New alternatives for improving and assessing the color of dark-cutting beef – a review

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Abstract: Myoglobin (Mb) is a sarcoplasmic heme protein present in muscle cells, which acts as a short-term oxygen (O₂) reserve in the muscle tissue. After slaughtering and exsanguination, Mb is the major pigment that provides the red color in meat. The concentration of Mb together with its redox state are two pivotal factors that determine meat color. The elevated pH of dark-cutting beef can affect both physical and biochemical properties resulting in decreased oxygenation. The darkening observed in high ultimate pH (pHu) beef concerns meat processors as color is the initial attribute that impacts on the purchase. Thus, any atypical meat color (i.e., loss of brightness) reduces consumer interest in the product. Several studies have demonstrated that immunological castration is effective in preventing both aggressive behavior and undesirable dark-cutting of bull meat. However, little information is available on the effects of processing techniques that limit the oxidation of ferrous iron (Fe²⁺), Mb or promote metmyoglobin (MMb) reduction in dark-cutting beef. Because of the importance of color to fresh beef marketability, this review aimed at overviewing the significance of pHu in beef color and color stability and to discuss new alternatives for improving and assessing the beef color of dark-cutting beef, especially in Nellore bulls and their crossbreds, which are widely used in beef cattle production in Brazil.

Keywords: MMb reductase enzyme, high pHu, ultimate pH, meat quality, bull

Introduction

Brazil is one of the world’s largest beef producers (Associação Brasileira das Indústrias Exportadoras de Carnes – ABIEC, 2019), producing a total of 8.23 million tons in carcass-weight equivalent in 2019 [Instituto Brasileiro de Geografia e Estatística – IBGE, 2019, 2020]. Of this total, 80.9% was destined for domestic consumption, providing an annual consumption of approximately 42 kg per capita (ABIEC, 2019). Bos indicus genetics, mainly the Nellore breed, have a considerable share of the Brazilian cattle herd [Mueller et al., 2019], in which males represent 59.1% of the total cattle slaughtered in Brazil during 2019 (IBGE, 2019, 2020).

The irritable behavior of bulls as well as other biological and environmental elements may impair proper muscle acidification until achieving the ultimate pHu (pH) of 5.3–5.8 (Ponnampalam et al., 2017), which is considered the normal pHu range (Ouali et al., 2006; Contreras-Castillo et al., 2016). Indeed, an ultimate pH above 5.8 is detrimental as it enhances the darkening of beef, which is known as dark-cutting beef (Tang et al., 2005; McKeith et al., 2016).

Dark-cutting beef results either from reduced myofibrillar shrinkage (i.e., lower light scattering) (Hughes, 2019) or from higher myoglobin (Mb) content (Ponnampalam et al., 2017). A high pHu then inhibits the Mb oxidation from developing a brown layer of metmyoglobin (MMb) due to improved metmyoglobin reducing activity (MRA) and consequent stability of the surface dark red color (Bekhit and Faustman, 2005).

Darkening is also attributable to increased mitochondrial oxygen consumption rate (OCR), which depletes the cellular oxygen (O₂) content and is amplified in high pHu muscles (Tang et al., 2005). Accordingly, higher pHu favors the formation of the purplish deoxymyoglobin (DMb) at the expense of the bright red oxymyoglobin (OMb) on the surface of the muscle (Mancini and Hunt, 2005). The negative consequence of this undesirable effect, dark cutting beef, is less marketable (Suman et al., 2014).

Since dark cutting beef has a negative impact on the fresh beef market and consumer acceptance, an understanding of the biochemical reactions and pathways is necessary in order to mitigate this effect. Thus, the objectives of this review were to provide an overview of the significance of pHu on beef color and color stability and to discuss new alternatives aimed at improving and assessing the beef color of dark-cutting beef, especially for Nellore bulls and their crossbreds, which are widely used in beef cattle production in Brazil.

Bever ultimate pH

After slaughtering, the pHu is reached after stabilization of the drop in pH. Although pH measurements at 24 h postmortem may be conducted on group beef muscles [Li et al., 2014; Ponnampalam et al., 2017; Pulford et al., 2008; Renerre, 1990], pHu seems to stabilize only after 48 h postmortem in longissimus lumborum muscles. In fact, a decline in pHu values was observed when measured repeatedly at 24 h and at 48 h postmortem; and this continuous drop profile until 48 h postmortem was presented by Ouali et al. (2006) and Mlynek et al. (2012).
There is a global incidence of beef muscle with pHu higher than the normal threshold established by experts and the beef industry. For instance, Mach et al. (2008) reported an incidence of 13.9 % within pH > 5.8, 24 h postmortem in Spain. According to Contreras–Barón [Personal communication], most of the cattle slaughtered in Brazil has shown pH above 5.8 48 h postmortem, which, to a certain extent, may impact the exporting of beef from Brazil. Nevertheless, there are still no records on the incidence of abnormal pH in meat in the Brazilian slaughter system.

To achieve a broader understanding on the influence of the high pHu on beef color, an initial overview on how pH drops to a determined pHu is required. Furthermore, the factors that can lead to a high pHu muscle and the definition of a pHu range to establish a cut-off standard are of pivotal importance.

Biochemical basis of ultimate pH

Living muscle has a neutral pH (7.2 – 7.4). After animal exsanguination, the delivery of O2 to muscle cells is interrupted. The depletion of O2 impairs the ability of cells to metabolize glucose aerobically by the tricarboxylic acid cycle and the respiratory chain. Glucose generated from muscle glycogen must be metabolized by glycolysis to generate lactate and H+ together with adenosine triphosphate (ATP), whose hydrolysis boosts the H+ ion concentration and thereby decreases the intracellular pH (England et al., 2016; Scheffler et al., 2015).

The pH drop profile does not present continuous behavior, but this profile passes through phases of transient pH stability until reaching the pHu (Ouali et al., 2006). Ouali et al. (2006) observed the gradual pH drop in two 19–month–old Charolais bulls’ longissimus muscle over the 0 – 10 h post–slaughter period. The authors associated these pH drop profiles with the phospholipid–dependent inversion of polarity in cellular membranes. This phenomenon involves the electronegative phosphatidylserine groups switching to the external leaflet of the membrane, while the electropositive phosphatidylcholine and phosphatidylethanolamine groups change to the internal leaflet when apoptosis occurs in muscle cells. This switching causes transient partial neutralization of protons formed from glucose by glycolysis (Ouali et al., 2006).

Muscle glycolytic potential is the molar sum of all substrates of glycolysis [glycogen, glucose and lactate]. Many researchers have demonstrated the negative correlation between the glycolytic potential and pHu (Holdstock et al., 2014; McKeith et al., 2016; Wulf et al., 2002). However, the type of fiber in the muscle may be a major determinant, as described in section 2.2.

Biological and environmental effects on pHu dropping

Several pre–harvest elements may determine the extension of pH drop. Physiological factors include the types of the myofiber and the muscle, as well as the testosterone level (Fink et al., 2018). Environmental factors involve the type of finishing, diet, and pre–slaughter stress, such as physical exercise, inadequate handling, and fighting between animals immediately prior to slaughter (Dunne et al., 2011; Ponnampalam et al., 2017).

Ultimate pH may vary according to the type of fiber that comprises a muscle. As stated by Patten et al. (2008) and England et al. (2014), glycolytic fibers, such as type IIB, present a fast glycolysis rate, which is able to produce a high concentration of ATP – substrate to H+ production. Greater concentration of glycogen on glycolytic fibers favors the acidification that provides sufficient amount of substrate. Therefore, in glycolytic muscles, the pH reduction is inversely correlated to the initial concentration of glycogen in the muscle (Lawrence et al., 2012).

Ultimate pH closer to the typical range [5.4 – 5.8] inhibits the activity of the glycolysis–regulatory enzyme phosphofructokinase, which ends the metabolic process, and then the pH drop in glycolytic fibers (England et al., 2014). Thus, glycolysis is active until the medium reaches an acidic pH. Conversely, oxidative fibers, such as type I, seem to have a shorter glycolysis pathway before reaching the necessary pH to inhibit the phosphofructokinase activity. Higher pHu in oxidative fibers may occur under two conditions: reduced glycogen content in the cell with low final ATP production, and a slow rate of glycolysis, which does not produce H+ ions given the extent to which the enzyme is inhibited at pH close to 5.8, even when there is an excess of glycogen (England et al., 2016).

As reviewed by Seideman et al. (1982) and reported by Węglarz (2010), bulls have been classically known to produce beef with higher pHu. The bull’s excitable temperament encompasses both aggressive and sexual activities, and it is linked to the testosterone level (Seideman et al., 1982). Testosterone stimulates the animal, which depletes the muscle content of glycogen prior to slaughter, thus interrupting the proper muscle acidification.

On the other hand, reports have shown that different categories of animal temperament do not result in different pHu values [King et al., 2006]. However, the degree of pre–slaughter stress should be taken into account because when stressful conditions are minimized or avoided, bulls do not produce elevated pHu. Even though the longissimus dorsi muscles from bulls have appeared darker compared to steers, DeVol et al. (1985) found no correlation between the excitatory effect of testosterone on muscle color because there was no pHu > 5.8 (5.66 for bulls and 5.58 for steers) nor any correlation between these pHu values. Therefore, this result was associated with pre–slaughter stress.

Extensive exercise and/or grass feeding is reported to enhance significant changes in muscle fiber
composition, as these factors increase the proportion of the oxidative fibers (MyHC–IIa) at the expense of the glycolytic type (MyHC–IIx) (Gagaoua et al., 2017). As previously described, oxidative fibers have a slower rate and limited extent of pH drop and, subsequently, greater production of dark meat. Although the high pHu after pasture has been associated with lower intracellular glycogen content and decreased glycolytic potential the shift towards oxidative fibers reveals that the lower energetic potential is not the preponderant factor in dropping the pH (Apaoblaza et al., 2020; Picard and Gagaua, 2020).

The nutrient restriction seems to increase the proportion of oxidative fiber type in the muscle and deplete the intracellular glycogen concentration [Bray et al., 1989; Kandeepan et al., 2009]. Both conditions lead to less H⁺ ions produced during the rigor mortis process (i.e., high pHu meat).

Understanding how pre-slaughter circumstances impact on the achievement of muscles with normal pHu and the economic consequences of postmortem biochemical reactions is a means of fomenting good animal welfare and ethical issues.

Animal stress may be avoided by a handler’s careful perception of animal behavior prior to and/or during slaughter, especially if there are signs of fear, pain, and distress (Grandin, 2010, 2019). According to Bomzon (2011), cattle may disguise signs of pain so as not to demonstrate weakness in front of a possible predator. However, changes in cattle mobility, behavior and/or appearance may indicate poor welfare, such as lameness, falling during handling, vocalization, apparent sclera (the white of the eye), struggling, agitation and slipping in the stunning box, and body lesion (Bomzon, 2011; Grandin, 2010, 2019).

Vocalization above a certain level is a clear indicator of pain or stress and can be perceived by the handler while moving cattle or holding them restrained during slaughter. However, certain bulls in a herd, normally vocalize at the lairage even in the absence of stressful conditions. Vocalizing animals show increased blood concentration of the stress biomarker cortisol, lactate, and glucose, indicating glycoenenolysis (Grandin, 2019). The neuropeptide P level also increases when painful procedures are inflicted (Coitzee et al., 2008).

The frequent and gentle animal handling on the farm as well as the previous experience of moving into a veterinary restraint encourage cattle to become accustomed to the presence of humans and to walking to the slaughter box at the abattoir (Probst et al., 2013; Grandin, 2019). When this approach is adopted, animals can be cajoled into a calmer and fearless behavior right up to the slaughter procedure itself.

**Ultimate pH ranges**

Meat pHu ranges from 5.3 to over 7.0, as reported by several authors (Abril et al., 2001; Contreras-Castillo et al., 2016; Hunt and Hedrick, 1977; Lawrie, 1958; McKeith et al., 2016; Pulford et al., 2008). Nevertheless, grouping pHu into ranges is a challenging task and there is no universal consensus, which results in several classifications and cut-off values.

Studies have focused on the meat tenderness of longissimus muscles, and a number of authors have grouped pHu into three ranges: normal (< 5.8) intermediate (5.9 - 6.2) and high (> 6.20) (Lomiwes et al., 2014; Wu et al., 2014, Contreras-Castillo et al., 2016). This can be attributable to the meat tenderness observed in the three groups, in which the intermediate range is tougher than the other ones (Lomiwes et al., 2013; Pulford et al., 2008). However, these ranges which assess meat tenderness cannot be applied to meat color studies.

As regards meat color, different classifications are proposed to group beef muscles according to their values of pHu. For instance, Viljoen et al. (2002) and Park et al. (2007) considered pHu > 5.8 as a single group to be studied. On the other hand, McKeith et al. (2016) split high pHu carcasses into four classes for longissimus thoracis muscles, according to their dark color: shady (pHu: 6.1 ± 0.03), mild (6.4 ± 0.03), moderate (6.6 ± 0.03), and severe (6.9 ± 0.03). Holdstock et al. (2014) used pHu 6.0 as the threshold for classifying beef muscles as normal (pHu < 6.0) and typically dark (pHu > 6.0).

One approach to reducing the human effect on pHu range classification is based on cluster analysis (Abril et al., 2001). This statistical technique involves analyzing the experimental data collected on the basis of their proximity – forming a group of samples with similar characteristics. This tool can show that data from the parameters assessed from three pHu muscle ranges can behave as two groups of pHu, which increases the credibility of data analysis. Abril et al. (2001) applied cluster analysis to all the color coordinates and divided the samples into two groups according to their pHu: pHu < 6.1 and pHu > 6.1. This difference was also observed in the reflectance spectra.

**Dark–cutting beef and color stability**

A typical denomination for beef darkening is dark-cutting beef. Certain authors have also referred it as DFD beef (Dark, Firm and Dry). Since the term firm might be interpreted as tough, it is important to note that not every high pHu muscle is tough. Because of enzymatic action by calpains proteases, meats with high pHu (especially above 6.3) can be as tender as those with normal pHu (Lomiwes et al., 2013). Therefore, the “dark-cutting” term seems more appropriate for studies focused on color.

Researchers have reported that pHu > 5.8 may result in dark meat (Abril et al., 2001; Ashmore et al., 1972; Holdstock et al., 2014; Hunt and Hedrick, 1977; Page et al., 2001; Stackhouse et al., 2016). Because of the
showed that 91.7% of the muscles with pHu ≥ 5.87 underwent minimal shrinkage in the myofibril structure. This alteration in the meat microstructure leads to an increase in hydration capacity by muscle proteins, also defined as water holding capacity (WHC). The greater content of water inside the muscle structure reduces the light scattered by the muscle fibers, which makes the meat surface look darker [Hughes et al., 2019].

Myofibrillar structural darkness has a great impact on beef marketability. However, the most significant attribute in purchasing is meat redness [Venturini et al., 2014]. The intensity of redness is dependent on the presence, concentration and chemical state of the central pigment of beef tissue: myoglobin [Mancini and Hunt, 2005].

### Influencing factors on myoglobin concentration in dark–cutting development

Similar to the determination of the pHu value, Mb concentration may be affected by both intrinsic (biological) and extrinsic elements. Endogenous components comprise bovine genetics, animal age, testosterone concentration, muscle type, and fiber involved. A noteworthy environmental influence on beef Mb concentration is the type of cattle, which dictates the frequency and extent of physical exercise.

King et al. [2010] demonstrated that genetic plays a role in Mb content in longissimus thoracis muscles. Seven steer breeds were investigated: Angus, Charolais, Gelbvie, Hereford, Limousin, Red Angus, and Simmental. Simmental showed the greatest concentration of Mb (3.71 mg g–1), whereas Charolais and Limousin had significantly lower Mb values (2.77 and 2.72 mg g–1, respectively).

DeVol et al. [1985] demonstrated the effect of age on the darkening of longissimus dorsi muscles from bulls and steers from Limousin and Angus genotypes. The authors reported that the muscles of bulls were visually darker [p < 0.01] with 3.25 mg g–1 of Mb content in fresh muscle, whereas the muscles of steers had 2.90 mg g–1 of Mb [p < 0.01]. As animals get older, the amount of Mb increases but the affinity of O2 for Mb decreases; thus, they need to synthesize more Mb to stock O2. This explains why meat from older animals looks darker than meat from younger animals.

Although stress can significantly trigger the dark-cutting, the high proportion of oxidative fiber types within muscles is the main factor that determines the susceptibility of animals to triggering this effect [Ponnampalam et al., 2017]. Based on the activity of myosin ATPase, fiber types within muscles can be classified as red, intermediate, and white fibers [Voisinet et al., 1997].

Red fibers, involved in oxidative metabolism, have higher Mb contents and capillary blood flow than white ones, which maintain the O2 supply to cells. These two characteristics allow red fiber to oxidize glucose via
aerobic pathways, which is corroborated by the larger number of mitochondria and tricarboxylic acid enzymes in muscle cells (Choi and Kim, 2009). Moreover, the increased content of Mb in oxidative fiber has been correlated with higher cytochrome c oxidase expression, which is present in the mitochondria and discernibly impacts on color stability (Apaoblaza et al., 2020).

Intermediate fibers have an oxidative–glycolytic metabolism and are intermediate between red and white fibers which have a glycolytic metabolism (Choi and Kim, 2009; Zerouala and Stickland, 1991). In other words, it is the ratio of red and intermediate fibers to white fibers that matters: muscles with high oxidative metabolic activity, where there is a predominance of β and α-red fibers, have a high risk of dark-cutting because their fibers tend to have greater affinity with circulating adrenaline which depletes glycogen more easily with the same level of activity or stress compared to an animal with α–white muscle fibers (McGilchrist et al., 2012).

White fibers have a glycolytic metabolism and the least Mb concentration compared to the other two fibers because of the reduced O2 demand by cells. The lower abundance is related to the reduced stress undergone by glycolytic muscles, which are used for fast bursts of energy (Choi and Kim, 2009; Wicks et al., 2019).

In general, muscles with high oxidative metabolic activity, where there is more than 40 % of slow oxidative red fibers, such as psoas major muscles, exhibit higher Mb concentration (4.66 ± 0.31 mg g⁻¹), higher content of mitochondria and thus more intense respiration, and are characterized as muscles with less color stability. In contrast, muscles comprising more than 40 % of fast glycolytic white fibers, such as longissimus lumborum, have lower Mb concentration (3.97 ± 0.12 mg g⁻¹), lower concentration of mitochondria and thus high color stability (Kirchofer et al., 2002; Salim et al., 2019).

Endogenous testosterone and steroidal compounds, such as trenbolone acetate, supplemented in diet or implanted in the animal trigger the growth of muscle fibers (Johnson and Chung, 2007). Fink et al. (2018) reported larger fiber cross-sectional area in bull muscles than in neutered males and females. Moreover, testosterone may also stimulate the prenatal enlargement and proliferation of the muscle cells, although postnatal cell formation is inhibited (Johnson and Chung, 2007; Wicks et al., 2019).

Nevertheless, hypertrophic pressure enlarges preferentially type IIB glycolytic fibers rather than type I oxidative fibers. Therefore, anabolic hormones and their analogues tend to generate muscles proportionally more glycolytic and thus lighter because of the lower Mb content (Wicks et al., 2019). Vaughn et al. (2019) found the growth of the glycolytic muscle longissimus was related to the extent of the proliferative capacity of satellite cells, which are muscle stem cells that continue to induce postnatal hypertrophy in myofibrils. Therefore, the darker color attributable to non-castrated cattle is likely related to the high pHu-dependent change in muscle microstructure rather than higher intracellular Mb concentration.

The shifting in the muscle fibers stimulated towards greater proportion in response to pasture-finishing boosts intracellular Mb concentration as a consequence (Dunne et al., 2011; Picard and Gagaoua, 2020). This conversion is also stimulated by long-term physical exercise, as observed by Dunne et al. (2005) as well as by the grass-fed nutritional plane (Apaoblaza et al., 2020; Wicks et al. 2019). The resultant increase in Mb content is a response to the high demand of O2 by the mitochondrial oxidative metabolism. As observed by Apaoblaza et al. (2020), even glycolytic muscles such as longissimus dorsi, show greater Mb concentration after grass feeding compared to grain feeding, with significant darkening [lower instrumental lightness – L* by Commission Internationale de l’Eclairage (CIE) in the longissimus muscle].

Myoglobin structure and ligands

Structurally, Mb is composed of a polypeptide chain and a group of Fe²⁺ protoporphyrin called heme. Heme consists of a tetapyrrolic ring with conjugated double bonds, which binds coordinately to an Fe²⁺ [American Meat Science Association – AMSA, 2012]. The resonant electronic distribution of these conjugated bonds can absorb visible light and enhance the Mb color. Hematinic iron forms six-coordinate bonds. Four of these bonds have heme pyrrole nitrogen molecules in the same plane. The fifth coordination involves the interaction between iron and proximal histidine, located at position 93 [His93] (Møller and Skibsted, 2006). The sixth site of heme is available for reversible binding with a small ligand, such as O₂, nitric oxide (NO) and carbon monoxide (CO). Fifth and sixth bonds are perpendicular to the heme ring. Hematinic iron may bind to O₂ as a result of Mb’s hydrophobic pocket, which is proper to the heme location. Inside this pocket, the accessibility of the heme group to solvent is highly restricted, which protects iron from oxidation. Bound to iron, oxygen forms a hydrogen bond to distal histidine [His64], which improves its bond (Mancini and Hunt, 2005).

The three states of Mb present in the fresh muscle interchange simultaneously under natural conditions – in the retail store or at home, and the conversion takes place in dynamic equilibrium. A higher proportion of one pigment is dependent on intrinsic and environmental factors, such as meat pHu, MRA, lipid oxidation, gas composition, temperature, and microbial growth (Mancini and Hunt, 2005).

Biochemical basis of color stability

The superficial dynamic equilibrium between DMB, OMB, and MMB over a period of time defines muscle color stability, which can be impaired depending on
internal and external conditions favoring the prevalence of one form on the muscle surface [Bekhit and Faustman, 2005].

Functionality of mitochondria plays a critical role in the intensity and stability of color from the postmortem aging muscle [Ma et al., 2017]. Respiratory consumption of \( \text{O}_2 \) generates reactive oxygen species (ROS) as by-products. Lipids and proteins are the main targets and are involved in the oxidation of \( \text{Fe}^{2+} - \text{Mb} \) to \( \text{Fe}^{3+} - \text{Mb} \) and thus increase muscle MMb content [Tang et al., 2005]. Nevertheless, the mitochondria also possess the ability to convert MMb back to DMB [Bekhit and Faustman, 2005].

High OCR in the initial periods of the postmortem is deleterious to the development of the surface bright red color since mitochondrial respiration outcompetes Mb for \( \text{O}_2 \) and ultimately results in dark colored muscle [Suman et al., 2014]. Over time, there is an improvement in color development because of the decrease in OCR. The depletion of substrates for OC, such as lactate, succinate, and the reduced state of nicotinamide adenine dinucleotide [NADH], during storage time makes \( \text{O}_2 \) diffuse more rapidly through the tissue in order to bind to Mb, which improves color development and stability [MacDougall, 1982; Mancini and Ramanathan, 2014].

The improvement in the superficial color of meat aged in the short-term can be attributed to a decrease in the OCR of substrates and enzyme activity in the mitochondria [Ramanathan et al., 2013; Seyfert et al., 2006]. On the other hand, long-term aging reduces the potential for oxygenation (blooming) by decreasing mitochondria-mediated MRA, when aged beef is subsequently displayed under retail light conditions [Mancini and Ramanathan, 2014].

Cytochrome c oxidase (complex IV) has been reported to be essential to OCR [Grabež et al., 2015; Seyfert et al., 2006]. This enzyme is located on the inner mitochondrial membrane and comprises the electron–transport chain along with complexes I (NADH–Q reductase), II (succinate–Q reductase), and III (cytochrome Q–cytochrome c oxidoreductase) [Arihara et al., 1995].

Under oxidative condition, MMb generation occurs beneath the surface, between the superficial OMb and inner DMB layers. In the intermediate layer, the insufficient availability of \( \text{O}_2 \) to oxygenate all available DMB creates a suitable condition for ROS molecules to initiate \( \text{O}_2 \) oxidation, which can propagate this chain reaction [O’Keefe and Hood, 1982]. MMb can be formed precisely in the intermediate layer; the subsurface layer of MMb increases and moves towards the surface as the OMb layer on the surface becomes thinner and is replaced by the MMb layer [Mancini and Hunt, 2005].

Oxidation requires the prior deoxygenation of OMb so that MMb can be obtained. This precondition is due to the high resonant structure of OMb providing it with great oxidative stability, which makes OMb oxidation thermodynamically unfavorable [Faustman and Cassens, 1990]. Once formed, MMb can be endogenously reduced to DMB either by enzymatic or non–enzymatic reactions and favors maintenance of ferrous forms of Mb in meat [Faustman et al., 2010]. However, the enzymatic system is considered prominent.

Metmyoglobin reducing activity is an intrinsic cellular ability that will reduce MMb to DMB, which can be oxygenated to OMb [Ramanathan et al., 2013]. Because of this reversible characteristic, MRA seems to be critically important to maintaining or prolonging surface color stability. The increase in MMb proportion throughout retail time implies a decrease in MRA [Bekhit and Faustman, 2005].

Enzymatic MRA involves the NADH–cytochrome b5 MMb reductase enzyme and is based on the transference of two electrons from an NADH–coenzyme to a ferrocytochrome b5, thus reducing it to ferrocytochrome b5. As an intermediate, ferrocytochrome b5 non–enzymatically reduces MMb to DMB, regenerating the content of oxidized cytochrome b5 and improving meat color [Arihara et al., 1995; Bekhit and Faustman, 2005; Zhai et al., 2019].

In non–enzymatic MMb reduction, an electron from DMB is transferred to MMb by an artificial electron carrier, such as NADH or NAPDH in the presence of ethylene c acidiaminetetraacetic (EDTA), cytochrome c, methylene blue, ascorbate, uridine diphosphate sugar (UDP–sugar), vitamin E, psychrotrophic bacteria and other systems [Bekhit and Faustman, 2005]. In addition to the enzymatic process, non–enzymatic MMb reduction replenishes NADH through the added oxidized form of nicotinamide adenine dinucleotide (NAD+) and Krebs cycle intermediate substrates. However, non–enzymatic processes depend on the \( \text{O}_2 \) intracellular content, which is inhibited under vacuum conditions [MacDougall and Mancini, 2018].

Denzer et al. [2020] studied how these electron donors and carriers act in non–enzymatic MMb reduction in vitro with equine muscle MMb at meat pH (5.2, 5.6, 6.0, and 6.4) and storage temperatures (4 and 25 °C). The authors used ascorbate and nicotinamide adenine dinucleotide, in a reduced form (NADH) as electron donors and methylene blue and cytochrome c as cofactors. Methylene blue in the presence of NADH or ascorbate was more reduced at 4 °C than at 25 °C, with ascorbate and NADH. Higher pH increased methemoglobin reduction with ascorbate and cytochrome c. These non–enzymatic data corroborated the greater enzymatic MMb reduction in high pHu range.

Despite Denzer et al. [2020] obtaining results from equine muscle MMb, Bechtold et al. [2019] found that the non–enzymatic MMb reduction was higher in bovine samples than in equine counterparts.

Metmyoglobin reducing activity and NADH can be gradually depleted throughout storage/display time [Mancini and Hunt, 2005]. Kim et al. [2009] observed a pattern for the decrease in lactate dehydrogenase (LDH)
activity, NADH concentration and the percentage of MRA during the retail time of longissimus lumborum muscle packaged under fluorescent light for seven days covered with a polyvinylchloride (PVC) film. Furthermore, MMB may also be reduced via electrons from the mitochondrial electron–transport chain to form DMb [Belskie et al., 2015; Tang et al., 2005].

Canto et al. (2016) evaluated MRA in Nellore longissimus lumborum and psoas major packaged on polystyrene trays with O₂–permeable film and stored refrigerated for nine days. The authors observed a progressive drop during all retail periods for both muscles. Longissimus lumborum showed more MRA than psoas major throughout the storage period. Similarly, other authors have reported an inverse correlation between MRA and meat discoloration (Bekhit and Faustman, 2005; Ledward, 1985; Mancini et al., 2008; Reddy and Carpenter, 1991; Seyfert et al., 2006).

Higher MMB reductase activity may explain the greater color stability shown by certain muscles, such as longissimus lumborum compared to psoas major, as observed by Canto et al. (2016) and Salim et al. (2019).

Several beef muscles were sorted by Renerre (1990) according to their oxidation stability: longissimus dorsi, obliquus externus and tensor fasciae latae were the most stable muscles. semi–membranosus had intermediate stability, while gluteus medius, supra–spinatus, psoas major and diaphragma medialis were the most prone to oxidation. Renerre’s classification agrees with previous studies (Ledward, 1971; O’Keeffe and Hood, 1980–81).

To date, the relationship between MRA and discoloration is not consensual. Although Kim et al. (2009) found a decrease in MRA; these authors did not find an increase in surface MMB accumulation [%] in beef steaks of longissimus lumborum stored for seven days at 1 °C. According to Sammel et al. (2002), the lack of uniformity in the MRA methodologies could explain these contradictory findings.

**Color stability of dark–cutting beef**

Higher postmortem muscle pH is the most important factor that can prolong mitochondrial functionality (Ramanathan et al., 2019). Previous reports state that the mitochondria from dark–cutting beef has progressively greater O₂ consumption and higher level of MMB reductase activity as the pH of dark–cutting meat increases [McKeith et al., 2016; Wu et al., 2020].

High pH₄ carcasses are reported to have greater mitochondrial OCR due to a higher activity in the enzyme cytochrome c oxidase (complex IV), which oxidizes O₂ into water, resulting in less surface OMb and a darker color [Bendall and Taylor, 1972; Tang et al. 2005; Renerre, 1990]. According to Bendall and Taylor (1972), mitochondrial O₂ consumption is approximately 50 – 75 % faster at pH 7.2 than at pH 5.8. As a result, the conversion to the DMb redox form is favored at high pH₄, giving the meat a purplish color.

Muscle structure also contributes to color development by affecting O₂ influx through the tissue, facilitating interaction between gas and Mb (Mancini and Hunt, 2005). A closed structure observed in high pH₄ muscles makes it difficult to diffuse O₂ into the swollen muscle fibers due to the higher WHC [Hughes et al., 2014b; Mancini and Hunt, 2005]. In addition, glycolytic fibers often have a smaller diameter, which is detrimental to O₂ diffusion [Choi and Kim, 2009; Kirchofer et al., 2002].

The decrease in O₂ concentration within the myofibrils favors the formation of DMb and the color of fresh beef turns to dark purple [Ramanathan et al., 2019]. Since the O₂ depth depends on the O₂ concentration in the medium, oxygenation by means of air-exposition (approximately 20 % O₂) would not result in a thick OMB layer on dark–cutting beef surfaces.

In addition to the O₂ concentration, oxygenation also depends on the time and the temperature of exposition, blooming takes place more efficiently when meat is exposed to O₂ at 0 – 2 °C for, at least, 30 min. Higher temperatures increase mitochondrial activity and thus the O₂ uptake, which reduces the O₂ available to bind to Mb [Bendall and Taylor, 1972; Renerre, 1990].

The opposite phenomenon is observed in high pH₄ meat compared to MRA. These muscles show a protective effect on the oxidation of Mb. As a result, dark–cutting beef is resistant to forming the brownish layer of MMB over time [Lu et al., 2020]. In contrast to OCR, the protective effect is endogenous and as regards the content of substrates, such as NADH, the MMB reductase activity decays over time.

Therefore, at a certain point, the oxidative pressure by fluorescent illumination, the amount of ROS produced in mitochondrial respiration, or other factors will promote Mb oxidation that will be noticeable on the surface of the muscle [Bekhit and Faustman, 2005; Ramanathan et al., 2019; Renerre, 1990] which leads to lower discoloration over the display time because of the accumulation of MMB on the meat surface [Renerre, 1990]. In short, low O₂ tension, pH₄ up to 5.8, high temperature, concomitant lipid oxidation and loss of MMB–reducing activity are amongst the factors that influence iron oxidation [Bekhit and Faustman, 2005; Mancini and Hunt, 2005]. To prevent oxidation of DMb to MMB, residual O₂ within vacuum packaging should be avoided. At low O₂ concentrations (< 7 mm Hg) DMb is susceptible to oxidation by ROS, forming MMB. At O₂ concentrations > 7 mm Hg, O₂ competes with the peroxide radicals for the DMb, inhibiting the formation of MMB (AMSA, 2012).

The role of pH in the rate of MMB formation has been the focus of research for quite some time. Chemically, iron–catalyzed oxidation has been reported to be more active under acidic conditions [George and Stratmann, 1952]. Enzymatically, Echevarne et al. (1990) showed that MMB reductase activity increases as a function of the medium pH, achieving its maximum
activity at pH 7.3. Ramanathan et al. (2012) reported that a mitochondrial NADH–dependent reductase reduced more MMb in the control group at pH 7.4 than at pH 5.6 \( (p < 0.05) \). The unbalanced equilibria among DMB, OMB, and MMb in the high pH \( \text{pH}_u \) muscles interfere with the visual and instrumentally measured surface color.

Table 1 summarizes the data on the effect of high pH \( \text{pH}_u \) on color attributes (instrumental, visual and color stability).

The most used instrumental colorimetric system to characterize color or evaluate color changes is the Commission Internationale de l’Eclairage (CIE) L*a*b*, where L* measures lightness \([0: \text{black}, 100: \text{white}] \), a* measures redness \([–60: \text{green}, +60: \text{red}] \), and b* measures yellowness \([–60: \text{blue}, +60: \text{yellow}] \). Another system available is the HunterLab, such as the RGB (red, green and blue) models.

Redness (a*) is reduced at a higher pH \( \text{pH}_u \) due to a decreased accumulation in OMB on the muscle surface because of more intense cellular respiration (Bendall and Taylor, 1972; Tang et al., 2005). Page et al. (2001) found that muscle pH \( \text{pH}_u \) was more correlated with a* and b* values than with L* values. The authors stated that

### Table 1 – Summary of studies on the effects of high ultimate pH on meat color.

| Experimental material | High pH \( \text{pH}_u \) | Findings | Publication |
|-----------------------|--------------------------|----------|-------------|
| LT from Bos taurus (Parda Alpina (15) and Pirenaica (16) breeds) | > 6.1 | CIE L*, a*, b*, Chroma, and hue were affected by \( \text{pH}_u \) \((p \leq 0.001)\). CIE b* is the best discriminant for \( \text{pH}_u \) groups. \( \text{pH}_u \) also had an effect on reflectance spectra during aging \((0 \text{ min}, 5 \text{ h}, 2 \text{ d}, \text{ and } 9 \text{ d}) \). There was a dependence between \( \text{pH}_u \) and the spectrophotometric indexes \((K/614 – K/632, K/630 – K/580)\) and \( R/632 – R/614 \) \((p \leq 0.001)\). | Abril et al. (2001). |
| LL from steers (680), heifers (315) and bullocks (5). | ≥ 5.87 | LL Beef from 24/1000 carcasses were grouped into \( \text{pH}_u > 5.87 \), with 22/22 carcasses classified as “dark cutters”. Muscle \( \text{pH}_u \) had positive correlation with dark-cutting \((r = 0.80)\) and negative correlation with L* \((r = –0.46)\), a* \((r = –0.58)\), and b* \((r = –0.56)\). | Page et al. (2001). |
| LD from beef carcasses | ≥ 5.80 | High \( \text{pH}_u \) samples had lower sensory scores related to color than normal \( \text{pH}_u \) samples \((p \leq 0.001)\). | Vlijgen et al. (2002). |
| LT from Beef carcasses (47) | av. 6.06 | High \( \text{pH}_u \) carcasses presented LT with lower values of CIE L*, a*, and b* than normal \( \text{pH}_u \) \((p < 0.05)\). DFD LT and MST after 7 days postmortem \((\text{pH}_u > 5.80)\) also showed lower CIE L*, a*, and b* than those of normal \( \text{pH}_u \) \((p < 0.05)\). | Wulf et al. (2002). |
| LL from Hanwoo steers and bulls (24) | > 5.80 | Positive correlation between beef color (lean meat) and \( \text{pH}_u \) \((r = 0.77)\). According to CIE a*, normal \( \text{pH}_u \) muscles were redder than the high \( \text{pH}_u \) group. | Park et al. (2007). |
| LT from young bulls (106), bulls (96), cows (317), and heifers (95) | ≥ 5.80 | \( \text{pH}_u \) muscles had lower L*, a* (CIEL*a*b), R, G, B (RGB), V, and L values than normal \( \text{pH}_u \) muscles \((p < 0.05)\). Total heme pigment content did not increase in the high \( \text{pH}_u \) group. \( \text{pH}_u \) showed high correlation with L* \((r = –0.80)\), R \((r = 0.79)\), G \((r = –0.69)\), B \((r = –0.68)\), V \((r = –0.79)\), and L \((r = –0.77)\), and moderate correlation with a* \((r = –0.44)\). | Weglarz (2010). |
| MS from bulls (36) and heifers (24). 13 – 24 months-old. | ≥ 5.80 | Muscles with \( \text{pH}_u > 6.0 \) were darker than those of the normal \( \text{pH}_u \) group. | Chmiel et al. (2012). |
| LT from 179 dark-cutting beef carcasses | > 6.0 | Muscles with \( \text{pH}_u > 6.0 \) were darker than those of the normal \( \text{pH}_u \) group based on JMGA \((p < 0.05)\). Atypical DC muscles \((\text{pH}u < 6.0)\) were also darker than those of the normal \( \text{pH}_u \) group. The correlation \((R^2)\) between \( \text{pH}u \) values and JMGA scores was 0.59. | Holdstock et al. (2014). |
| LL from beef carcasses (10) aged during 62 d under vacuum (dark at 2 ºC) | av. 6.4 (all aging) | High \( \text{pH}_u \) muscles showed higher OCR and MRA, and lower CIE L* and surface OMB (during oxygenation period) for all aging times than those of the normal \( \text{pH}_u \) muscles. | English et al. (2016). |
| LL from steers and heifers. 9 – 30 months-old. | av. 6.1/6.4/6.6/6.9 | Animal selection based on dark-cutting, with subsequent grouping into four sub–groups, with average \( \text{pH}_u \) from 6.1 to 6.9. Instrumental color (L*, a*, b*, Chroma, and hue) and color stability parameters (IMF and bloomed OMB) decreased as \( \text{pH}_u \) increased \((p < 0.05)\). | McKeith et al. (2016). |
| LL from beef carcasses (9) | av. 6.65 | Non-enhanced dark-cutting muscles showed higher \( \text{pH}u \) than those of the normal \( \text{pH}_u \) group (USDA choice). High \( \text{pH}_u \) muscles (non-enhanced) were darker L* and visual color), less red and had lower hue and Chroma values than non-enhanced normal \( \text{pH}_u \) muscles on initial retail time. Color parameters were more stable during retail time for high \( \text{pH}_u \) than those for the normal \( \text{pH}_u \) group. | Stackhouse et al. (2016). |

Muscles: LL = longissimus lumborum; LT = longissimus thoracis; LD = longissimus dorsi; LTD = latissimus dorsi; RA = rectus abdominis; MSM = semimembranosus; MST = semitendinosus; \( \text{pH}_u \) = ultimate pH; DC = dark cutting; DFD = dry, firm, and dark meat; RGB = computer vision system \((\text{R} = \text{red}, \text{G} = \text{green}, \text{B} = \text{blue})\); V and L = value and lightness software models for color analysis; CIE L* (lightness); a* (redness); b* (yellowness); av. = average; K/S = ratio of absorptive (K) and scattering (S) color properties; \( R_\infty \) = reflectivity; OMB = oxymyoglobin; OCR = oxygen consumption rate; MRA = metmyoglobin reducing activity; IMF = initial metmyoglobin formed; JMGA = Japanese meat grading association; USDA = The United States Department of Agriculture.
\( \text{pH}_u \) affected the muscle color by changing the hue angle. Reduction in hue angle values moves color attributes closer to the \(+a^*\) axis (red) and farther from the \(+b^*\) axis (yellow). It was concluded that the closer the hue angle was to 0° and thus to \(b^* = 0\), the less the muscle was discolored. A number of authors have observed this decrease in angle and chroma as the \( \text{pH}_u \) increased (Mckeith et al., 2016; Stackhouse et al., 2016; Wglarz, 2010; Abril et al., 2001).

**Improving the color of dark-cutting beef**

Currently, vacuum-packaged meat is becoming more common because of its convenience for processors and consumers. However, vacuum-packaged muscles are dark and purplish in color due to the DMb layer on the surface, which can be easily oxygenated after adequate exposition to \(O_2\). A high DMb concentration is also observed in muscle immediately after cutting, since \(O_2\) penetration does not persist through the entire meat (Mancini and Hunt, 2005). The absence of DMb on a steak immediately after cutting is particularly useful for OCR analysis to measure \(O_2\) intake by mitochondria. The higher \(O_2\) partial pressure found in high oxygen modified atmosphere packaging (HiOx MAP) increases the depth of penetration of \(O_2\) through the muscle microstructure, which results in a thicker OMb layer on the muscle surface. Therefore, packaging dark-cutting beef in HiOx MAP is a strategy which can induce a bright red surface on the muscle despite the high \(\text{pH}_u\) (McMillin, 2008; Renerre, 1990). Moreover, HiOx retards uprising in the MMb layer on the muscle surface (Mancini and Hunt, 2005), whereas steaks packaged with \(O_2\) permeable films can maintain the cherry red color for hours or a few days. Thus, beef with HiOx MAP can be displayed for 6–10 days only (McMillin, 2008), because of microbial spoilage (Sun and Holley, 2012).

Another strategy for improving the quality of dark-cutting meat is the use of carbon monoxide modified atmosphere packaging (CO–MAP). Beef with high \(\text{pH}_u\) was shown to be lighter and redder in CO–MAP, but resulted in a higher surface MMb content than beef in HiOx MAP, which indicates that the relationship between MRA and MMb content in CO–MAP was different compared to that found in HiOx MAP (Zhang et al., 2018).

The use of carbon dioxide (CO\(_2\)) can inhibit the growth of food pathogens and Gram-negative aerobic bacteria, since they are more sensitive to CO\(_2\) compared to Gram-positive bacteria (Daniels et al., 1985). A higher proportion of CO\(_2\) can be used to package meat and prevent aerobic deterioration with an increase in microbial control (Sun and Holley, 2012).

Although the use of HiOx MAP increases consumer acceptance of meat color (O’Sullivan et al., 2015), this modified atmosphere has been associated with some loss in overall meat quality because of the presence of off-odors and off-flavors attributable to lipid oxidation (Jayasingh et al., 2002). Seyfert et al. (2006) also found an increase in thiobarbituric acid reactive substance (TBARS) values of beef packaged in HiOx MAP after storage and display.

According to Ramanathan et al. (2012) and Suman et al. (2014), beef Mb is greatly susceptible to nucleophile attack by ROS and aldehydes generated from peroxidation, such as 4-hydroxynonenal, which increases the proportion of MMb (Fe\(^{3+}\)) on the beef surface. However, fresh steaks from Nellore bulls showed extremely low TBARS values in both HiOx MAP and CO–MAP (Santos et al., 2016). Fiber IIB exhibit glycolytic metabolism and pasture-fed zebu steers (longissimus lumborum muscles), and contains natural antioxidants, which may explain these results (Canto et al., 2016).

Research indicates that the addition of various glycolytic and tricarboxylic acid metabolites, such as succinate, lactate, and malate can regenerate NADH through NADH–dependent reducing systems to enhance MMb reduction and the color stability of whole-muscle beef cuts (Table 2).

Furthermore, the ability of pyruvate and succinate to minimize lipid oxidation has been reported (Ramanathan et al., 2011). Kim et al. (2006) reported that color stabilization by lactate enhancement is related to MRA, by NADH replacement via LDH, reducing NAD+ both enzymatically and non-enzymatically. However, NADH can also promote mitochondrial \(O_2\) consumption resulting in darkened muscle because of the lowering of Mb oxygenation (Ramanathan et al., 2019). Lactate also results in color stabilization of cooked beef (Knock et al., 2006). In addition, the application of solutions containing lactate improve the meat’s juiciness, tenderness, taste, shelf life, and yield (Lawrence et al., 2003).

The addition of potassium lactate increased the color stability of steaks packaged in HiOx MAP (Kim et al., 2006). Additional research indicated the effect of lactate enhancement on beef steaks is packaging-dependent (Suman et al., 2009): lactate can be utilized for improving the color stability of beef steaks in HiOx MAP, but has no effect on the beef color stored in vacuum packaging and CO–MAP. Many advances have been made in the field of active packaging, such as the detection of microbial spoilage and safety within different MAP systems, to which dark-cutting has increased susceptibility (Holman et al., 2018). Sensors for real-time monitoring of beef freshness and quality can be used as an attractive and effective tool for assessing the microbial quality of packaged fresh meat (Kuswandi and Nurfaawaidi, 2017; Shukla et al., 2015).

**Color methods applied in dark-cutting beef fresh muscle**

Because of the difference in \(\text{pH}_u\), certain research methods for evaluating meat color in high \(\text{pH}_u\) beef samples require adaptations or a new approach to obtain...
The color of dark–cutting beef

Table 2 – Summary of results from studies that have examined the relationship between injection–enhanced beef with various Krebs cycle intermediates and the color stability.

| Results                                                                 | References                                      |
|------------------------------------------------------------------------|-------------------------------------------------|
| A combination of substrates relevant to mitochondrial oxygen consumption, improved meat color stability forms in aerobic and Bjelanovic et al. (2016). anaerobic packaging systems. |                                                 |
| An acceptable beef color resulted from the infusion of beef muscles with a phosphate and lactate blend. Erikson et al. (2018). |                                                 |
| Lactate and rosemary in beef enhancement solutions for improving strip loin steer color stability during display in modified atmosphere packages (HiOx MAP 80 % O2/20 % CO2). Mancini et al. (2005). |                                                 |
| Beef steaks enhanced with lactate, pyruvate, and succinate were less discolored than control steaks in PVC and high oxygen. Ramanathan et al. (2011a). |                                                 |
| Succinate had the greatest and pyruvate had the least metmyoglobin–reducing activity. |                                                 |
| Lactate–enhanced steaks had the least overall surface reflectance and the darkest surface color (lower L*). Ramanathan et al. (2010). |                                                 |
| Lactate can improve the color stability of lamb, possibly by increasing both oxygen consumption and metmyoglobin reducing activity. Ramanathan et al. (2011b). |                                                 |
| Effects of lactate on myoglobin are temperature and pH dependent. Suman et al. (2014). |                                                 |
| Calcium lactate/phosphate enhancement have beneficial effects on lipid stability, surface color, and sensory attributes of beef round cuts under HiOx MAP. Cruzen et al. (2015). |                                                 |
| The glutamate, succinate and citrate combinations acted as pro–oxidants that promoted lipid oxidation in minced beef under both high and low oxygen conditions. Yi et al. (2015). |                                                 |
| Lactate enhancement on beef steaks is packaging–dependent. Suman et al. (2010). |                                                 |
| Enhancing beef with lactate replenishes NADH via increased LDH activity, ultimately resulting in greater meat color stability. Kim et al. (2006). |                                                 |

PVC = polyvinylchloride; NADH = reduced form of nicotinamide adenine dinucleotide; LDH = lactate dehydrogenase.

reliable data. Herein, we discussed a number of methods employed in color stability research and compare them with standard methods applicable to normal pHu beef muscles. It is important to note that these methods are in–line with the Meat Color Measurement Guidelines issued by AMSA (2012): Mb quantification, MRA, and OCR.

There are two protocols for quantifying total Mb of fresh and cooked meat in AMSA’s guidelines (AMSA, 2012): the first based on the isobestic spectrophotometric point and the second, by reducing all Mb forms to DMB. Both strategies can be used for meat with high pHu. However, the extraction of the pigment with neutral buffer (40 mM potassium phosphate, pH 6.8) is hampered in high pH meat, which results in incomplete total pigment extraction and, thus, in a pink color remaining in the centrifuge pellet after one extraction. In order to improve the pigment extraction, an acidic buffer adapted from Poel’s Mb quantification method can be deployed (DeDuve, 1948; Hunt and Hedrick, 1977). The use of an acidic buffer [0.01 N, Hunt and Hedrick (1977), or 800 mM, McKeith et al. (2016), sodium acetate, pH 4.5] compensates for the native higher pH of the muscle, also observed in pre–rigor muscle (abdominal muscle from biopsy) as shown by DeDuve (1948). The repetition of the extraction step ensures the removal of Mb from the tissue, as followed by Hunt and Hedrick (1977) and McKeith et al. (2016).

The MRA method is based on the complete oxidation of Mb in sodium nitrite solution [0.3 % for 20 min at 25 °C] with subsequent Mb reduction under vacuum conditions at 20 – 30 °C for 2 h. The estimation of the proportion of each Mb redox form is achieved by scanning samples with a Hunter Miniscan colorimeter with settings previously described that had been calibrated through the O2 impermeable film of a vacuum bag. Complete oxidation may be estimated by the ratio of specific wavelengths [572 nm/525 nm], being the proportion of each Mb redox form confirmed by the 630 nm/580 nm ratio.

Another protocol for estimating the MMb redox form on the surface of the samples is based on the creation of reference standards for 100 % MMb and DMb. Thus, the proportion of surface MMb is obtained by dividing the difference between MMb for 100 % DMb and for samples by the difference between MMb for 100 % DMb and 100 % MMb. However, Mb does not oxidize completely in pHu meat > 5.8, and does not result in 100 % MMb samples prior to vacuum–reduction. There is no method in the literature for achieving total Mb oxidation. Therefore, the initial MMb formation is used by a number of authors, such as McKeith et al. (2016). The initial MMb formation represents the proportion of surface MMb formed after Mb oxidation in sodium nitrite solution.

Instead of forcing Mb oxidation/reduction, as seen in MRA analysis, OCR methodology is based on the assessment of the Mb oxygenation/deoxygenation stimulated. In OCR protocol, described by Madhavi and Carpenter (1993), muscle samples are air–oxygenated for 2 h at 2 °C before vacuum–packaging (reading 1) and subsequent incubation for 20 – 30 min at 25 – 30 °C (reading 2). Two readings assess the surface OMb proportion in order to evaluate mitochondrial O2 consumption during thermal incubation.

McKeith et al. (2016) observed that performing OCR in high pHu, beef dark–cutting samples had lower initial blooming (before incubation) than normal pHu samples, failing to achieve 100 % of OMb on the surface.
of the muscle – as was also observed by Krzywicki (1979). Therefore, similar to the behavior of MRA in high pHu samples, the evaluation of the proportion of the deoxygenated OMb may not be reached. Therefore, the use of the proportion of OMb formed during the blooming step is an interesting alternative for comparing OCR between samples.

Calculating MMb, OMb, and DMb proportions on the beef surface has been conducted by Krzywicki (1979). However, assessment of the Mb states in high pHu muscle is challenging when disabling a full conversion of these forms, further to this pHu range changing muscle structure and WHC. English et al. [2016] estimated DC beef pigments by their reflectance values and demonstrated that using the K/S ratios at isobestic points were useful. K/S improves pigment proportion quantification by making data more linear accounting for absorptive (K) and scattering (S) color properties; and its formula is \(1-R^2/(2R)\), where R is the reflectance obtained using a spectrophotometer [AMSA, 2012].

Although Ramanathan et al. [2010] have not worked with high pHu, the authors showed that lactate-enhancement in beef had changed the overall percentage reflectance. Nonetheless, 525 nm remained the isobestic point for MMb, OMb, and DMb, while 572, 610, and 473 nm also remained isobestic for MMb, OMb, and DMb, respectively. Therefore, these wavelengths are still useful for calculating surface pigment in high pHu beef using the following formulae: K/S 572 ÷ K/S 525 (MMb), K/S 610 ÷ K/S 525 (OMb), and K/S 473 ÷ K/S 525 (DMb) [AMSA, 2012].

Novelties in meat color studies of dark-cutting beef

Molecular profiling techniques including metabolomic and proteomic, have been increasingly applied as targeted or non–targeted approaches to exploring biochemical changes in postmortem muscle and its influence on meat quality characteristics [Ma et al., 2020]. Mass spectrometry–based metabolomic and proteomic studies have been carried out to evaluate modifications in the metabolite and protein profile as proteins and metabolites are directly involved in molecular mechanisms related to color stability, lipid oxidation, WHC and tenderness in fresh meats [Kim et al., 2016; Li et al., 2018; Ma et al., 2017; Subbaraj et al., 2016].

On the subject of meat color, a proteomic study indicated that the sarcoplasmic proteome of color–stable longissimus lumborum beef muscle has higher levels of soluble antioxidant proteins (thioredoxin, peroxiredoxin–2, and peptide methionine sulfoxide reductase) and chaperones (heat–shock protein–27 kDa) compared to the color–labile psoas major muscle [Joseph et al., 2012]. In turn, Mato et al. (2019) assessed phosphoproteomic differences between dark–cutting and normal beef in response to pre–slaughter stress. These authors found that protein phosphorylation levels were three times higher in dark–cutting beef compared to normal beef. This effect was mainly observed in proteins with biological functions related to structural–contractile properties, actin polymerization, stress response, metabolism and electron transport chain.

As postmortem glycolysis and pH decline in muscle are associated with protein phosphorylation, a previous gel–based phosphoproteomic study hypothesized that phosphorylation modifies the Mb structure and its susceptibility to oxidation, thereby influencing meat color stability [Li et al., 2018]. This study identified that the phosphorylation of color stability–related proteins regulates the activity of glycolytic enzymes, thus influencing meat discoloration.

In metabolomic studies, Ma et al. [2017] stated that metabolic pathways influencing both color and lipid oxidative stability in beef are dependent on the muscle type as well as the postmortem aging period. In fact, psoas major muscle was more susceptible to discoloration, lower free radical scavenging activity, higher non–heme iron content and lipid oxidation compared to semimembranosus and longissimus lumborum muscles, which were more stable. These authors reported that metabolites such as the NAD/NADH ratio, acyl carnitines, free amino acids, nucleotides, nucleosides, and glucuronides play an important role in the oxidative stabilization of beef muscles. These metabolites can be potential biomarkers for further validation studies.

To identify differentially abundant metabolites related to color stability from longissimus lumborum and psoas major muscle, a gas chromatography–mass spectrometry (GC–MS) based non–targeted metabolomic approach was used by Abraham et al. (2017). The Longissimus lumborum muscle had higher levels of pyruvic acid, glucose 6–phosphate, fructose and citric acid compared to psoas major muscle. Additionally, key regulatory metabolites can increase MRA and mitochondrial activity. The malonic acid levels were higher in psoas major muscle when compared to the values found for longissimus lumborum. In fact, malonic acid is a complex II inhibitor recognized by promoting NADH oxidation, thus negatively affecting meat color [Abraham et al., 2017; Ramanathan et al., 2019].

In loins obtained from lamb carcasses and exposed to different storage conditions and display times, the metabolite profile was studied by a hydrophilic interaction liquid chromatography–mass spectrometry (HILIC–MS)–based metabolomic approach. The study identified metabolites including malic acid, NADH, and guanosine levels as being significantly higher in color–stable samples than in color–labile samples during aging [Subbaraj et al., 2016].

Moreover, as regards the differences in MRA and OCR, the influence of sarcoplasmic proteome and metabolome on the differential color stability of beef
has been recently reported [Ramanathan et al., 2019; Mancini et al., 2018]. These results indicated that muscle-specific differences in mitochondrial activity may partially contribute to variations in the color stability of longissimus lumborum and psoas major beef muscles.

**Final Remarks**

Dark-cutting beef has been cataloged as a multifactorial phenomenon dependent on ante and postmortem factors. Although not the exclusive factor, high pHu was the major influencing element to produce dark-cuts and their color stability, even though there was, at the time, no definitive cut-off pH for determining a carcass as dark-cutting beef. Animal genotype was observed as being one factor among several others that affect pHu drop and beef color stability. However, there was little information available regarding an important breed in Brazilian herds: Nellore bulls and their crossbreeds.

The significant color stability found in the muscles with high pHus was outshone by the darker appearance which, closely associated with the increased mitochondrial OCR, reduces beef marketability. Post-slaughter interventions, such as improvements in the use of organic acid salts through multi-needle injection followed by HiOx MAP either limited the oxidation of Fe2+-Mb or promoted reductions in M Mb.

Future studies should focus on the use of accurate techniques to quantify tissue OCR and M RA and their relationships to color stability of fresh dark-cutting meat, such as the –omics techniques, especially in Nellore bulls in order to substantiate the effect of animal phenotype on beef color.

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