Abstract: Color is the main aspect that defines a food’s quality and that most influences consumer choice. Sensorial quality, especially color and appearance of meat, can be affected by both internal and external factors.

Chromoproteins, carotenes, and carotenoproteins are important in meat color. Carotenes are responsible for the color of beef fat, poultry meat and skin, fish, and shellfish. Of the hemoglobins present postmortem in the muscle, myoglobin is the one mainly responsible for color. Cytochromes are metalloproteins with a heme group with a role in meat coloration. The principal role of fat is in the brightness of meat products.

The color of meat may be altered by several factors, including exposure to light, microbial growth, rancidity, and exposure to oxygen.

Keywords: Muscle-based food color, carotenes, hemoproteins, myoglobin, cytochromes, fat color, melanosis, discoloration, premature browning, shelf life

1.1 GENERAL ASPECTS OF MUSCLE-BASED FOOD COLOR

The first impression that a consumer receives concerning a food product is established visually, and among the properties observed are color, form, and surface characteristics. Color is the main aspect that defines a food’s quality, and a product may be rejected simply because of its color, even before other properties, such as aroma, texture, and taste, can be evaluated. This is why the appearance (optical properties, physical form, and presentation) of muscle-based products at the sales point is of such importance (Lanari et al. 2002).

Regarding the specific characteristics that contribute to the physical appearance of meat, color is the quality that most influences consumer choice (Krammer 1994). The relationship between meat color and quality has been the subject of study since the 1950s, indeed, since Urbain (1952) described how consumers had learned through experience that the color of fresh meat is bright red, and any deviation from this color (nonuniform or anomalous coloring) is unacceptable (Diestre 1992). The color of fresh meat and associated adipose tissue is, then, of great importance for its commercial acceptability, especially in the cases of beef and lamb (Cornforth 1994), and in certain countries, for example, the USA and Canada, there have been many studies to identify the factors controlling its stability. Adams and Huffman (1972) affirmed that consumers relate the color of meat to its freshness. In poultry, the consumers of
many countries also associate meat color with the way in which the animal was raised (intensive or extensive) and fed (cereals, animal feed, etc.).

Color as a quality factor for meat can be appreciated in different ways in different countries; for example, in Denmark, pork meat color holds fifth place among qualities that affect consumers’ purchase decisions (Bryhni et al. 2002). The sensorial quality, especially color and appearance (Brewer & Mckeith 1999), of meat can be affected by both internal and external factors. In the case of internal factors, in fish, for example, a particular problem that has been encountered in rearing some Pargus species is the darkening of the body after the capture of wild fish and during farming. During farming and marketing, the skin color (silver-red) turns dark gray (especially the tail and fins) (Kentouri et al. 1995; Lin et al. 1998). In the case of farmed salmon, too, feeding fish with carotenoid pigments is regarded as the most important management practice for marketing (Moe 1990), because without them, flesh and skin color would be less visually attractive, and therefore would be less valued as a food (Baker 2002).

Food technologists, especially those concerned with the meat industry, have a special interest in the color of food for several reasons—first, because of the need to maintain a uniform color throughout processing; second, to prevent any external or internal agent from acting on the product during its processing, storage, and display; third, to improve or optimize a product’s color and appearance; and, lastly, to attempt to bring the product’s color into line with what the consumer expects. Put simply, the color of meat is determined by the pigments present in it. These can be classified into four types: (1) biological pigments (carotenes and hemopigments), which are accumulated or synthesized in the organism antemortem (Lanari et al. 2002); (2) pigments produced as a result of damage during manipulation or inadequate processing conditions; (3) pigments produced postmortem (through enzymatic or nonenzymatic reactions) (Montero et al. 2001; Klomklao et al. 2006); and (4) pigments resulting from the addition of natural or artificial colorants (Fernández-López et al. 2002).

As a quality parameter, color has been widely studied in fresh meat (MacDougall 1982; Cassens et al. 1995; Faustman et al. 1996) and cooked products (Anderson et al. 1990; Fernández-Ginés et al. 2003; Fernández-López et al. 2003). However, dry-cured meat products have received less attention (Pérez-Alvarez 1996; Pagán-Moreno et al. 1998; Aleson et al. 2003) because in this type of product, color formation takes place during different processing stages (Pérez-Alvarez et al. 1997; Fernández-López et al. 2000); recently, a new heme pigment has been identified in this type of product (Parolari et al. 2003; Wakamatsu et al. 2004a, b).

From a practical point of view, color plays a fundamental role in the animal production sector, especially in meat production (beef and poultry, basically) (Zhou et al. 1993; Esteve 1994; Verdoes et al. 1999; Irie 2001), since in many countries of the European Union (e.g., Spain and Holland) paleness receives a wholesale premium.

For fish, skin and flesh discoloration is a very important problem, especially in highly appreciated species. Since the skin and flesh color must be very vivid, many efforts have been directed at improving color, mainly through dietary control (carotene-enriched diets) (Fujita et al. 1983; Mori 1993). Without these pigments, the aquaculture industry would find it hard to undertake the production of some species because fish demand is driven through consumer demand for quality products (Baker 2002). In fish, consumer preference is often influenced by body pigmentation. Fish flesh color is an important quality parameter for most farmed fish, especially with salmonids (salmon, rainbow trout), (Francis 1995; Hyun et al. 1999), in which the pink or red color of fillets is an important feature (Sigurgisladottir et al. 1994; Sigurgisladottir et al. 1997). For example, a uniform red color in rainbow trout is considered to indicate a high-quality product and is a reason for its acceptability, while for the tuna fish industry, it is very important to avoid discoloration in fresh and processed meat and to increase its shelf life.
(Goodrick et al. 1991; Tze et al. 2001). Fish nutrition has an important impact on several parameters that directly influence the quality of fish, some of which are color and appearance. The color of salmonid flesh is one of the most important quality parameters, because consumers have a preference for red- or pink-colored products in the case of salmonids. This is the reason for using carotenoids in aquaculture.

1.2 CHEMICAL AND BIOCHEMICAL ASPECTS OF MUSCLE-BASED FOOD COLOR

Of the major components of meat, proteins are the most important since they are only provided by essential amino acids, which are very important for the organism’s correct functioning; proteins also make a technological contribution during processing, and some are responsible for such important attributes as color. These are the so-called chromoproteins, and they are mainly composed of a porphyrinic group conjugated with a transition metal, principally iron metallocorphyrin, which forms conjugation complexes (heme groups) (Whitaker 1972) that are responsible for color. However, carotenes and carotenoproteins (organic compounds with isoprenoid-type conjugated systems) exist alongside chromoproteins and also play an important part in meat color. There are also some enzymatic systems whose coenzymes or prosthetic groups possess chromophoric properties (peroxidases, cytochromes, and flavins) (Faust-man et al. 1996). However, their contribution to meat color is slight. Below, the principal characteristics of the major compounds that impart color to meat are described.

1.3 CAROTENES

Carotenes are responsible for the color of beef fat, poultry meat and skin, fish, and shellfish; in the last two cases, these are of great economic importance. The color of the fat is also important in carcass grading. Furthermore, carotenoids can be used as muscle-based food coloring agents (Verdoes et al. 1999). An important factor to be taken into account with these compounds is that they are not synthesized by the live animal but are obtained by assimilation (Pérez-Alvarez et al. 2000), for instance, in the diet. Salmonids, for example, obtain carotenines in the wild in their preys, but in intensive fish culture, carotenoids must be added to the diet. Farmed fish, especially colored fish (e.g., salmon and rainbow trout), are now a major industry. For example, Norway exports a great part of its salmon production. Carotenoid pigments have been used in aquafeed for many years in order to impart the desired flesh color in farmed salmonids (Baker 2002). Astaxanthin has been the main flesh-coloring pigment of choice in most trout- and salmon-farming industries. The type of carotene used in animal feed is very important because the fish farmer may find that pigmentation takes on a heterogeneous appearance, which is contrary to general consumer acceptance (Yanar et al. 2006). The preferred pigments used in the Canadian aquaculture industry are synthetic canthaxanthin (Cx) and synthetic astaxanthin (Ax) (Higgs et al. 1995). In fats, the fatty acid composition can affect their color. When the ratio of cis-monounsaturated to saturated fatty acids is high, the fat exhibits a greater yellow color (Zhou et al. 1993). In the case of the carotenines present in fish tissues, these come from the ingestion of zooplankton, algae, and crustacean wastes (Ostermeyer & Schmidt 2004), and the levels are sometimes very high. This is possible because fish have the capacity to transport and deposit this pigment to specific sites in their muscles (Baker 2002). The deposition of Ax is higher in dark muscle than in light muscle (Ingemansson et al. 1993). The shells of many
crustaceans, for example, lobster (*Panulirus argus*), also contain these compounds. Carotenoids have been extracted from crustacean wastes with organic solvents, but in many of the methods pigment degradation occurs (Charest *et al.* 2001).

The pigments responsible for color in fish, particularly salmonids (trout and salmon, among others), are Ax and Cx, although they are also present in tunids and are one of the most important natural pigments of marine origin. In the case of shellfish, their color depends on the so-called carotenoproteins, which are proteins with a prosthetic group that may contain various types of carotene (Minguez-Mosquera 1997), which are themselves water soluble (Shahidi & Matusalach-Brown 1998). Henmi *et al.* (1990a) reported that carotenoid–protein interaction in the salmon muscle is weak, and that Ax and Cx have a trans configuration in vivo. Henmi *et al.* (1990b) also reported that the actomyosins from salmonids showed a higher affinity for ketocarotenoids than those of other fish, except common mackerel. These authors also described correlations between the surface hydrophobicity of actomyosins and the combination of Ax and/or canthaxanthin with actomyosins. From a chemical point of view, astaxanthin or canthaxanthin bind via a beta-ionone ring to a hydrophobic binding site on actomyosin; the hydroxyl and keto end groups of the beta-end group of carotenoids intensify binding to actomyosin. Salmon actomyosin forms complexes with free Ax, astaxanthin monoester, canthaxanthin, echinenone, zeaxanthin, and beta-carotene, but not astaxanthin diester (in which a long-chain fatty acid residue may cause steric hindrance). The lipids in the actomyosin complex have no effect on the binding of carotenoids (Henmi *et al.* 1989). They are distributed in different amounts in the flesh, head, and carapace of crustaceans; for example, astaxanthin and its esters are the major carotenoids found in the extracts from different species of shrimp (*Penaeus monodon*, *Penaeus indicus*, *Metapenaeus dobsonii*, *Parapenaeopsis styliifera*) (Sachindra *et al.* 2005a), but there are different types of carotenes, depending on whether the crustaceans are marine or fresh water (Sachindra *et al.* 2005b). Another difference is that the concentration of unsaturated fatty acids in its carotenoid extracts was found to be higher than that of saturated fatty acids. In raw muscle, the main carotenoid concentration was strongly correlated with some color attributes (hue, chroma, and lightness) (Choubert *et al.* 1992). Torrisen *et al.* (1989) reported that a level of 4 mg/kg in fish fillets is regarded as a minimum acceptable carotenoid concentration in marketable-farmed salmon. Sex also affects carotene concentration: female muscles, which contain much more carotenoid, are more strongly colored than male muscles (Norris & Cunningham 2004).

As suggested by Torrisen *et al.* (1989), the rate of carotenoid deposition in salmonids is curvilinear throughout the life of the fish. As the growth rate is obviously under strong genetic control, the genetic correlation between the growth rate and color is high. It must be taken into account that carotenoids migrate from the muscle to the gonads. Carotene-type deposition in salmonid species differs; for example, Ax is more efficiently deposited than Cx in rainbow trout (Storebakken & Choubert 1991; Torrissen 1986), but this pattern is not the same for Atlantic salmon. These differences may be due to genetic background and/or environment (Baker 2002). Choubert *et al.* (1997) reported that in rainbow trout there is an unequal distribution of carotenoids so that the color of the muscle lightens from the head toward the tail and from the midline of the fish toward the dorsal and ventral external area of the fish.

From a chemical point of view, carotenoids are organic molecules that contain a conjugated carbon–carbon double bond system, which is responsible for their color. But this can be a problem during processing, because a high number of conjugated double bonds may be subject to oxidation, which can lead to discoloration of the carotenoids (Liaaen-Jensen 1971; Choubert & Baccanaua 2006). As carotenoids are lipid soluble compounds, it might be thought that increasing dietary fat would increase carotene absorption and deposition, but this is not necessarily the case for all salmonids. The retention of carotenoids in the flesh is relatively poor,
with only 10%–18% of pigment obtained from the diet being retained (Nickell & Bromage 1998). Astaxanthin can be found in its free, mono-, or diesterified forms. In processed shrimps, eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) are the principal fatty acids esterified with the portion of astaxanthin linked to chitin in the carapace (Guillou et al. 1995). b-carotene and Ax are fat-soluble pigments found in squid oil. However, technological processes, such as refining, can remove Ax completely (Hwei & Min 1994).

In fish-derived products, the carotene content has previously been used as a quality parameter on its own; however, it has been demonstrated that this is not appropriate, and that other characteristics may influence color (Little et al. 1979). The carotene content and its influence on color is perhaps one of the characteristics that has received most attention (Swatland 1995). In the case of meat, especially beef, an excess of carotenoids may actually lower the quality (Irie 2001), as occurs sometimes when classifying carcasses. The Japanese system for beef carcass classification identifies acceptable fats as white, slightly off-white, or slightly reddish white in color, while pink-yellowish and dark yellow are unacceptable (Irie 2001). It is precisely the carotenoids that are responsible for these last two colorations. However, in other animal species, such as chicken (Castaneda et al. 2005), the opposite effect is observed, since a high carotene (xanthophyll) concentration is much appreciated by consumers (Esteve 1994), yellow being associated with traditional or “home-reared” feeding (Pérez-Alvarez et al. 2000). The use of the carotenoid canthaxanthin as a coloring agent in poultry feeds is designed to result in the desired coloration of poultry meat skins. The carotenoids used include citranaxanthin, capsanthin, and capsorubin, but Cx shows superior pigmentation properties and stability during processing and storage (Blanch 1999). To improve its color and brilliance, 0.004–0.04 wt% proanthocyanidin is added to fish feed containing carotenoids (Sakiura 2001). For rainbow trout, carotenoid concentrations could be 10.7 or 73 ppm Cx, or 47 or 53 ppm Ax.

### 1.4 HEMOPROTEINS

Of the hemoproteins present, postmortem in the muscle, myoglobin (Mb) is the one mainly responsible for color, since hemoglobin (Hb) arises from the red cells that are not eliminated during the bleeding process and are retained in the vascular system, basically in the capillaries (incomplete exsanguination; the average amount of blood remaining in meat joints is 0.3%) (Warris & Rodes 1977). However, the contribution of red cells to color does not usually exceed 5% (Swatland 1995). There is wide variation in the amounts of Hb from muscle tissue of bled and unbled fish. Mb content is minimal compared with the Hb content in fish light muscle and white fish whole muscle. Hb made up 65% and 56% by weight of the total heme protein in dark muscle from unbled and bled fish, respectively (Richards & Hultin 2002). Mb, on average, represents 1.5% by weight of the proteins of the skeletal muscle, while Hb represents about 0.5%, the same as the cytochromes and flavoproteins combined. Mb is an intracellular (sarcoplasmic) pigment apparently distributed uniformly within muscles (Ledward 1992; Kanner 1994). It is red in color and water soluble, and it is found in the red fibers of both vertebrates and invertebrates (Knipe 1993; Park & Morrissey 1994), where it fulfills the physiological role of intervening in the oxidative phosphorylation chain in the muscle (Moss 1992).

#### 1.4.1 Structure of myoglobin

Structurally, Mb can be described as a monomorphic globular protein with a very compact, well-ordered structure that is specifically, almost triangularly, folded and bound to a heme group (Whitaker 1972). It is structurally composed of two groups: a proteinaceous group and a heme
group. The protein group has only one polypeptidic chain composed of 140–160 amino acid residues, measuring 3.6 nm and weighing 16,900 Da in vertebrates (Lehninger 1981). It is composed of eight relatively straight segments (where 70% of the amino acids are found), separated by curvatures caused by the incorporation into the chain of proline and other amino acids that do not form alpha-helices (such as serine and isoleucine). Each segment is composed of a portion of alpha-helix, the largest of 23 amino acids and the shortest of 7 amino acids, all dextrogyrating. Mb’s high helicoidal content (forming an ellipsoid of $44 \times 44 \times 25\,\text{Å}$) and lack of disulfide bonds (there is no cysteine) make it an atypical globular protein. The absence of these groups makes the molecule highly stable (Whitaker 1972). Although the three-dimensional structure seems irregular and asymmetric, it is not totally anarchic, and all the molecules of Mb have the same conformation. One very important aspect of the protein part of Mb is its lack of color. However, the variations presented by its primary structure and the amino acid composition of the different animal and fish species destined for human consumption are the cause of the different colorations of meat and their stability when the meats are displayed in the same retail illumination conditions (Lorient 1982; Lee et al. 2003a). The heme group of Mb (as in Hb and other proteins) is, as mentioned above, a metalloporphyrin. These molecules are characterized by their high degree of coloration as a result of their conjugated cyclic tetrapyrrolic structure (Kalyanasundaram 1992). The heme group is composed of a complex, organic annular structure, protoporphyrin, to which an iron atom in ferrous state is united (Fe II). This atom has six coordination bonds, four with the flat protoporphyrin molecule (forming a flat square complex) and two perpendicular to it. The sixth bond is open and acts as a binding site for the oxygen molecule.

Protoporphyrin is a system with a voluminous flat ring composed of four pyrrolic units connected by methyl bridges (=C–). The Fe atom, with a coordination number of 6, lies at the center of the tetrapyrrole ring and is complexed to four pyrrolic nitrogens. The heme group is complexed to the polypeptidic chain (globin) through a specific histidine residue (imidazolic ring) occupying the fifth position of the Fe atom (Davidsson & Henry 1978). The heme group is bound to the molecule by hydrogen bridges, which are formed between the propionic acid side chains and other side chains. Other aromatic rings exist near and almost parallel to the heme group, which may also form π bonds (Stauton-West et al. 1969).

The Hb contains a porphyrinic heme group identical to that of Mb and equally capable of undergoing reversible oxygenation and deoxygenation. Indeed, it is functionally and structurally paired with Mb, and its molecular weight is four times greater since it contains four peptidic chains and four heme groups. The Hb, like Mb, has its fifth ligand occupied by the imidazole group of a histidine residue, while the sixth ligand may or may not be occupied. It should be mentioned that positions 5 and 6 of other hemoproteins (cytochromes) are occupied by R groups of specific amino acid residues of the proteins and therefore cannot bind to oxygen ($O_2$), carbon monoxide (CO), or cyanide (CN–), except a1, which, in its biological role, usually binds to oxygen.

One of the main differences between fish and mammalian Mb is that fish Mb have two distinct endothermic peaks, indicating multiple states of structural unfolding, whereas mammalian Mb followed a two-state unfolding process. Changes in alpha-helix content and tryptophan fluorescence intensity with temperature are greater for fish Mb than for mammalian Mb. Fish Mb shows labile structural folding, suggesting greater susceptibility to heat denaturation than that of mammalian Mb (Saksit et al. 1996).

The helical contents of frozen-thawed Mb were practically the same as those of unfrozen Mb, regardless of pH. Frozen-thawed Mb showed a higher autoxidation rate than unfrozen Mb. During freezing and thawing, Mb suffered some conformational changes in the nonhelical region, resulting in a higher susceptibility to both unfolding and autoxidation (Chow et al. 1996).
In tuna fish, Mb stability followed the order bluefin tuna (Thunnus thynnus), yellowfin tuna (Thunnus albacares), and bigeye tuna (Thunnus obesus); autoxidation rates were in the reverse order. The pH dependency of Mb from skipjack tuna (Katsuwonus pelamis) and mackerel (Scomber scombrus) were similar. Lower Mb stability was associated with higher autoxidation rates (Chow 1991).

1.4.2 Chemical properties of myoglobin

The chemical properties of Mb center on its ability to form ionic and covalent groups with other molecules. Its interaction with several gases and water depends on the oxidation state of the Fe of the heme group (Fox 1966), since this may be in either its ferrous (Fe II) or its ferric (Fe III) state. Upon oxidation, the Fe of the heme group takes on a positive charge (Kanner 1994) and, typically, binds with negatively charged ligands, such as nitrites, the agents responsible for the nitrosation reactions in cured meat products.

When the sixth coordination ligand is free, Mb is usually denominated deoxymyoglobin (DMb), which is purple in color. However, when this site is occupied by oxygen, the oxygen and the Mb form a noncovalent complex, denominated oxymyoglobin (OMb), which is cherry or bright red (Lanari & Cassens 1991). When the oxidation state of the iron atom is modified to the ferric state and the sixth position is occupied by a molecule of water, the Mb is denominated metmyoglobin (MMb), which is brown. There are several possible causes for MMb generation, and these may include the ways in which tunids, meat, and meat products are obtained, transformed, or stored (MacDougall 1982; Lee et al. 2003b; Mancini et al. 2003). Among the most important factors are low pH, the presence of ions, and high temperatures during processing (Osborn et al. 2003); the growth and/or formation of metabolites from the microbiota (Renerre 1990); the activity of endogenous reducing enzymes (Arihara et al. 1995; Osborn et al. 2003); and the levels of endogenous (Lanari et al. 2002) or exogenous antioxidants, such as ascorbic acid or its salts, tocopherols (Irie et al. 1999), or plant extracts (Xin & Shun 1993; Fernández-López et al. 2003; Sánchez-Escalante et al. 2003). The pH, which may be altered depending on postslaughter metabolism and on ingredient addition, can affect the stability of the central iron atom in Mb and Hb. At high pH, the heme iron is predominantly in the Fe21 state; low pH accelerates Fe21 conversion to Fe31 (Zhu & Brewer 2002, 2003). While oxygen can bind to Fe21 only, many other ligands (CN, nitric oxide [NO], CO) can bind to either Fe21 or Fe31 so producing a variety of colors. This change in the oxidation state of the heme group will result in the group being unable to bind with the oxygen molecule (Arihara et al. 1995). DMb is able to react with other molecules to form colored complexes, many of which are of great economic relevance for the meat industry. The most characteristic example is the reaction of DMb with nitrite, since its incorporation generates a series of compounds with distinctive colors: red in dry-cured meat products or pink in heat-treated products. The products resulting from the incorporation of nitrite are denominated cured, and such products are of enormous economic importance worldwide (Pérez-Alvarez 1996). The reaction mechanism is based on the propensity of nitric oxide (NO, generated in the reaction of nitrite in acid medium, readily gives up electrons) to form strong coordinated covalent bonds; it forms an iron complex with the DMb heme group independent of the oxidation state of the heme structure. The compound formed after the nitrification reaction is denominated nitrosomyoglobin (NOMb). As mentioned above, the presence of reducing agents such as hydrogen sulfide acid (H2S) and ascorbates leads to the formation of undesirable pigments in both meat and meat products. These green pigments are called sulfomyoglobin (SMb) and colemyoglobin (ColeMb), respectively, and are formed as a result of bacterial activity and an excess of reducing agents in the medium. The formation of SMb is reversible, but that of...
ColeMb is an irreversible mechanism, since it is rapidly oxidized between pH 5 and 7, releasing the different parts of the Mb (globin, iron, and the tetrapyrrolic ring).

From a chemical point of view, it should be borne in mind that the color of Mb, and therefore of the meat or meat products, depends not only on the molecule that occupies the sixth coordination site, but also on the oxidation state of the iron atom (ferrous or ferric), the type of bond formed between the ligand and the heme group (coordinated covalent, ionic, or none), and the state of the protein (native or denatured form), not to mention the state of the porphyrin of the heme group (intact, substituted, or degraded) (Pérez-Alvarez 1996).

During the heat treatment of fish flesh, the aggregation of denatured fish proteins is generally accompanied by changes in light-scattering intensity. Results demonstrate that changes in relative light-scattering intensity can be used for studying structural unfolding and aggregation of proteins under thermal denaturation (Saksit et al. 1998). When fatty fish meat like Trachurus japonicus was heat treated, the MMb content increased linearly, and the percentages of denatured Mb and apomyoglobin increased rapidly when mince was exposed to heat; however, when the temperature reached 60°C, the linearity was broken. The results indicated that MMb color stability was higher than that of Mb and that the thermal stability of heme was higher than that of apomyoglobin (Hui et al. 1998). Both Mb and ferrous iron accelerated the lipid oxidation of cooked, water-extracted fish meat. Ethylenediaminetetraacetic acid (EDTA) inhibited the lipid oxidation accelerated by ferrous iron, but not that accelerated by Mb. Also, with cooked, nonextracted mackerel meat, EDTA noticeably inhibited lipid oxidation. Nonheme iron catalysis seemed to be related in part to lipid oxidation in cooked mackerel meat. The addition of nitrite in combination with ascorbate resulted in a marked inhibition of lipid oxidation in the cooked mackerel meat. From these results, it was postulated that nitric oxide ferrohemochromogen, formed from added nitrite and Mb (present in the mackerel meat) in the presence of a reducing agent, possesses an antioxidant activity, which is attributable in part to its function as a metal chelator (Ohshima et al. 1988).

Tuna fish meat color can be improved when the flesh is treated or packaged with a modified atmosphere in which CO is included. Normally, the rate of penetration of CO or carbon dioxide (CO₂) in fish meat such as tuna, cod, or salmon, under different packaging conditions, is measured by monitoring pressure changes in a closed constant volume chamber with constant volume and temperature. Alternatively, however, the specific absorption spectrum of carboxymyoglobin (MbCO), within the visible range, can be obtained and used as an indicator of MbCO formation. Mb extracts from tuna muscle treated with CO exhibited higher absorbance at 570 than at 580 nm. Therefore, the relationship between absorbance at 570 nm and absorbance at 580 nm could be used to determine the extent of CO penetration of tuna steaks placed in a modified atmosphere in which CO was included. The penetration of CO into tuna muscle was very slow. After approximately 1–4 h, CO had penetrated 2–4 mm under the surface, and after 8 h, CO had penetrated 4–6 mm (Chau et al. 1997).

In products with added nitrite or nitrate, the complex nitrosylmyoglobin (MbFe[II]NO) is the main contributor to the characteristic color of cooked cured ham, and brine-cured and dry-cured meat products. Meat and meat products without nitrite/nitrate addition will normally attain a dull brown color or a gray color in heated products, which influences consumer acceptance negatively (Adamsen et al. 2005). In dry-cured meat products such as Parma ham produced without nitrite or nitrate addition, the characteristic bright red color (Wakamatsu et al. 2004a) is caused by a Zn-protoporphyrin IX (Zn-pp) complex, a heme derivative. Adamsen et al. (2005) showed that the use of nitrite as a curing ingredient inhibits the formation of Zn-pp. In the same work, the author described that this color compound is present in other meat products like Iberian ham, although in a lower concentration.

Virgili et al. (1999) reported that this color may be due to the action of low-molecular weight compounds containing electron-donating atoms, formed during maturation, in particular
basic peptides or amino acids resulting from an external proteolysis, which may play a role as Fe ligands in Mb. Wakamatsu et al. (2004b) reported that anaerobic conditions favor the formation of Zn-pp and that endogenous enzymes as well as microorganisms may also be involved. There are several hypotheses that try to explain the formation of this compound. Wakamatsu et al. (2004b) described three possible substitution patterns: (1) a nonenzymatic reaction in which Zn(II) substitutes Fe(II) under anaerobic conditions, with concomitant dissociation of the heme; (2) a bacterial enzymatic reaction, whereby bacterial growths naturally degrade the meat proteins including the pigment; and (3) an enzymatic reaction where an endogenous ferrochelatase interchanges the two metals. However, Adamsen et al. (2005) described this process as having the three following mechanisms to explain the metal substitution: (1) a nonenzymatic enzymatic reaction driven by the binding of iron in the high chloride meat matrix; (2) a bacterial enzymatic reaction; and (3) an endogenous enzymatic reaction.

Also, spectroscopic studies of Parma ham during processing revealed a gradual transformation of muscle Mb, initiated by salting and continuing during aging. Using electron spin resonance spectroscopy, Moller et al. (2003) have shown that the Parma ham pigment is different from MbFe(II)NO and is not a nitric oxide complex such as that found in brine-cured ham and Spanish Serrano hams. These authors also establish that the heme moiety is present in the acetone-water extract and that Parma ham pigment is gradually transformed from a Mb derivative into a nonprotein heme complex, which is thermally stable in an acetone-water solution. Adamsen et al. (2003) also demonstrated that the heme moieties of Parma ham pigments have antioxidative properties. Pigments became increasingly lipophilic during processing, suggesting that a combination of drying and maturing yields a stable red color (Parolari et al. 2003).

1.4.3 Cytochromes

Cytochromes are metalloproteins with a prosthetic heme group, whose putative role in meat coloration is undergoing revision (Boyle et al. 1994; Faustman et al. 1996). Initially, they were not thought to play a very important role (Ledward 1984). These compounds are found in low concentrations in the skeletal muscle, and in poultry, they do not represent more than 4.23% of the total hemoproteins present (Pikul et al. 1986). It has now been shown that the role of cytochrome (especially its concentration) in poultry meat color is fundamental when the animal has been previously exposed to stress (Ngoka & Froning 1982; Pikul et al. 1986). Cytochromes are most concentrated in cardiac muscle so that when this organ is included in meat products, the heart’s contribution to color, not to mention the reactions that take place during elaboration processes, must be taken into consideration (Pérez-Alvarez et al. 2000).

1.5 COLOR CHARACTERISTICS OF BLOOD

Animal blood is little used in the food industry because of the dark color it imparts to the products to which it is added. For solving the negative aspects of blood incorporation, specifically food-color-related problems, several different processes and means have been employed, but they are not always completely satisfactory. The addition of 12% blood plasma to meat sausages leads to pale-colored products. The addition of discolored whole blood or globin (from which the Hb’s heme group has been eliminated) has also been used to address color problems. Natural red pigments can be obtained from blood without using coloring agents such as nitrous acid salts; these pigments have Zn-pp as the metalloporphyrin moiety and can be used to produce favorably colored beef products, whale meat products, and fish products (including fish pastes) (Numata & Wakamatsu 2003). There was wide variation in amounts of
Hb extracted from the muscle tissue of bled and unbled fish, and the residual level in the muscle of bled fish was substantial. Mb content was minimal as compared with Hb content in mackerel light muscle and trout whole muscle. Hb made up 65% and 56% by weight of the total heme protein in dark muscle from unbled and bled mackerel, respectively. The blood-mediated lipid oxidation in fish muscle depends on various factors, including Hb concentration, Hb type, plasma volume, and erythrocyte integrity (Richards & Hultin 2002). The presence of blood, Hb, Mb, Fe12, Fe13, or Cu12 can stimulate lipid oxidation in the fillets of icefish (Rehbein & Orlick 1990; Richards & Li 2004). Kanner et al. (1987) reported that Hb, Mb, copper, and iron have the potential to promote lipid oxidation in muscle foods. Since iron can be released from Hb during storage, it is difficult to ascertain whether the intact heme protein, dissociated heme, or released iron is responsible for the bulk of lipid oxidation that occurs during storage. For this reason, Svingen et al. (1979) used the term “low molecular weight iron” instead of “free iron” since iron binds to other low molecular weight compounds to gain solubility and hence potential reactivity. Ferrous and ferric forms of iron can promote lipid oxidation processes (Gutteridge 1986; Tadolini & Hakim 1996). Iron shows a high reactivity with reactants such as hydrogen peroxide and lipid peroxides (Kanner & Harel 1987).

Mitochondria are a source of reactive oxygen species that could confound lipid oxidation reactions due to added Hb. During fish processing (e.g., tuna fish), the loss of redness can be a good indicator that lipid oxidation processes mediated by Hb are progressing. Just after death, Hb in muscle tissue is primarily in the reduced state (i.e., oxyhemoglobin [OHb] and deoxyhemoglobin [deoxyHb]).

This mixture of OHb and deoxyHb has a red color. With increased postmortem aging, Hb autoxidizes to methemoglobin (MHb), a brown pigment. MHb is considered more prooxidative than reduced Hb due to its less tightly bound heme group and its reactivity with hydrogen peroxide and lipid peroxides to form hypervalent Hb catalysts (Everse & Hsia 1997).

From a technological point of view, during meat or fish processing, rapid chilling may alter oxygen solubility in tissues resulting in less available oxygen to oxygenate either OMb or Hb. The conversion of OMb to MMb, which is brown and unattractive, occurs under conditions of very low oxygen tension as well (Nicolalde et al. 2005).

Field et al. (1978) describe how bone marrow is high in Hb, while muscle has a high Mb content. As with other meats, its color and Hb stability depend on packaging and storage conditions. Good temperature control and modified atmosphere packaging (MAP) with high oxygen atmospheres (80%) are often used to extend both microbiological and color shelf life (Nicolalde et al. 2005).

### 1.6 FAT COLOR

From a technological point of view, fat fulfills several functions, although, regarding color, its principal role is in the brightness of meat products. Processes such as “afinado” during the elaboration of dry-cured ham involve temperatures at which fat melts so that it infiltrates the muscle mass and increases its brilliance (Sayas 1997). When the fat is finely chopped, it “dilutes” the red components of the color, thus decreasing the color intensity of the finished product (Pérez-Alvarez et al. 2000). However, fats do not play such an important role in fine pastes since, after emulsification, the fat is masked by the matrix effect of the emulsion so that it contributes very little to the final color. The color of fat basically depends on the feed that the live animal received (Esteve 1994; Irie 2001). In the case of chicken and ostrich, the fat has a “white” appearance (common in Europe) when the animal has been fed with “white” cereals or other ingredients not containing xanthophylls, since these are accumulated in subcutaneous fat.
and other fatty deposits. However, when the same species are fed maize (rich in xanthophylls), the fatty deposits take on a yellow color. Beef or veal fat that is dark, hard (or soft), excessively bright, or shiny lowers the carcass and cut price. Fat with a yellowish color in healthy animals reflects a diet containing beta-carotene (Swatland 1988). While fat color evaluation has traditionally been a subjective process, modern methods include such techniques as optical fiber spectrophotometry (Irie 2001). Another factor influencing fat color is the concentration of the Hb retained in the capillaries of the adipose tissues (Swatland 1995). As in meat, the different states of Hb may influence the color of the meat cut. OMB is responsible for the yellowish appearance of fat, since it affects different color components (yellow-blue and red-green).

The different states of Hb present in adipose tissue may react in a similar way to those in meat so that fat color should be measured as soon as possible to avoid possible color alterations. When the Hb in the adipose tissue reacts with nitrite incorporated in the form of salt, nitrosohemoglobin (NOHb), a pigment that imparts a pink color to fat, is generated. This phenomenon occurs principally in dry-cured meat products with a degree of anatomical integrity, such as dry-cured ham or shoulder (Sayas 1997). When fat color is measured, its composition should be borne in mind since its relation with fatty acids modifies its characteristics, making it more brilliant or duller in appearance. The fat content of the conjunctive tissue must also be borne in mind—collagen may present a glassy appearance because, at acidic pH, it is “swollen,” imparting a transparent aspect to the product.

1.7 ALTERATIONS IN MUSCLE-BASED FOOD COLOR

The color of meat and meat products may be altered by several factors, including exposure to light (source and intensity), microbial growth, rancidity, and exposure to oxygen. Despite the different alterations in color that may take place, only a few have been studied; these include the pink color of boiled uncured products, premature browning (PMB), fish skin discoloration, and melanosis in crustaceans.

1.8 PINK COLOR OF UNCURED MEAT PRODUCTS

The normal color of a meat product that has been heat treated but not cured is “brown,” although it has recently been observed that these products show an anomalous coloration (red or pink) (Hunt & Kropf 1987). This problem is of great economic importance in “grilled” products since this type of color is not considered desirable. This defect may occur both in meats with a high hemoprotein content, such as beef and lamb (red), and in those with a low hemoprotein concentration, including chicken and turkey (pink) (Conforth et al. 1986). One of the principal causes of this defect is the use of water rich in nitrates, which are reduced to nitrites by nitrate-reducing bacteria, which react with the Mb in meat to form NOMb (Nash et al. 1985). The same defect may occur in meat products containing paprika, which according to Fernández-López (1998) contains nitrates that, once incorporated in the product, may be similarly reduced by microorganisms. Conforth et al. (1991) mention that several nitrogen oxides may be generated in gas and electric ovens used for cooking ham and that these nitrogen oxides will react with the Mb to generate nitrosohemopigments. CO is also produced in ovens, which reacts with Mb during thermal treatment to form a pink-colored pigment, carboxyhemochrome. It has also been described how the use of adhesives formed from starchy substances produces the same undesirable pink color in cooked products (Scriven et al. 1987). The same anomalous pink color may be generated when the pH of the meat is high (because
of the addition of egg albumin to the ingredients) (Froning et al. 1968) and when the cooking temperature during processing is too low. These conditions favor the development of a reducing environment that maintains the iron of the Mb in its ferrous form, imparting a reddish/pink color (as a function of the concentration of hemopigments) instead of the typical grayish brown color of heat-treated, uncured meat products.

Cooking uncured meat products, such as roast beef, at low temperatures (less than 60°C) may produce a reddish color inside the product, which some consumers may like. This internal coloring is not related to the formation of nitrosopigments, but results from the formation of OMb, a phenomenon that occurs because there exist in the muscle MMb-reducing enzymatic systems that are activated at temperatures below 60°C (Osborn et al. 2003). Microbial growth may also cause the formation of a pink color in cooked meats since these reduce the oxidoreduction potential of the product during their growth. This is important when the microorganisms that develop in the medium are anaerobes, since they may generate reducing substances that decrease the heme iron. When extracts of Pseudomonas cultures are applied, the MMb may be reduced to Mb (Faustman et al. 1990).

1.9 MELANOSIS

Melanosis, or blackspot, involving the appearance of a dark, even black, color, may develop postmortem in certain shellfish during chilled and frozen storage (Slattery et al. 1995). Melanosis is of huge economic importance since the coloration may suggest a priori in the eyes of the consumer that the product is in bad condition, despite the fact that the formation of the pigments responsible involves no health risk. Melanosis is an undesirable surface discoloration of such high-value shellfish as lobsters that takes place immediately after harvesting since it starts with oxygen contact (López-Caballero et al. 2006). Blackspot is caused by enzymatic formation of the precursors of phenolic pigments (Williams et al. 2003). Blackspot is a process regulated by a complex biochemical mechanism, whereby the phenols present in a food are oxidized to quinones in a series of enzymatic reactions caused by polyphenol oxidase (PPO) (Ogawa et al. 1984). This is followed by a polymerization reaction, which produces pigments of a high molecular weight and dark color. Melanosis is produced in the exoskeleton of crustaceans, first in the head and gradually spreading toward the tail. Melanosis of shell and hyperdermal tissue in some shellfish, such as lobsters, has been related to stage of molt, since the molting fluid is considered to be the source of the natural activator(s) of pro-PPO.

PPO (catechol oxidase) can be isolated from shellfish cuticle (Ali et al. 1994) and is still active during iced or refrigerated storage. Some authors have found a connection between melanosis and microbial growth in crustaceans. Thus, color formation (melanin) due to strains of P. fragi may occur if prawns are not properly chilled (Chinivasagam et al. 1998). In this respect, López-Caballero et al. (2006) reported that the presence of microorganisms (e.g., Proteus spp., Pseudomonas, etc.) and the H2S produced reacted with metals of the lobster shell resulting in melanosis. Sulfites can be used to control the process (Ferrer et al. 1989; Gomez-Guillen et al. 2005), although their use is prohibited in many countries. It is well known that the inhibitory effect of blackspot is specific for each species, requiring adequate doses and formulations (Montero et al. 2001). The effective dose of 4-hexylresorcinol differs depending on the physiological state, season, method of application, etc., although the species being treated is one of the most important of these factors. Montero et al. (2004) and López-Caballero et al. (2006) found that, regardless of the season, a concentration of 0.25% 4-hexylresorcinol was effective in extending the shelf life of pink shrimp. Ficin (Taoukis et al. 1990) and 4-hexylresorcinol also functioned as a blackspot inhibitor, alone and in combination with l-lactic acid (Benner et al. 1994).
1.10 FISH SKIN DISCOLORATION

In fish and other vertebrates, in which the pigmentation of the skin can be changed by hormonal stimulation, the color of the background and illumination are determining factors for the intensity and/or the pattern of skin fish pigmentation (Duray et al. 1996; Crook 1997; Sugimoto 1997; Healey 1999; Papoutsoglou et al. 2000; Rotllant et al. 2003). In addition, temperature may also have an impact on color (Fernandez & Bagnara 1991). In some types of fish, especially those with a red skin, the color tone becomes dark immediately after killing, reducing the commercial value of the fish. Most of the color changes in fish are often related to stress. It is generally accepted that melanophores play an important role in the rapid color change of certain fish (Fujii 1969). These changes are related to hormonal (α-melanocyte-stimulating hormones) responses causing dispersion of the melanin granules in melanophores and are responsible for skin darkening (Green & Baker 1991; Lamers et al. 1992; Gröneveld et al. 1995; Arends et al. 2000; Burton & Vokey 2000).

In the case of Red Sea Bream, the rapid skin-color changes after killing of cultured fish is thought to be mainly due to the rapid dispersion of chromatosomes in melanophores elicited by handling and killing stresses. Potassium ions through the noradrenaline pathway can induce aggregation of chromatosomes in melanophores (Kumazawa & Fujii 1984). In fish skin, besides melanophores there are other chromatophores, such as xanthophores and erythrophores, the latter mainly contributing to the red color of the fish skin.

1.11 PREMATURE BROWNING

Hard-to-cook patties show persistent internal red color and are associated with high pH (>6) raw meat. Pigment concentration affects red color intensity after cooking (residual undenatured Mb), so this phenomenon is often linked to high pH dark cutting meat from older animals. Premature browning (PMB) is a condition in which ground beef (mince) looks well done at a lower than expected temperature (Warren et al. 1996). PMB of ground beef is a condition in which Mb denaturation appears to occur on cooking at a temperature lower than expected; it may indicate falsely that an appropriate internal core temperature of 71°C has been achieved (Suman et al. 2004). The relationship between cooked color and internal temperature of beef muscle is inconsistent and depends on pH and animal maturity. Increasing the pH may be of benefit in preventing PMB, but it may increase the incidence of red color in well-cooked meat (cooked over an internal temperature of 71.1°C) (Berry 1997). When pale, soft, exudative (PSE) meat was used in patty processing, patties containing OMb easily exhibited PMB. One reason for this behavior is that the percentage of Mb denaturation increased as cooking temperature rose (Lien et al. 2002).

1.12 COLOR AND SHELF LIFE OF MUSCLE-BASED FOODS

Meat and meat products are susceptible to degradation during storage and throughout the retail process. In this respect, color is one of the most important quality attributes for indicating the state of preservation in meat. Any energy received by food can initiate its degradation, but the rate of any reaction depends on the exact composition of the product (Jensen et al. 1998), environmental factors (light, temperature, presence of oxygen), and the presence of additives. Transition metals such as copper and iron are very important in the oxidative/antioxidative
balance of meat. When the free ions of these two metals interact, they reduce the action of certain agents, such as cysteine, ascorbate, and alpha-tocopherol, oxidizing them and significantly reducing the antioxidant capacity in muscle (Zanardi et al. 1998). Traditionally, researchers have determined the discoloration of meat using as criterion the brown color of the product, calculated as percent MMb (Mancini et al. 2003). These authors demonstrated that in the estimation of the shelf life of beef or veal (considered as discoloration of the product), the diminution in the percent of OMb is a better tool than the increase in percentage of MMb. Occasionally, when the meat cut contains bone (especially in pork and beef), the hemopigments (mainly Hb) present in the medulla lose color because the erythrocytes are broken during cutting and accumulate on the surface of the bone Hb. When exposed to light and air, the color of the Hb changes from bright red (oxyhemoglobin [OHb]), the characteristic of blood, to brown (methemoglobin [MHb]) or even black (Gill 1996). This discoloration basically takes place during long periods of storage, especially during shelf-life display (Mancini et al. 2004). This characteristic is aggravated if the product is kept in a modified atmosphere rich in oxygen (Lanari et al. 1995). These authors also point out that the effect of bone marrow discoloration is minimized by the effect of bacterial growth in MAP. As in the case of fresh meat, the shelf life of meat products is limited by discoloration (Mancini et al. 2004). This phenomenon is important in this type of product because they are normally displayed in illuminated cabinets. Consequently, the possibility of photooxidation of NOMb needs to be taken into account.

During this process, the molecule is activated because it absorbs light; this may subsequently deactivate the NOMb and give the free electrons to the oxygen to generate MMb and free nitrite. In model systems of NOMb photooxidation, the addition of solutions of dextrose, an important component of the salts used for curing cooked products and in meat emulsions, can diminish the effect of NOMb photooxidation. When a meat product is exposed to light or is stored in darkness, the use of ascorbic acid or its salts may help stabilize the product’s color. Such behavior has been described both in model systems of NOMb (Walsh & Rose 1956) and in dry-cured meat products (e.g., longanizas, Spanish dry-fermented sausage). However, when sodium isoascorbate or erythorbate is used in longanizas production, color stability is much reduced during the retail process (Ruiz-Peluffo et al. 1994).

The discoloration of white meats such as turkey is characterized by color changes that go from pink-yellow to yellow-brown, while in veal and beef, the changes go from purple to grayish brown. In turkey, it has been demonstrated that the presence or absence of lipid oxidation depends on, among other things, the concentration of vitamin E in the tissues. The color and lipid oxidation are interrelated since it has been seen that lipid oxidation in red and white muscle depends on the predominant form of catalyzing iron, Mb, or free iron (Mercier et al. 1998). Compared with red meat, tuna flesh tends to undergo more rapid discoloration during the refrigerated storage. Discoloration due to the oxidation of Mb in red fish presented a problem, even at low temperatures. This low color stability might be related to the lower activity or poorer stability of MMb reductase in tuna flesh (Ching et al. 2000). Another reason for the low color stability is that aldehydes produced during lipid oxidation can accelerate tuna OMb oxidation in vitro (Lee et al. 2003a). Tuna flesh could be immersed in an MMb reductase solution to extend the color stability of tuna fish. Also, the use of this enzyme can reduce MMb formation during the refrigerated storage of tuna (Tze et al. 2001). Yellowtail (Seriola quinqueradiata) fillets stored in gas barrier film packs filled with nitrogen (N2) and placed in cold storage at 0–5°C stayed fresh for 4–7 days. N2 or CO2 packaging did not prevent discoloration in frozen tuna fillets; better results were achieved by thawing the frozen tuna meat in an O2 atmosphere (Oka 1989). Packaging in atmospheres containing 4% or 9% O2 was inferior to packaging in air, as these atmospheres promoted MMb formation. Packaging in 70% O2 maintained the fresh red color of tuna dorsal muscle for storage periods less than 3 days.
To change the dark brown color to a bright red color, processors sometimes treat tuna with 100% carbon monoxide (CO) during MAP. Since Mb can react with CO rapidly even at low CO concentrations (Chi et al. 2001), MAP with 100% CO may result in high CO residues in the flesh, which may cause health problems.

### 1.13 MICROORGANISMS AND MUSCLE-BASED FOOD COLOR

Although the real limiting factor in the shelf life of fresh meat is the microbial load, consumers choose fresh meat according to its color. The bacterial load is usually the most important cause of discoloration in fresh meat and meat products (sausages and other cooked products), and slaughter, cutting, and packaging must be strictly controlled. Bacterial contamination decisively affects the biochemical mechanisms responsible for the deterioration of meat (Renerre 1990). Is it important to take into account that, just as with the bacterial load, the effect of discoloration on meat is more pronounced in meats that are more strongly pigmented (beef) than in less pigmented meats such as pork and chicken (Gobantes & Oliver 2000). Another variable affecting color stability in meat is the quantity of microorganisms present (Houben et al. 1998); concentrations in excess of 106/g have a strong effect. Although antioxidants, such as ascorbic acid, slow lipid oxidation and consequently improve color stability, these substances have little effect when bacterial growth is a problem (Zerby et al. 1999).

### REFERENCES


Choubert, G. & Baccamoua, M. (2006) Colour changes of fillets of rainbow trout (Oncorhyncus mykiss W.) fed astaxanthin or canthaxanthin during storage under controlled or modified atmosphere. LWT—Food Science and Technology, 39(10), 1203–1213.


