



The effect of extended post-mortem ageing on the Warner–Brazler shear force of *longissimus thoracis* from beef heifers from two sire breeds, slaughtered at 20 or 25 mo of age

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Abstract

The effects on tenderness of extended ageing of *longissimus thoracis* (LT, striploin) muscle that differed in structure and composition were examined. Spring-born Angus × Holstein-Friesian heifers ($n = 48$) and Belgian Blue × Holstein-Friesian heifers ($n = 48$) were slaughtered, within sire breed, at 20 or 25 mo of age. Approximately 48 h post-mortem, LT steaks (2.5 cm) were removed, and either stored at -20°C for chemical analysis or vacuum-packed, stored at 2°C for 7, 14 or 28 d post-mortem and then at -20°C pending Warner–Brazler shear force (WBSF) analysis. Muscle from Angus-sired heifers had higher ($P < 0.001$) intramuscular fat (IMF) concentration, lower ($P < 0.001$) proportion of type IIX muscle fibres and higher ($P < 0.001$) proportion of type IIA and type I muscle fibres compared to muscle from Belgian Blue-sired heifers. Collagen characteristics did not differ between sire breeds. Later slaughter increased ($P < 0.001$) IMF concentration and decreased ($P < 0.001$) total and insoluble concentrations and collagen solubility. There were no interactions between the main effects for WBSF and no difference between sire breeds. Later slaughter and increasing the duration of ageing decreased ($P < 0.05$) WBSF. Based on threshold WBSF values in the literature, all samples would be considered tender ($<39\text{ N}$) after 7 d ageing. Untrained consumers are likely to detect the decrease in WBSF from 7 to 14 d ageing but not due to further ageing. Within the production system examined and based on WBSF data, extending LT ageing to 28 d is not necessary to ensure consumer satisfaction.

Keywords

Beef • breed • instrumental texture • wet ageing

Introduction

Tenderness is generally considered the major influence on consumer satisfaction when eating beef (Miller *et al.*, 2001). Post-mortem proteolysis results in the degradation of myofibrillar and cytoskeletal proteins which enhances beef tenderness (Ouali *et al.*, 2006). This process, termed maturation or ageing, most commonly involves removal of individual muscles from the carcass, vacuum-packaging (wet ageing) and storage at $2\text{--}4^{\circ}\text{C}$. While it is accepted that the time taken to reach a final tenderness endpoint is muscle dependent and that the rate of tenderisation decreases as ageing proceeds, the influence of production factors on tenderness evolution is unclear. Nevertheless, an ageing period is included in many “quality” beef specifications. In the United Kingdom (UK), to comply with a quality standard specification, “primals used for frying, roasting and grilling must be subject to a minimum 14 d maturation (from slaughter to the final consumer)” (EBLEX,

2012). In the United States (US), the earlier recommendation was for *longissimus thoracis* (LT) (commercial name = striploin) to be aged for at least 14 d (Mies *et al.*, 1998). More recently, Gruber *et al.* (2006) concluded that post-mortem ageing should be managed with respect to US Department of Agriculture carcass grade with striploins from carcasses that graded “upper two-thirds Choice” achieving 95% of the ageing response in 14 d compared to 66% for striploins from carcasses that are graded “Select”. In Australia, all beef that has the Meat Standards Australia (MSA) label is aged for a minimum of 5 d (MLA, 2018). However, the MSA grading model predicts little increase in striploin quality score, of which tenderness represents 40%, beyond 14 d. Recently in Ireland there has been a move towards extended ageing of up to 30 d for striploin, particularly for “premium” branded products (Appendix 1). This is based presumably on a perception that

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this results in enhanced quality due to an increase, at least in part, in tenderness. From a commercial perspective, the cost incurred in extended ageing must be judged against the additional increase in tenderness or overall palatability. There is a paucity of information as to whether extended ageing of striploin from cattle within Irish beef production systems significantly, either statistically or commercially, increases tenderness. Therefore, the objective of this study was to determine the interaction between modifications within one typical heifer beef production system (Teagasc, 2015) and post-mortem ageing to 28 d on the tenderness, measured instrumentally as toughness, of striploin muscle.

Materials and methods

Experimental design and animal management

The study was carried out under license from the Irish Government Department of Health and Children and all procedures used complied with national regulations concerning experimentation on farm animals. Spring-born early-maturing sired (Aberdeen Angus) × Holstein-Friesian heifers (AAH, $n = 48$) and late-maturing sired (Belgian Blue) × Holstein-Friesian heifers (BBH, $n = 48$) were reared from 4 mo of age according to the heifer grass-based dairy calf to beef blueprint described by Keane & Drennan (2008). Within breed, the animals were blocked on bodyweight and within block assigned at random to early slaughter or late slaughter. The animals grazed a perennial ryegrass-dominant sward from July 1, and were offered a daily allowance of grass dry matter (DM) sufficient to ensure a consumption of 2.5% bodyweight (assuming 75% utilisation) for the whole group. They were housed on November 30 and offered unwilted grass silage *ad libitum* and 1.5 kg of concentrates per animal daily until turnout to pasture on April 1 when they again grazed a perennial ryegrass-dominant sward. At the end of October, animals previously assigned to early slaughter, were weighed on consecutive days, transported approximately 90 km to a commercial abattoir and slaughtered (Halal procedure, Farouk *et al.* (2014)) within 45 min of arrival. Carcasses were subjected to low voltage electrical stimulation (24 V, 30 s). Animals previously assigned to late slaughter were housed on November 15 and offered unwilted grass silage *ad libitum* and 5.0 kg of concentrates per animal daily. At the end of March, they were weighed and slaughtered as described for animals slaughtered the previous October. Thus, within sire breed, there were two slaughter weight/age categories, that is, young/light and older/heavier.

Carcass trait measurements and sample collection

At <45 min post slaughter, a sample (*ca* 20 g) of LT muscle tissue (from the 10th rib position) was taken, snap frozen in

liquid nitrogen and maintained at -80°C for metabolic enzyme activity and fibre typing analyses. The carcasses were then weighed and graded for conformation (5-point scale, classes E [highest] to P [lowest], E is 5) and fatness (5-point scale, scores 5 [highest] to 1 [lowest]) according to the EU Beef Carcass Classification Scheme (EC, 2006) and chilled under factory conditions. After approx. 24 h the right side of each carcass was cut at the 5th/6th rib interface ("pistola" cut) and the LT muscle, between the 10th and 13th ribs (three rib striploin cut), was excised, vacuum packed, transferred to the Teagasc Food Research Centre, Ashtown, Dublin and stored at 2°C until approximately 48 h post-mortem.

Muscle sampling and measurement

At approximately 48 h post-mortem, the pH of the LT muscle (pH_u) was measured by making a scalpel incision at the 10th rib end and inserting a glass electrode (Model EC-2010-06; Amagrass Electrodes Ltd., Westport, Co. Mayo, Ireland) attached to a portable pH meter (Model no. 250A; Orion Research Inc., Boston, MA, USA) approximately 2.5 cm into the muscle. Five LT steaks of 2.5 cm thickness were then removed and vacuum-packed (Multivac[®], model A300/16; Multivac, Ltd., Swindon, UK). Two steaks were immediately frozen for chemical composition and collagen analysis, respectively. Three steaks were randomly assigned within carcass to storage at 2°C for 7, 14 or 28 d post-mortem after which they were stored at -20°C .

Thawed samples of LT (4°C overnight) with external fat and connective tissue removed were homogenized using a Robot coupe blender (R301 Ultra; Robot Coupe SA, Vincennes, France) and total intramuscular fat (IMF) content measured as described by Folch *et al.* (1957). Total and heat-soluble (77°C , 65 min) collagen concentrations were determined by quantitative determination of hydroxyproline by a colorimetric reaction as described by Listrat & Hocquette (2004).

The primary references for the analysis of the activity of glycolytic enzymes (phosphofructokinase [PFK, EC 2.7.1.11], lactate dehydrogenase [LDH, EC 1.1.1.27]) and oxidative enzymes (isocitrate dehydrogenase [ICDH, EC 1.1.1.42], citrate synthase [CS, EC 4.1.3.7] and cytochrome c oxidase [COX, EC 1.9.3.1]) were given in Moran *et al.* (2017). Muscle fibre types (Type I, IIA, IIX and IIB) were identified by separating myosin heavy chain (MyHC) isoforms (MyHC I, IIA, IIX and IIB) using high-resolution mini-gel electrophoresis as described by Picard *et al.* (2011).

For instrumental texture analysis, frozen vacuum-packed steaks were thawed in circulating water at 20°C . External fat and connective tissue surrounding the muscle were removed and the steaks were conditioned for 15 min at 20°C . The steaks were subsequently cooked in vacuum pack bags to an internal temperature of 70°C , by immersing in a water bath (Model Y38; Grant Instruments Ltd. Royston, UK) at

72°C. The internal temperature of the steaks was measured using a digital thermometer (HI 904; Hanna Instruments Ltd, Leighton Buzzard, Bedfordshire, UK). After cooking, the juices were removed, the steaks were cooled to room temperature and stored overnight in a closed bag at 4°C, for subsequent Warner–Bratzler shear force (WBSF) analysis according to the procedure of Shackelford *et al.* (1994). Briefly, six cores (1.25 cm diameter) parallel to the direction of the muscle fibres were obtained and sheared using an Instron Universal testing machine, model 5543 (Instron Corporation, High Wycombe, Buckinghamshire, UK) equipped with a Warner–Bratzler shearing device. The crosshead speed was 5 cm/min. Instron Series IX Automated Materials Testing System software for Windows (Instron Corporation, High Wycombe, Buckinghamshire, UK) was employed in the analysis.

Statistical analysis

Animal, carcass and LT data (measured once), were subjected to analysis of variance using a model that had block, sire breed, slaughter time (early or late) and sire breed × slaughter time as sources of variation. Warner–Bratzler shear force data were analysed as a randomised block, split-plot design with the sources of variation described already in the main plot and time of ageing and all time-related interactions in the sub-plot. Where significant effects were detected (F-test), means were separated using a Tukey's test. Data (measured once) were also subjected to multiple analysis of variance to calculate partial correlation coefficients (ρ) from the error sum of squares and cross products matrix.

Results

Carcass and LT muscle characteristics are summarised in Table 1. There was no interaction between sire breed and slaughter time for any of the variables listed. Carcasses from AAH heifers were lighter, fatter (higher fat score) and had a poorer conformation ($P < 0.001$) than those from the BBH heifers. Later slaughter increased ($P < 0.001$) carcass weight and carcass fatness and improved carcass conformation. Muscle from AAH heifers had higher ($P < 0.001$) IMF content and lower ($P < 0.05$) LDH activity, whether expressed on a tissue or protein basis, compared to muscle from BBH heifers but other enzyme activities did not differ between sire breeds. Muscle from AAH heifers had a lower ($P < 0.001$) proportion of type IIX muscle fibres and a higher ($P < 0.001$) proportion of type IIA and type I muscle fibres compared to muscle from BBH heifers, but collagen characteristics did not differ between sire breeds. Later slaughter resulted in a higher muscle pH at 75 min and 48 h post-mortem, increased ($P < 0.001$) IMF concentration and the activities of LDH and PFK but decreased the activity

of CS ($P < 0.05$), total and insoluble concentrations and collagen solubility.

The WBSF data are summarised in Table 2. There were no interactions between sire breed and slaughter time and there was no difference between sire breeds for WBSF. Later slaughter decreased ($P < 0.05$) WBSF as did the duration of ageing ($P < 0.001$) with both linear and quadratic terms being significant.

Partial correlation coefficients between characteristics of carcass or LT muscle and WBSF are shown in Table 3. The IMF content was negatively correlated with WBSF after 7 ($P < 0.05$), 14 ($P < 0.05$) and 28 ($P < 0.1$) d ageing. ICDH activity, expressed on a protein basis, and LDH activity, expressed on a tissue basis, were negatively correlated ($P < 0.05$) with WBSF after 7 d ageing.

Discussion

The primary objective of this study was to determine the effects on tenderness of extended ageing of Irish striploin muscle. A larger project concerned with the enhancement of the nutritional quality of beef (Moloney *et al.*, 2008) provided an opportunity to examine the influence of two important factors within beef production systems on the tenderness response to ageing. These factors were sire breed and time of slaughter, which were expected to change muscle structure and composition and possibly influence the tenderness response to extended ageing. It is recognised that within the production system chosen, slaughtering animals early, at a younger age, results in a lighter carcass and that the effects of slaughter time, age and carcass weight are therefore confounded. "Premium" striploins are often also branded as being from the progeny of early maturing breed sires such as Angus steers (Appendix 1). Comparison with the late maturing Belgian Blue breed was therefore of commercial interest. In this study, striploins were aged in vacuum sealed air-tight bags. This "wet" ageing is the most common commercial practise in Ireland. Extended ageing of intact joints in controlled atmospheric conditions termed "dry" ageing, is also used primarily for speciality products with a particular focus on flavour. This is considered a different post-mortem technology in the context of the present study and is not discussed further. An objective, standardised instrumental approach was used to directly measure toughness to allow comparison with the literature but also because of the practical constraints to carrying out trained sensory panel or consumer assessment of the large sample set used.

As expected, the sire breeds resulted in differences in carcass characteristics which are qualitatively similar to those reported for dairy origin steers sired by Angus or Belgian Blue bulls (Keady *et al.*, 2017). The size of the difference between

Table 1: Characteristics of the carcass and *longissimus* muscle from Angus and Belgian Blue sired Holstein-Friesian heifers slaughtered at 20 (early) or 25 (late) mo of age

Slaughter time (S)	Sire breed (B)				sed	Significance ¹	
	Angus		Belgian Blue			B	S
	Early	Late	Early	Late			
No of animals	24	24	24	24			
Final weight (kg)	472	588	478	603	10.1		***
Age (mo)	20.1	24.7	20.1	24.6	0.09		***
Carcass (kg)	234	298	250	321	5.91	***	***
Fat score ²	3.4	4.2	2.5	3.6	0.19	***	***
Conformation score ³	2.3	2.8	2.8	3.2	0.13	***	***
<i>Longissimus</i> muscle							
Fat (g/kg)	51.3	67.8	35.7	42.5	4.33	***	***
Protein (g/kg)	212	212	212	212	4.91		
Enzymatic activities							
ICDH ⁴	1.31	1.25	1.30	1.20	0.090		
ICDHp ⁵	6.20	5.91	6.14	5.67	0.437		
LDH ⁴	1120	1141	1155	1205	24.4	**	*
LDHp ⁵	5283	5412	5452	5700	134.5	*	*
PFK ⁴	70.7	95.3	66.4	91.7	14.48		*
PFKp ⁵	335	455	313	434	68.9		*
COX ⁴	14.3	13.8	13.2	13.9	0.92		
COXp ⁵	67.7	65.1	62.2	66.2	4.41		
CS ⁴	6.66	6.45	7.52	6.21	0.438		*
CSp ⁵	31.5	30.4	35.5	29.5	2.09		*
pH (75 min)	6.13	6.50	6.18	6.55	0.56		*
pH (48 h)	5.52	5.50	5.57	5.49	0.031		*
Muscle fibre types (%)							
IIX	45.7	46.9	54.0	54.5	2.42	***	
IIA	34.8	35.0	29.7	30.1	2.20	***	
I	19.4	18.0	16.3	15.4	1.07	***	
Collagen content							
Insoluble collagen (% DM) ⁶	2.38	1.81	2.35	2.05	0.130		***
Total collagen (% DM)	3.31	2.36	3.23	2.70	0.196		***
Collagen Solubility (%)	27.1	23.3	26.7	23.4	1.92		*

¹There were no sire breed × slaughter weight interactions.

²Fat score; 5 = fattest, 1 = leanest.

³Conformation score; 5 = best, 1 = worst.

⁴Enzymatic activity expressed as $\mu\text{mol}/\text{min}$ per g of fresh muscle.

⁵Enzymatic activity expressed as $\mu\text{mol}/\text{min}$ per g of protein in muscle.

⁶DM = dry matter.

sire breeds in LT IMF content across both studies reflects the different genders and experimental designs (59.6 vs. 39.1 in the present study and 74.5 vs. 36.4 g/kg in Keady *et al.*

(2017) for Angus and Belgian Blue sires, respectively). The higher glycolytic enzyme activity (LDH) and the trend towards lower oxidative enzyme activities (ICDH and COX) for BBH

Table 2: Warner Bratzler shear force (WBSF) of *longissimus* muscle from Angus and Belgian Blue sired Holstein-Friesian heifers slaughtered at 20 (early) or 25 (late) mo of age

Ageing (d)	Sire breed (B)				Mean ¹
	Angus		Belgian Blue		
	Early	Late	Early	Late	
WBSF (N)					
7	34.0	31.6	35.6	33.7	33.7 ^a
14	30.4	25.8	30.0	28.5	28.7 ^b
28	26.2	22.9	27.0	23.6	24.9 ^c
Significance					
B					
Slaughter time (S)	*				
Days (D)	***(L, Q) ²				
B × S					
B × D					
S × D					
B × S × D					
Sed (B × S × D)	1.92				

¹Means within a column with a different superscript differ significantly ($P < 0.05$).

²L and Q are linear and quadratic effects, respectively of duration of ageing.

heifers was similar to that seen in the comparison of early and late maturing bulls in Mezgebo *et al.* (2017). The difference in the distribution of muscle fibres between sire breeds was also similar to that reported by Mezgebo *et al.* (2017) and interpreted from the data reported by Maltin *et al.* (2001) for a comparison of purebred Angus and Charolais steers using a different methodology to measure muscle fibres. The lack of difference in collagen characteristics between sire breeds in the present study and also reported by Maltin *et al.* (1998) contrasts with that of Christensen *et al.* (2011) who reported lower collagen solubility for Angus bulls compared to Charolais bulls although total collagen concentration did not differ in that study. Similar to the present study, Cuvelier *et al.* (2006) found no difference in WBSF in meat from Angus (IMF concentration = 24.0 g/kg) and Belgian Blue bulls (IMF concentration = 6.5 g/kg) aged for 2 d and 8 d, respectively. Chambaz *et al.* (2003) compared the WBSF of LT from Angus, Charolais, Limousin and Simmental steers slaughtered at a common IMF concentration (32.5 g/kg) and found no difference. In contrast to the present study, Keady *et al.* (2017) reported lower WBSF for LT from Angus-sired steers than from Belgian Blue sired steers after 14 d ageing, which likely reflects the greater difference in IMF content in the two studies.

The effect of later slaughter (increasing slaughter age/weight) on IMF content and collagen solubility was as expected (Maltin *et al.*, 1998; Mezgebo *et al.*, 2019). The time interval in this

study and that of Maltin *et al.* (1998) was not sufficient to alter the muscle fibre distribution seen in Mezgebo *et al.* (2019). The increase in glycolytic enzyme activity (LDH and PFK) and decrease in oxidative enzyme activity (CS) due to later slaughter which was not seen in Mezgebo *et al.* (2019) may reflect the higher energy density in the pre-slaughter diet of the heavier/older heifers. In support of this suggestion, when late maturing bulls were slaughtered at the same age from pasture or a high energy ration, carcass weight differed but the LT from the heavier animals had a more glycolytic and less oxidative metabolism (Moloney *et al.*, 2017). The decrease in WBSF (increase in tenderness) with later slaughter in the present study is consistent with Lively *et al.* (2005) and McNamee *et al.* (2014) and is likely due to the increase in IMF content within the relatively narrow range in age. However, if animals were slaughtered at an older age, it is likely that the increase in IMF would not prevent an increase in WBSF due to a further decrease in collagen solubility as was seen by Mezgebo *et al.* (2019) in late maturing breed sired suckler bulls slaughtered between 19 and 24 mo of age. The age at which this transition would occur for heifers remains to be determined but slaughtering heifers older than 25 mo of age is unlikely to be commercially attractive.

In summary, the biochemical and structural characteristics of LT from the four categories of animal in this study were generally as expected based on the literature but the variation between the groups was very modest.

Table 3: Partial correlation coefficients between characteristics of carcass or *longissimus* muscle and Warner Bratzler shear force (WBSF) after 7, 14 or 28 d ageing post mortem

Ageing (d)	WBSF		
	7	14	28
Final weight (kg)	-0.171	-0.149	-0.141
Age (mo)	-0.137	-0.112	-0.037
Carcass (kg)	-0.106	-0.110	-0.071
Fat score ¹	0.078	0.007	0.120
Conformation score ²	0.144	-0.000	0.016
Longissimus muscle			
Fat (g/kg)	-0.220*	-0.251*	-0.213*
Protein (g/kg)	0.245*	0.181	0.163
Enzymatic activities			
ICDH ³	-0.126	-0.035	0.059
ICDHp ⁴	-0.219*	-0.098	-0.003
LDH ³	0.226*	0.074	0.035
LDHp ⁴	-0.053	-0.122	-0.140
PFK ³	0.129	0.037	0.079
PFKp ⁴	0.092	0.002	0.045
COX ³	0.136	0.087	0.053
COXp ⁴	0.039	0.022	-0.011
CS ³	0.005	-0.014	0.061
CSp ⁴	-0.083	-0.076	-0.010
Muscle fibre types (%)			
IIX	0.060	-0.017	0.017
IIA	-0.015	0.014	-0.049
I	-0.108	0.008	0.061
Collagen content			
Insoluble collagen (% DM) ⁵	0.017	0.085	-0.059
Total collagen (% DM)	-0.020	0.103	-0.046
Collagen solubility (%)	-0.096	0.034	0.029

¹Fat score; 5 = fattest, 1 = leanest.

²Conformation Score; 5 = best, 1 = worst.

³Enzymatic activity expressed as $\mu\text{mol}/\text{min}$ per g of fresh muscle.

⁴Enzymatic activity expressed as $\mu\text{mol}/\text{min}$ per g of protein in muscle.

⁵DM = dry matter.

It is generally accepted that ageing *per se* decreases muscle WBSF. The lack of an interaction between sire breed and slaughter time with ageing time for WBSF is indicative of a consistent pattern of ageing for the four animal type combinations examined in this study and reflects the small differences in muscle structure observed. The effect of ageing on muscle tenderness has been the subject of many

studies and the findings are equivocal (below). The variation between studies likely reflects the animal/carcass type used, the ageing conditions, that is, whether as joint/partial carcass or individual muscle, whether vacuum packaged or not and the ageing temperature, the power of individual experiments and the instrument/shear force measurement protocol employed. Thus, Smith *et al.* (1978) observed no decrease in WBSF of LT aged up to 28 d compared to 11 d. Similarly, Jaborek *et al.* (2019) and Harris *et al.* (1992) concluded that compared to ageing for 14 d, ageing for 28 d did not result in a significant decrease in WBSF of LT. In addition, ageing striploin for 42 d compared to 14 d (Colle *et al.*, 2015) or for 35 d compared to 28 d (Laster *et al.*, 2008; Juarez *et al.*, 2010) did not decrease WBSF. In contrast, Jennings *et al.* (1987) observed a decrease in shear force of LT between 10 and 20 d ageing while Huff and Parrish (1993) observed a decrease in WBSF of LT between 14 and 28 d ageing (2.8 vs. 2.3 kg) as did Adcock *et al.* (2015). No further decrease was observed in the latter study when ageing was extended to 42 d. While Lepper-Blilie *et al.* (2016) reported a linear decrease in WBSF of striploin when aged up to 49 d, WBSF at 28 d was lower than that at 14 d (2.48 vs. 2.80 kg) but not subsequently. Across studies, the incremental decrease in WBSF with prolonged ageing (after 14 d) is small. Gruber *et al.* (2006) examined the effects of post-mortem ageing for 2, 4, 6, 10, 14 or 28 d on 17 individual muscles from two USDA grades of carcass. They reported that for the higher quality grade of carcass (upper two-thirds USDA Choice, marbling score 575), LT WBSF did not decrease beyond 21 d (3.96, 3.66 and 3.55 kg for 14, 21 and 28 d ageing, respectively). By contrast, for the lower grade carcasses (USDA Select, marbling score 351), LT WBSF decreased from 21 to 28 d ageing (5.02, 4.52 and 4.21 kg for 14, 21 and 28 d ageing, respectively). Similarly, using the same procedure as in the present study, Moloney *et al.* (2018) showed that WBSF of LT from steers (IMF = 76 g/kg) did not decrease between 14 and 21 d ageing while WBSF of LT from bulls of a similar genotype (IMF = 43 g/kg) decreased by 4.8 N. These data are consistent with the data from the present study, that is, LT with the higher IMF concentration had lower WBSF and on average, WBSF was lower after 28 d ageing compared to 14 d. The small negative partial correlation supports the influence of IMF content on WBSF.

While the lower WBSF value for 28 compared to 14 d aged LT in the present study was statistically significant, the actual difference was small (3.72 N). Several studies have sought to interpret WBSF values in the context of consumer acceptability and the ability of consumers to detect differences. Miller *et al.* (2001) reported that WBSF values of <3.0 (29.4 N), 3.4 (33.4 N) and 4.0 (39.2 N) would result in 100, 99 and 94% US consumer satisfaction for beef tenderness. Similarly,

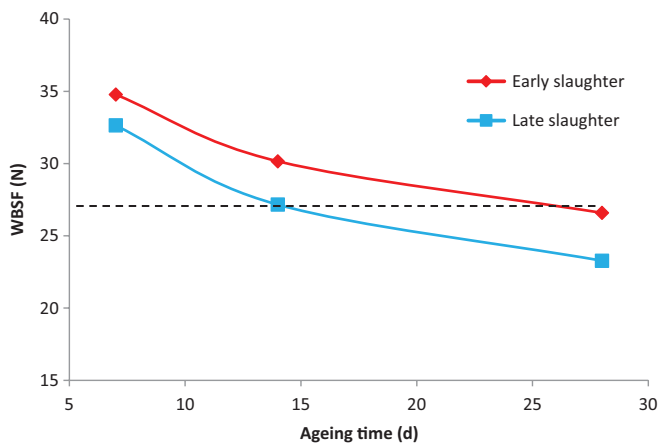


Figure 1. Warner Bratzler shear force (WBSF) during post-mortem ageing of longissimus muscle from heifers slaughtered at 20 (early) or 25 (late) mo of age.

Belew *et al.* (2003) concluded that “beef with WBSF values <3.2 kg (31.4 N) would be rated very tender by trained panellists irrespective of muscle source”. Destefanis *et al.* (2008) indicated that European consumers would consider LT with a WBSF value <42.87 N to be tender. Based on these data, on average all samples in the current study would be considered tender even after 7 d ageing. From the data of Miller *et al.* (2001) a difference in WBSF of approximately 4.9 N can be detected by a consumer. Therefore, untrained consumers are likely to detect the decrease in WBSF from 7 to 14 d ageing (5.48 and 4.62 N for LT from late and early slaughtered heifers, respectively) but not due to further ageing (3.88 and 3.57 N for LT from late and early slaughtered heifers, respectively). It is noteworthy that in the present study, WBSF of LT from carcasses from the later slaughtered heifers aged for 14 d was similar to the LT from carcasses from the early slaughtered heifers aged for 28 d (Figure 1) indicating that a particular WBSF target is influenced by both carcass type and the duration of ageing. In this regard, Lepper-Bililie *et al.* (2016), who examined the effect of extended ageing on both WBSF and sensory characteristics of striploin concluded that “while WBSF values declined linearly from day 14 to day 49 of ageing, economic and flavour attribute data suggest the small improvement in WBSF value beyond 21 d cannot be used to justify the extended ageing period”. Similarly, Adcock *et al.* (2015) concluded that for Certified Angus muscle despite the decrease in WBSF from 14 to 28 d ageing, “consumers signified only minute differences (in sensory palatability) following 14 d of ageing”.

It is concluded that within the production system examined in the present study, and based on WBSF data, extending LT ageing to 28 d is not necessary to ensure consumer satisfaction.

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Appendix 1: Declared aging time on packaged striploins in the major Irish retailers

Retailer	Product name	Ageing time (d)
Dunnes Stores	“Simply Best” Irish Angus Striploin	28
Dunnes Stores	Irish Beef Striploin Steaks	21
Tesco	“Tesco Finest” Irish Angus Striploin	30
Tesco	Irish Beef Striploin	28
Aldi	“Butchers Selection” Irish Matured Striploin	30
Aldi	Irish Beef Striploin	28
Aldi	Irish Black Angus Striploin	30
Lidl	“Deluxe” Irish Hereford Striploin	28
Lidl	Inisvale 100% Irish Striploin	28
Lidl	Inisvale Irish Striploin	25
Supervalu	“Signature Tastes” Striploin	21

Survey carried out 6 and 7 July 2019.