1	Effect of finishing diet and duration on the sensory quality and volatile profile of lamb
2	meat
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Abstract

Animal production factors diet can affect the sensory quality of lamb meat. The study 35 investigated the effect of diet composition and duration of consumption on theproximate 36 analysis, volatile profile and sensory quality of lamb meat. Ninety-nine male Texel × Scottish 37 Blackface lambs were raised at pasture for 10 months before being assigned in groups of 11 38 to one of the following treatments: 100% Silage (S) for 36 (S36), 54 (S54) or 72 (S72) days; 39 50% Silage 50% - 50% Concentrate (SC) for 36 (SC36), 54 (SC54) or 72 (SC72) days; 100% 40 Concentrate (C) for 36 (C36) or 54 (C54) or 72 (C72) days. A trained sensory panel found 41 42 Intensity of Lamb Aroma, Dry Aftertaste and Astringent Aftertaste to be higher in meat from lambs on the concentrate diet. Discriminant analysis showed that the volatile profile enabled 43 discrimination of lamb based on dietary treatment but the volatile differences were 44 45 insufficient to impact highly on sensory quality. Muscle from animals in the S54 group had higher Manure/Faecal Aroma and Woolly Aroma than the SC54 and C54 groups, possibly 46 47 related to higher levels of indole and skatole. Further research is required to establish if these small differences would influence consumer acceptability. 48

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50 Keywords: Animalfeed, Silage, Concentrate, Discriminant analysis, Palatability,
51 SPME/GC/MS

53 **1 Introduction**

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The main feedstuffs consumed by sheep for meat production are derived from cereal 54 55 grains and pasture (either grazed or ensiled grass), with combinations of both feed sources often in use over the lifetime of animal (Almela et al., 2010). The growth rates of sheep 56 57 receiving solely grass-based diets are lower and ultimate carcass weights may also be lower (Murphy, Loerch, McClure, & Solomon, 1994; Priolo, Micol, Agabriel, Prache, & 58 Dransfield, 2002); thus, grain-based concentrates, which are more energy dense, are often 59 used to shorten the time to slaughter, increase dressing percentage, and improve carcass 60 quality (De Brito, Ponnampalam, & Hopkins, 2017; Jaborek, Zerby, Moeller, & Fluharty, 61 62 2017). In addition to the effects of diet on production parameters (De Brito et al., 2017), 63 dietary constituents may also have a considerable effect on meat quality (Kitessa et al., 2009). 64 There are differences in the consumer acceptability of meat from grain-fed and grass-fed 65 sheep (Font i Furnols et al., 2006; Sanudo et al., 2007) attributable to, among other factors, 66 variation in the level of intramuscular fat (IMF) and subcutaneous fat and their fatty acid 67 composition (Howes, Bekhit, Burritt, & Campbell, 2015). Consumer assessment of lamb 68 69 meat is influenced by the taste and/or aroma deriving from volatile compounds, which are known to be affected by the relative proportions of fatty acids in the meat (Ponnampalam, 70 Sinclair, Egan, Ferrier, & Leury, 2002). With regard to flavour specifically, the extent to 71 which flavour intensity is altered depends on the types of both forage and grain consumed 72 (Duckett & Kuber, 2001). Meat from sheep receiving primarily grass-based diets (pasture or 73 74 grass silage) is reported to have a pastoral (grassy) flavour (Young, Lane, Priolo, & Fraser, 2003). In this context, nutritional strategies may be used to modulate the sensory quality of 75

lamb ultimately affecting consumer preference (Almela et al., 2010); There are other

study (<u>Gkarane et al., 2017</u>), we reported less favourable sensory attributes in lamb from rams compared to castrates; in this instance a modification to the diet might be useful in overcoming undesirable sensory attributes. The objective of the current study was to test the hypothesis that different proportions and durations of feeding cereal concentrate and silagebased diets would affect the sensory quality and volatile profile of lamb meat from rams.

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84 **2 Materials and methods**

85 2.1 Animal husbandry, slaughter and sampling

All animal procedures used in this study were conducted under experimental license 86 from the Irish Health Products Regulatory Authority (HPRA) in accordance with the 87 European Union (Protection of Animals used for Scientific Purposes) Regulations 2012 (S.I. 88 No. 543/2012). Ninety-nine ram lambs (Texel × Scottish Blackface) were sourced from Irish 89 farms in March 2015. Lambs were raised at pasture from birth (March 2015) and were 90 weaned at 130 d of age after which they were transported to the Teagasc Sheep Research 91 92 Centre, Athenry, Co. Galway, Ireland (Claffey et al., 2018). Lambs were maintained at 93 pasture until selected for commencement of an intensive indoor finishing period. Lambs were allocated to the following nine dietary treatments consisting of three grass silage:concentrate 94 95 ratios (100:0 (S), 50:50 (DM basis) (SC), 0:100 (C)) with each diet being fed for three preslaughter feeding durations (36, 54 and 72 d) to give the following dietary treatments: S36, 96 S54, S72, SC36, SC54, SC72, C36, C54, C72. The grass silage was predominantly Lolium 97 perenne L. and the concentrate diet consisted of 30% maize, 30% barley, 16.5% soya hulls 98 99 and 15.5% soybean meal. In line with commercial practice, lambs were selected for treatment 100 based on initial live weight and predicted growth rate on the assigned diets to yield lambs with similar weights at slaughter. Thus, the lightest lambs were assigned to the C72 treatment 101 and the heaviest to the S36 treatment. For the indoor finishing period (36, 54 or 72 d) lambs 102

were individually penned in metal floor feeding pens (182 cm \times 122 cm). At the end of the 103 finishing period, lambs were transported to a commercial abattoir (Gillivan's, Moate, Co. 104 Westmeath, Ireland) for slaughter. The mean ages in days $(\pm SD)$ of the animals at slaughter 105 were 252 (±6.4), 260 (±3.7), 273 (±6.0), 248 (±3.8), 254 (±4.8), 271 (±5.3), 248 (±6.1), 258 106 (±5.0), 266 (±4.3) for the S36, S54, S72, SC36, SC54, SC72, C36, C54, C72 treatments, 107 respectively. After slaughter, carcasses were chilled overnight and transported to Teagasc 108 109 Food Research Centre, Ashtown, Dublin 15, Ireland, for dissection. Ultimate pH (pHu) of M. longissimus thoracis et lumborum (LTL) was measured at 25 h post slaughter at the 13th rib 110 111 using a SympHony SP70P hand-held pH meter (VWR, Dublin, Ireland). Both LTL muscles were excised from each carcass, cut into 2.5 cm thick steaks, vacuum packed, aged for 8 d at 112 4 °C and frozen at -20 °C until required for analysis. 113

114 **2.2 Compositional analysis**

Samples of LTL were thawed overnight at 4 °C and homogenized using a Kenwood 115 CH180 Compact Mini Chopper (Kenwood, Hampshire, UK). Moisture and intramuscular fat 116 (IMF) contents were determined using the SMART Trac Rapid Fat Analyzer (CEM 117 Corporation, NC, USA) according to AOAC Methods 985.14 and 985.26 (AOAC, 1990), 118 respectively. Protein concentration was determined using a LECO FP328 (LECO Corp., MI, 119 USA) protein analyzer based on the Dumas method and according to AOAC method 992.15 120 (AOAC, 1990). Ash was determined following incineration of samples overnight in a furnace 121 at 540 °C. 122

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124 **2.3 Reagents and fibres for volatile analysis**

Volatile standards, the alkane mixture (C7 - C30), methanol (for preparation of stock
solutions of the standards), and sodium sulfate were supplied by Sigma-Aldrich Ireland Ltd

(Arklow, Co. Wicklow, Ireland). The volatile standards hexanoic acid and α-terpineol were
supplied from VWR International Ltd (Blanchardstown, Dublin 15, Ireland) while 1pentadecanol was supplied from Fisher Scientific Ireland Ltd (Blanchardstown, Dublin 15,
Ireland). Solid phase microextraction (SPME) fibres (50/30 µm CAR/DVB/PDMS fibre; 1
cm length) were supplied by Agilent Technology (Part Number: SU57298U; Unit 3, Euro
Business House, Cork, Ireland). All reagents and chemicals were of chromatographic quality.

134 **2.4 Sample preparation and volatile analysis**

Before analysis LTL samples were thawed by immersion of frozen vacuum packed 135 samples in water at room temperature for 20 min. Thawed steaks were grilled with the fat 136 attached, using a clamshell grill until an internal temperature of 70 °C was reached 137 (monitored using a hand-held digital thermometer; Eurolec, Dublin, Ireland). Subcutaneous 138 fat was removed and 7 g from the core was weighed and homogenized with 7 g Na₂SO₄ using 139 140 a Kenwood CH180 Compact Mini Chopper (Kenwood, Hampshire, UK). A 5 \pm 0.05 g 141 sample of the mixture was placed in a 20 ml glass headspace vial sealed with a polytetrafluoroethylene (PTFE)-faced silicone septum (VWR, Dublin, Ireland). The vial 142 containing the sample was equilibrated in a water bath set at 90 \pm 2 °C for 20 min and the 143 fibre was exposed to the headspace over the sample for a further 20 min. These SPME 144 conditions (adopted based on maximizing the number of compounds detected, the total peak 145 area and the detection of BCFAs) were considered optimum as previously described in 146 Gkarane et al. (2018). After adsorption, the fibre (50/30 µm CAR/DVB/PDMS) was removed 147 148 from the vial and immediately inserted into the injection port of the GC. Analysis of the volatile compounds was carried out using a Varian 3800 GC coupled to a Varian Saturn 2000 149 150 ion trap mass spectrometer (Varian Chromatography Systems, Walnut Creek, CA, USA). 151 Volatile extraction, adsorption and injection were performed manually. The injector,

operating in splitless mode, was set at 250 °C and the desorption time was 8 min. Helium was 152 used as carrier gas with a constant flow rate of 1.0 ml/min. Volatile compounds were 153 separated using an Agilent ZB-5MS column (30 m length, 0.25 mm internal diameter, 0.25 154 µm film thickness) (Agilent Technologies, Palo Alto, CA, USA). The GC oven temperature 155 was programmed as follows: 40°C for 5 min, increasing to 230 °C at 4 °C/min and holding 156 for 5 min, with a total acquisition time of 57.5 min. The GC/MS transfer line was heated at 157 158 280°C. Acquisition was performed in electron impact (EI) mode (70 eV) at 10 microscans/s, scanning the mass range 33–230 m/z. Saturated n-alkanes (C7 - C30) injected directly (1 µl) 159 160 onto the column were run under the same GC-MS conditions (at split ratio of 1:50) to obtain linear retention index (LRI) values for the volatile compounds detected. Compounds were 161 identified by comparing their mass spectra with those of spectra from the NIST/EPA/NIH 162 Mass Spectral Database (Version 2.0 g, 2011), those of authenticated standards and linear 163 retention indices matching those of published values (Gkarane et al., 2018). Individual 164 animals were considered as experimental units and one meat sample from each animal was 165 subjected to analysis using a randomized block design to avoid experimental bias. Integration 166 of the peak areas of the volatile compounds used specific ion identification for each molecule 167 (to deal with co-elution of some compounds). An external standard (bromobenzene (10 ppm)) 168 was run daily under the same SPME and GC-MS conditions as the samples. For volatile 169 analysis, the peak area (PA) of each volatile was first normalised against bromobenzene 170 171 before adding a constant (+1) and being logarithmically transformed to achieve a normal distribution. The amount of each volatile was expressed as logarithmically transformed PA 172 for that compound. 173

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175 **2.5 Lamb meat preparation for sensory analysis**

The LTL muscle from the left side of each carcass was used for sensory analysis. On 176 the day of sensory testing, packaged frozen steaks were thawed by immersion in water at 177 room temperature for 45 min. Steaks were grilled, with subcutaneous fat attached, to an 178 internal temperature of 70 °C, using a clamshell grill. On reaching 70 °C (monitored using a 179 hand-held digital thermometer (Eurolec, Dublin, Ireland)) the steaks were removed from the 180 grill, wrapped with aluminium foil and allowed to rest for 3 min. Each steak was unwrapped 181 182 and following removal of the subcutaneous fat, cut into 8 pieces of approximately 2 cm³. Samples were re-wrapped with foil, assigned a random three-digit code, held in an oven set at 183 184 60 °C and served to the panellists within 20 min.

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186 **2.6 Panel training**

187 Staff at Teagasc Food Research Centre, Ashtown, participated as sensory panellists in 188 16 training sessions prior to participating in sensory testing. Training sessions included: lamb 189 meat tasting to generate descriptors for aroma, flavour, texture/mouthfeel, taste and aftertaste; 190 spiking sessions using lamb flavour/aroma compounds; and training using physical and 191 chemical reference standards. A detailed procedure for the panel training is described in 192 <u>Gkarane et al. (2017)</u>.

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194 **2.7 Quantitative descriptive analysis**

Quantitative descriptive analysis (QDA) was performed on one day per week over 8 weeks with two sensory sessions per day (morning and afternoon). In each session, six samples were assessed using a balanced and randomized design. Panellists were asked to rate attributes (generated during the training) for each sample, by marking a point on a 100 mm unstructured line scale. Unsalted crackers and water at room temperature were given to panellists to cleanse the palate between samples. The sensory attribute definitions, agreed
during the training sessions (<u>Gkarane et al., 2017</u>), were available to each panellist during
tasting. Panellist evaluations were recorded using Compusense 5 (v4.4, Compusense Inc.,
Guelph, Ontario, Canada).

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205 2.8 Statistical analyses

Proximate and sensory analysis data were tested for the normality of the residuals for each variable. In the case of non-normal distribution, data were transformed using the Box-Cox transformation. All data were analysed using a mixed model with diet, duration and diet x duration as fixed effects (SAS (v9.4)). For the sensory data, the sensory analysis session and carcass weight were considered as random effects. Analysis was conducted in the MIXED procedure of SAS (v9.4). All data were presented as least square means.

Principal component analysis (PCA) of the sensory and volatile data for the nine treatments was performed using XLSTAT®statistical software (Version 19.01.41647; Addinsoft, Paris, France). Associations between sensory attributes and diets, and volatile compounds and diets were also investigated using Discriminant Analysis (DA) performed using XLSTAT®statistical software (Version 19.01.41647; Addinsoft, Paris, France).

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218 **3. Results and discussion**

219 **3.1 Proximate analysis**

There was no difference in muscle fat content among dietary treatments or finishing periods (Table 1). Other authors have reported that lambs receiving concentrate diets generally have higher growth rates (Fraser & Rowarth, 1996) and IMF than lambs receiving pasture-based diets (De Brito et al., 2017). However, Crouse et al. (1978) found no difference in fat thickness or percentage carcass fat of lambs fed low, medium or high energy diets and
slaughtered at constant weights. Similarly, <u>Aurousseau et al. (2007)</u> detected no differences
in the lipid content of *M. longissimus thoracis* of lambs raised and finished on pasture only,
raised on grass and finished in stalls for 22 or 41 d, or raised and finished indoors (in stalls)
on concentrates and hay only. They attributed the lack of differences between treatments to
similarity in energy expenditure between animals and a higher rate of gain from good quality
grass.

231 For protein, there was a diet \times duration interaction whereby the muscle from the S 232 group had lower protein content than that of the SC and C groups at 54 d and 72 d, but there were no differences due to diets at 36 d (Supplementary Table S1). The lower protein content 233 of the lamb muscle from the S group may be explained by the fact that concentrate diets have 234 higher dry matter and crude protein content than silage-based diets (Warren et al., 2008); 235 however, this was more noticeable when the feeding duration increased to 54 and 72 days. In 236 addition, there were differences due to duration in the C group, whereby the muscle of the 54 237 d and 72 d groups had higher protein content than the 36 d group. In general, concentrate-238 based diets favour the production of propionate leading to increased insulin secretion and 239 stimulation of protein and fat synthesis in muscle (Weekes, 1986). Muscle from lambs 240 receiving the experimental diets for 36 and 54 d duration had higher muscle ash content (P 241 <.05) than lambs fed for the 72 d duration, although there was a diet \times duration interaction 242 whereby the SC group at 54 d had higher ash content than the S and C groups. 243

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245 **3.2 Effect of diet on the sensory and volatile profiles of lamb meat**

In general, a limited effect of the different dietary treatments on the 38 sensory descriptors was noted (only seven were significantly affected; P < .05) (Table 2). For three of these (*Animal/Farm Smell, Woolly Aroma* and *Fattiness*) there were diet \times duration

interactions which are discussed in the next section (3.3). Intensity of Lamb Aroma, Dry 249 Aftertaste and Astringent Aftertaste scored higher (P < .05) in the C group compared to the S 250 251 and SC groups. Farmyard Flavour scored lower (P < .05) in the SC group compared to the C group, but was similar to S group. Although significant effects on sensory descriptors were 252 few, lamb from animals fed the SC group received lower scores (P = .015-0.078) for 253 attributes that may be considered hedonically negative by some consumers (i.e. Animal/Farm 254 255 Smell, Woolly Aroma, Manure/Faecal Aroma, Off-flavours) (Table 2) although no consumer evaluation was performed in this study. Similar conclusions regarding lamb meat assessed by 256 257 European consumers was reported by Font i Furnols et al. (2009) where meat from lambs fed concentrate or a mixture of pasture and concentrate was more acceptable compared to meat 258 from lambs at pasture. Specifically, the meat from lambs fed a mixture of pasture (6% of live 259 260 weight, LW) and concentrate (1.2% of LW) was the most acceptable. Arsenos et al. (2002) showed that meat from lambs fed lucerne hay with low and medium levels of concentrate was 261 preferred more than meat from lambs fed high levels of concentrates. Other studies have 262 reported bigger differences when comparing grass-based systems with concentrate-based 263 system (Priolo et al., 2002; Resconi, Campo, Furnols, Montossi, & Sanudo, 2009), with 264 concentrate-fed lambs having more intense lamb odour and/or flavour than grass or forage-265 fed lambs but also higher acceptability (Borton, Loerch, McClure, & Wulf, 2005; Resconi et 266 al., 2009; Schreurs, Lane, Tavendale, Barry, & McNabb, 2008). 267

The volatile analysis showed that only ten volatile compounds were significantly (P <.05) affected by diet (Table 3), seven of which showed diet \times duration interactions (described in section 3.3). The SC and C groups had higher (P < .05) values for dimethyl sulphide (formed through Strecker degradation of methionine (Bailey, Rourke, Gutheil, & Wang, 1992)), than the S diet. Levels of hexanal (a compound that derives from oxidation of linoleic acid in muscle (C18:2n-6) (Elmore et al., 2005)), increased gradually with increasing

dietary concentrate although only the C and S groups were significantly different from each 274 other (P < .05). This could be due to the higher proportion (%) of C18:2n-6 in the C group 275 compared to the other groups (Rowe, Macedo, Visentainer, Souza, & Matsushita, 1999). 276 Muscle from lambs fed the S diet had higher values (P < .05) for skatole than the SC and C 277 diets. Skatole (which has a "faecal/manure aroma") derives from the degradation of dietary 278 tryptophan and since lush pasture is a source of more readily degradable protein than cereal 279 280 concentrates, it is also a possible source of tryptophan (Tavendale, Lane, Schreurs, Fraser, & Meagher, 2006). In addition, pasture-based diets have a high ratio of protein to readily 281 282 fermentable carbohydrate (Schreurs et al., 2008; Young et al., 2003). This may explain the higher levels of skatole in muscle from animals on the S group compared the other groups. 283 Priolo et al. (2004) reported differences in p-cymene and eight sesquiterpenes among lambs 284 fed either on grass or on concentrates for different periods while <u>Resconi et al. (2010)</u> found 285 that lambs fed only on pasture had lower levels of carbonyl compounds (alkanals, 286 alkadienals, ketones, strecker aldehydes) than those fed on grass with a concentrate 287 supplement, or only with concentrate. 288

Multivariate analysis techniques were applied to investigate potential differences 289 between groups and associations with the sensory and volatile data. Following discriminant 290 analysis of the sensory data, the first component (F1) explained 58.87% of the variation and 291 the second component (F2) explained 41.13% of the variation (Fig. 1). The centroids of the 292 dietary treatments were placed in different quadrants (Fig. 1a), revealing some associations 293 with some sensory attributes (Fig. 1b). The factor loadings of the sensory attributes that were 294 considered significant were higher than 0.30. In general, the overlapping of the groups 295 confirmed that the sensory profile of the lambs fed on different diets was similar. Also, the P 296 values from Wilk's Lambda test, Pillai's trace test and Roy's greatest root test showed that the 297 mean vectors only approached significance (range P = .06-0.10). Nevertheless, the C group 298

(centroid located in the upper right quadrant) was more associated with Dry Aftertaste and 299 Astringent Aftertaste. The S group (centroid located in the upper left quadrant) was more 300 301 associated with *Fattiness*. For the SC group (centroid in the bottom left quadrant), although visually it was associated with Juiciness, Intensity of Roast Meat Aroma and Intensity of Roast 302 *Meat Flavour*, the factor loadings of these attributes were ≤ 0.30 . However, it is clear that the 303 SC group was not associated with attributes that may be viewed as undesirable (i.e. 304 305 Manure/Faecal Aroma, Animal/Farm Smell, Off-flavours, Farmyard Flavour; factor loadings >0.30 for F2). 306

307 The discriminant analysis plot of the volatile data (Fig. 2) showed that the three groups (S, SC and C) were clearly separated. The first component (F1) explained 73.04% of 308 the variation and the second component (F2) explained 26.96 % of the variation. The factor 309 loadings of the volatile compounds that were considered significant were equal or higher than 310 0.30. The P values from Wilk's Lambda test, Pillai's trace test and Roy's greatest root test (P 311 <.001) indicate that at least one of the groups was different from another, whereby according 312 to the Fisher distances test the C group differed from the S group (P < .001) and from the SC 313 group (P = .001). For F1, the S and SC groups (both placed on the left side of the plot) were 314 separated from the C group. The compounds that contributed to this separation were 2,5-315 dimethylpyrazine, 2,6-dimethylpyrazine, 2-ethyl-3,6-dimethylpyrazine, (*E*,*Z*)-2,6-nonadienal, 316 pentadecane, hexadecane, and pentadecanol (factor loadings ≥ 0.3 for F1, data not shown). 317 The slight overlap of the S and SC groups indicated that their volatile profile had some 318 similarities. The results are in accordance with Vasta et al. (2011) who, through discriminant 319 analysis, showed that the volatile profile of meat from animals fed silage-based diets was 320 different from those on a concentrate-based diet suggesting that this could be due to the 321 presence of compounds in silage-based diets arising from bacterial fermentation of herbage 322 that makes the "volatile fingerprint" different. The second component separated the SC from 323

the C and S groups and the compounds that contributed to the variation were dimethyl sulfide 324 and indole (factor loadings ≥ 0.3 for F2). The differences in the volatile profile (Fig. 2) show 325 326 that both S and SC groups differed from the C group; however, the differences were not reflected in the sensory quality to a large extent as few differences were detected (Table 2). 327 The explanation could be that, while the volatile analysis showed 10 compounds to be 328 significantly affected, only seven (dimethylsulfide, hexanal, 2,6-nonadienal, indole, skatole, 329 330 2,5-dimethylpyrazine and 2-ethyl-3,6-dimethylpyrazine) have low odour thresholds and have been reported to be odour-active in previous lamb meat flavour studies (Gkarane et al., 2018). 331 332 Furthermore, only three out of the seven compounds (dimethylsulfide, hexanal and skatole) had a "clear" diet effect since the others had an interaction with duration. These compounds, 333 even if present at concentrations above the odour threshold, may not be adequate to elicit 334 significant sensory differences among diets which could explain the similarity in the sensory 335 profiles of the lambs on different diets. Another hypothesis is that the panellist's sensitivity 336 was insufficient to detect the differences in the aroma or that even if they detected them they 337 didn't score them very differently on the magnitude scale of 0-100. Thus, while the 338 discriminant analysis separated the lamb based on diets, it seems that there are limitations 339 that should be considered regarding the compounds that could ultimately influence flavour. 340

The fact that only few effects of dietary treatment on the volatile and sensory profiles 341 of lamb were noted in the present study is surprising given that differences in the fatty acid 342 343 profile of the lambs due to the different dietary treatments were present (unpublished results). For example, the C18:3 content was higher (P < .001) and the C18:2 content lower (P < .001) 344 in LTL from the S treatment compared to the C treatment while LTL from the SC treatment 345 had intermediate values (unpublished results). However, the lack of differences in IMF in this 346 study could explain the lack of differences in the volatile profile due to diets. According to 347 Vasta, D'Alessandro, Priolo, Petrotos, and Martemucci (2012) and Frank, Kaczmarska, 348

Paterson, Piyasiri, and Warner (2017) most of the odour-impact volatiles in meat systems are 349 lypophilic and their accumulation in animal tissue is correlated with the level of 350 intramuscular fat deposition. Furthermore, differences in flavour volatiles and/or fatty acid 351 composition following diet modification do not always have a major effect on sensory quality 352 as reported by Kitessa et al. (2009) and Muir, Deaker, and Bown (1998). It is also important 353 to recognise that the volatiles extracted by a static method headspace such as SPME may not 354 355 be representative of the headspace volatiles (considering that many factors (Jelen, Majcher, & Dziadas, 2012) influence the extracted compounds). Finally, the compounds detected by 356 357 SPME may not be perceived by trained panellists and the perception of trained panellists can't be equated to the perception of consumers (Munoz, 1998). 358

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360 3.3 Effect of finishing duration on the sensory and volatile profiles of lamb meat

Sensory analysis showed that only two attributes (*Animal/Farm Smell* and *Woolly Aroma*) were affected by finishing duration, both of which had a diet × duration interaction which will be described later in this section (Table 2). A recent study (Guerrero et al., 2018) also reported that feeding duration (30, 50 or 70 d) had a minor impact on sensory attributes of dry cured ham from culled ewes.

The volatile analysis showed that seven volatile compounds were affected (P < .05) by 366 the finishing duration, regardless of the finishing diet (Table 3). For four compounds (octanal, 367 nonanal, 1-octanol, and nonanoic acid) the 54 d group had higher levels (P <.05) than the 368 other two groups (36 d and 72 d) which did not differ from each other. For two compounds 369 370 (dodecanal and tridecanal) the 54 d group was higher (P < .05) than the 72 d group but both were similar to the 36 d group. For one compound (2-pentylfuran) values for the 54 and 72 d 371 372 groups were both higher (P < .05) than the 36 d group. This quadratic pattern (i.e. an increase 373 to 54 d and a decrease thereafter) could be attributed to a number of factors including the

different average daily gains and feed conversion efficiencies of the lambs. In the current 374 study the average live weights (and ages) of lambs assigned to the experimental diets (S, SC 375 and C) were 41.9 ± 2.4 kg (214 ± 5 d), 39.0 ± 5.2 kg ($204 \pm 5d$) and 38.9 ± 5.9 kg (197 ± 8 d) 376 for the 36, 54 and 72 d groups, respectively. These differences in maturity and associated 377 differences in average daily gain (ADG) and feed conversion efficiency (FCE), on 378 assignment to the experimental diets, may have contributed to the minor differences in 379 380 sensory character and volatiles. Similarly Arsenos et al. (2002) reported that lambs slaughtered at similar target slaughter weights may have differences in degree of maturity 381 382 which may impact on meat quality and consumer acceptability.

Studies indicate that regardless of diet there is a limit to daily intake in ruminants 383 (Allison, 1985; Caton & Dhuyvetter, 1997) after a defined period on a diet. A multitude of 384 factors can affect feed palatability in ruminants and, thus, voluntary feed intake and rate of 385 passage through the gut, including interactions between environmental conditions, animal 386 requirements (physiological or metabolic demands), physical characteristics of the diet 387 (composition, digestibility, energy density) and amount of protein which bypasses the rumen, 388 efficiency of microbial growth and extent of methane loss (Baumont, 1996; Caton & 389 Dhuyvetter, 1997; Decruyenaere, Buldgen, & Stilmant, 2009; Okine, Mathison, Kaske, 390 Kennelly, & Christopherson, 1998). These factors may in turn be influenced by feeding 391 duration with an ultimate effect on the lamb's metabolism and meat quality. 392

There were some interactions between diet and duration with respect to their effects on sensory and volatile profiles. The sensory analysis showed differences among groups at 54 d, whereby *Manure/Faecal Aroma* scores for the S group were higher (P < .05) than the scores for the SC and C groups, but there were no differences among groups at the other two feeding durations (Supplementary Table S2). In the S group specifically, scores of *Manure/Faecal Aroma* and *Woolly Aroma* for 54 d were higher (P < .05) than for 36 d and 72 399 d (P <.05) whereas for *Animal /Farm Smell* scores for 54 d were higher (P <.05) than 36 d but 400 similar (P >.05) to 72 d. For *Fattiness/Greasiness*, scores from the S group were higher (P 401 <.05) than the scores of SC and C groups only at 72 d.

There were ten significant (P < .05) diet \times duration interactions in the volatile analysis 402 (Supplementary Table S3). For (Z)-4-heptenal there were no differences due to duration in 403 the S group; however, in the SC group the 54 d value was higher (P < .05) than the 72 d value, 404 405 neither of which differed (P >.05) from the 36 d value, while in the C group the 36 d value was higher (P <.05) than both the 54 and 72 d values. In addition, there were differences due 406 407 to diet in the 72 d period, with S group having higher values than the C group and similar to the values of the SC group. For (E,Z)-2,6-nonadienal there were differences due to duration 408 only in the SC and C groups, whereby the 36 and 54 d values, which did not differ, were 409 410 higher (P <.05) than the 72 d values. A difference due to diet was found only for the 72 d group, whereby the S group had higher (P < .05) values than both the SC and C groups which 411 did not differ. These two aldehydes derive from linolenic acid (C18:3n-3) (Elmore et al., 412 2005), associated with grass-based diets (Enser et al., 1998), which could explain why levels 413 were lower with inclusion of concentrates for the longer (i.e. 72 d) finishing duration. 414

For 1-pentadecanol, the S and SC groups had higher (P < .05) values at 36 and 54 d, 415 which did not differ, compared to the C group; at 72 d values decreased (P <.05) from S to 416 SC to C group. Long-chain fatty alcohols, like pentadecanol, derive from wax ester 417 hydrolysis and are considered as diet biomarkers; notable differences in the alcohol content 418 of wax are found mainly among grasses and legumes (Kelman, Bugalho, & Dove, 2003), 419 which could explain the higher levels in muscle from the S and SC groups compared with the 420 C group, regardless of the finishing duration. For 2-heptanone, differences due to diet were 421 observed; thus, at 36 d values were lower (P <.05) in the S group than either the SC or C 422 groups, which did not differ. This compound was generally present at higher levels (although 423

424 not significant) in muscle from the SC and C groups at all finishing durations, probably 425 because it derives from C18:2n-6 (Elmore et al., 2005), which is associated with grain-based 426 diets (Enser et al., 1998). Differences in 2-heptanone due to feeding duration were significant 427 only in the S group, whereby values at 36 d were lower (P < .05) than either 54 or 72 d, which 428 did not differ.

Indole was detected at each duration of feeding in the S group, but only detected at 54 429 430 d in the SC group and at 72 d in the C group (Supplementary Table S3). The frequency of detection was higher in muscles from the S group since it derives from tryptophan 431 432 degradation in the rumen mainly of grass-fed lambs and has been identified with pastoral flavours (Schreurs et al., 2008). The higher scores for Woolly Aroma and Manure/Faecal 433 Aroma in muscle from the S54 group could be due to the higher levels of indole and skatole 434 (faecal, mothball-like aroma) compared to SC54 and C54 groups (although for skatole the 435 diet \times duration effect approached significance (P <.1)). 436

437 For 2,5-dimethyl pyrazine, values at 36 d in the S group were higher (P < .0.5) than in the C group, neither of which differed from the SC group; there were no statistical significant 438 differences (P > .05) due to dietary treatment at the other durations of feeding despite the fact 439 440 that the trend was similar (Supplementary Table S3). Differences due to feeding duration were significant only in the S group, whereby values at 36 d were higher (P < .05) than at 72 441 d, neither of which differed from 54 d. For 2-ethyl-3,6-dimethylpyrazine, values at 54 d in the 442 S group and SC groups were higher (P < .0.5) than those of the C group, while there were no 443 differences (P > .05) due to dietary treatment at the other durations of feeding (although a 444 similar trend was observed). Similar to 2,5-dimethylpyrazine, the S group, had higher (P 445 <.05) values at 36 d than 72 d, neither of which differed from values at 54 d. Muscle from 446 animals fed the S and SC diets had numerically higher levels for some pyrazines than the C 447 diet. This could be due to a possibly higher content of specific amino acids (e.g. cysteine, 448

glycine), that contribute to the Maillard reaction, in muscle from animals fed silage-based 449 diets as reported by other authors (Farmer, 1994). Koutsidis et al. (2008) reported a 450 significant effect of the diet (grass silage vs concentrate) on the concentration of free amino 451 acids (which can participate in the Maillard reaction) in bovine muscle, with animals fed 452 grass silage having higher levels than animals fed a concentrate diet. In addition, Tai and Ho 453 (1997) found that an oxidized cysteine/glucose reaction model produced more pyrazines and 454 455 furans as opposed to a non-oxidized cysteine/glucose reaction model that produced more sulphur compounds; thus, differences in susceptibility of muscle to oxidation may contribute 456 457 to differences in pyrazine formation.

For pentadecane there were differences among diets at all feeding durations whereby 458 the S group and SC groups, which did not differ, had higher (P <.05) values than the C group 459 460 (Supplementary Table S3). For hexadecane, differences among diets were found for all finishing durations whereby at 36 and 54 d the S group had higher levels (P < .05) than the C 461 group but both were similar (P >.05) to the SC group, while at 72 d the S and SC groups, 462 which did not differ, had higher values than the C group. Hydrocarbons like pentadecane and 463 hexadecane, are lipid oxidation compounds, and have been characterised as tracers of a 464 pasture diet in lamb (Sivadier, Ratel, & Engel, 2010); this could explain why levels were 465 lower with concentrate feeding at all durations. For 4-methyloctanoic acid differences among 466 diets were detected only at the 36 d of feeding duration with the S and SC groups, which did 467 not differ, having higher values (P < .05) than the C group. 468

In general, the majority of the aroma and flavour attribute scores as well as volatile compounds followed a quadratic pattern, i.e. values increased from 36 to 54 d and decreased from 54 to 72 d, mainly in S and SC groups (Supplementary Table S2 and Table S3). The PCA plot (Supplementary Fig. S1) for all nine groups (using only the aroma and flavour attributes and selected volatiles) explained 46.55% of the variance, whereby the first

component separated the three groups of 54 days duration (located on the right side of the 474 plot) from the other six groups (left side of the plot). The plot showed that S54 group was 475 characterised by the attributes "Manure/Faecal Aroma" and "Rancid Aroma", clustered with 476 skatole, indole, p-cresol, 4-heptenal, 2-nonenal, 2,6-nonadienal 2,5-dimethylpyrazine and 2-477 ethyl-3,6-dimethylpyrazine (factor loadings 0.6-0.8 on PC2). Previous studies have shown 478 that phenols and indoles (associated with animal-like odours) as well as 4-heptenal (Young, 479 480 Berdagué, Viallon, Rousset-Akrim, & Theriez, 1997; Young et al., 2003) and pyrazines (Bueno et al., 2013) have low odour thresholds and may be causally involved in lamb meat 481 482 aromas perceived by trained panellists. The SC54 group was characterised by "Animal/Farm Smell", "Woolly Aroma" and "Sweaty Aroma" (factor loadings 0.6-0.8 on PC1) (this 483 association is more meaningful when comparing SC54 group with SC36 or SC72 groups; See 484 supplementary Table S2). The compounds which may have contributed to these attributes 485 (Factor loadings 0.6-0.9 on PC1) were mainly lipid oxidation compounds (heptanal, 1-486 hexanol, 1-heptanol, octanal, 2-octenal, 1-octanol, nonanal, decanal, 2-decenal, 2,4-487 decadienal, 2-octen-1-ol) and other compounds e.g. a-terpineol, 2-pentylfuran, nonanoic acid, 488 benzaldehyde and phenylacetaldehyde, dimethyldisulfide and dimethyltrisulfide. 489

The results of the PCA plot could also be explained in part by the numerically higher (although not significant) proportions of C18:3n-3, eicosapentaenoic acid (EPA; C20:5n-3) and n-3 fatty acids in the S54 group compared to S36 and S72 groups, the higher proportion of arachidonic acid (C20:4n-6) of the SC54 compared to SC36 and SC72 groups and the higher level of PUFA of C54 group compared to C36 and C72 group (unpublished results).

495

496 **4. Conclusion**

497 When lambs receive different proportions of silage and concentrates for durations up 498 to approximately ten weeks pre-slaughter, effects on the sensory quality (and flavour volatiles) of lamb meat are relatively few. Some sensory attributes with potentially negative connotations (*Animal/Farm Smell*, *Manure/Faecal Aroma*, *Farmyard Flavour*, *Off-flavours*) appear to be lower when a mixed diet of silage and concentrate is fed. With regard to the duration of feeding, a diet composed of silage only, fed for an intermediate period, appears to be associated with less desirable sensory aroma attributes (*Manure/Faecal Aroma*, *Rancid Aroma*) which could be due to indoles or lipid oxidation compounds. Further research is required to establish if these small differences would influence consumer acceptability

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516

517 **Conflict of interest**

518 The authors have no conflicts of interest to declare.

519

520 Authors contributions

521 V. Gkarane conducted the experimental work, collected and statistically analysed the data
522 and drafted the manuscript. N. Brunton contributed to the method development for volatile
523 analysis and interpretation of the volatile results. P. Allen contributed to the study design and

524 sensory analysis. R. Gravador contributed to the sensory analysis and conducted the proximate analysis. N. Claffey contributed to the animal management and sample collection. 525 M. Diskin contributed to the study design and animal management oversight. A. Fahey 526 527 contributed to the study design and the univariate statistical analysis. L. Farmer contributed to the method development for volatile analysis and manuscript revision. A. Moloney 528 contributed to the study design. M. Alcalde contributed to the sensory analysis. P. Murphy 529 contributed to the multivariate statistical analysis. F. Monahan had overall responsibility for 530 the project, contributed to the study design, method development and interpretation of the 531 532 results. All authors read the manuscript and contributed to manuscript revisions.

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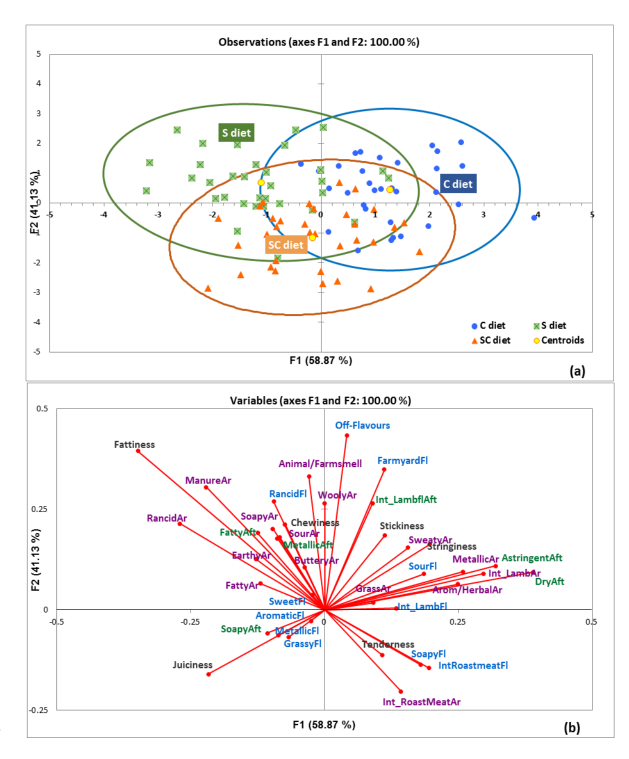


Fig. 1 Discriminant analysis (DA) plot (a) and variable loadings plot (b) of the sensory attributes of
lamb meat as affected by dietary treatments. Plot (a): S diet refers to the silage diet; SC diet refers to
the diet consisting of 50:50 silage:concentrate; C diet refers to the concentrate diet. Plot (b): "Ar",
"Fl" and "Aft" refer to Aroma, Flavour and Aftertaste attributes, respectively.

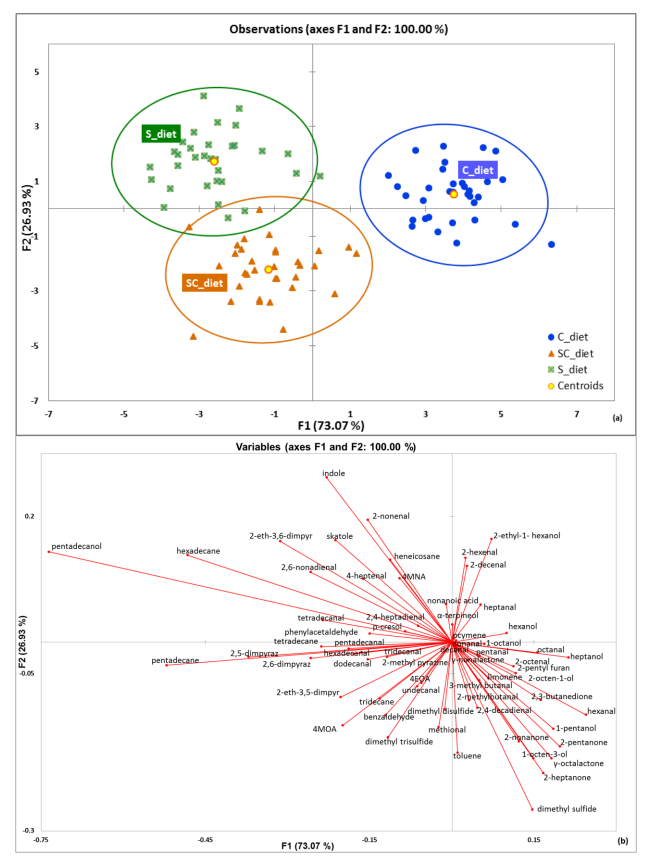


Fig. 2 Discriminant analysis (DA) plot (a) and variable loadings plot (b) of the volatile compounds as affected
by dietary treatment. Plot (a): S diet refers to the silage diet; SC diet refers to the diet consisting of 50:50
silage:concentrate; C diet refers to the concentrate diet. Plot (b) 4-MOA, 4-methyloctanoic acid in muscle; 4MNA, 4-methylnonanoic acid in muscle, 4-EOA; 4-ethyloctanoic acid in muscle; 2-eth-3,6-dimpyr, 2-ethyl-3,6dimethylpyrazine; 2-eth-3,5-dimpyr, 2-ethyl-3,5-dimethylpyrazine; 2,5-dimethylpyrazine; 2,6dimpyraz, 2,6-dimethylpyrazine.

692 Table 1 Least square mean values for proximate analysis and ultimate pH (pHu) in *longissimus thoracis et lumborum* (LTL) muscle fed three different diets

		Diet		Fe	eding durati	on	SEM -	Significance			
	S	SC	С	36	54	72	SEM	Diet	Duration	Diet x Duration	
Moisture	73.9 ^b	73.1 ^a	73.3 ^{ab}	73.5	73.6	73.2	0.14	0.041			
Protein	21.1ª	21.9 ^b	22.0 ^b	21.5	21.6	21.8	0.10	< 0.001		0.001	
Fat	3.71	3.94	3.79	3.77	3.78	3.88	0.13				
Ash	1.05	1.09	1.05	1.10 ^b	1.08 ^b	1.01 ^a	0.01		0.001	0.023	
pHu	5.73	5.71	5.78	5.69	5.78	5.76	0.02				

693 (100% Silage (S); 50% Silage: 50% Concentrate (SC); 100% Concentrate (C)) for three durations of feeding (36, 54, 72 days).

695 a,b Within row, means assigned different superscripts indicate differences among diets (S vs SC vs C) or durations (36 vs 54 vs 72 days).

697	Table 2 Least square mean scores for sensory attributes in grilled LTL muscle as affected by diet (100% Silage (S); 50% Silage: 50% Concentrate (SC);
698	100% Concentrate (C)) and duration of feeding (36, 54, 72 days).

		Diet			Duration		SEM		Significance	l
Sensory	Silage (S) 100%	50% (S) - 50% (C)	Concentrate (C) 100%	36	54	72		Diet	Duration	Diet × Duratior
Aroma										
Intensity of roast meat aroma	41.3	44.8	43.8	44.4	41.7	43.8	0.93			
Intensity of lamb aroma	39.8 ^a	40.1 ^a	43.1 ^b	40.4	40.5	42.0	0.71	0.036		
Grassy	7.5	7.6	7.9	7.8	7.3	8.0	0.28			
Aromatic/herbal	11.8	12.0	13.3	13.2	11.3	12.7	0.38			
Metallic/bloody	14.0	14.0	15.3	14.2	14.0	15.1	0.34			
Animal/farm Smell	15.5 ^b	12.8^{a}	15.1 ^b	12.1ª	16.6 ^b	14.7 ^b	0.60	0.039	0.007	0.032
Woolly	14.3 ^b	12.5 ^a	14.4 ^b	11.9ª	16.2 ^b	13.0 ^b	0.56	0.045	0.007	0.038
Buttery	7.0	6.8	7.1	6.8	7.3	6.8	0.30			
Fatty	8.2	8.0	7.9	7.6	8.2	8.3	0.33			
Rancid	8.0	6.2	6.3	6.6	7.5	6.4	0.43			
Manure/faecal	9.8	6.8	7.7	6.9	10.1	7.4	0.55			0.016
Sour	7.8 ^b	6.6^{a}	7.2 ^{ab}	6.0	8.7	7.0	0.43	0.078*		
Sweaty	14.9	14.5	16.2	14.0	16.5	15.1	0.49			
Soapy	3.7	3.2	3.4	3.5	3.3	3.5	0.16			
Earthy	10.5	9.9	10.0	9.8	10.2	10.4	0.27			
Flavour										
Intensity of roast meat flavour	36.9	39.3	39.4	39.5	37.0	39.0	0.78			
Intensity of lamb flavour	r 42.9	43.5	42.9	43.9	42.8	44.1	0.70			
Grassy	8.3	8.4	8.0	7.8	8.2	8.6	0.24			
Metallic/bloody	20.2	20.6	19.8	20.3	20.7	19.6	0.49			
Aromatic/herbal	9.4	9.3	9.2	8.8	9.1	10.0	0.27			

Soapy	5.2	6.2	6.3	5.3	6.6	5.9	0.28		
Rancid	8.5	6.8	7.8	7.0	8.5	7.6	0.41		
Farmyard	8.9^{ab}	7.3 ^a	9.9 ^b	8.3	8.9	8.9	0.47	0.015	
Sour	7.9	8.2	9.5	9.4	8.3	7.9	0.45		
Sweet	11.4	11.3	11.4	10.8	11.2	12.2	0.39		
Off-flavours	19.6 ^b	15.8 ^a	19.7 ^b	18.9	18.7	17.5	0.67	0.066*	
Texture									
Tenderness	54.4	58.0	57.5	56.7	57.7	55.6	1.57		
Juiciness	48.4	49.1	45.7	47.8	46.1	49.3	0.81		
Chewiness	51.9	46.7	49.5	49.8	47.6	50.7	1.47		
Fattiness/greasiness	30.7	25.5	26.4	27.1	28.4	27.2	0.65	0.003	0.044
Stringiness/fibrousness	33.8	32.5	37.7	36.7	34.1	33.3	1.29		
Stickiness	26.8	25.7	27.9	27.2	27.2	25.9	0.63		
Aftertaste									
Intensity of lamb aftertaste	34.1	32.9	34.7	34.6	33.5	33.7	0.43		
Soapy	9.3	9.5	8.9	8.6	9.4	9.6	0.31		
Metallic/ bloody	20.8	19.1	19.9	19.6	19.4	20.7	0.49		
Fatty/ greasy	17.7	15.9	16.5	16.7	17.4	15.9	0.48		
Dry	11.3ª	11.8 ^a	13.5 ^b	12.6	12.2	11.7	0.34	0.009	
Astringent	7.2ª	7.6 ^a	9.3 ^b	7.6	8.3	8.2	0.35	0.030	

699 a,b within row, different superscripts indicate differences among diets (S vs SC vs C) or durations (36 vs 54 vs 72 days).

¹Probability of significance for the main effects of diet, duration and diet x duration tested using the MIXED model (P < .05).

*P <.1.

Table 3 Least square mean values for logarithmically transformed peak areas of aroma compounds detected in the headspace of grilled *longissimus thoracis et*

lumborum (LTL) muscle fed three different diets (100% Silage (S); 50% Silage: 50% Concentrate (SC); 100% Concentrate (C)) for three durations (36, 54,

706 72 days).

Volatile compound	LRI ¹	Ions	Method of		Diet		Feeding	g Duration	(days)	SEM		Significance	
Ĩ		Used ²	Identification ³	S	SC	С	36	54	72	-	Diet	Duration	Diet × Duration
Sulphur													
compounds													
Dimethyl sulfide		63,62,61	NIST, Std, LRI	1.97ª	2.90 ^b	2.70 ^b	2.73	2.63	2.22	0.151	0.029		
Dimethyl disulfide	719	94,79	NIST, Std, LRI	2.34	2.71	2.41	2.42	2.89	2.16	0.176			
Dimethyl trisulfide	963	126	NIST, Std, LRI	4.27	4.36	4.21	4.25	4.39	4.21	0.037			
Aldehydes													
2-Methylbutanal		39,41,57	NIST, Std, LRI	4.17	4.36	4.29	4.40	4.28	4.15	0.095			
3-Methylbutanal		41,43,58	NIST, Std, LRI	4.33	4.47	4.47	4.57	4.47	4.24	0.097			
Pentanal		43,44,58	NIST, Std, LRI	4.26	4.32	4.36	4.39	4.42	4.13	0.082			
(E)-2-Hexenal	849	39,41,55	NIST, Std, LRI	3.01	2.65	2.97	3.12	2.70	2.82	0.139			
Hexanal	800	39,41,56	NIST, Std, LRI	5.26ª	5.37 ^{ab}	5.45 ^b	5.32	5.43	5.33	0.031	0.044		
Methional	905	48,104	NIST, Std, LRI	3.82	4.13	3.86	4.13	3.93	3.76	0.117			
(<i>E</i> , <i>E</i>)-2,4-	1008	81,53	NIST, Std, LRI	2.20	2.04	1.91	2.23	2.25	1.66	0.194			
Heptadienal													
(Z)-4-Heptenal	898	67,39,55	NIST, Std, LRI	4.08	4.00	3.96	4.08 ^b	4.06 ^b	3.90ª	0.029		0.013	0.007
Heptanal	900	39,41,70	NIST, Std, LRI	5.33	5.30	5.35	5.32	5.40	5.26	0.025			
(E)-2-Octenal	1056	39,55,83	NIST, Std, LRI	4.40	4.44	4.48	4.40	4.51	4.41	0.027			
Octanal	1002	41,67,69	NIST, Std, LRI	5.41	5.44	5.50	5.42 ^a	5.55 ^b	5.39ª	0.024		0.012	
(<i>E</i> , <i>Z</i>)-2,6-	1150	41,69,70	NIST, Std, LRI	4.10 ^b	4.01 ^b	3.93 ^a	4.06	4.09	3.89	0.028	0.026	0.007	0.003
Nonadienal													
(E)-2-Nonenal	1158	29,41,55	NIST, Std, LRI	4.99	4.86	4.86	4.92	4.96	4.82	0.028			
Nonanal	1101	69,81,57	NIST, Std, LRI	6.02	6.03	6.03	6.01 ^a	6.13 ^b	5.95ª	0.023		0.006	
(E,E)-2,4-	1315	81,67	NIST, Std, LRI	3.98	4.05	4.03	3.97	4.09	4.01	0.031			
Decadienal													
(E)-2-Decenal	1260	39,81,55	NIST, Std, LRI	4.47	4.40	4.46	4.42	4.53	4.37	0.029			
Decanal	1204	41,67,55	NIST, Std, LRI	4.83	4.83	4.84	4.82	4.89	4.79	0.021			
Undecanal	1306	41,67,81	NIST, Std, LRI	3.90	4.04	3.78	3.66	3.97	4.09	0.126			
Dodecanal	1406	41,67,81	NIST, Std, LRI	4.43	4.43	4.36	4.39 ^a	4.48 ^b	4.35 ^a	0.022		0.044	
Tridecanal	1510	41,67,81	NIST, LRI	4.48	4.47	4.40	4.46 ^a	4.52 ^b	4.37ª	0.024		0.042	

Tetradecanal Pentadecanal Hexadecanal	1705	41,67,81 41,67,81 41,67,81	NIST, LRI NIST, LRI NIST, LRI	4.97 5.07 5.67	4.93 5.05 5.65	4.85 4.96 5.55	4.90 5.02 5.62	4.97 5.09 5.66	4.87 4.97 5.59	0.023 0.026 0.030			
Alcohols 1-Pentanol	800	41,55,70	NIST, Std, LRI	3.24	3.73	3.88	3.67	3.63	3.55	0.132			
1-Hexanol		41,55,70	NIST, Std, LRI	5.24 4.35	4.35	5.88 4.40	3.07 4.35	3.03 4.44	3.33 4.31	0.132			
1-Heptanol		41,56,59	NIST, Std, LRI	4.55 4.48	4.55 4.52	4.40 4.60	4.55 4.51	4.44 4.60	4.31 4.49	0.028			
1-Octen-3-ol		43,57,69	NIST, Std, LRI	4.40 4.81	4.92	4.00	4.31	4.00	4.49	0.023			
2-Octen-1-ol		41,57,67	NIST, Std, LRI	3.97	4.00	4.01	3.93	4.05	4.01	0.027			
2-Ethyl-1-hexanol		41,57,67	NIST, Std, LRI	4.43	4.00	4.46	4.27	4.48	4.01	0.024			
1-Octanol		41,55,69	NIST, Std, LRI	4.43 5.11	5.12	4.40 5.14	4.27 5.10 ^{ab}	4.48 5.22 ^b	4.41 5.04 ^a	0.033		0.008	
α-Terpineol		41,55,69 93,59,121	NIST, Std, LRI	4.91	4.89	4.91	4.87	3.22 4.95	4.88	0.023		0.008	
1-Pentadecanol		69,83,97	NIST, Std, LRI	4.91 5.57°		4.91 5.00 ^a	5.31	5.31	4.88 5.29	0.030	< 0.001		< 0.001
		, ,	, ,										
Ketones													
2-Pentanone		43,71,86	NIST, Std, LRI	0.70	1.42	1.59	1.24	1.45	1.02	0.166			
2,3-Butanedione		43	NIST, Std, LRI	2.59	3.13	3.42	2.99	3.35	2.80	0.202			
2-Heptanone	887	43,58	NIST, Std, LRI	3.73	4.04	4.03	3.74	4.07	3.99	0.062			0.029
2-Nonanone		43,58	NIST, Std, LRI	3.91	4.01	4.00	3.92	4.04	3.95	0.027			
γ-Octalactone		85,57	NIST, Std, LRI	1.11	1.82	1.89	1.15	1.87	1.80	0.153			
γ-Nonalactone	1356	85,29	NIST, Std, LRI	3.06	3.12	3.13	2.91	3.22	3.18	0.089			
Terpenes													
p-cymene	1020	119,91	NIST, Std, LRI	2.81	2.80	2.82	2.51	3.06	2.87	0.122			
Limonene		67,68,93	NIST, Std, LRI	4.27	4.31	4.32	4.25	4.38	4.27	0.029			
Phenols													
p-Cresol	1071	107,108	NIST, Std, LRI	3.32	3.19	2.99	3.46	3.17	2.88	0.171			
p-cressi	1071	107,100	NIST, SIG, LICI	5.52	5.17	2.))	5.40	5.17	2.00	0.171			
Indoles													
Indole		117,89	NIST, Std, LRI	0.64 ^b		0.07ª	0.09	0.54	0.17	0.086	0.006		0.021
Skatole (3-methyl	1379	130,131	NIST, Std, LRI	1.11 ^b	0.51 ^{ab}	0.34 ^a	0.59	1.05	0.32	0.137	0.048	0.080*	0.063*
indole)													
Pyrazines													
2-Methyl pyrazine	822	94,67	NIST, Std, LRI	1.09	1.07	0.65	0.94	1.23	0.64	0.183			
2,5-Dimethyl	909	108,42	NIST, Std, LRI	2.44 ^b	2.14 ^b	0.68ª	2.27	1.55	1.45	0.222	0.002		0.022
pyrazine													

2,6-Dimethyl	909	108,42	NIST, Std, LRI	4.27 ^b	4.11 ^{ab}	3.09ª	3.58	4.05	3.85	0.216	0.055*		
pyrazine 2-Ethyl-3,5-	1071	135,134	NIST, Std, LRI	4.86	4.93	4.54	4.83	4.81	4.69	0.084			
dimethyl-pyrazine	1071	155,154	MIST, SIG, LKI	4.80	4.95	4.34	4.85	4.81	4.09	0.084			
2-Ethyl-3,6-	1083	135,136	NIST, Std, LRI	2.69 ^b	1.66 ^{ab}	1.03ª	2.45 ^b	1.92 ^{ab}	1.01 ^a	0.213	0.003	0.012	0.003
dimethyl-pyrazine													
Benzenoid													
compounds													
Benzaldehyde	957	105,77	NIST, Std, LRI	6.19	6.25	6.12	6.15	6.29	6.12	0.034			
Phenyl	1039	91,92	NIST, Std, LRI	4.97	4.94	4.86	4.90	4.99	4.89	0.033			
acetaldehyde													
Toluene	748	91,92	NIST, Std, LRI	4.68	4.82	4.73	4.72	4.83	4.67	0.038			
Furans													
2-Pentylfuran	987	81,138,53	NIST, Std, LRI	4.03	4.25	4.43	3.66ª	4.46 ^b	4.60 ^b	0.139		0.010	
Hydrocarbons													
Tridecane		41,57,71	NIST, Std, LRI	4.52	4.55	4.46	4.49	4.57	4.47	0.028			
Tetradecane		41,57,71	NIST, Std, LRI	4.57	4.54	4.44	4.48	4.60	4.47	0.025			
Pentadecane		41,57,71	NIST, Std, LRI	4.88 ^b	4.84 ^b	4.62 ^a	4.77	4.82	4.75	0.023	< 0.001		0.000
Hexadecane		41,57,71	NIST, Std, LRI	4.57 ^b	4.46 ^b	4.33ª	4.47	4.49	4.41	0.021	< 0.001		0.000
Heneicosane		41,57,71	NIST, Std, LRI	0.98	0.59	0.58	0.52	0.96	0.67	0.122			
BCFAs													
4-Methyloctanoic	1232	55,57,73	NIST, Std, LRI	1.31	1.58	0.76	1.18	1.44	1.04	0.157			0.048
acid		, ,	, ,										
4-Ethyloctanoic	1313	55,57,71	NIST, Std, LRI	1.94	2.12	1.81	2.34	1.42	2.11	0.168			
acid													
4-Methylnonanoic	1323	55,57,71	NIST, Std, LRI	1.26	0.89	0.86	0.98	0.76	1.27	0.149			
acid													
Organic acids													
Nonanoic acid	1275	60	NIST, Std, LRI	3.74	3.67	3.71	3.58ª	3.89 ^b	3.65 ^{ab}	0.051		0.029	
^{a,b} Within row means	assigne	d different su	persorints indicate diffe	arences a	mong diet	s (S vs SC vs C) or duration	ne (36 ve 5/	ve 72 dave)				

^{a,b} Within row, means assigned different superscripts indicate differences among diets (S vs SC vs C) or durations (36 vs 54 vs 72 days).

 ¹ Linear retention indices (LRI) calculated from the n-alkanes (C7-C30) run under the same GC-MS conditions as LTL muscle samples.
 ² Specific ions used for volatile identification and peak area integration.
 ³ Method of identification: NIST (NIST library), Std (authentic standard) and LRI.

710

* P <.1 712 Supplementary Table S1 Least square mean values for proximate analysis and ultimate pH (pHu) in *longissimus thoracis et lumborum* (LTL) muscle fed three

713 different diets (100% Silage (S); 50% Silage: 50% Concentrate (SC); 100% Concentrate(C)) for three durations of feeding (36, 54, 72 days).

714

		100 % \$	5		SC			100% C	,		p-values			
	Fee	eding dur	ation	Fe	eding dur	ation	Fee	eding dura	ation					
	36	54	72	36	54	72	36	54	72	SEM	Diet	Duration	Diet x Duration	
Moisture	73.4	74.3	74.1	73.4	72.9	72.9	73.7	73.6	72.8	0.14	0.045			
Protein	21.3	20.9 ^x	21.1 ^x	21.9	21.7 ^y	22.1 ^y	21.4 ^a	22.2 ^{by}	22.3 ^{by}	0.10	< 0.001		0.001	
Fat	4.04	3.48	3.6	3.51	4.32	3.98	3.75	3.54	4.07	0.13				
Ash	1.09	1.05 ^x	1.02	1.14	1.10 ^y	1.03	1.08	1.08 ^{xy}	1.00	0.01		0.001	0.023	
pH_{u}	5.61	5.84	5.75	5.68	5.75	5.69	5.77	5.74	5.82	0.02				

715

^{a,b,c} Within row, means assigned different superscripts indicate differences between durations within each diet (i.e. S36 *vs* S54 *vs* S72, SC36 *vs* SC54 *vs* SC72
 and C36 *vs* C54 *vs* C72).

718 ^{x,y,z} Within row, means assigned different superscripts indicate differences between diets with finishing duration 36 or 54 or 72 days (i.e. S36 vs SC36 vs C36 \sim SC36 \sim SC36 vs C36 \sim SC36 \sim SC36

719 or S54 *vs* SC54 *vs* C54 or S72 *vs* SC72 *vs* C72).

50% Supplementary Table S2 Least square mean scores for sensory attributes in grilled LTL muscle as affected by (100% Silage (S); 50% Silage: 50%

722 Concentrate (SC); 100% Concentrate (C)) and duration of feeding (36, 54, 72 days).

723

		100 % S		5	0%S:50%	ъC		100% C		SEM	Significance ¹		
	Fee	ding dura	tion	Fee	ding dura	tion	Feed	ling durat	tion			-	
	36	54	72	36	54	72	36	54	72		Diet	Duration	Diet x Duration
Aroma													
Intensity of roast meat aroma	43.6	41.6	38.9	46.3	43.3	44.8	43.9	39.6	47.9	0.93			
Intensity of lamb aroma	39.9	39.7	39.9	40.8	39.4	40.1	41.0	42.1	46.1	0.71	0.036		
Grassy	7.4	6.6	8.6	8.3	6.5	8.0	7.6	8.7	7.4	0.28			
Aromatic/herbal	13.2	9.2	12.9	12.5	12.0	11.5	13.2	12.8	13.8	0.38			
Metallic/bloody	14.4	13.0	14.5	13.3	13.5	15.3	14.9	15.5	15.6	0.34			
Animal/farm Smell	12.7 ^a	18.9 ^b	14.9 ^{ab}	10.7	14.4	13.3	13.2	16.3	15.8	0.60	0.039	0.007	0.032
Woolly	12.7ª	17.3 ^b	13.0 ^a	11.4	14.6	11.5	12.0	16.6	14.6	0.56	0.045	0.007	0.038
Buttery	6.2	8.0	6.8	6.4	7.1	7.0	8.0	6.8	6.5	0.30			
Fatty	7.5	8.9	8.4	7.7	8.9	7.5	7.8	6.8	9.1	0.33			
Rancid	8.0	8.6	6.5	5.8	7.4	5.6	5.9	6.5	6.3	0.43			
Manure/faecal	8.5 ^a	14.1^{by}	7.0 ^a	5.4	7.3 ^x	7.9	7.4	8.4 ^x	7.3	0.55			0.016
Sour	7.0	10.2	6.3	5.6	7.6	6.8	5.9	7.8	8.1	0.43	0.078*		
Sweaty	14.3	16.5	14.1	13.1	16.0	14.3	14.8	16.9	16.9	0.49			
Soapy	4.1	3.3	3.6	2.9	7.6	6.8	3.4	7.8	8.1	0.16			
Earthy	10.6	9.9	11.0	9.5	10.4	9.8	9.0	10.3	10.6	0.27			
Flavour													
Intensity of roast meat flavour	37.4	36.7	36.7	39.9	36.8	41.2	41.8	37.5	38.9	0.78			

Intensity of lamb flavour	43.5	42.1	43.1	44.4	41.5	44.5	43.8	44.8	44.7	0.70		
Grassy	7.8	8.1	9.0	8.1	8.1	9.1	7.7	8.2	8.3	0.24		
Metallic/bloody	21.8	19.3	19.5	19.8	21.5	20.5	18.9	21.2	19.3	0.49		
Aromatic/herbal	9.1	8.6	10.4	8.5	9.1	10.3	8.6	9.6	9.3	0.27		
Soapy	4.9	5.3	5.4	5.4	7.2	6.2	5.3	7.2	6.4	0.28		
Rancid	8.2	8.5	8.6	5.5	7.8	7.1	7.2	9.2	7.0	0.41		
Farmyard	7.9	10.5	8.4	7.4	7.0	7.6	10.4	9.0	10.4	0.47	0.015	
Sour	7.8	8.2	7.7	9.1	7.8	7.7	11.4	8.9	8.3	0.45		
Sweet	11.9	10.4	12.0	10.7	10.9	12.4	9.6	12.3	12.3	0.39		
Off-flavours	19.9	21.2	17.8	15.7	15.5	16.2	21.2	19.3	18.5	0.67	0.066*	
Texture												
Tenderness	58.4	59.1	45.6	54.9	56.0	63.0	56.7	57.8	58.1	1.57		
Juiciness	49.8	45.1	50.3	50.6	46.5	50.2	43.0	46.6	47.3	0.81		
Chewiness	49.1	48.3	58.3	51.9	44.6	43.7	48.4	50.0	50.1	1.47		
Fattiness/greasiness	29.6	31.1	31.5 ^y	25.4	26.1	25.1 ^x	26.0	28.1	25.0 ^x	0.65	0.003	0.044
Stringiness/	34.2	30.5	36.6	36.7	32.8	28.1	39.1	38.9	35.0	1.29		
fibrousness	777	27.2	25 4	26.6	22.0	267	27.5	20.5	25.7	0.62		
Stickiness	27.7	27.2	25.4	26.6	23.9	26.7	27.5	30.5	25.7	0.63		
Aftertaste												
Intensity of lamb aftertaste	34.9	33.1	34.1	33.8	32.6	32.3	34.8	34.7	34.7	0.43		
Soapy	8.3	9.8	9.8	9.0	9.2	10.2	8.7	9.2	8.8	0.31		
Metallic/ bloody	21.8	18.0	22.4	18.3	19.6	19.3	18.0	20.7	20.7	0.49		
Fatty/ greasy	16.8	18.3	17.8	16.2	16.2	15.4	16.9	18.2	14.2	0.48		
Dry	11.7	12.2	9.9	12.0	11.6	11.9	14.6	12.7	13.4	0.34	0.009	
-												

Astringent 6.3 8.1 7.3 7.0 8.0 8.0 9.7 8.8 9.3 0.35 0.	.030
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¹Probability of significance for the main effects of diet, duration and diet x duration tested using the MIXED model (P < 0.05)

^{a,b,c} within row, different superscripts indicate differences between durations within each diet (i.e. S36 vs S54 vs S72, SC36 vs SC54 vs SC72 and C36 vs C54 vs C72) (P < 0.05).

x,y,z within row, different superscripts indicate differences among diets with finishing duration 36 or 54 or 72 days (i.e. S36 *vs* SC36 *vs* C36 or S54 *vs* SC54 *vs* SC54 *vs* SC54 *vs* SC54 or S72 *vs* SC72 *vs* C72) (P < 0.05).

Supplementary Table S3 Least square mean values for logarithmically transformed peak areas of aroma compounds detected in the headspace of grilled
 longissimus thoracis et lumborum (LTL) muscle fed three different diets (100% Silage (S); 50% Silage: 50% Concentrate (SC); 100% Concentrate (C)) for

three durations of feeding (36, 54, 72 days).

	LRI	LRI 100% S			50	0%S:50%	С	100% C			SEM		Significand	ce
		Feeding Duration			Feed	ding Dura	tion	Feeding Duration						
		36	54	72	36	54	72	36	54	72		Diet	Duration	Diet × Duration
Sulphur compounds														
Dimethyl sulfide		2.45	1.89	1.57	2.63	3.12	2.95	3.10	2.87	2.14	0.151	0.029		
Dimethyl disulfide	719	2.67	2.39	1.97	2.60	3.55	2.00	1.99	2.73	2.50	0.176			
Dimethyl trisulfide	963	4.33	4.26	4.22	4.28	4.56	4.25	4.14	4.34	4.15	0.037			
Aldehydes														
2-Methylbutanal		4.42	4.13	3.96	4.40	4.22	4.47	4.38	4.48	4.01	0.097			
3-Methylbutanal		4.60	4.37	4.02	4.53	4.36	4.53	4.58	4.67	4.16	0.095			
Pentanal		4.33	4.50	3.95	4.41	4.15	4.39	4.43	4.60	4.06	0.082			
(E)-2-Hexenal	849	3.48	2.66	2.91	2.67	2.53	2.76	3.22	2.91	2.78	0.139			
Hexanal	800	5.17	5.31	5.31	5.34	5.47	5.31	5.46	5.50	5.39	0.031	0.044		
Methional	905	4.05	3.93	3.49	4.30	3.94	4.16	4.04	3.93	3.63	0.117			
(<i>E</i> , <i>E</i>)-2,4- Heptadienal	1008	1.70	2.19	2.71	2.94	2.14	1.03	2.06	2.41	1.25	0.194			
(Z)-4-Heptenal	898	4.03	4.12	4.09 ^y	4.05 ^{ab}	4.12 ^b	3.83 ^{xya}	4.17 ^b	3.94 ^a	3.78 ^{xa}	0.029		0.013	0.007
Heptanal	900	5.26	5.40	5.32	5.31	5.43	5.18	5.38	5.37	5.29	0.025			
(E)-2-Octenal	1056	4.36	4.43	4.41	4.44	4.54	4.33	4.39	4.55	4.48	0.027			
Octanal	1002	5.36	5.50	5.38	5.40	5.60	5.32	5.48	5.55	5.46	0.024		0.012	
(<i>E</i> , <i>Z</i>)-2,6- Nonadienal	1150	4.07	4.13	4.11 ^y	4.06 ^b	4.14 ^b	3.81 ^{xa}	4.03 ^b	3.99 ^b	3.76 ^{xa}	0.028	0.026	0.007	0.003

(E)-2-Nonenal	1158	4.96	5.05	4.96	4.89	4.97	4.71	4.92	4.87	4.80	0.028			
Nonanal	1101	5.99	6.10	5.99	6.00	6.20	5.89	6.03	6.09	5.97	0.023		0.006	
(<i>E</i> , <i>E</i>)-2,4- Decadienal	1315	3.93	4.02	3.99	4.07	4.12	3.98	3.91	4.13	4.05	0.031			
(E)-2-Decenal	1260	4.44	4.57	4.39	4.41	4.53	4.27	4.42	4.50	4.47	0.029			
Decanal	1204	4.77	4.90	4.81	4.84	4.91	4.75	4.84	4.86	4.80	0.021			
Undecanal	1306	3.51	3.52	4.23	3.95	3.98	4.18	3.52	3.97	3.84	0.126			
Dodecanal	1406	4.43	4.48	4.40	4.43	4.52	4.34	4.31	4.46	4.33	0.022		0.044	
Tridecanal	1510	4.51	4.51	4.43	4.47	4.59	4.36	4.40	4.46	4.33	0.024		0.042	
Tetradecanal	1607	4.99	4.98	4.95	4.93	5.02	4.84	4.80	4.92	4.83	0.023			
Pentadecanal	1705	5.11	5.09	5.01	5.06	5.14	4.95	4.90	5.03	4.95	0.026			
Hexadecanal	1818	5.73	5.62	5.66	5.66	5.72	5.57	5.45	5.65	5.56	0.030			
Alcohols														
1-Pentanol	809	3.91 ^x	2.50	3.31	3.38	4.20 ^y	3.60	3.73	4.19 ^y	3.73	0.132			
1-Hexanol	868	4.27	4.42	4.36	4.36	4.46	4.24	4.42	4.44	4.34	0.026			
1-Heptanol	969	4.43	4.54	4.47	4.52	4.63	4.40	4.57	4.63	4.60	0.025			
1-Octen-3-ol	980	4.77	4.79	4.87	4.92	5.03	4.85	4.83	5.02	4.94	0.027			
2-Octen-1-ol	1066	3.93	3.95	4.03	3.93	4.13	3.96	3.94	4.07	4.03	0.024			
2-Ethyl-1-hexanol	1027	4.36	4.47	4.46	3.98	4.50	4.33	4.48	4.46	4.44	0.053			
1-Octanol	1069	5.06	5.20	5.06	5.12	5.27	4.96	5.13	5.18	5.11	0.023		0.008	
α-Terpineol	1191	4.89	4.93	4.92	4.87	4.97	4.83	4.86	4.96	4.89	0.036			
1-Pentadecanol	1766	5.52 ^y	5.53 ^y	5.67 ^z	5.32 ^y	5.45 ^y	5.27 ^y	5.08 ^x	4.97 ^x	4.94 ^x	0.03	< 0.001		< 0.001
Ketones														
2-Pentanone		0.64	1.18	0.28	1.21	1.51	1.55	1.88	1.65	1.23	0.166			
2,3-Butanedione		2.50	3.43	1.85	2.34	3.15	3.91	4.15	3.48	2.63	0.202			
2-Heptanone	887	3.24 ^{xa}	3.96 ^b	3.98 ^b	4.01 ^y	4.13	3.98	3.96 ^y	4.11	4.02	0.062			0.029
2-Nonanone	1089	3.84	3.93	3.95	3.99	4.10	3.93	3.95	4.08	3.98	0.027			

γ-Octalactone γ-Nonalactone	1251 1356	0.83 2.73	1.44 3.43	1.07 3.02	1.17 2.99	2.53 3.13	1.77 3.26	1.47 3.01	1.65 3.11	2.56 3.27	0.153 0.089			
Terpenes p-cymene Limonene	1020 1024	2.72 4.24	2.45 4.31	3.24 4.25	2.37 4.29	3.38 4.40	2.67 4.24	2.43 4.21	3.35 4.45	2.69 4.30	0.122 0.029			
Phenols p-Cresol	1071	3.84	3.15	2.98	3.27	3.36	2.95	3.27	3.00	2.71	0.171			
Indoles Indole Skatole (3-methyl indole)	1287 1379	0.27 0.94	1.36 ^y 2.00	0.28 0.38	0.00 0.59	0.26 ^x 0.65	0.00 0.31	0.00 0.24	0.00 ^x 0.50	0.22 0.27	0.086 0.137	0.006 0.048	0.080*	0.021 0.063*
Pyrazines 2-Methyl pyrazine 2,5-Dimethyl pyrazine 2,6-Dimethyl pyrazine 2-Ethyl-3,5- dimethyl-pyrazine 2-Ethyl-3,6- dimethyl-pyrazine	822 909 909 1071 1083	1.21 3.60 ^{yb} 5.10 3.43 5.06 ^b	1.28 2.09 ^{ab} 3.66 3.13 4.56 ^{yab}	0.79 1.65 ^a 4.07 1.50 4.97 ^a	0.80 2.43 ^{xy} 3.31 1.99 4.87	1.67 2.08 5.05 2.25 5.09 ^y	0.75 1.91 3.96 0.75 4.83	0.80 0.79 ^x 2.34 1.94 4.55	0.75 0.48 3.44 0.38 4.80	0.39 0.78 3.51 0.76 4.27 ^x	0.183 0.222 0.216 0.213 0.084	0.002 0.055* 0.003	0.012	0.022
Benzenoid compounds Benzaldehyde Phenyl acetaldehyde	957 1039	6.20 4.98	6.18 4.97	6.18 4.97	6.21 4.92	6.41 5.04	6.12 4.85	6.05 4.79	6.27 4.96	6.05 4.84	0.034 0.033			

Toluene	748	4.69	4.69	4.65	4.77	4.95	4.72	4.69	4.85	4.65	0.038			
Etherocyclic compounds 2-Pentylfuran	987	3.72	3.79	4.59	3.41	4.79	4.56	3.84	4.79	4.66	0.139		0.010	
Hydrocarbons														
Tridecane		4.48	4.49	4.59	4.59	4.65	4.42	4.40	4.58	4.39	0.028			
Tetradecane		4.57	4.60	4.53	4.53	4.64	4.46	4.36	4.55	4.41	0.025			
Pentadecane		4.89 ^y	4.89 ^y	4.87 ^y	4.81 ^y	4.92 ^y	4.79 ^y	4.62 ^x	4.66 ^x	4.59 ^x	0.023	< 0.001		0.000
Hexadecane		4.60 ^y	4.56 ^y	4.56 ^y	4.48 ^{xy}	4.52 ^{xy}	4.39 ^y	4.34 ^x	4.39 ^x	4.28 ^x	0.021	< 0.001		0.000
Heneicosane		1.03	0.92	0.99	0.53	0.73	0.50	0.00	1.23	0.51	0.122			
BCFAs														
4-Methyl octanoic acid	1232	1.45 ^y	1.95	0.54	2.08 ^y	1.19	1.48	0.00 ^x	1.18	1.11	0.157			0.048
4-Ethyl octanoic acid	1313	2.21	1.20	2.42	2.98	1.91	1.47	1.81	1.16	2.45	0.168			
4-Methyl nonanoic acid	1323	0.98	1.46	1.35	0.81	0.24	1.61	1.15	0.59	0.85	0.149			
Organic acids														
Nonanoic acid	1275	3.66	3.90	3.66	3.38	3.96	3.68	3.70	3.82	3.60	0.051		0.029	
735														

^{a,b,c} Within row, means assigned different superscripts indicate differences between durations within each diet (i.e. S36 vs S54 vs S72, SC36 vs SC54 vs SC72 and C36 vs C54 vs C72).

^{x,y,z} Within row, means assigned different superscripts indicate differences among diets at finishing durations of 36 or 54 or 72 days (i.e. S36 vs SC36 vs C36 or S54 vs SC54 vs C54 or S72 vs SC72 vs C72).

740 * P <.1.