role in the strength of acid gels as the storage modulus of acid gels increases with decreasing measurement temperature (Lucey, 2003). Cooling gels results in an increase in the storage modulus, probably as a result of the swelling of casein particles (caused by the weaker hydrophobic interactions) and an increase in the contact area between particles (Lucey, 2003). With increasing ionic strength, the charged groups on casein are screened, thereby weakening interactions between casein particles.

Milk has been reversibly acidified by means of carbonation—injecting pressurized CO₂ as the acidifying agent—in order to reduce the pH (usually done at low temperature). Neutralization is obtained by pressure release followed by degassing under vacuum. Upon CO₂ treatment, the zeta potential and the size of the casein micelles were restored although the total amount of CCP was not restored (Raouche et al., 2007). The rheological properties of acid gels (made using GDL) from CO₂-treated milk were similar to those of acid gels from untreated milk (Raouche et al., 2007).

Some factors influencing the texture of yoghurt gels

It is well established that the way in which the milk is handled or prepared, including the processing conditions used in yoghurt manufacture, greatly influences the gel texture, strength and stability (Lucey and Singh, 1998; Walstra, 1998; Tamime and Robinson, 1999; Jaros and Rohm, 2003a, 2003b), and that these factors include: (a) fortification level and material(s) used in the mix, (b) stabilizer type and usage levels, (c) fat content and homogenization conditions, (d) milk heat treatment conditions, (e) starter culture (type, rate of acid development and production of exopolysaccharides), (f) incubation temperature (influences growth of starter cultures, gel aggregation, bond strength), (g) pH at the breaking of the gel (stirred) and/or the start of cooling (set), (h) cooling conditions (when cooling is started, rate of cooling) and (i) post-manufacture handling of the product, e.g. physical abuse (e.g. vibration) and temperature fluctuations (i.e. if the product is not maintained at $\approx 5^\circ$C).
Homogenization and fat globule surface material

The fat globules in milk are surrounded by membrane proteins and, unless homogenized, fat acts as an inert filler in milk gels. Cho et al. (1999) prepared fat globules with different surface materials and studied the effects of these surface materials on the rheological properties of acid milk gels. Gels containing fat globules stabilized by non-interacting materials (“structure breaker”) (i.e. Tween and unheated whey protein concentrate [WPC]) had low storage moduli compared with interacting surface materials (“structure promoter”) (skim milk powder, sodium caseinate and heated WPC).

Milk for the manufacture of yoghurt is normally homogenized (15–20 MPa) to increase the yoghurt consistency and to decrease whey separation during storage (Tamime and Robinson, 1999). High-pressure homogenization has a similar principle to conventional homogenization but works at significantly higher pressures (up to 400 MPa). Milk given a high-pressure (>200 MPa) treatment gave firmer yoghurt gels than milk heat treated (90°C for 90 s) and traditionally homogenized at 15 MPa (Serra et al., 2007); presumably this effect reflects a combination of the creation of very small fat globules, whey protein denaturation and possible modification to the CCP content (Huppertz and de Kruif, 2006; López-Fandiño, 2006).

High hydrostatic pressure

High-hydrostatic-pressure treatment of milk enhances the mechanical properties of yoghurt gels (Needs et al., 2000). The storage moduli of gels made from high-pressure-treated milk were considerably higher than those of gels made from heat-treated milk (85°C for 20 min), although the yield stress and the yield strain were lower in the pressure-treated gel (Needs et al., 2000). The combined use of high thermal treatment and high hydrostatic pressure results in extensive whey protein denaturation and casein micelle disruption respectively (Harte et al., 2003). The net effect of the combined high hydrostatic pressure and thermal treatments was an improvement in the yield stress of the yoghurt and a reduction in syneresis (Harte et al., 2003). High-pressure treatment up to 600 MPa (for 20 min) improved the viscosity of stirred yoghurt, which had similar rheological properties to yoghurt made from milk heated at 90°C for 30 min (Knudsen et al., 2006).

Enzymatic modification of proteins

Acid-induced gelation of TGase-cross-linked casein resulted in increased gel firmness, lower permeability, finer protein networks and improved whey drainage (Færgemand and Qvist, 1997; Færgemand et al., 1999; Schorsch et al., 2000). Lauber et al. (2000) reported that even a very small amount of casein cross-linking, due to the action of TGase, is capable of inducing significant changes in yoghurt texture (i.e. a large increase in gel strength). A slightly slower acidification rate by the starter culture was observed in yoghurts made from TGase-treated milk; possibly there was a reduction in availability of the low molecular weight peptides required by Streptococcus thermophilus as a result of the cross-linking reaction (Færgemand et al., 1999; Ozer et al., 2007).
Cross-linking of caseins restored the sensory texture profile of a lower protein yoghurt to be comparable with that of a higher protein yoghurt, suggesting that TGase could be used instead of some of the milk solids currently used in yoghurt fortification (Færgemand et al., 1999). Excessive protein cross-linking increased the gel firmness but the yoghurt became grittier than the control samples (Færgemand et al., 1999). TGase is capable of cross-linking caseins even under high pressure (Lauber et al., 2001).

When TGase treatment was performed during high-hydrostatic-pressure treatment, a markedly higher final storage modulus was observed in acid milk gels compared with gels with only pressure treatment or when a separate TGase treatment was performed (Anema et al., 2005). Anema et al. (2005) proposed that there is an increase in cross-linking of the whey proteins and an increase in cross-linking between the whey proteins and casein when TGase treatment is performed under high pressure.

**Heat treatment**

Acid gels formed from unheated milk are very weak and this may arise, at least partly, because the interparticle contact area is still dominated by the presence of the \( \kappa \)-casein hairs (GMP), which have collapsed but are still present (Li and Dalgleish, 2006). The \( \kappa \)-casein hairs are rich in hydroxylated amino acids, some of which are glycosylated, and also acidic and basic residues. Thus, the interface between the aggregating particles will tend to be highly hydrated and attractive interactions will be partly offset by the hydrophilic tendency of the \( \kappa \)-casein hairs (Li and Dalgleish, 2006).

There has been considerable recent research on the topic of how whey proteins influence yoghurt texture. Native whey proteins in unheated milk are inert fillers in yoghurt (Lucey et al., 1999). Added whey proteins alter yoghurt gelation and texture as long as the mix is given a sufficiently high heat treatment to denature the whey proteins and cause them to associate with the casein micelles (Lucey et al., 1999). Commercially, WPC is often used to increase the solids content of yoghurt and to give improved viscosity and lower whey drainage. High-heat treatment causes considerable whey protein denaturation (e.g. 85ºC for 15 min results in >80% \( \beta \)-lactoglobulin denaturation). As a result, \( \beta \)-lactoglobulin becomes mostly attached to the \( \kappa \)-casein of the casein micelles or forms soluble complexes (with serum casein), depending on the heating conditions (i.e. pH) (Lucey et al., 1998b).

Denatured whey proteins (DWP) attached to the surface of casein micelles during heating (i.e. bound DWP) are a critical factor in the increased stiffness of yoghurt gels made from heated milk. DWP cause micelles to aggregate at higher pH because of the higher isoelectric pH (\( \approx 5.3 \)) of the main whey protein, \( \beta \)-lactoglobulin, than that of caseins (Lucey et al., 1997; Guyomarc’h et al., 2003). An alternative view is that the DWP associated with the micelles alter the hydrophobic interactions between heated micelles, which facilitates gelation at higher pH values (although there is greater electrostatic repulsion at higher pH) (Jean et al., 2006). More cross-linking of gels by bound DWP increases the gel strength. Soluble DWP are not able to increase the gel stiffness of milk in which there are no bound DWP present, i.e. the micelle surface does not contain any “bound” DWP, which can be created experimentally (Lucey et al., 1998b) (Figure 16.3).
In industrial practice, heating milk always creates some bound DWP, which allows soluble DWP to become attached to the micelles and to contribute to the gel strength. The pH at heating influences the association of DWP with casein micelles. At pH 6.5, most DWP are associated with micelles (e.g. >70% for milk heated at 90°C for 30 min). At higher pH (e.g. 7.0), fewer DWP are associated with micelles as more κ-casein dissociates from the micelles to interact with β-lactoglobulin during heating.

The gel strength of acid gels made from milk heated at high pH is higher than that of acid gels made from milk heated at the natural pH of milk (Lucey et al., 1998b; Anema et al., 2004); this may not be valid for situations in which there is a lot of added whey protein. At high pH values there is an increase in the concentration of CCP (additional cross-linking) in milk (McCann and Pyne, 1960), which could potentially increase the stiffness of acid gels made from high-pH milk. Increasing the pH of heat treatment of the milk from 6.5 to 7.0 should also alter protein unfolding and disulfide bond formation, involving β-lactoglobulin, as the pK value of its free thiol group is 9.35 (Kella and Kinsella, 1988a). The creation of additional covalent disulfide bonds that involve whey protein and caseins should increase the strength of the yoghurt gel.

Irrespective of the pH of the milk at heating, DWP (i.e. those designated as “soluble” and “bound” at the pH of heat treatment) are insoluble at low pH and should associate with casein at the pH values involved in yoghurt fermentation. As the pH decreases during fermentation virtually all the residual soluble complexes become attached to caseins via the bound DWP. The rate of acidification and the gelation temperature may also influence how these complexes associate with the caseins during acidification. The extent of denaturation of the whey proteins is often determined by their loss of solubility at pH 4.6 (de Wit, 1981), so that all the DWP should precipitate as the pH approaches pH 4.6.
The addition of WPC to milk that was then given a high-heat treatment resulted in an increase in the pH of gelation, an increase in gel stiffness and a reduction in fracture strain compared with gels made from heated milk without added WPC (Lucey et al., 1999). If WPC was added to heated milk and this mixture was not given any further heat treatment, the acid gels formed after acidification were weaker than those made from heated milk without WPC. This suggests that any added whey proteins must be denatured in order to reinforce the network, even when DWP are already present in the milk.

Schorsch et al. (2001) examined the effect of heating whey proteins in the presence or absence of casein micelles on the subsequent acid gelation properties of milk. The acid-induced gelation occurred at a higher pH (around pH 6.0) and in a shorter time when the whey proteins (concentration of 1 g whey protein/kg) were denatured separately from the casein micelles than when the whey proteins were heated in the presence of the casein micelles. However, the gels formed were very weak, probably because of the formation of a weak network in which whey proteins entrapped caseins.

Various studies have shown some conflicting results about the relative importance of the soluble and bound DWP fractions to the texture of acid milk gels (Lucey et al., 1998b; Guyomarc’h et al., 2003; Anema et al., 2004). Differences in the proportions of soluble and bound DWP fractions in these studies could have contributed to these conflicting results. Guyomarc’h et al. (2003) had only a small proportion (10–15%) of β-lactoglobulin in the bound DWP fraction whereas Lucey et al. (1998b) had around 80% β-lactoglobulin in the bound DWP fraction. Guyomarc’h et al. (2003) suggested that differences in the quantitative amounts of aggregates (and the total amount of DWP) present in the systems, independently of whether they were soluble or not, could be the reason for some of the conflicting results reported by the different groups.

In gels made from heated milk, because of the high gelation pH, the gel goes through a period of solubilization of the CCP that is present within casein particles that are already part of the gel network (this event is responsible for the maximum in the loss tangent during gelation) (Lucey et al., 1997). This process loosens the interactions between caseins in the gel network, and the higher bond mobility in yoghurt gels during this period has been associated with whey separation (Lucey, 2001).

The rheological changes during the acid-induced gelation (with GDL) of unheated and heated milk at 30°C are shown in Figure 16.4. Note the much shorter gelation time, the large increase in the storage modulus and the maximum in the loss tangent (as indicated by the hatched region between the two arrows, region A) in the heated milk sample. As the low gelation pH (4.8) of the unheated milk gel occurs after most or all of the CCP is already solubilized, there is no maximum in the loss tangent in this type of gel. When acid-induced gelation of heated milk occurs rapidly at high temperature, a plateau in the storage modulus, which corresponds to the region where there is a maximum in the loss tangent, can be observed (Horne, 2001).

Bikker et al. (2000) reported that the addition of β-lactoglobulin variant B or variant C to the milk prior to heating and acidification caused a larger increase in the storage modulus of acid gels than the addition of β-lactoglobulin variant A.
Soluble whey protein polymers have been used as ingredients for yoghurt applications (Britten and Giroux, 2001). The use of whey protein polymers to standardize the protein content of milk increased the yoghurt viscosity to about twice that obtained using skim milk powder at the same protein concentration. The water-holding capacity of yoghurt standardized with whey protein polymers was considerably higher than that of yoghurt standardized with skim milk powder (Britten and Giroux, 2001).

**Incubation temperature**

Although 42°C is a commonly used fermentation temperature for yoghurt, the use of slightly lower incubation temperatures (e.g. 40°C) leads to slightly longer gelation times, but firmer and more viscous gels that are less prone to whey syneresis are formed (Lee and Lucey, 2004). At a lower incubation temperature, there is an increase in the size of the casein particles because of a reduction in hydrophobic interactions which, in turn, leads to an increased contact area between the casein particles (Lee and Lucey, 2004); a similar trend occurs when the gels are cooled. A high incubation temperature also makes the gel network more prone to rearrangements (more flexible) during gelation and these changes can lead to greater whey separation (Lucey, 2001; Mellema *et al.*, 2002).

**Whey protein gels**

As whey is usually obtained as a by-product of cheesemaking (although recent developments in membrane technology mean that, in future, “whey” will come not
necessarily from a cheese vat but as “native” whey directly from milk prior to cheese-making), its composition depends on the cheesemaking conditions, e.g. acid whey derived from cottage cheese has different mineral (ash), lactic acid and pH values from whey derived from rennet-coagulated cheeses such as Cheddar (Table 16.1).

Whey products are widely used as food ingredients because of their excellent functional and nutritional properties. Various types of whey products are made commercially, ranging from dried whey to WPC (WPC has protein contents ranging from \( \approx 35\% \) to \( 80\% \)) to whey protein isolate (WPI) (protein contents \( \geq 90\% \)) (Table 16.2). Membrane filtration, i.e. ultrafiltration (UF) and diafiltration (DF), is used to concentrate the protein fraction before spray drying into WPC. Two different approaches are used to produce WPI: (a) membrane filtration (microfiltration, UF and DF) and (b) ion-exchange chromatography coupled with UF/DF. These two approaches result in WPI with different protein profiles (Table 16.3; Wang and Lucey, 2003). Many serum proteins take part in heat-induced gelation whereas GMP and proteose peptones do not (Walstra et al., 2005).

Whey proteins are globular proteins and heating induces denaturation and aggregation. At sufficiently high protein levels (usually \( \geq 6\% \), except for purified individual whey

<table>
<thead>
<tr>
<th>Whey ingredient</th>
<th>Moisture (%)</th>
<th>Fat (%)</th>
<th>Protein (%)</th>
<th>Lactose (%)</th>
<th>Ash (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sweet whey</td>
<td>3–5</td>
<td>1.1–1.5</td>
<td>11–14.5</td>
<td>75</td>
<td>8–10</td>
</tr>
<tr>
<td>Acid whey</td>
<td>3.5</td>
<td>0.5–1.5</td>
<td>11–13.5</td>
<td>70</td>
<td>10–12</td>
</tr>
<tr>
<td>WPC35</td>
<td>3–4.5</td>
<td>3–4.5</td>
<td>34–36</td>
<td>48–52</td>
<td>6.5–8</td>
</tr>
<tr>
<td>WPC80</td>
<td>3.5–4.5</td>
<td>6–8</td>
<td>80–82</td>
<td>4–8</td>
<td>3–4</td>
</tr>
<tr>
<td>WPI</td>
<td>4–5</td>
<td>&lt;1.0</td>
<td>90–92</td>
<td>(&lt;1.0)</td>
<td>2.5–3.5</td>
</tr>
</tbody>
</table>
Milk protein gels

proteins; this depends on many factors especially pH), gelation occurs during heating or cooling. The formation and the properties of whey protein gels are influenced by many factors:

- pH;
- protein content;
- ionic strength;
- rate and temperature/time of heating;
- types and ratios of the serum proteins;
- concentration of divalent ions (e.g. Ca\(^{2+}\));
- concentration of sugars;
- concentration of lipids including phospholipids.

There have been several reviews of the gelation of globular proteins (Oakenfull, 1987; Clark, 1992, 1996, 1998; Doi, 1993; Gosal and Ross-Murphy, 2000), as well as reviews of the thermal denaturation and gelation of whey proteins (Mulvihill and Kinsella, 1987; Mangino, 1992; Aguilera, 1995; Singh and Havea, 2003; Foegeding, 2006).

### Thermal denaturation of whey proteins

Many studies on the denaturation of whey proteins have been conducted (see the review by Mulvihill and Donovan, 1987), especially \(\beta\)-lactoglobulin as this is the major whey protein and its behavior dominates the gelation behavior of whey protein products. Denaturation has been used to describe both the loss of native structure (conformational change) and the loss of solubility (e.g. at pH values close to the isoelectric point). At around neutral pH values, denaturation becomes irreversible above about 65°C (Holt and Sawyer, 2003); with a decrease in the pH, the denaturation temperature increases (Kella and Kinsella, 1988b). Disulfide bond formation is favored as the pH is increased towards the pK value of the thiol group on \(\beta\)-lactoglobulin (9.35; Kella and Kinsella, 1988b).

Denaturation, i.e. conformational change, can be reversible and, for whey proteins, the cause of irreversibility is often the formation of new covalent (mostly disulfide) bonds. Various mechanisms for the thermal denaturation/aggregation of \(\beta\)-lactoglobulin (at neutral pH) have been proposed, in which the basic steps are: (a) the dissociation

<table>
<thead>
<tr>
<th>Protein type</th>
<th>Membrane filtration</th>
<th>Ion exchange chromatography</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\beta)-lactoglobulin</td>
<td>48–55%</td>
<td>60–73%</td>
</tr>
<tr>
<td>(\alpha)-lactalbumin</td>
<td>15–22%</td>
<td>12–25%</td>
</tr>
<tr>
<td>Bovine serum albumin and</td>
<td>4–7%</td>
<td>6–16%</td>
</tr>
<tr>
<td>Immunoglobulins</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glycomacropeptide</td>
<td>17–26%</td>
<td>0.2–1.4%</td>
</tr>
</tbody>
</table>
of the dimer into monomers and a conformational change leading to the exposure of Cys121, which initiates sulphydryl–disulfide interchange reactions, (b) an endothermic transition to a “molten globule” state, and (c) the unfolding of the protein and a second, high-temperature endothermic transition (Holt and Sawyer, 2003).

The reactive monomers formed during the denaturation process initially form dimers and trimers via the thiol–disulfide exchange reaction and the conversion of dimer to trimer is considered to be the rate-limiting step in the aggregation process (Prabakaran and Damodaran, 1997). Patel et al. (2006) proposed that the following reactions occur when milk is heated at $\approx$85°C; the major whey proteins ($\beta$-lactoglobulin and $\alpha$-lactalbumin) alter their structures and the free cysteine (CysH121) of $\beta$-lactoglobulin initially reacts reversibly with the adjacent Cys106–Cys119 disulfide bond to give a free CysH119, which, in turn, reacts with the Cys66–Cys160 disulfide bond of the same or another $\beta$-lactoglobulin molecule to give a free CysH160. CysH160 is mobile and free to move because it is so close to the C-terminus of the molecule. Thus, it reacts with disulfide bonds in other proteins, allowing a chain reaction with other $\beta$-lactoglobulin or casein molecules to occur (Patel et al., 2006).

A possible model of these reactions during the denaturation and aggregation of $\beta$-lactoglobulin is shown in Figure 16.5. In the presence of different types of whey proteins, various heteropolymers (e.g. $\beta$-lactoglobulin–$\alpha$-lactalbumin or $\beta$-lactoglobulin–bovine serum albumin) are formed during heating (Havea et al., 2001).

During the heating of $\beta$-lactoglobulin, the loss of native structure occurs via both disulfide-linked aggregate formation and non-covalently linked aggregates (e.g. hydrophobic interactions) (McSwiney et al., 1994). When $\beta$-lactoglobulin was heated at 75°C, gelation was not observed until most of the protein had aggregated (McSwiney et al., 1994). Pure $\alpha$-lactalbumin is very heat stable (because it does not have a free thiol group), although it does undergo a reversible transition at
64°C (Ruegg et al., 1977). In the presence of β-lactoglobulin, it undergoes irreversible aggregation via the thiol–disulfide exchange reaction as well as other types of interactions (Elfagm and Wheelock, 1978).

During the heating of β-lactoglobulin, most of the helical conformation is lost by about 65°C; with increasing temperature there is progressive loss of β-sheet structure (Qi et al., 1997). However, in β-lactoglobulin, a considerable amount of secondary structure, particularly β-sheet, still remains intact even at 90°C (Bhattacharjee et al., 2005). Aggregation of globular proteins starts when heat causes some unfolding of the molecule, which exposes reactive groups or sites (e.g. hydrophobic regions) that favor intermolecular interactions (Foegeding, 2006). Gupta et al. (1999), using Monte Carlo computer simulations, indicated that protein-like molecules need only partially unfold before they are susceptible to aggregation. Aggregation ultimately results in gelation if the protein concentration and other gelling conditions are favorable. This aggregation process is governed by a balance between attractive hydrophobic and repulsive electrostatic interactions.

Fractal aggregation theory has been applied to the aggregation and formation of whey protein gels (Vreeker et al., 1992; Ikeda et al., 1999). Euston (2004) argued that theories of fractal aggregation are not necessarily a good representation of protein gel structure as they treat the aggregating protein as a rigid particle and ignore any structural changes that occur in the protein during denaturation and aggregation. This criticism could be particularly important for the gelation of globular proteins, such as β-lactoglobulin.

A gel is formed when the extent of aggregation exceeds a critical level for the formation of a self-supporting network that is able to entrap the solvent.

Types and properties of whey protein gels

Different types of gel networks can be formed by globular proteins, such as whey proteins. The network structure in a heat-induced globular protein gel is strongly dependent on the balance between attractive and repulsive forces among (partially) denatured protein molecules during the aggregation process. As whey proteins have isoelectric points (pI) in the vicinity of pH 5, they are negatively charged at neutral pH values. In whey protein solutions, the ionic strength is important as it regulates the amount of ions available for the screening of charged groups on the proteins. At neutral pH values and under low ionic strengths, there is intermolecular repulsion. Aggregation of denatured proteins proceeds via hydrophobic sites and this leads to the formation of fine-stranded gels (with a transparent or translucent appearance and strands that are often 10–20 nm in thickness) (Stading and Hermansson, 1991).

Intermolecular repulsion can be reduced by increasing the ionic strength or by adjusting the pH to be closer to the isoelectric point of the whey proteins (≈5). Under gelation conditions of high ionic strength or pH values close to 5, whey proteins form opaque or particulate or turbid gels. The particles/clusters in this type of gel are in the micron size range. This type of gel structure has a poorer water-holding capacity than fine-stranded gels (Bottcher and Foegeding, 1994). Particulate gels break down rapidly during mastication to yield a homogeneous distribution of small
particles, whereas fine-stranded gels break down into large, inhomogeneous particles with irregular shapes (Foegeding, 2006).

A fine-stranded gel formed at neutral pH is rubbery and deformable to a large strain with a small fracture stress (Stading and Hermansson, 1991). At acidic pH values, intermolecular disulfide bonding is unlikely to occur and the fine-stranded networks formed at very low pH values (e.g. 3) are brittle. Particulate gels normally fracture at a small strain, but the stress required to reach the fracture strain is relatively large (Stading and Hermansson, 1991; Bottcher and Foegeding, 1994; Foegeding et al., 1995). After heat-induced gelation, cooling results in strengthening of the network because of hydrogen bond formation.

Heat-induced β-lactoglobulin gels exhibit the characteristics of a “strong gel”, i.e. they have a low frequency dependence on the storage modulus (the linear slope, $n$, of the plot of log frequency versus log storage modulus is <0.06) (Stading and Hermansson, 1990). The slope $n$ is slightly higher for particulate gels than for fine-stranded gels (Stading and Hermansson, 1990).

At pH values around 2 and low ionic strengths, whey protein gels that have some similarities in structure to β-amyloid fibrils are formed (Gosal et al., 2004; Bolder et al., 2006). Fibrils are usually rigid, non-branching and filamentous structures, around 8 nm (or larger) in width (for β-lactoglobulin) and often more than 1 μm long, that arise from linear aggregation of partly unfolded proteins (Gosal et al., 2004). α-Lactalbumin and bovine serum albumin can also form fibrils during heating at pH 2 (Goers et al., 2002; Veerman et al., 2003).

Other factors influencing properties of whey protein gels

pH and ionic strength greatly impact on the type of gel formed and its properties. The strength of whey protein gels increases with protein content. The minimum protein content needed for gelation depends on whether an individual whey protein (e.g. β-lactoglobulin) or a commercial mixture (e.g. WPC) is used as well as the gelation conditions (e.g. pH, heat treatment, ionic strength). Pure solutions of β-lactoglobulin can form a self-supporting gel at 5% protein content when tested at pH 8.0 and a heat treatment of 90°C for 15 min (100 mM Tris-HCl buffer) (Matsudomi et al., 1991).

The protein profile is important for whey gelation, e.g. higher gelling whey products can be made by increasing the proportion of β-lactoglobulin and decreasing the proportion of GMP. As α-lactalbumin is a poorer gelling protein than β-lactoglobulin, increasing the proportion of β-lactoglobulin to α-lactalbumin also increases the gelation properties of whey products (Hines and Foegeding, 1993); commercial whey products with a higher ratio of β-lactoglobulin to α-lactalbumin are available (e.g. WPI made by ion-exchange chromatography compared with WPI made by membrane filtration, or acid whey WPC; both have little or no GMP).

Salts have a major effect on the type of whey protein gel formed as a result of heat treatment and its mechanical/sensory properties. It is generally recognized that the addition of NaCl or CaCl$_2$ to dialyzed samples of WPC or WPI results in an increase in gel strength. Above a level of 10–20 mM CaCl$_2$ and 100–200 mM NaCl, the gel firmness starts to decrease (Schmidt et al., 1979; Kuhn and Foegeding,
Excessive calcium has been speculated to cause rapid protein aggregation, which limits protein unfolding and network formation (Mangino, 1992). Caussin et al. (2003) reported that the addition of calcium to whey proteins resulted in the formation of very large protein aggregates during heating. Most commercially available WPC products probably have calcium contents that are greater than that required for optimal gel strength (Mangino, 1992).

There is considerable variability in the thermal aggregation behavior of commercial whey products and some of these differences could be removed by dialysis of these samples to a common ionic strength (McPhail and Holt, 1999). The concentrations of divalent cations are higher in WPC made from cheese whey than in WPC made from acid whey and these cations are not easily removed by dialysis, suggesting some binding with the whey proteins (Havea et al., 2002). Although acid whey starts with a higher calcium content than cheese whey (Table 16.1), it is presumably easier to remove these salts in the manufacture of acid whey WPC than in the manufacture of cheese whey WPC.

Acid whey WPC is known as a superior heat-gelling product compared with cheese whey WPC (Veith and Reynolds, 2004). These differences could be due to the absence of GMP and the low calcium concentration in acid whey WPC. It has been reported that polyphosphates have been added to WPC to improve the gelling properties (Veith and Reynolds, 2004). There are various possible processing approaches to reduce the calcium/mineral content of cheese whey WPC (e.g. electrodialysis, addition of chelating agents, low-pH UF/DF) in order to improve its gelling properties.

The gelling time is also dependent on temperature, with the time required for gelation decreasing with increasing temperature although, at very high temperatures, gelation may occur only during the subsequent cooling stage (Hillier and Cheeseman, 1979). Many reports show that, when all other factors are kept constant, the gel strength increases with increasing temperature (presumably reflecting greater unfolding and reactivity of the proteins) (Mulvihill and Kinsella, 1987). The presence of lipids and lactose impairs the gelation of whey proteins (Mulvihill and Kinsella, 1987). Sugars, such as lactose, are known to protect the protein against loss of solubility during heat treatment and increase the thermal denaturation temperature of whey proteins (de Wit, 1981; Jou and Harper, 1996). Possibly, lipids might interfere with the hydrophobic interactions that play a role in the aggregation of partly unfolded whey proteins during heat treatment.

**Cold gelation of whey proteins**

Gels can also be produced using a two-step process that involves heat treatment at low ionic strength and/or far from the isoelectric point, followed by an increase in ionic strength and/or an adjustment in pH (Barbut and Foegeding, 1993; Britten and Giroux, 2001). These gels are labelled as cold-set gels, as the initial heat treatment produces a polymerized solution, with gelation occurring during the subsequent cold-set conditions through screening of the repulsive forces. To obtain gels via the cold-set gelation method, it is necessary to prepare a heat-denatured solution, with a protein concentration below the critical gelation concentration. Gelation can then be induced at low temperatures by the addition of mono- or polyvalent cations (e.g. Ca^{2+}).
Britten and Giroux (2001) acidified whey protein polymers to pH 4.6 with GDL and formed opaque particulate gels. The storage modulus and the firmness of the gels were affected by the conditions used to prepare the protein polymers.

**Enzymatic modification of whey protein (for gelation purposes)**

Extensive hydrolysis of whey protein using proteinases results in gelation mainly via hydrophobic interactions, with hydrogen bonding and electrostatic interactions also playing a minor role (Otte et al., 1996; Doucet et al., 2003).

The casein fractions in milk are more susceptible to TGase cross-linking than the globular whey protein fractions (Jaros et al., 2006). Some unfolding of β-lactoglobulin improves the extent of cross-linking with TGase (Færgemand et al., 1997; O’Sullivan et al., 2002a). Cold-set whey protein gels at low pH have been cross-linked with the TGase enzyme under either low pH or alkaline conditions (Eissa et al., 2004; Eissa and Khan, 2005).

One approach involved two steps, firstly cross-linking whey proteins with TGase at pH 8 and 50°C and secondly cold-set acidifying the resulting solution using GDL (Eissa et al., 2004). During the first step, the whey proteins undergo enzyme-catalyzed ε-(γ-glutamyl)lysine bond formation with a substantial increase in viscosity. Enzyme-cross-linked gels had significantly higher yield/fracture stress and strain than cold-set gels prepared without TGase enzyme or conventional heat-set gels. In addition, the elastic modulus of the enzyme-catalyzed gel was higher than that of its non-enzyme-treated counterpart.

**Mixed gels made with rennet and acid**

Milk coagulation can be induced by the combined action of acid and enzyme (i.e. mixed gels). The study of mixed milk coagulation has received very little attention when compared with rennet- or acid-induced coagulation, although there have been several recent studies (Roefs et al., 1990; Lucey et al., 2000, 2001; Tranchant et al., 2001).

Cottage cheese is generally manufactured by acid coagulation of pasteurized skim milk and a small concentration of rennet is sometimes added after the starter has been allowed to develop some acidity (i.e. at pH around 5.5) (Castillo et al., 2006). The use of rennet in combination with acid development initiates gelation at a high pH and the gel can undergo a “weakening” stage (as indicated by a decrease/plateau of the storage modulus, a decrease in the light backscatter ratio or an increase in the loss tangent).

This weakening is more pronounced with unheated milk gels and where there have been very high levels of κ-casein hydrolysis prior to acidification (Li and Dalgleish, 2006). This “weakening” stage is related to rearrangements caused by CCP demineralization of the casein particles in the gel network because this CCP solubilization occurs after gelation (gelation is initiated at a high pH in mixed gels) (Lucey et al., 2000). The final storage modulus of mixed gels can be considerably higher than that of acid gels made without rennet. Mixed gels made from heated milk formed firmer
gels, as they were cross-linked by denatured whey proteins and underwent fewer large-scale rearrangements (Lucey et al., 2000).

The rheological and microstructural properties of mixed gels are complex and these properties can be adjusted by varying the rennet level or the acidification rate (Tranchant et al., 2001). The use of low rennet levels during the fermentation of milk resulted in a coarser acid gel network and higher syneresis (Aichinger et al., 2003). Micelle fusion was faster in gels with rennet added because of the removal of the κ-casein hairs (Aichinger et al., 2003).

Gastaldi et al. (2003) studied the acid-induced gelation of milk samples in which chymosin was used to vary the degree of κ-casein hydrolysis prior to acidification (further chymosin activity during acidification was blocked using an inhibitor). The gelation pH increased and the gelation time decreased with an increasing degree of κ-casein hydrolysis. Gels with much higher storage moduli were formed as a result of partial κ-casein hydrolysis prior to gelation, although the loss tangent and the serum-holding capacity were lower (Gastaldi et al., 2003). Presumably, partial κ-casein hydrolysis prior to acid gelation facilitated greater rearrangements/fusion of casein, which was responsible for the increase in the storage modulus but also increased the serum separation (Lucey et al., 2001).

Conclusions

The formation and the physical properties of milk protein gels have been the subject of intense study because of the great economic impact of these gels for dairy products such as cheese, yoghurt and heat-set whey gels. There is a growing recognition that the internal structure of casein micelles plays an important role in the structural properties of rennet, acid and mixed gels. These gels are dynamic in nature and undergo rearrangements.

Technologists have recently studied the impact of high pressure and enzymatic cross-linking of these proteins to modify their functionality. The interaction between DWP and caseins has received a lot of attention and this interaction has been used to alter the texture of acid gels, although there is disagreement about the exact mechanism(s) involved. DWP polymers have been used for making cold-set gels and they have interesting possible applications in various milk gels/products. Fine-stranded whey proteins made at very low pH values have been shown to be similar in structure to amyloid fibrils. From an industrial viewpoint, these fine-stranded fibril types of gels might have some useful applications because they gel at low protein concentrations.

References


Abstract

Milk proteins have a central role to play in the development of functional foods—foods that have targeted physiological effects in the body over and above the normal effects of food nutrients. Milk proteins contain high amounts of bioavailable amino acids making them ideal ingredients for the manufacture of nutritionals—foods designed for specific nutritional purposes. Certain amino acids (e.g., tryptophan as a precursor of serotonin or leucine in the regulation of muscle metabolism) have specific physiological roles and some isolated milk proteins have particularly high concentrations of these amino acids, allowing foods to be developed to target physiological end points.

Milk proteins, and especially whey protein and glycomacropeptide, have an application in inducing satiety in humans and the relatively low yield of ATP per unit amino acid in comparison with glucose or fatty acids means that milk proteins are ideal ingredients for weight-loss foods.

Finally, milk proteins are known to be a rich source of bioactive peptides, released in the gut naturally during digestion. These peptides have a plethora of physiological effects and notable effects locally at the gut level. This chapter discusses the multiple nutritional and physiological properties of milk proteins and peptides in the context of functional foods.
Introduction

Over the last century, scientists have gradually come to better understand the complexity of foods. In addition to delivering nutrients, satisfying hunger and providing pleasure, food components are now known to have a role in directly influencing physiological processes in the body. In particular, certain food components may assist in maintaining or promoting health and well-being and preventing the development of disease. These components may be non-nutrients (e.g. the antioxidant effect of polyphenols or the blood-cholesterol-lowering effect of plant sterols) or nutrients (e.g. the effects of short-chain volatile fatty acids in modulating gut development and function). In fact, the often multiple effects of some chemical compounds released from foods during digestion, traditionally considered to have a sole role in nourishment, call into question the very definition of “a nutrient”.

It is also now widely appreciated that body growth and maintenance processes are a subtle interaction between nutrient supply and genetically regulated metabolism. The assimilation and metabolism of food compounds is subject to genetic and epigenetic control, which will vary among individuals and diverse populations, leading to important gene–nutrient interactions. In turn, nutrients and food non-nutrients also greatly influence gene expression, with effects once again varying among individuals; the complexity and the subtlety of nutrigenomics and nutrigenetics are really only beginning to be understood.

Over the next 50 years, it is expected that great advances will be made in understanding how food influences gene expression and how genetic regulation influences the assimilation and utilization of nutrients, and how the individual’s genome explains differences among individuals in their physiological and nutritional response to different foods under different conditions. Such understanding will pave the path towards personalized nutrition and personalized health foods and dietary/exercise regimens.

Milk is an excellent example of a food having both nutritional and non-nutritional physiological roles in the human diet. Milk and, in particular, milk proteins not only supply the body with amino acids necessary for the maintenance and growth of body protein, but also give rise, during food manufacture and/or food digestion, to a myriad of protein fragments and large and small peptides that have distinct biological functions (Ward and German, 2004).

Also, certain amino acids (e.g. leucine, tryptophan) released during digestion have regulatory functions or act as precursors for the synthesis of key non-protein metabolites. Such compounds are a rich source of bioactive components for the development of functional foods. Although these compounds are undoubtedly of major significance to the suckled infant, whereby milk should be regarded as a biological fluid specifically designed by nature for optimal growth and development, they are also probably of importance in the adult diet, where milk has long been an important constituent.
Functional foods

A functional food has been defined by Diplock et al. (1999) as:

A food can be regarded as functional if it has beneficial effects on target functions in the body beyond nutritional effects in a way that is relevant to health and well-being and/or the reduction of disease.

In a sense, and following this definition strictly, most if not all foods could be described as “functional” and perhaps the definition needs to be expanded to include the notion that the “beneficial effect on target functions” occurs at a meaningful level. Minor beneficial effects on target functions, which would be consequent upon the ingestion of many mainstream foods, may be relevant in the long run, in the context of a balanced diet, to health, well-being and disease prevention, but such foods are not widely regarded as functional foods.

Under the latter broader definition of a functional food, foods may be functional naturally (e.g. oily fish containing high amounts of omega 3 fatty acids) or may be rendered functional usually by adding a bioactive component to a food or by removing some component that is inhibiting bioactivity. Foods may also be enriched with a given bioactive component or components, through conventional animal or plant breeding practices, by genetic engineering or by manipulation of the feeding and nutrition of the plant or animal.

Functional foods is a rapidly growing sector of the international food industry, with development spurred by a number of technical, social and economic drivers. Firstly, there is, today, a high degree of awareness of the link between diet and health established largely through well-publicized epidemiological studies. This knowledge has been expanded by the more cogent “proof of cause and effect”, from human intervention studies. Well-educated consumers, aware of the importance of diet to health, are demanding healthy and functional foods and are prepared to pay a price premium for such foods. Escalating healthcare costs, which are a major concern to governments, encourage “disease prevention rather than cure” community health strategies.

Completion of the human genome project with a rapid accumulation of knowledge concerning single and multiple gene effects (heralding pre-symptomatic testing for disposition to particular conditions), coupled with a better understanding of inter-generational nutritional effects (Barker hypothesis) on predisposition to chronic disease states, is likely to lead in the near future to personalized nutrition strategies and further demand for specific functional foods tailored to the requirements of the individual (mass customization).

If the functional foods industry is to achieve its full potential, however, it will be critical that regulatory bodies have a clear, consistent and rigorous approach to safety, labeling and health claims issues and that food manufacturers contract reputable science providers to independently establish “proof of concept” around their products (Roberfroid, 2000). There will also need to be a considerable investment in research and development to clearly establish cause-and-effect relationships between
food compounds and targeted physiological end points and recognized disease risk factors.

Consumer confidence can quickly be eroded by conflicting messages received from the scientific community, underlining the critical role of regulatory authorities in ensuring that sufficient adequate information is available and is correctly analyzed using meta-analysis techniques, to make sustainable health claims, both qualified and approved claims. The burden of proof for generic health claims and for claims on the efficacy of specific food products needs to be considerable, such that consumers can have high levels of confidence.

The need for adequate evidence to substantiate claims and the need for regulatory authorities to be cautious in allowing claims is borne out by the recently published conclusion of the American Heart Association (Sacks et al., 2006) that:

earlier research indicating that soy protein has clinically important favorable effects [decreased blood LDL cholesterol] as compared with other proteins has not been confirmed.

The US Food and Drug Administration (FDA) had previously (October, 1999) approved labeling for foods containing soy protein as protective against heart disease, which in the light of fuller information does not appear to be supported. There has been widespread acceptance in the food industry and among consumers of the cholesterol-lowering properties of soy protein, but it would now appear that such an effect is meagre even at high daily intakes of soy protein. There are many other examples of conflicting information. The food industry must be careful that, while forging ahead with new product development and associated implied or stated health claims, long-term consumer confidence is not eroded.

Foods undoubtedly have a major role to play in preventing disease and ensuring health and vitality. However, if functional foods are to achieve their potential as part of an overall lifestyle stratagem towards healthfulness, then consumers must be guided by the highest quality information and distilled findings that have a strong likelihood of remaining substantiated over time. The food industry, science providers and government bodies all have a responsibility to ensure that the functional foods movement is led by ethical informed decision making. The recent Institute of Food Technologists (IFT, USA) Expert Report on Functional Foods (2005), although recognizing the enormous potential for functional foods, stated:

But that is not to say that IFT believes that all foods on the market for which claims are being made are being properly represented based on science and proper regulatory policies. IFT does not support some claims on foods marketed today because they are not supported by today’s science.

Clearly there is a challenge! The IFT Expert Panel goes on to recommend basing structure/function health claims on broad-based scientific criteria that address the underlying link between health and nutrition and meet the need for sound scientific substantiation supporting the structure/function effect. The panel discusses principles around ensuring the safety of functional foods and has introduced the concept of GRAE (generally recognized as efficacious), analogous to GRAS (generally recognized as safe), to encourage public confidence in the labeling of functional foods.
The dairy industry is in an excellent position to take advantage of the trend towards more healthy diets, more healthy foods and functional foods. Because milk proteins are readily available sources of amino acids and give rise to many bioactive compounds, they have a central place in the development of both specialized nutritionals and functional foods.

**Milk proteins as a source of amino acids**

Milk and milk proteins have long been regarded as a rich source of readily digestible and nutritionally available amino acids.

**Specialized nutritionals**

Early *in vivo* determinations of protein digestibility were based on fecal measurement, which is now known to be flawed given the significant degree of colonic microbial metabolism known to occur and that amino acids either are not absorbed as such from the large intestine or are absorbed to only a very limited extent (Moughan, 2003). The preferred accurate method for determining amino acid digestibility is to determine unabsorbed amino acids at the end of the small bowel (terminal ileum). This can be achieved in humans using naso-intestinal intubation or through the co-operation of ileostomates. Alternatively, animal models can be used with rat and pig ileal digestibility assays, both being suitable (Moughan *et al*., 1994).

Where a protein has undergone structural alteration due to processing or storage (especially at high temperatures), conventional digestibility measures are inappropriate for some amino acids and in particular the often first-limiting amino acid, lysine. A new lysine bioavailability assay (based on the collection of ileal digesta and application of the guanidination reaction) has been developed; it can usefully be applied to damaged proteins (Moughan, 2003). When digestibility determinations are based on the sampling of ileal digesta, it is important to recognize that digesta contain copious quantities of endogenous (of body origin) protein in addition to undigested protein. This endogenous protein component needs to be taken into account (Moughan *et al*., 1998), to yield “true” rather than “apparent” estimates of digestibility.

There are very few published data on true ileal protein digestibility as determined using human subjects. A comprehensive set of studies in humans (Gaudichon *et al*., 1994, 1995, 1996; Mahé *et al*., 1995, 1996) demonstrated a high true ileal digestibility of protein in milk proteins of around 95%. Comparable values, using the same methodology, for soy and pea proteins were 91 and 89% respectively. Sandstrom *et al*. (1986) gave soy- and meat-based diets to ileostomates and reported true ileal digestibility coefficients for total nitrogen in the range 80–85%. The naso-intestinal intubation method with normal volunteers has also been used to determine digestibility coefficients for individual amino acids (Gaudichon *et al*., 2002).

For cow’s milk, true ileal digestibility ranged from 92% for glycine to 99% for tyrosine, whereas, for soy bean, digestibility ranged from 89% for threonine to 97% for tyrosine. In our own laboratory, true ileal amino acid digestibility determined
using ileostomates ranged from 98% for aspartate to 100% for cysteine in sodium caseinate; from 93% for threonine to 99% for cysteine in whey protein concentrate; from 95% for glycine to 99% for arginine in soy protein isolate; and from 91% for cysteine to 100% for arginine in soy protein concentrate (Moughan et al., 2005a).

There are more comprehensive data on the true ileal digestibility of amino acids in various milk proteins, which have been obtained using animal models for digestion in humans (Rutherfurd and Moughan, 1997). Table 17.1 shows ileal digestibility data obtained using the laboratory rat for selected amino acids in soy protein concentrate, soy protein isolate, lactic casein, sodium caseinate, whey protein concentrate, α-lactalbumin and milk protein concentrate. These data confirm the very high digestibility of milk-derived proteins to the end of the ileum in simple-stomached mammals. The dietary amino acids are virtually completely digested.

Almost all dairy proteins have been subjected to some type of processing during their manufacture and, given that milk products often contain the reducing sugar lactose, they are susceptible to damage to the amino acid lysine. A specific assay designed to allow an accurate determination of lysine bioavailability in processed foods (Moughan and Rutherfurd, 1996) has recently been applied to a range of commercially available milk-protein-based products (Table 17.2), once again underscoring the high bioavailability of milk proteins and the limited amount of lysine damage incurred by proteins with modern controlled processing. In contrast, when the same bioassay was applied to grain-based processed foods, including cereals for children, substantial amounts of lysine damage were found (Table 17.3). Milk and milk-based products have an important role in complementing cereal foods and in supplying available lysine.

Figure 17.1 highlights the substantial differences that exist in the amounts of digestible amino acids supplied by plant proteins (e.g. soy) and milk proteins (e.g. α-lactalbumin). The often first-limiting amino acids (lysine and methionine plus cysteine) are found in much higher concentrations in the milk proteins, making them excellent sources of amino acids and very important dietary constituents to afford a balanced dietary protein intake.

Because of their relatively high levels of nutritionally important amino acids, milk proteins are utilized efficiently by humans, when given as a sole protein source. Tomé and Bos (2000) reported net post-prandial protein utilization values of 80% and 72% for milk protein and soy protein respectively, measured over 8 h after the ingestion of standard meals by healthy human subjects.

### Table 17.1 Mean true ileal digestibility of selected amino acids in a range of soy and dairy products

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>Soy protein concentrate</th>
<th>Soy protein isolate</th>
<th>Lactic casein</th>
<th>Sodium caseinate</th>
<th>Whey protein concentrate</th>
<th>α-Lactalbumin</th>
<th>Milk protein concentrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lysine</td>
<td>97.3</td>
<td>98.5</td>
<td>98.8</td>
<td>98.0</td>
<td>98.2</td>
<td>94.7</td>
<td>98.3</td>
</tr>
<tr>
<td>Methionine</td>
<td>95.3</td>
<td>100.0</td>
<td>100.0</td>
<td>99.6</td>
<td>100.0</td>
<td>99.2</td>
<td>100.0</td>
</tr>
<tr>
<td>Cysteine</td>
<td>86.9</td>
<td>95.3</td>
<td>99.2</td>
<td>93.0</td>
<td>99.6</td>
<td>96.1</td>
<td>97.8</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>96.4</td>
<td>96.8</td>
<td>94.8</td>
<td>90.6</td>
<td>98.1</td>
<td>95.4</td>
<td>94.9</td>
</tr>
<tr>
<td>Leucine</td>
<td>95.7</td>
<td>95.3</td>
<td>99.1</td>
<td>97.6</td>
<td>99.1</td>
<td>96.1</td>
<td>98.9</td>
</tr>
</tbody>
</table>

*Source: Adapted from Rutherfurd and Moughan (1997), with permission of the publisher*
Given the high bioavailability of amino acids in milk proteins and their abundant supply, it is hardly surprising that milk proteins are commonly used for the manufacture of so-called nutritionals, i.e. foods designed for a specific nutritional purpose (e.g. infant, sports, elderly formulas).

In the future, with increasing human population growth and greater pressure on food supplies, it is likely that milk proteins will play an ever more important role as protein “balancers” in plant-based processed foods.

### Specific physiological roles

Amino acids may have physiological roles that are unrelated to their direct involvement in protein synthesis. These include their roles as neurotransmitters (e.g. glutamate, aspartate and glycine) and as precursors for the synthesis of other molecules.
involved in neuromuscular function (e.g. creatine and taurine) and in host defences (e.g. glutathione and nitric oxide). Tryptophan is a precursor for the synthesis of serotonin, potentially impacting mood control (van de Poll et al., 2006) and appetite regulation (Fernstrom and Wurtman, 1972; Fernstrom and Fernstrom, 1995), whereas the nitrogen-rich amino acid arginine leads to the production of nitric oxide (Wu and Morris, 1998), which is considered to have a significant role in cell signalling and the control of endothelial tone. Depending on its site of release, nitric oxide exerts several known functions, including stimulation of the pituitary gland, vasodilation, neurotransmission and immune modulation.

Another example of an amino acid with a specific metabolic role is the branched-chain amino acid leucine, which has a unique role in the regulation of muscle protein synthesis (Kimball and Jefferson, 2001). Interestingly, leucine stimulates protein synthesis directly in skeletal muscle but not in liver. The other branched-chain amino acids—iso-leucine and valine—are less effective in stimulating muscle protein synthesis compared with leucine. Leucine supplementation has been shown to stimulate recovery of muscle protein synthesis during food restriction and after endurance exercise (Gautsch et al., 1998; Anthony et al., 2000). It has also been suggested (Layman, 2002, 2003) that leucine has a potential regulatory role in glycaemic control.

It has been known for many years that glutamine, glutamate and aspartate are preferred oxidative fuels for the gut—a highly metabolic organ. Consequently, many studies...
with humans and animals have been undertaken to investigate the effects, especially of dietary glutamine, on intestinal mucosal integrity, glutathione synthesis and immune function. This has led to debate as to whether glutamine should be regarded as a “conditionally essential” dietary amino acid (Grimble, 1993). In the traumatized patient, dietary glutamine may be needed to maintain immune responsiveness and for maintenance of the mucosal barrier against bacterial action and endotoxins.

There are also amino acids not found in proteins (i.e. non-protein amino acids) with specific physiological functions. The classic example is taurine (β-aminoethanesulfonic acid), which is synthesized by the body from cysteine or methionine and is essential for the production of conjugated bile salts (taurocholic acid). Taurine is found in milk, normally in the free form. It is recognized that cow’s milk has low concentrations of taurine relative to human milk, raising the question as to whether cow’s-milk-based infant formulas should contain added taurine.

The above are examples of specific physiological functions of amino acids; there are many others. It is anticipated that, over the next decade, our understanding of the physiological roles of individual amino acids will increase, leading to opportunities to develop functional foods containing higher or lower amounts of certain amino acids. van de Poll et al. (2006) have provided a useful summary of current evidence for proven functional effects (clinical benefits) in humans consequent upon dietary supplementation with specific amino acids.

Arginine has been widely used in supplemental nutrition for surgical patients and patients with burns, to modify the inflammatory response, to enhance organ perfusion and to stimulate wound healing, but the benefits accruing from arginine supplementation are not uniformly proven and accepted. There is some evidence that taurine supplementation improves retinal development in premature babies receiving parenteral nutrition. Taurine is often added to feeding formulas for infants and growing children. Glutamine is one of the more extensively studied amino acids and has been used in the preparation of medical foods. There is evidence that glutamine supplementation may reduce infectious morbidity and the length of hospital stay in surgical patients. Phenylalanine-free preparations have application in phenylketonuria.

Etzel (2004) highlighted an opportunity for the dairy industry, whereby a number of refined high-quality proteins are produced and marketed. The diverse amino acid compositions of these proteins can be exploited. The mixtures of proteins in milk and whey may be fractionated to give isolated proteins (α-lactalbumin, β-lactoglobulin, immunoglobulins, bovine serum albumin, the caseins, lactoferrin, lactoperoxidase and the peptide glycomacropeptide that is cleaved from κ-casein by chymosin) and blends of proteins with unique amino acid patterns. Etzel (2004) compared the amino acid compositions of several milk proteins with the composition of a theoretical “average” protein. The amino acid composition of the theoretical “average” protein was calculated from the frequency of occurrence of each amino acid in 207 unrelated proteins of known sequence.

Table 17.4 provides an abridged version of the Etzel (2004) data set. Some interesting comparisons can be made. Firstly, glycomacropeptide is completely devoid of cysteine, histidine, phenylalanine, tyrosine and tryptophan. Cysteine content is relatively high in α-lactalbumin, whereas glutamine has a relatively high concentration
Milk proteins: a cornucopia for developing functional foods

in β-lactoglobulin and glutamic acid is found at a high concentration in three of the dairy products. Leucine content is some twofold higher in β-lactoglobulin compared with the “average” protein, and the threonine content of glycomacropeptide is extraordinarily high. The branched-chain amino acids as a group are higher in concentration than that found for the “average” protein. It is clear from this type of comparison that milk-based products can be developed with amino acid compositions targeting particular physiological end points.

Role in providing calories and in promoting satiety

In addition to their role as a substrate for body protein synthesis and for the synthesis of various non-protein nitrogenous compounds, amino acids may also be used as a source of dietary energy; and the interaction between dietary protein and energy has long been understood.

However, the fact that different dietary macronutrients give rise, biologically, to quite different amounts of free energy (ATP) per unit gross energy (bomb calorimeter) is often overlooked. It is often argued that a “calorie is a calorie” regardless of the macronutrient giving rise to the energy. This is true but, what often fails to be appreciated, is that usually the numbers of calories in a food deemed to be derived from the respective macronutrients are estimates, not absolute measures. Conversion factors such as the Atwater factors are applied to determined amounts of macronutrients in a food and “available” energy is estimated. The point is that the conversion factors are not completely accurate and thus the “estimated calories” will not be completely accurate. This has particular relevance in the case of amino acids.

Atwater factors attempt to take into account the loss of energy due to incomplete absorption of the amino acid during digestion and the loss of energy in excreted urinary metabolites, post catabolism. However, the net yield of ATP during the catabolism of the amino acid and the ATP cost of synthesizing urea are not accounted for. The capture

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>β-Lactoglobulin</th>
<th>α-Lactalbumin</th>
<th>Glycomacropeptide</th>
<th>Whey protein isolate</th>
<th>“Average” protein a</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cysteine</td>
<td>2.8</td>
<td>5.8</td>
<td>0</td>
<td>1.7</td>
<td>2.6</td>
</tr>
<tr>
<td>Glutamine</td>
<td>6.3</td>
<td>4.5</td>
<td>3.8</td>
<td>3.4</td>
<td>4.6</td>
</tr>
<tr>
<td>Glutamic acid</td>
<td>11.3</td>
<td>7.3</td>
<td>15.5</td>
<td>15.4</td>
<td>7.3</td>
</tr>
<tr>
<td>Histidine</td>
<td>1.5</td>
<td>2.9</td>
<td>0</td>
<td>1.7</td>
<td>2.6</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>6.2</td>
<td>6.4</td>
<td>11.9</td>
<td>4.7</td>
<td>4.8</td>
</tr>
<tr>
<td>Leucine</td>
<td>13.6</td>
<td>10.4</td>
<td>1.7</td>
<td>11.8</td>
<td>7.8</td>
</tr>
<tr>
<td>Valine</td>
<td>5.4</td>
<td>4.2</td>
<td>8.9</td>
<td>4.7</td>
<td>6.2</td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>3.2</td>
<td>4.2</td>
<td>0</td>
<td>3.0</td>
<td>4.7</td>
</tr>
<tr>
<td>Tryptophan</td>
<td>2.0</td>
<td>5.3</td>
<td>0</td>
<td>1.3</td>
<td>1.9</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>3.6</td>
<td>4.6</td>
<td>0</td>
<td>3.4</td>
<td>5.2</td>
</tr>
<tr>
<td>Threonine</td>
<td>4.4</td>
<td>5.0</td>
<td>16.7</td>
<td>4.6</td>
<td>5.5</td>
</tr>
</tbody>
</table>

aBased on amino acid compositions of 207 unrelated sequenced proteins
Source: Adapted from Etzel (2004), with permission of the publisher
of net energy as ATP for an amino acid is less efficient than for other nutrients such as glucose and fatty acids, with an accompanying higher dietary-induced thermogenesis. That amino acids are used less efficiently energetically (i.e. have a higher dietary-induced thermogenesis) compared with glucose and fatty acids has important implications for designing weight-loss diets. Foods containing high amounts of protein will provide less “available energy” (i.e. ATP) per unit dry matter or gross energy, compared with foods high in available carbohydrate and/or fat.

Dairy proteins are a highly versatile source of amino acids, for the design of weight-loss foods, and more care should be taken in describing the caloric values especially for functional foods designed for weight loss. Protein has a further advantage for the formulation of weight-loss foods. It is now widely accepted that protein is a satiating nutrient and is effective relative to carbohydrate and fat in suppressing voluntary food intake independent of its caloric value. The role of dietary protein in the regulation of food intake and body weight in humans, and underlying mechanisms, has been the subject of recent reviews (Anderson and Moore, 2004; Westerterp-Plantenga and Lejeune, 2005). There is strong evidence that the protein content of a food is a determinant of short-term satiety and of how much food is eaten. The role of protein in the regulation of long-term food intake and body weight is less clear, because of a paucity of relevant experimental observations.

The role of protein in body weight regulation, in comparison with other macronutrients, is considered to consist of several often-related but different aspects: satiety, thermogenesis, metabolic energy efficiency and body composition. As stated, protein appears to increase satiety and therefore helps to sustain reduced-energy-intake diets.

Firstly, the highly satiating effect of protein has been observed both post-prandially and post-absorptively. Secondly, and also as discussed, high-protein diets are associated with a high dietary-induced thermogenesis, which could be related to the satiety effect of proteins. Thirdly, high-protein diets assist to maintain or increase fat-free body mass and the maintenance of a higher lean mass is costly energetically (i.e. a higher resting energy expenditure), leading to a lower associated metabolic efficiency of energy utilization.

Of particular interest to the dairy industry is the observation that protein source per se may be a factor influencing short-term satiety in humans. Whey protein has been identified as a candidate protein that may be highly effective in promoting satiety (Vandewater and Vickers, 1996; Portman et al., 2000; Hall et al., 2003). A basis for differences in satiety related to source of protein may be found in amino acid composition (e.g. a high leucine content stimulating protein synthesis and altering body energetics), in bioactive peptides released from the protein during digestion (refer to the following section), in different kinetics of protein digestion and, in the case of whey, in the presence of glycomacropeptide.

**Milk proteins as a source of bioactive peptides**

Milk is known to contain proteins (e.g. lactoferrin, lactoperoxidase, immunoglobulins) and free peptides having specific non-nutritional physiological functions. These
compounds are undoubtedly important in the case of the human infant and may also have a functional role in adults. Of potentially greater significance, however, are the many small (from 3 to 20 amino acids) bioactive peptides encrypted in food protein amino acid sequences and released during digestion. Bioactive peptides are specific protein fragments that influence body function. These peptides are inactive within the sequence of the parent protein and can be released during proteolysis or fermentation. Bioactive peptides may act as physiological modulators locally in the gut and, potentially, systemically. Most, if not all, proteins appear to contain bioactive sequences, although the majority of research to date has been conducted with milk proteins.

An opioid activity of peptides derived from partial enzymatic digestions of milk proteins and wheat gluten was reported in the literature as early as 1979 (Brantl et al., 1979; Zioudrou et al., 1979). Since then, a considerable body of research has been undertaken, many different bioactive amino acid sequences have been discovered and physiological functions have been defined. The potential for bioactive peptides in the development of functional foods is great. It is now appreciated that bioactive peptides have a wide range of physiological effects, some of which are listed in Table 17.5. Specific bioactive peptides and protein hydrolysates can now be produced commercially, allowing for dietary supplementation and protein fortification. Casein-derived peptides are already in commercial use as food supplements (e.g. phosphopeptides) and as pharmaceuticals (Meisel, 1997).

The remainder of this section focuses on the first two functions listed in Table 17.5 (i.e. gut function), as an example of the potential of food-derived peptides as natural bioactive peptides. Several studies have described the involvement of bioactive peptides (exorphins) in regulating stomach emptying rate, gastrointestinal motility and gut secretory activity in mammals (see Rutherfurd-Markwick and Moughan, 2005). A role for bioactive peptides in influencing gut function is not to be unexpected, as the effects may be mediated both directly and hormonally, involving receptor sites in the gut without the need for absorption and systemic uptake of the peptide.

Our own studies within the Riddet Institute at Massey University highlight the potential importance, quantitatively, of the net effect of food-derived peptides on overall gut metabolism. The gut is a highly metabolic organ with changes in the rate of organ metabolism having significant implications for total body energetics, protein dynamics and amino acid and other nutrient requirements. In our series of studies, terminal ileal digesta amino acid or nitrogen flow was determined as an indicator of overall protein dynamics in the upper digestive tract consequent upon the ingestion

<table>
<thead>
<tr>
<th>Table 17.5 Reported effects of natural food-derived peptides</th>
</tr>
</thead>
<tbody>
<tr>
<td>● Modulation of gastrointestinal motility</td>
</tr>
<tr>
<td>● Stimulation of secretory processes</td>
</tr>
<tr>
<td>● Mineral binding</td>
</tr>
<tr>
<td>● Antimicrobial properties</td>
</tr>
<tr>
<td>● Immunomodulation</td>
</tr>
<tr>
<td>● Antithrombotic activity</td>
</tr>
<tr>
<td>● Inhibition of angiotensin-converting enzyme (ACE) in the control of hypertension</td>
</tr>
<tr>
<td>● Analgesic (pain relief) and other neuroactive effects</td>
</tr>
</tbody>
</table>

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of a meal and is reflective of the various cellular and dietary controls on the protein secretion and amino acid reabsorption processes.

Endogenous amino acid flows (the net result of secretion and reabsorption) at the end of the ileum were determined following the provision of semi-synthetic corn-starch-based diets, differing in the source of dietary nitrogen (protein-free, synthetic amino acids, protein, hydrolyzed protein). A range of methods (Moughan et al., 1998) to determine endogenous (of body origin) as opposed to exogenous (diet origin) nitrogen were applied.

Table 17.6 gives results for endogenous lysine (marker for total protein) flow at the end of the small bowel from a representative study from our series of experiments using the pig as a generalized mammalian model. The results clearly demonstrate that, when amino acids were present in the gut (either directly from the hydrolyzed casein or after being released from the digestion of dietary zein), endogenous protein loss at the end of the small bowel was significantly enhanced. The peptides led to an enhanced secretion of protein into the gut lumen and/or a reduced reabsorption of endogenous amino acids.

In any case, the loss of extra protein into the colon, whereupon the amino acids are not salvageable, represents a considerable loss of amino acids and is associated with a high metabolic energy cost. Further work has demonstrated that the quite dramatic effect (an almost 60% increase in flow for the hydrolyzed casein) of dietary peptides is dose dependent (Hodgkinson et al., 2000; Hodgkinson and Moughan, 2006).

These results combined with those of several other similar studies provide compelling evidence for a significant influence of dietary peptides on gut protein dynamics and overall metabolism. Little is known about how these effects are mediated or how the magnitude of the effect is influenced by the source of dietary protein. Claustre et al. (2002) have recently shown that casein and lactalbumin hydrolysates (but not egg or meat hydrolysate) greatly stimulate mucin secretion in rat jejunum. The casein-hydrolysate-mediated effect was blocked by the administration of naloxone (an opioid antagonist), and β-casomorphin-7, an opioid peptide released from β-casein during digestion, also induced mucin secretion. The peptide effect was also inhibited by naloxone. It may be that the effects are more pronounced with milk proteins.

The co-operation of ileostomates has allowed our results obtained from animal studies to be confirmed in humans (Moughan et al., 2005b; Table 17.7).

<table>
<thead>
<tr>
<th>Diet</th>
<th>Significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>PF</td>
<td>SAA a</td>
</tr>
<tr>
<td>Lysine loss (mg/kg dry matter intake)</td>
<td>252</td>
</tr>
</tbody>
</table>

a Devoid of lysine, with intravenous lysine infusion
b Digesta were centrifuged and ultrafiltered (10 000 Da molecular weight cut-off)
c Naturally devoid of lysine, with intravenous lysine infusion
Adapted from Butts et al. (1993), with permission of the publisher
Bioactive peptides, and it would seem particularly those arising from the digestion of milk proteins, have been shown to have multiple physiological effects often at modest dietary intakes. As more is understood about these effects, there will be the possibility to develop novel functional foods. The potential to develop protein hydrolysates, peptide fractions and commercially synthesized peptides, to target physiological end points associated with gut motility, digestion, energetics and satiety, is particularly promising.

Conclusions

In this chapter, a case has been made for the central place of dairy proteins in the development of functional foods and specialized nutritionals. Dairy proteins are a source of highly bioavailable amino acids and offer a diverse range of amino acid patterns and specific amino acid concentration ratios. Certain amino acids, having direct physiological as opposed to nutritional functions in humans, are found in some milk proteins in high concentrations. Milk proteins may have antimicrobial and immunomodulatory functions and induce and maintain satiety.

Additionally, amino acids have a relatively high dietary thermogenesis; thus dairy proteins are ideal for the formulation of specialized weight-loss foods. Almost every week, new information is reported concerning bioactive peptides, which are released in the gut during the natural digestion of milk proteins. Milk is indeed a veritable cornucopia for developing functional foods.

References


Milk proteins: the future

Mike Boland

Abstract

This final chapter contemplates future trends and their likely impact on the production and use of milk proteins. We consider first global issues, including energy consumption, the global water economy and specific issues for dairy relating to greenhouse gases. We then review current and emerging trends in consumer demands and how they might impact on the market for milk proteins. Important factors are expected to be food safety and traceability, as well as an increasing concern for the effect of food on health and an increasing importance of nutrig-enomics and personalized nutrition. Finally we consider some emerging technologies and how they might affect the future of milk protein production and processing.

Introduction

As a wrap-up of our journey from expression to food, this chapter takes a look at the possible future of food, especially as it relates to milk proteins. Global macro-environmental factors are considered first, then we examine consumer demands and trends and the likely impact of new technologies.

Global issues for food

Global issues that can be expected to have a predominant impact on future food production (and hence production of milk proteins) include: energy—primarily...
because of the greenhouse gas implications of energy use, but also because of the rising cost of energy; the effect of the water economy; and the consequence of methane emissions from cows on global warming. Fifty-year predictions for the USA include a 50% loss of available water, as well as land, and a 50% reduction in animal agriculture.

**Milk and energy**

Milk is energetically very expensive. Milk is an animal product: to produce it requires the cow to eat vegetable material that has already been produced in a nutritional format. However, milk is the most efficiently produced of the animal-produced foods—largely because the animal itself is not consumed.

It has been estimated that production of 50 kg of milk protein in the USA requires $7 \times 10^6$ kcal of feed energy (i.e. 585 kJ/kg), an energy efficiency of 30:1 (Pimentel and Pimentel, 1979). In contrast, the total energy input per kilogram for production of corn or soy protein in the USA calculates out to 58 kJ/kg (data calculated from Pimentel and Pimentel, 1979). These figures do not take into account the uptake of direct solar energy through photosynthesis as the crops grow, or the opportunity cost in energy for other use of the land used to grow these products.

These figures mean that dairy has a strong sensitivity to energy and energy-related costs. As energy costs rise, dairy protein products will increasingly be restricted to use in high-value or luxury foods and substitution by vegetable proteins will increase, particularly in the area of nutritional ingredients.

Early signs of energy sensitivity in the market are coming through the use of “food miles”—an inappropriately named measure of the carbon footprint (i.e. the energy cost) expended in producing and distributing foods. These measures can be expected to become more accurate and more stringently applied in future, but also will be potentially misused as non-tariff barriers in some jurisdictions. We note that, because most food products are shipped by sea from remote markets, the greenhouse gas component of shipping is small compared with production costs, even when food is shipped halfway across the world, such as from New Zealand to the UK: the contribution of CO$_2$ from shipping was estimated at 125 kg CO$_2$/tonne milk solids out of a total of 1422 kg CO$_2$/tonne, which in turn compared favorably with the figure of 2921 kg CO$_2$/tonne milk solids for the locally produced equivalent in the UK (Saunders et al., 2006).

**Milk and the water economy**

Increasingly, international attention is being paid to the “water economy” as water becomes a limiting resource in many regions. The amount of “virtual water” in a product means the amount of water required to produce it throughout the production chain. The amount of virtual water in a range of products is given in Table 18.1.

Most of the virtual water in these products arises from on-farm activities, with processing water a minor component. Hence, protein product values have been calculated here by simply adjusting for the amount of protein in the parent product, without
adjustment for processing water or credit for the water value of any co-products. The key point is that, as with energy, the cost of water for producing milk-origin products is several-fold higher than for producing similar plant-origin products. This means that only countries that are very water-rich can ever contemplate producing animal-based products. This will impact in future as water distribution changes with climate change, but also threatens production in some parts of the world where existing water use is unsustainable, such as parts of Australia where water offtake has led to saline ingress into soils (Anderies et al., 2006). A full discussion of the implications of climate change on dairy production is beyond the scope of this chapter, but recent droughts in Australia, leading to a downsizing of the dairy herd, may be portents of the future.

**Implications of dairy methane production**

Methane is worthy of special mention as a greenhouse gas because emissions from cows contribute substantially to the greenhouse gas load as a by-product of rumen digestion. Methane is recognized as a greenhouse gas and is rated as having a global warming potential 21 times that of the equivalent amount of carbon dioxide on a 100-year timescale. It has been estimated that methane is second in effect only to carbon dioxide and is responsible for around 10–15% of the present greenhouse gas effect in the atmosphere. Globally, ruminant livestock produce about 28% of methane emissions from human-related activities. A single adult cow is a relatively minor contributor, emitting only 80–110 kg of methane but, with about 100 million cattle in the USA alone and 1.2 billion large ruminants in the world, ruminants are one of the largest methane sources. In the USA, cattle emit about 5.5 million tonnes of methane per annum into the atmosphere, accounting for 20% of US methane emissions, with dairy cattle producing around one-quarter of the total (see http://www.epa.gov/methane/rlep/faq.html).

Most governments recognize the need to limit greenhouse gases and current international negotiations are expected to impose penalties on greenhouse gas producers. This will have serious implications for the dairy industry and penalties, or costs of compliance, may become prohibitively high in some jurisdictions. Research efforts to specifically target the removal of methanogenic organisms from the rumen are important for the future economic viability of the industry. Because methanogens are believed to have an important role in managing the hydrogen concentration in

<table>
<thead>
<tr>
<th>Product</th>
<th>Virtual water (m³/ton)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Milk</td>
<td>990</td>
<td>Hoekstra and Chapagain (2007)</td>
</tr>
<tr>
<td>Milk Powder</td>
<td>4602</td>
<td>Hoekstra and Chapagain (2007)</td>
</tr>
<tr>
<td>Milk Protein Powders</td>
<td>18,400</td>
<td>Calculated from above</td>
</tr>
<tr>
<td>Soy Beans</td>
<td>1,789</td>
<td>Hoekstra and Chapagain (2007)</td>
</tr>
<tr>
<td>Soy Protein</td>
<td>5,400</td>
<td>Calculated from above</td>
</tr>
</tbody>
</table>
the rumen, it may be necessary to find or create a micro-organism that can transfer hydrogen into a product other than methane.

**Consumer demands and trends for food and ingredients**

**Food safety and traceability**

Throughout the world, awareness of foodborne disease has risen in response to the high level of publicity that such outbreaks receive. The toll exacted in human and economic terms is considerable. Notable dairy outbreaks in recent years include *Salmonella* in ice cream (USA, 1994: 224000 cases of illness) and staphylococcal enterotoxin in milk (Japan, 2000: 15000 cases). Contaminated soft cheeses and raw milk are often in the news. Most dairy products, processed to modern standards of hygiene, have an excellent safety record, but consumers are demanding increased surveillance and control of all foods, including dairy. The contamination of animal feed with dioxin in Belgium in 1999 highlighted the importance consumers place on the absence of toxic chemicals in their food. There will be no lessening in the demands on food producers to control risks and deliver assurances of safety. The increased costs from providing this assurance through effective process control will become the norm for dairy businesses in the future.

In recent times, increasing attention has been paid to traceability, so that any food safety issue can be quickly traced to its origin and other food from the same batch can be quickly quarantined. Traceability can also be important because of consumers’ desire for products that are sustainably produced or have other connotations of quality (such as organically produced products).

Traceability is usually managed through labeling and tracking of products through manufacture and distribution, usually by means of labels on the packaging. This is usually well handled and food manufacturers and distributors are good at it. There have, however, been attempts at “false-flagging” products in the past and this will no doubt continue. For products containing milk proteins, it is often possible to obtain an internal check on the origin of the product: dairy herds in different countries and regions tend to have a rather unique mixture of breeds and genetics. This is reflected in the distribution of polymorphisms of the proteins, which can be relatively simply analyzed using gel electrophoresis and/or mass spectroscopy. Additional information about processing can be gained from mass spectroscopic analysis of post-production changes in the chemistry of milk proteins (see Chapter 10 earlier in this volume).

**Food and health**

Consumers are being increasingly sensitized to the effects of diet on health (and appearance). The success of diet clinics attests to this. The occurrence of (and attention being paid to) current high levels of obesity in affluent societies is spurring interest in diet at all levels of society, from individual to government. Food products on
supermarket shelves are increasingly differentiated by the presence of (omega-3 fats, antioxidants) or absence of (fat-free, gluten-free) components believed to affect health.

**Nutrigenomics**

The combination of the availability of individual genetic data on a scale never before seen with a detailed understanding of nutrition has led to the field of “nutrigenomics”: the study of the relationship between a person’s genetic makeup and their individual nutritional needs. The relationship between the human body and food is now understood to be far more complex than simple nutrition.

Food contains bioactive materials that can interact with the body to stimulate or inhibit the activity of enzymes, the immune system and even the expression of genes. Individuals have differences in the genes that code for synthesis or control the expression of particular enzymes and other proteins called polymorphisms. The simplest and most common of these differ in only a single base pair in the DNA and are called single nucleotide polymorphisms, or SNPs. Individuals with different SNPs will respond differently to some food components and nutrigenomics is the science of how nutrition interacts with different individuals’ genomes, with a view to creating a healthier diet for each individual.

Some experts distinguish between nutrigenomics—where the study is based on a whole genome and systems biology approach—and nutrigenetics, which involves hypothesis-driven investigation around specific known genetic variations (for example Mutch *et al.*, 2005). “Nutrigenomics” is the term widely used to cover both aspects of nutritional science and it is used in that sense here. Whole-genome nutrigenomics is still many years away and will not be possible until all the common SNPs are mapped and their interaction with food is understood. Meantime, nutritional advice based around a series of known SNPs that do interact with diet is being provided by several companies, based on specific DNA tests for these SNPs. Such advice is usually general nutritional advice, modified to take into account risk factors associated with “bad” SNPs. This is called personalized nutrition and is dealt with in the following section.

Nutrigenomics is expected to have a wide effect on the food market in affluent countries over the next 10–20 years (Oliver, 2005) and will provide both opportunities and challenges for the food manufacturer using dairy, and most other ingredients. While there are a few known SNPs that are likely to lead to effects from milk consumption (most notably those affecting the ability to metabolize lactose, leading to lactose intolerance), there is little or no specific information relating to milk proteins, apart from the “A2” milk case discussed below.

**Personalized nutrition**

Individuals can now obtain information about their own genetic profile, with respect to known genetic polymorphisms related to health and metabolism. In the USA, companies such as Sciona and Genelex will provide a mail-order analysis of key genetic polymorphisms together with advice about diet and lifestyle.
Personalized nutrition is a nutritional response to differences between individuals, whether from a nutrigenomics input or through other identified needs and preferences, and attempts to balance an individual’s diet to their specific individual and situational needs. Nutrition today is not just about balance of macro- and micronutrients: a plethora of “functional” (bioactive) food components are also known to affect health in ways that extend far beyond the simple supply of nutrients, and they can be modifiers of nutrient uptake and usage, thus modifying the effect of nutritional balance as seen by the body’s metabolism. The kinetics of nutrient uptake are just as important as overall absolute uptakes of nutrients. Personalized nutrition attempts to take this into account, to provide optimal customized nutrition for the individual.

In sophisticated markets today, there is increasing acceptance that nutrition has a profound effect on health and wellness and, as individuals become more aware of their specific nutritional needs, the demand for personalized nutrition is set to increase.

The impact of all this on milk proteins has to date been minimal. However, three aspects are notable.

- Allergies to milk, particularly in infants, have been attributed to β-lactoglobulin in cows’ milk (although recent work has cast some doubt on this [Brix et al., 2003]). This protein is not produced in human milk and is the dominant whey protein in bovine milk (see Chapter 1 of this volume). Whey proteins are important nutritionally, as they are a valuable source of essential amino acids. β-lactoglobulin is a particularly important source of branched chain amino acids. So-called hypoallergenic products are therefore produced by hydrolyzing milk proteins, more particularly whey proteins, so that fragments are sufficiently small to be non-antigenic.

- There is some literature that suggests a weak correlation between consumption of milk containing the β-casein A1 variant and some diseases, notably type 1 diabetes (Elliott et al., 1997) and ischaemic heart disease (McLachlan, 2001). Further studies on diabetes proved to be inconclusive (Beales et al., 2004) and the heart disease data do not stand up to scrutiny; furthermore, other epidemiological data show that the A2 hypothesis does not hold up (Truswell, 2005). Notwithstanding this, a New Zealand-based company, the A2 Corporation, has been formed to produce and market a niche milk product, called A2 milk, from cows that do not carry the A1 gene. The milk has, to date, been sold mostly in Australia and the company is very careful not to make claims about any specific health benefits after having been prosecuted and fined $15,000 in Queensland in 2004 for making such claims. This product is now being sold in New Zealand and is shortly to be released in the USA—it will be interesting to see how it fares.

- There is increasing evidence that milk proteins and peptides have physiological functionality, in particular effects on cardiovascular health, immune modulation and anti-cancer effects. The validity of these effects remains to be fully proven, but in time may lead to new functional foods based on milk proteins and their products.
New technologies and their possible effect on milk protein ingredients and products

A range of new technologies has the potential to affect dairy production and processing in the near future. They include gene technologies that could lead to new, different milk proteins, new kinds of processing that can produce novel milk protein materials and products containing them, and new analytical techniques that have the potential to improve processing and place ever more stringent requirements on product quality.

Genetic modification

Milk proteins have been genetically modified and expressed in non-bovine animals (e.g. Bleck et al., 1998) and in cows (Brophy et al., 2003). However, it seems unlikely that transgenic modification of milk proteins for functional or nutritional purposes will occur widely in the foreseeable future. There are several reasons for this (Creamer et al., 2002):

- Consumer acceptance of genetically modified (GM) foods is still variable, throughout the world, with most countries now having strict labeling requirements. Because milk is a liquid product handled in large volumes during processing, maintenance of batch identity and keeping GM milk separate are more problematic than with discrete products, although recent efforts with organic milk have proven that this is possible.
- Milk is an animal product that is strongly targeted at the health of babies and young people. This has been identified in consumer surveys as a very sensitive area (compared, for example, with the acceptability of GM fruit and vegetables) and milk will probably be one of the last foods for which genetic modification is acceptable to most consumers.
- The cost of producing herds of GM cows will be very high and developing herds will be very slow unless expensive cloning and embryo transfer methods are used. This is not justified by a small premium for improved nutrition or functionality arising from genetic modification.
- More importantly, a switch to genetic modification will severely limit genetic gain, because the gene pool will be restricted to the genetics of the donor animals for the original GM parents. This segregation from the global bovine gene pool will prevent, or severely limit, participation in the ongoing genetic improvement of the species, currently occurring at about 2% per annum.

Notwithstanding these points, if a strong nutraceutical or pharmaceutical component were to be identified, enhanced expression through genetic modification would not be out of the question. However, much-touted “gene-pharming” in dairy animals has not yet been notably successful commercially.
In contrast to milk proteins, competitive plant-origin proteins are well advanced in improvement using genetic modification. GM soy beans are now predominant in world soy bean crops, covering 54.4 million hectares in 2005, which makes up 60% of world-wide soy bean production (http://www.gmo-compass.org/eng/agri_biotechnology/gmo_planting/194.global_growing_area_gm_crops.html). Soy beans can be genetically modified to remove undesirable proteins such as trypsin inhibitor, soy haemagglutinin and allergens (e.g. Friedman et al., 1991), while at the same time soy proteins can be modified to provide a more favorable nutritional balance of essential amino acids (Mandal and Mandal, 2000). The more efficient production of soy proteins in terms of energy and water, coupled with these improvements from genetic modification, means that soy proteins will increasingly out-compete dairy proteins as bulk nutritional and functional food ingredients.

**Novel processing**

High-pressure processing was originally developed in the late nineteenth century (Hite, 1899) but did not find application in food processing until the 1990s, when new materials enabled the development of production-scale processing equipment. High-pressure processing has been used commercially as an alternative method for preservation, particularly for acidic foods (Dunne and Kluter, 2001).

When milk is subjected to high pressure, the casein micelle undergoes dramatic non-reversible changes, leading to a smaller micelle that is less opaque (see Chapters 5 and 7 earlier in this volume). It has also been reported that high-pressure processing can alter the functionality of whey proteins (Patel et al., 2005). Whether these novel modified proteins will find application in new foods remains to be seen.

**New analytical methods**

Recent years have seen a range of new and improved analytical methods that have potential to improve process control and tighten product specification. Particularly important are methods that can control product safety (particularly microbiology), as well as nutritional and functional properties.

One of the weaknesses in product safety in the past has been the need to grow up samples on Petri dishes to test for the presence of undesirable microbial species. This process is time-consuming and laborious and can identify issues with process and product only well after they have occurred. A range of novel microbial detection methods is emerging; they have the potential to allow at-line detection of microbiological problems, or conversely to provide early assurance of food safety. For example, use of flow cytometry was able to reduce times for measuring bacterial numbers from the 3 days required for the traditional plate count to 2 h (Flint et al., 2006). This could be particularly important for proteins manufactured using ultrafiltration, such as whey protein concentrates and milk protein concentrates, because the ultrafiltration step co-concentrates any microbial contaminants that may be present.

Modern electrospray mass spectrometric analysis has enhanced our ability to understand and control processing effects that can alter the nutritional value of milk proteins, particularly the loss of bioavailable lysine due to processing and storage effects.
(see Chapter 10 earlier in this volume). Similarly, once the relationship between functionality and protein chemistry is well understood, the same techniques will allow better management of functional properties. Novel in-line and at-line methods are becoming possible through a range of techniques such as nuclear magnetic resonance, Fourier transform infrared spectroscopy and surface plasmon resonance analysis, for example, to measure water activities inside packaging and to predict the flavor of cheeses at maturation.

Materials science and nanotechnology

Food structure is important at all dimensional scales for the sensory properties of food (including texture, mouthfeel and flavor release) and may have an important effect on nutrient release and bioavailability (Parada and Aguilera, 2007). Increasingly, attention is being paid to materials science approaches to understanding and potentially managing these effects. An example is the physics of soft materials being applied to food (Mezzenga et al., 2005).

The potential application of nanotechnology and nanoscience to food is likely to become an important area, for the reasons outlined above. Much of the higher dimensional structure of food is a consequence of nanostructures. It is unlikely that nano robotics will be applied to food in the foreseeable future: regardless of the considerable technical challenges, public acceptance can be expected to be a major barrier. Notwithstanding this, nanotechnology is having a considerable impact on food science, in part through the use of new improved instrumentation becoming available to support nanotechnology research (Foegeding, 2006; Weiss et al., 2006).

One of the important features in nanotechnology is the occurrence of self-assembling molecular superstructures (nanostructures). It turns out that foods naturally contain many such systems, examples being the actin–myosin complex in the muscle fibers of meat, starch granules in plant foods and the casein micelle in milk. Whey proteins have also been shown to form self-assembling systems under certain conditions (Bolder et al., 2006; Graveland-Bikker and de Kruijff, 2006).

Conclusions

This brief chapter has provided a glimpse of some of the global issues and new technologies that may or may not influence the future development and use of dairy protein products. It is clear that milk proteins are expensive food ingredients and will increasingly be restricted to niche applications, and that global trends such as rising energy costs, scarcity of water and effect of greenhouse gases will increase the cost of production and will restrict land areas where they can be sustainably produced.

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Plate 2.6  Microarray analysis of genes coding for secreted proteins in the tammar wallaby mammary gland during lactation. The phases of lactation are described in Figure 2.2. A total of 75 gene transcripts are shown (see also Figure 2.6).
Plate 2.11  Histological sections of the mammary gland from Cape fur seals: (a) lactating while nursing on-shore and (b) lactating while foraging at sea. Sections are stained with hematoxylin and eosin. Magnification x 100. (c) Milk protein gene expression. β-Casein expression during Cape fur seal lactation cycle. Analysis of expression using canine Affymetrix chips hybridized to cDNA probes generated from RNA from pregnant (placental gestation and non-lactating, \( n = 2 \)), lactating on-shore (\( n = 2 \)) and lactating at sea (\( n = 1 \)) (animals in embryonic diapause) Cape fur seals. (d) Cluster analysis of gene expression profiles from the Cape fur seal mammary gland during different stages of lactation. A total of 1020 Cape fur seal mammary messenger RNA (mRNA) transcripts were identified with expression levels above an intensity of 250 in any sample type. Hierarchical clustering was conducted using Euclidean distance. Pregnant and on-shore lactating data represent an average of two animals. Off-shore data represent a single sample. Reprinted from *Current Topics in Developmental Biology*, 72, G. P. Schatten, snr ed., J. A. Sharp, K. N. Cane, C. Lefevre, J. P. Y. Arnold, and K. R. Nicholas, *Fur Seal Adaptations to Lactation: Insights into Mammary Gland Function*, pp. 276–308, New York: Academic Press. Copyright 2006, with permission from Elsevier (see also Figure 2.11).
Plate 6.1 Diagram of the dimeric structure of bovine β-Lg A looking down the two-fold axis. The co-ordinates are taken from the structure of β-Lg A in the trigonal Z lattice with 12-bromododecanoic acid bound (PDB code: 1bso). The strands that form the β barrel are labeled A to H. The I strand, together with part of the AB loop, forms the dimer interface at neutral pH. The locations of the sites of difference between the A and B variants are also shown. The structure is rainbow colored, beginning with blue at the N-terminus and ending with red at the C-terminus. Ser21, which shows conformational flexibility, and the 12-bromododecanoate anion are shown as spheres. Figure drawn with PyMOL (Delano, 2002) (see also Figure 6.1).

Plate 5.4 Representations of casein micelle structures at various pH values as indicated. The pale chains indicate protein molecules, where they cross being a hydrophobic interaction junction. The small black circles are the calcium phosphate nanoclusters that are solubilized when the pH is lowered. The outer circle is indicative of the range of steric repulsion generated between micelles and preventing interaction of the surface protein chains (see also Figure 5.4).
Plate 6.2  Ligand-binding sites on β-Lg as inferred from NMR measurements of binding of small (<12 atoms) ligands to β-Lg at acidic pH (Luebke et al., 2002). The binding site of 12-bromododecanoic acid is shown for reference (Qin et al., 1998b). (a) View into the calyx showing the primary binding site of fatty acids at pH =7 and of flavor components at pH =2, highlighted in yellow. (b) Secondary binding site for flavor components at pH 2 at N-terminal ends of strands A, B, C and D, and the C-terminal strand, highlighted in pink. (c) Secondary binding site for flavor components at pH 2 adjacent to the three-turn helix and strand G, highlighted in cyan. To show more clearly the attachment of side chains to the main chain, loops and strands have not been smoothed. The pH-sensitive EF loop is colored in magenta. Figure drawn with PyMOL (Delano, 2002) using co-ordinates with PDB code 1bso (see also Figure 6.2).
Plate 6.3  (a) Structure of bovine α-La showing the Ca$^{2+}$ ion binding site (PDB code: 2yfd). The peptide chain is rainbow colored, beginning at the N-terminus in blue and progressing to the C-terminus in red, in order to show the assembly of the sub-domains. The Ca$^{2+}$ ion is seven co-ordinate. Loop 79–84 provides three ligands, two from main-chain carbonyl oxygen atoms of Lys79 and Asp84 and one from the side chain of Asp82. Co-ordination about the Ca$^{2+}$ ion is completed by carboxylate oxygen atoms from Asp87 and Asp88 at the N-terminal end of the main four-turn helix and by two water molecules. The four disulfide bonds are shown in ball-and-stick representation (one in the helical domain is obscured and the two linking the helical domain and the Ca$^{2+}$ ion binding loop to the β domain are on the left half of the panel). (b) The lactose synthase complex formed from bovine α-La (yellow) with β-1,4-galactosyltransferase (gray) (PDB code: 1f6s). Several substrate molecules are observed, together with the cleaved nucleotide sugar moiety (cyan sticks). The Mn$^{2+}$ ion is shown as a pink sphere and the Ca$^{2+}$ ion is shown as a gray sphere. For clarity, loop regions are given a smoothed representation. Figure drawn with PyMOL (Delano, 2002) (see also Figure 6.3).
Plate 6.4  Structure of HSA complexed with halothane (slate/purple) partially occupying seven distinct sites and myristic acid (yellow/red) fully occupying five distinct sites (PDB code: 1e7c). Domain IA (residues 5–107) is shown in blue; domain IB (residues 108–196) is shown in light blue; domain IIA (residues 197–297) is shown in green; domain IIB (residues 297–383) is shown in light green; domain IIIA (residues 384–497) is shown in red; and domain IIIB (residues 498–582) is shown in light red. The single cysteine, Cys34, is labeled (Bhattacharya et al., 2000). The 17 disulfide bonds, which tie together individual sub-domains, are represented in stick format. Figure drawn with PyMOL (Delano, 2002) (see also Figure 6.4).
Plate 6.5  (a) Schematic of the general structure of Igs. Reproduced from Gapper et al. (2007) with permission. The different classes are distinguished by the constant or Fc regions of the heavy and light chains. (b) The X-ray structure of the human IgG1 molecule (PDB code: 1hzh). The heavy chains are in blue and green; the light chains are in magenta and pink. The asparagine N-linked glycan is shown in stick representation. The lack of two-fold symmetry indicates the extreme flexibility of the domains with respect to one another and the consequent sensitivity to crystal-packing effects. Disulfide bridges are shown as spheres. Each domain has a disulfide bridge joining the two sheets; additional disulfide bridges link the two heavy chains and the light chains to the heavy chains. The N-terminus of each chain is at the top left and top right of the diagram; the C-termini of the heavy chains are at the base of the molecule. Structures (e.g. PDB code: 1wej, 3hfm) where antigens are bound at the light chain–heavy chain interface indicate that binding of antigen occurs across the top of the molecule with little embedding of antigen between the domains, contrary to the mode implied by (a). Figure drawn with PyMOL (Delano, 2002) (see also Figure 6.5).
Plate 6.6  Structure of Lf. (a) Human apo-Lf (PDB code: 1cb6), showing the domain structure. The N lobe (blue for sub-domain N-I$_N$ and cyan for sub-domain N-I$_C$) is in the open conformation; the C-lobe (magenta for sub-domain C-I$_N$ and pink for sub-domain C-I$_C$), despite no metal ion present, is in the closed conformation. The helix connecting the two lobes is shown in yellow. Metal-binding ligands are shown as spheres. (b) Bovine holo-(Fe(III))-Lf (PDB code: 1blf; the human analog, 1b0l, is structurally very similar). The polypeptide is rainbow colored, blue at the N-terminus to red at the C-terminus, to highlight the manner in which the sub-domains N-I$_N$ and C-I$_N$ are formed from residues ≈1–90 and ≈250–320 (N-I$_N$) and ≈350–440 and ≈560–680 (C-I$_N$); for reference, these sub-domains are shown in approximately the same orientation in frames (a) and (b). The iron-binding ligands, including the synergistic bicarbonate ion in (b), are shown in stick form, the iron atom is shown as a red sphere and the cysteines are shown as orange spheres (16 Cys forming eight disulfides for human Lf and 34 Cys forming 17 disulfides for bovine Lf). Loops are not smoothed. Figure drawn with PyMOL (Delano, 2002) (see also Figure 6.6).