Delayed carcass chilling improves tenderness of the beef *gluteus medius* muscle

**Abstract** – The objective of this work was to evaluate the effects of a slower chilling rate on the physical traits of 14-day aged top sirloin (*gluteus medius*) from F1 Angus x Nellore young bulls. Slight and lean beef carcasses (n = 30) were split in half and selected for control (2°C for 24 hours) and treatment (10°C for 10 hours followed by 2°C for 14 hours). Carcass temperature and pH decline were monitored in the *longissimus dorsi* (LD). Sarcomere length, color, and instrumental tenderness (Warner-Bratzler shear force and Volodkevich bite jaws) were measured on the *gluteus medius* (GM). pH 6.0 was reached when the LD temperature was very low, 3°C for the control and 6°C for the treatment. Differences in color and cooking losses were not significant. A trend was noticed for shorter sarcomere length and greater Warner-Bratzler shear force in the GM control. However, a lower value for Volodkevich bite jaws was found in the GM subjected to the treatment. The cooling regimes adopted were not sufficiently different to cause changes in color traits; however, the slow chilling of carcass improved tenderness and can be a good alternative to produce high-value cuts.

**Index terms**: beef carcass, chilling rates, sarcomere length, top sirloin, Volodkevich.
Introduction

Consumer expectations of acceptable beef are one of the most crucial factors affecting the meat supply chain. When buying meat, they take into account color, amount of visible fat, price, and cut, and, when consuming it, tenderness, flavor, and juiciness (Robbins et al., 2003).

In fully developed meat markets, the beef top sirloin, known as “alcatra” in Brazil, is a cut of highly acceptable appearance, which is suitable for grilling, roasting or frying. However, it is rarely the first option in the retail market or restaurants, and it is usually sold as a cost-effective alternative. According to some authors, this cut, which is mainly formed by the gluteus medius (GM), has problems related to the consistency of palatability traits, as tenderness and juiciness (McKeith et al., 1981; Wheeler et al., 1990; Neely et al., 1998; Savell et al., 1999).

In postmortem muscles, the degree of rigor shortening and toughening depends largely on cooling temperature. Cold shortening, the rapid chilling of beef carcasses immediately after slaughter, has been intensively used by the meat industry due to its favorable influence on meat weight loss and storage life (James & Bailey, 1986); however, it has negative impacts on the quality of the consumed meat, becoming more severe with decreasing temperatures (Locker & Hagyard, 1963). Simmons et al. (2008) suggested that toughness by cold shortening could be related to a delayed aging due to fast chilling.

The effects of early-postmortem chilling rate on beef quality have been shown in several research works, particularly regarding the longissimus dorsi (Lochner et al., 1980; Marsh et al., 1981; Koh et al., 1987; Olsson et al., 1994; Rhee & Kim, 2001; Sørheim et al., 2001; Hannula & Puolanne, 2004; Yu et al., 2008; Wolcott et al., 2009). This type of information, however, is not easily available for the GM. Stolowski et al. (2006) reported that temperature and pH decline were muscle dependent and that, of the evaluated beef muscles, the GM, electrically stimulated or not, had the fastest rate of pH decline and highest temperatures during chilling, making it more tender than the others. This seems to be the case for heavier and fatter top sirloins, highly demanded by the “churrascarias” (barbecue-style restaurants) in Brazil, but not for lighter and leaner cuts, currently more frequent.

The obtained results are indicative that cooling carcasses too slowly or too quickly can damage meat quality. Therefore, special care must be taken to ensure that the cooling process be favorable to all aspects regarding quality and safety in meat production (Braden, 2013).

The objective of this work was to evaluate the effects of a slower chilling rate on the physical traits of 14-day aged top sirloin from F1 Angus x Nellore young bulls.

Materials and Methods

Thirty F1 Angus x Nellore young bulls of approximately 13 months of age and 440 kg live weight were selected for this experiment out of 200 heads of cattle that had been raised on a feedlot for 107 days. The animals were slaughtered at a commercial beef plant in the municipality of Lins, in the state of São Paulo, Brazil. Pre-slaughtering handling was carried out according to good animal welfare practices, and slaughtering procedures followed the regulation for sanitary and industrial inspection of products of animal origin (Brasil, 1997). All carcasses were visually classified for fatness and weighed on the kill floor after kidney knob, channel, and inguinal fat removal. Average hot carcass weight was 240 kg, and fatness was slight (1–3-mm fat on the twelfth rib). Half carcasses were used: the right sides, as a control subjected to conventional chilling; and the left ones, for a treatment with a distinct chilling rate. The half carcasses were moved into a chilling room within 30 min of slaughter, and this time was established as the 0 hour postmortem.

Control half carcasses were chilled according to the system used in modern packing plants, where a cooler is programmed to maintain constant air temperature at 2°C for 24 hours. For the treatment half carcasses, another cooler was programmed for an initial temperature of 10°C for the first 10 hours, and then the temperature was adjusted to 2°C for 14 hours.

Half carcass temperature and pH were manually monitored in the longissimus dorsi (LD), at the last two ribs, using the LoT406-M6-DXXK-57/25 portable pH meter (Mettler Toledo Ind. e Com. Ltda., Barueri, SP, Brazil) with electrode and temperature probe, which was inserted approximately 5 cm deep, at 4, 7, 9, and 11 hours postmortem.
pH and temperature were determined in the Longissimus muscle because these traits are always measured in this position in the meat industry. However, since, in the present study, the aim was to evaluate the quality of another muscle, the Longissimus was considered as the reference to monitor pH and temperature decline.

At 24 hours postmortem, the boneless top sirloin (MLA, 2003) of each half carcass was completely trimmed, skinned, and vacuum packaged in high-barrier Cryovac bags (Sealed Air: Food Care, Charlotte, NC, USA). These cuts were aged for 14 days, frozen at -18°C, and transported to the meat laboratory at Universidade Estadual de Campinas for analysis.

Later, meat cuts were thawed at 4°C, and divided in two portions of 11 cm. Measurements of pH, sarcomere length, instrumental color, and shear force were taken. A total of 3 g muscle were homogenized in 20 mL distilled water for 15 s to determine pH, which was obtained three times using the DM-22 pH meter (Digmed: Analytical Instrumentation, São Paulo, SP, Brazil) with a combined glass electrode.

Color was measured using the L*, a*, and b* color space (also referred to as the CIELAB color space) in three positions in the middle of each GM half carcass, after a 30 min bloom time at 4°C, using the CM-508d portable spectrophotometer (Konica Minolta Sensing Americas, Ramsey, NJ, USA) and the MiniScan XE (Hunter Associates Laboratory, Inc., Reston, VA, USA) with the illuminant D-65, 10ºC standard observer angle, and an aperture size of 3.5 cm. The unit was calibrated using a black and a white standard plate, as specified by International Commission on Illumination (CIE, 1986).

GM sarcomere length was determined by the neon-laser diffraction method described by Cross et al. (1981), modified to evaluate a fresh piece of meat (0.5 x 0.5 cm x 100 µm) removed parallel to the orientation of the muscle fibers. This piece was placed on microscope slides with cover slips and exposed to the 155SL helium-neon laser beam (Spectra-Physics, Santa Clara, CA, USA) of 632.8 nm. The lengths of ten diffraction patterns from each sample were measured, and sarcomere length was determined by averaging these measurements.

Each middle sample of the GM was weighed (approximately 400 g), and then six samples were cooked at a time in a Série 8 – 4000 W conventional oven (Imeque, Alto da Lapa, SP, Brazil), equipped with upper and lower electrical resistances. The oven was preheated with a thermostat adjusted to 170°C, and each roast was placed on a metal rack over an aluminum tray. Thermocouples were used to individually monitor the internal temperatures of the roasts, which were removed from the oven when they reached the internal temperature of 71°C (AMSA, 1995).

After cooking, the roasts were weighed to calculate cooking losses and then were kept at room temperature for 2 hours, packed in plastic bags, and chilled at 4°C overnight. Cooked samples were used to measure instrumental tenderness by Warner-Bratzler shear force (WBSF) and Volodkevich bite jaws.

Three 1.27-cm cores were taken from two 2.5-cm thick slices, and three 1x1-cm parallelepipeds were cut parallel to the muscle fibers. Each core and sample were sheared once in the perpendicular direction of the muscle fibers using the TA-XT2i texture meter (Texture Technologies Corp. and Stable Micro Systems, Ltd. Hamilton, MA, USA) equipped with a 1.06-mm thick Warner-Bratzler blade or Volodkevich bite jaws, respectively.

Data, including basic statistics and tables, were analyzed using the Statistica, version 7.0, software (TIBCO Software Inc., Palo Alto, CA, USA), in order to determine means, standard errors, correlation coefficients, and t-test for dependent samples to compare treatment means. Results were considered significant at 5% probability.

Results and Discussion

In the LD, temperature showed a fast drop in the first 4 hours of chilling, for both control and treatment half carcasses, reaching 14.4°C. At 7 hours of chilling, the temperature of the control was already 8°C; however, that of the treatment half carcasses, reached 11°C. At 9 hours, the temperature of the control was 6°C, but it remained the same as at 7 hours for the treatment half carcasses. At 11 hours postmortem, temperature was already below 4°C and a little above 6°C in the control and treatment half carcasses, respectively. Therefore, control and treatment means differed (p<0.05) at 7, 9, and 11 hours (Figure 1). Although the muscle chilling rate often influences the glycolytic rate (Pike et al., 1993; Yu et al., 2008), this was not observed in the
present study since differences in LD temperature were slight.

The pH means of 6.62, 6.29, 6.23, and 6.04 were not different (p>0.05) at 4, 7, 9, and 11 hours postmortem, respectively (Figure 1). Prado & Felício (2010) also found no difference in the pH decline curve when studying conventional and slow-air and spray chilling. However, Mohrhauser et al. (2014) reported that delayed chilling caused a faster pH decline, resulting in lower pH values at 6, 12, and 24 hours. The same trend was verified by Li et al. (2012).

Measurements of pH at 48 hours showed no difference between the control and treatment half carcasses (p>0.05; 5.54). This is in alignment with other researchers who found no significant differences in pH values at 24 or 48 hours due to cooling rates (Koh et al., 1987; White et al., 2006; Kahraman et al., 2011, 2014; Li et al., 2012).

pH 6.0 was reached at 11 hours postmortem when the LD temperature was below 7 and 4°C, for the treatment and control half carcasses, respectively, which can be interpreted as a fall into the cold shortening zone of the superficial muscles of both carcass sides (Locker & Hagyard, 1963; Bendall, 1973; Olsson et al., 1994).

No significant differences were observed between the control and treatment means in the GM regarding L*, a*, b*, and Chroma values (Table 1), which is in agreement with the results of Yu et al. (2008) and Mao et al. (2012) for the longissimus lumborum (LL), but differs from those of Farouk & Lovatt (2000) for the Semitendinosus and Biceps femoris, and of Janz et al. (2000) for the LL, who reported higher L*, a*, and b* values with increasing rigor temperature. Young et al. (1999) concluded that beef going into rigor at 9°C had the best bloomed color at 2 and 8 weeks of aging; however, in the present study, no color difference was detected at 14 days of aging. The obtained results suggest that the differences between the two cooling regimes evaluated here may have not been great enough to cause any effects on instrumental color.

There were no significant differences regarding the means of sarcomere length for the GM control (1.61±0.10 µm) and treatment (1.65±0.07 µm) (Table 2). The average sarcomere length of 1.63 µm for the GM in the present study was similar to that of 1.66 µm reported by Rhee et al. (2004), but notably shorter than those of 1.90, 1.91, and 1.77 µm, respectively, obtained by McKeith et al. (1985), Harris et al. (1992), and

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**Figure 1.** Temperature and pH decline of the control and treatment of Longissimus muscle half carcasses during chilling. Control, constant air temperature of 2°C for 24 hours; and Treatment, initial temperature of 10°C for the first 10 hours followed by 2°C for 14 hours.
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Stolowski et al. (2006), and longer than those of 1.50 and 1.58 µm by King et al. (2009a, 2009b).

The means found for sarcomere length in the present study are indicative that the GM, subjected to both the treatment and control, likely suffered cold shortening, since it was much shorter than those described by McKeith et al. (1985), Harris et al. (1992), and Stolowski et al. (2006). Moreover, the chilling rate was too fast, because, at 11 hours postmortem, the LD temperature was below 7°C, which can be interpreted as a fall into the cold shortening zone for superficial muscles, as the GM (Locker & Hagyard, 1963; Bendall, 1973; Olsson et al., 1994).

Rhee et al. (2004) reported an average range of 1.3 µm (1.6–2.9 µm) for sarcomere length between 11 muscles. The Psoas major had the longest sarcomere length, and the GM the shortest. According to these authors, there is a high degree of variation in sarcomere length both among and within GM steaks. However, Hostetler et al. (1972) found a mean sarcomere length of 2.0 µm for the GM in a condition of supposedly high fatness (only shown in pictures), obtaining sarcomere measurements of ≥ 1.9 µm for all ten studied muscles.

It should be noted that sarcomere shortening can be reduced by changing the way the carcass is suspended. Hip suspension or tenderstretch results in a natural position of the muscles and keeps sarcomeres longer, which improves meat tenderness (Ahnström et al., 2012).

The results for cooking losses, Volodkevich bite jaws, and WBSF for the GM are shown in Table 2. There were no differences for means of cooking losses between cooling regimes, but there was a tendency for greater (p=0.08) WBSF in the control (53.32±14.73 N) than in the treatment (49.03±9.29 N). Rhee et al. (2004) reported WBSF mean values of 42.22 to 44.59 N, depending on the location within the GM. In addition, Honikel (2009) concluded that muscle pH at 24 hours postmortem can affect losses during cooking.

In this experiment, means of cooking losses were higher in both control and treatment half carcasses (29.70±2.94 and 28.94±2.08%), compared with the mean of 23.6% obtained by Rhee et al. (2004) for the same trait and muscle. However, Luchak et al. (1998) found a very similar mean of 29.9%, when cooking steak at 74°C internal temperature, and George-Evins et al. (2004) reported 31% cooking losses.

When measured by WBSF, no difference was observed for tenderness of the GM from carcasses that were slowly chilled (49.03 N), compared with the control ones (53.32 N). However, when determined by Volodkevich bite jaws, the treatment GM (43.1 N) was more tender than that the control one (46.8 N). According to Pomponio & Ertbjerg (2012), calpastatin activity is lower at higher temperatures, which can contribute to increase tenderness of beef subjected to slow chilling processes.

In the present study, the GM did not show good enough WBSF values, with an average of 51.2 N, probably due to the short sarcomere lengths in both the control and treatment half carcasses. Trying to investigate the impacts of shorter sarcomeres on GM tenderness, Hostetler et al. (1972) observed that WBSF, converted to Newton, was 48.02 N when the carcass was hung by the Achilles tendon, but 36.26 N when

### Table 1. Effect of cooling regime on the mean values of top sirloin (gluteus medius) instrumental color.

<table>
<thead>
<tr>
<th>Instrumental color</th>
<th>Control (n = 30)</th>
<th>Treatment (n = 30)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lightness</td>
<td>34.18 2.65</td>
<td>34.24 2.89</td>
<td>0.83</td>
</tr>
<tr>
<td>Redness</td>
<td>18.01 1.34</td>
<td>18.19 0.97</td>
<td>0.34</td>
</tr>
<tr>
<td>Yellowness</td>
<td>17.67 1.40</td>
<td>17.80 1.21</td>
<td>0.46</td>
</tr>
<tr>
<td>Chroma</td>
<td>25.24 1.83</td>
<td>25.45 1.46</td>
<td>0.40</td>
</tr>
</tbody>
</table>

1Control, constant air temperature of 2°C for 24 hours. 2Treatment, initial temperature of 10°C for the first 10 hours followed by 2°C for 14 hours. SD, standard deviation. 3Statistical comparisons were performed using Student’s t-test, at 5% probability.

### Table 2. Effect of cooling regime on the mean values of top sirloin (gluteus medius) ultimate pH, sarcomere length, cooking loss, and instrumental tenderness analysis by Warner-Bratzler shear force (WBSF) and Volodkevich bite jaws (VBJ).

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Control (n = 30)</th>
<th>Treatment (n = 30)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sarcomere (µm)</td>
<td>1.61 0.10</td>
<td>1.65 0.07</td>
<td>0.08</td>
</tr>
<tr>
<td>pH (48 hours)</td>
<td>5.53 0.12</td>
<td>5.56 0.12</td>
<td>0.10</td>
</tr>
<tr>
<td>Cooking loss (%)</td>
<td>29.70 2.94</td>
<td>28.94 2.08</td>
<td>0.17</td>
</tr>
<tr>
<td>WBSF (N)</td>
<td>53.32 14.74</td>
<td>49.03 9.29</td>
<td>0.08</td>
</tr>
<tr>
<td>VBJ (N)</td>
<td>46.8 13.2</td>
<td>43.1 12.1</td>
<td>0.02</td>
</tr>
</tbody>
</table>

1Control, constant air temperature of 2°C for 24 hours. 2Treatment, initial temperature of 10°C for the first 10 hours followed by 2°C for 14 hours. SD, standard deviation. 3Statistical comparisons were performed using Student’s t-test, at 5% probability.
hung by the pelvic bone due to muscle stretching, which increased sarcomere length from 2.0 to 2.8 μm. 

Derbyshire et al. (2007) found that sarcomere lengths were significantly increased by hip suspension, but were not affected by electrical stimulation. However, both of these treatments were able to improve meat tenderness, showing that other traits, besides sarcomere length, can affect tenderness.

McKeith et al. (1985), aiming to prevent cold shortening at pH 5.5 and 15.5°C, obtained 1.90 μm for sarcomere length and 34.10 N for WBSF for the GM. Harris et al. (1992) reported similar sarcomere length of 1.91 μm and WBSF of 41.25 N. According to George-Evins et al. (2004) and Gruber et al. (2006), the WBSF of the GM can improve the quality grade (carcass fatness) used to rank meat by United States Department of Agriculture; for example, for 14 day-aged GM, these authors found values of 44.39 and 53.99 N for Choice and Select grades, respectively. In Brazil, a current commercial practice is to select top sirloin cuts out of heavy and fat carcasses for the “churrascarias”.

**Conclusion**

The slower cooling conditions in the first 10 hours postmortem do not affect top sirloin (*gluteus medius*) color, but are effective for improvement of instrumental tenderness, adding more value to this cut.

**Acknowledgments**

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