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1           **CONSUMER PALATABILITY SCORES AND VOLATILE BEEF FLAVOR**  
2           **COMPOUNDS OF FIVE USDA QUALITY GRADES AND FOUR MUSCLES**

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## Abstract

Proximate data, consumer palatability scores and volatile compounds were investigated for four beef muscles (*Longissimus lumborum*, *Psoas major*, *Semimembranosus* and *Gluteus medius*) and five USDA quality grades (Prime, Upper 2/3 Choice, Low Choice, Select, and Standard). Quality grade did not directly affect consumer scores or volatiles but interactions ( $P < 0.05$ ) between muscle and grade were determined. Consumer scores and volatiles differed ( $P < 0.05$ ) between muscles. Consumers scored *Psoas major* highest for tenderness, juiciness, flavor liking and overall liking, followed by *Longissimus lumborum*, *Gluteus medius*, and *Semimembranosus* ( $P < 0.05$ ). Principal component analysis revealed clustering of compound classes, formed by related mechanisms. Volatile *n*-aldehydes were inversely related to percent fat. Increases in lipid oxidation compounds was associated with *Gluteus medius* and *Semimembranosus*, while greater quantities of sulfur-containing compounds was associated with *Psoas major*. Relationships between palatability scores and volatile compound classes suggests that differences in the pattern of volatile compounds may play a valuable role in explaining consumer liking.

*Keywords:* beef; flavor; GC-MS; HS-SPME; Muscle; USDA Quality Grade

## 1. Introduction

Beef palatability is often believed to be most dependent on tenderness (Miller, Carr, Ramsey, Crockett, & Hoover, 2001; Miller, *et al.*, 1995; Savell, *et al.*, 1987). However, flavor is also considered a primary palatability factor and is shown to be of great importance when tenderness is acceptable (Behrends, *et al.*, 2005a, 2005b; Goodson, *et al.*, 2002; Killinger, Calkins, Umberger, Feuz, & Eskridge, 2004). Flavor has been identified as the single most important factor in determining consumer acceptability when meat was prepared at home (Huffman, Miller,

47 Hoover, Wu, Brittin, & Ramsey, 1996). Beef flavor is a combination of taste and odor. While  
48 taste is generally detected on the tongue as sweet, sour, salty, bitter or other taste sensations such  
49 as “umami”, odor or aroma is detected in the nose and plays a large role in flavor perception.  
50 Numerous volatile compounds have been identified from beef, including: sulfur-containing  
51 compounds, furan thiols, disulfides, aldehydes, ketones and other heterocyclic compounds (Cerny  
52 & Grosch, 1992; Farmer & Patterson, 1991; Gasser & Grosch, 1988; Mottram, 1991).

53 Consumers have associated increased flavor desirability with increased intramuscular fat  
54 (O’Quinn *et al.*, 2012; Smith, Savell, Cross, & Carpenter, 1983). However, laboratory studies  
55 have repeatedly found that increased intramuscular fat rarely produces increases in volatile flavor  
56 compounds (Cross, Berry, & Wells, 1980; Mottram & Edwards, 1983; Mottram, Edwards, &  
57 MacFie, 1982). Evidence from studies on meat products suggests that fat acts as a solvent for  
58 volatile compounds, thus delaying flavor release (Chevance, Farmer, Desmond, Novelli, Troy, &  
59 Chizzolini, 2000). Documentation of the effect of USDA quality grade among multiple beef  
60 muscles upon volatile flavor compounds was not found in the literature.

61 Research regarding differences in flavor among muscles has focused on flavor intensity and  
62 the presence of off-flavors. Calkins and Hodgen (2007) have summarized muscle rankings based  
63 on flavor intensity and off-flavors. In most cases flavor intensity and off-flavors were correlated  
64 with each other. Volatile compounds associated with lipid oxidation have been reported to vary  
65 between muscles of the chuck and round influencing perceived flavor (Hodgen, Cuppett, &  
66 Calkins, 2006). Recently a beef flavor lexicon of beef attributes was used to determine  
67 differences between top loin, top sirloin, tenderloin, and inside round steaks (Adhikari &  
68 Chambers, 2010; Miller, 2010).

69 To date, no studies have assessed the palatability and volatile profile of multiple beef muscles  
70 in various quality grades. The objective of this study was to determine the effects of USDA  
71 quality grade and muscle on consumer palatability perception and volatile beef flavor  
72 compounds.

## 73 **2. Materials and Methods**

### 74 *2.1. Product procurement and preparation*

75 Boneless striploins [Institutional Meat Purchase Specifications (IMPS) 180, North American  
76 Meat Processers Association (NAMP)], tenderloins (IMPS 189, NAMP), inside rounds (IMPS  
77 169, NAMP), and top sirloins (IMPS 184, NAMP) were collected from three ‘A’ maturity (9 to  
78 30 month animals at harvest) carcasses representing each of five USDA quality grades (Prime,  
79 Upper 2/3 Choice, Low Choice, Select, and Standard) at a commercial beef processing facility in  
80 the Midwest region of the United States. Carcasses were selected by trained individuals who  
81 assessed the amount of visual intramuscular fat of the ribeye face at the 12<sup>th</sup> and 13<sup>th</sup> rib along  
82 with lean color and skeletal ossification (USDA, 1997). Subprimals of the selected carcasses  
83 were vacuum packaged and transported to the Gordon W. Davis Meat Laboratory where they  
84 were stored at 2 to 4 °C in the absence of light, and aged to 21 days postmortem prior to  
85 fabrication. Steak cutting, selection and cooking followed Meat Standards Australia (MSA)  
86 protocols (Watson, Gee, Polkinghorne, & Porter, 2008). The muscles, *Longissimus lumborum*,  
87 *Psoas major*, *Semimembranosus*, and *Gluteus medius* (from striploin, tenderloin, inside round,  
88 and top sirloin subprimals, respectively) were denuded of all epimysium and fat.  
89 *Semimembranosus* and *Gluteus medius* muscles were sectioned parallel with muscle fibers in  
90 order to allow steak cutting across the grain. *Longissimus lumborum* and *Psoas major* muscles  
91 were cut perpendicular to the length of each muscle having some grain angle, specifically in

92 *Longissimus lumborum* steaks. All muscles were cut into 25 mm thick steaks approximately 10  
93 cm x 5 cm in length and width, starting at the anterior end of the muscle or muscle section. The  
94 resulting steaks were individually wrapped in plastic, vacuum packed in sets of five, identified  
95 with a unique sample code and frozen (-20 °C). Frozen wrapped steaks were later sorted into  
96 predetermined groups of 10 steaks, each being a single steak from 10 of the original sample  
97 codes, representing a cooking round and re-vacuum packaged. This re-sorting was determined by  
98 MSA protocols and related software routines to produce a six by six latin square presentational  
99 order in which six test products were arranged so that each product was cooked and served an  
100 equal number of times in each of six presentational orders (serving rounds two to seven) and  
101 served before and after each other product an equal number of times. The first cooking and  
102 serving round utilized a common presumed mid position “starter” served to all consumers. The  
103 five individual steaks from each original sample were placed and served in five different rounds  
104 to counter potential order effects.

## 105 2.2. *Consumer palatability scores*

106 Consumer palatability scoring was conducted in accordance with MSA protocols (Watson *et*  
107 *al.*, 2008). Steaks were thawed at 2 to 5 °C for 24 hours prior to cooking. All steaks were cooked  
108 using a Silex clamshell grill (model S-143k, Silex Grills Australia Pty. Ltd., Marrickville,  
109 Australia). Plate surface temperature was set at 225 °C and preheated 45 min prior to panels.  
110 Each panel session was conducted using a count up timer and timed schedule. Each session  
111 commenced with cooking of a warm up load to stabilize grill recovery temperatures prior to the  
112 seven cooking rounds. Loading and unloading of both the warm up and subsequent six test  
113 rounds was conducted in accordance with the time schedule as was serving of test samples.  
114 During panels steaks were loaded on the grill in seven designated groups (rounds) of 10. The

115 grill surface was scraped, cleaned and greased with non-flavored cooking spray (Pam<sup>®</sup> Original  
116 Non-Stick Cooking Spray, ConAgra Foods, Inc., Omaha, NE, USA) between rounds. Steaks  
117 were cooked 5 min with the grill closed, removed at the designated time and allowed to rest for 3  
118 min. During resting three 1.27 cm diameter cores were removed across the center line of selected  
119 steaks for volatile analysis by coring through the thickness of steaks perpendicular to cut surfaces  
120 in order to produce cores of similar volume (approximately 2.5 cm in length and 1.27 cm in  
121 diameter). After the resting period each steak was cut into two pieces (across the cored section),  
122 and immediately served to two designated consumers.

123 Sessions were conducted in evenings by paid consumers (n=278) recruited from Lubbock,  
124 TX, USA and the surrounding area. Consumers were recruited from various community and  
125 charity groups with the group paid for attendance as a fund raiser rather than paying individuals.  
126 Consumers were screened to include only regular beef eaters that preferred “medium doneness.”

127 Each consumer was assigned to a numbered booth containing a ballot, plastic knife, plastic  
128 fork, toothpicks, napkins, a cup of water, an expectorant cup, and between sample palate  
129 cleansers (a 10% apple juice, 90% water solution and unsalted crackers). Panelists were verbally  
130 instructed to utilize the provided plastic utensils to cut steaks into bite sizes similar to their  
131 normal beef consumption habits.

132 Groups of 20 consumers each evaluated seven steaks, the first a standard “starter”, chosen to  
133 be of a mid-range quality, to acclimate consumers, followed by one from each of six product  
134 groups encompassing a wide quality range derived from multiple muscles and USDA quality  
135 grade. Each steak was rated on a 100-mm continuous line scale for tenderness, juiciness, flavor  
136 liking and overall liking. On the scale, zero was verbally anchored as “not tender,” “not juicy,”  
137 “dislike flavor extremely,” and “dislike overall extremely.” Conversely, 100 was verbally

138 anchored as “very tender”, “very juicy”, “like flavor extremely”, and “like overall extremely”.  
139 The MSA “MQ4” score was calculated as a weighted consumer score between one and 100,  
140 using the standard MSA weightings of 30% for tenderness, flavor and overall liking and 10% for  
141 juiciness.

### 142 *2.3. Volatile compound evaluation*

143 Volatile compound collection and gas chromatography-mass spectrometry (GC-MS) analysis  
144 was conducted on selected steaks from those that were grilled and served to consumers during  
145 each evening’s consumer panel. Samples for volatile collection were collected from the selected  
146 steaks, once removed from the grill, by obtaining three 1.27-cm diameter cores from the center  
147 line of selected steaks during the resting period and before the remaining steak was cut into two  
148 portions and served to two consumers. Each core was then cut again perpendicular to the muscle  
149 fibers to enable the six pieces to be placed into a 15 mL clear glass vial (Supelco, Bellefonte, PA,  
150 USA; preconditioned in an oven held at 95 °C). Preheated (60 °C) vials and screw caps  
151 containing a polytetrafluoroethylene septum were then closed. The vial was then placed in a 65  
152 °C water bath (Thermo Scientific, Waltham, MA, USA) and allowed to equilibrate for 5 min.  
153 Volatiles were extracted by solid phase microextraction (SPME) using an 85 µm film thickness  
154 carboxen polydimethylsiloxane fiber in a manual SPME needle and holder (Supelco, Bellefonte,  
155 PA, USA). Following equilibration, a SPME fiber was placed in the headspace above the sample  
156 for 10 min. After collection, samples were withdrawn into the SPME needle, capped using an  
157 inert GC septum (LB-2, Supelco, Bellefonte, PA, USA) and placed in a glass test tube with a  
158 PTFE-lined lid (all preheated in an oven at 95 °C). The SPME fibers with collected volatiles  
159 were held at 2 to 4 °C for up to a maximum of 24 hours, prior to analysis. Collection and holding



160 was required as multiple volatile samples were collected simultaneously during consumer  
161 palatability scoring sessions.

162 An Agilent 6890 series GC (Agilent Technologies, Santa Clara, CA, USA) equipped with a  
163 5975 MS detector (Agilent Technologies, Santa Clara, CA, USA) was used for separation and  
164 detection of volatile compounds. Extracted volatile compounds were desorbed from SPME fibers  
165 at the GC-MS inlet at 250 °C in splitless mode. Cryogenic focusing was conducted by placing  
166 the front of the GC column into a bed of dry ice (solid CO<sub>2</sub>). A loop of the front end of the  
167 column (approximately 100 mm), between the injector and the remaining portion of the column,  
168 was placed into the dry ice for a period of 5 min prior to injection. The software program was  
169 then loaded and prepared to start and the SPME fiber was injected and desorbed for 5 min while  
170 the column remained in the dry ice. After 5 min the column was removed from the dry ice and  
171 the oven method was started. The SPME fiber remained exposed within the inlet for the first 3  
172 min of the oven method to ensure all volatile compounds had been desorbed.

173 Compounds were separated using a BPX-5 capillary column (25 m × 0.32 mm, 0.25 µm film  
174 thickness; SGE, Austin, TX, USA) with helium as the carrier gas at 1 mL per min. The oven  
175 method used included an initial 5 min at 35 °C, followed by an 8 °C per min ramp to 220 °C, then  
176 a 20 °C per min ramp to 290 °C, and finally a 5 min hold period at 290 °C. The total run time was  
177 37 min. The inlet was operated in splitless mode for the first 3 min followed by a 10:1 split.

178 The MS detected ions within 33-500 m/z range in the electron impact mode at 70 eV.  
179 Chromatography data was collected in the selective ion monitoring/scan mode (SIM/Scan;  
180 Agilent MSD Chemstation D.03.00.611 software, Agilent Technologies, Santa Clara, CA, USA).  
181 Ions were selected based on the presence of three primary ions from compounds of interest.

182 *2.4. Mass spectral identification of volatile compounds*

183 A solution of *n*-alkanes (C<sub>8</sub>-C<sub>22</sub>, Supelco, Bellefonte, PA, USA; 1 ng/μL) was run each day of  
184 analysis and linear retention indices (LRI) were calculated with reference to the *n*-alkanes  
185 (Goodner, 2008). The calculated LRI were used to determine retention times of compounds of  
186 interest. Volatile compound identity was confirmed by comparison of the ion fragmentation  
187 patterns and the LRI with that of the authentic compounds. Three target ions were selected for  
188 the comparisons between sample and standard runs with one quantitative ion and two qualifying  
189 ions being selected for each compound of interest. A single-point external standard method was  
190 used for quantitation. External standard reference compounds (Sigma Aldrich, Saint Louis, MO,  
191 USA) were delivered in solutions (1 ng/μl) of pentane (later eluding compounds) or toluene  
192 (early eluting compounds) in splitless-mode. Quantitative ion abundances of sample runs were  
193 compared with quantitative ion abundances of standard runs of known concentration.

194 Compounds not detected in sample runs were treated as zero

#### 195 2.5. Proximate Analysis

196 Proximate analysis of raw steaks was conducted by an AOAC official method (2007.04;  
197 Anderson, 2007) using a near infrared spectrophotometer (FoodScan, FOSS NIRsystems, Inc.,  
198 Laurel, MD, USA). Chemical percentages of fat, moisture, protein, and total collagen were  
199 determined for each muscle within each USDA quality grade, as described previously (O'Quinn  
200 *et al.*, 2011).

#### 201 2.6. Statistical analysis

202 Statistical analysis was conducted based on a generalized linear mixed model, using the Proc  
203 Glimmix procedure of SAS (Version 9.3, Cary, NC). Two-way analysis of variance was used to  
204 evaluate the fixed effects of USDA quality grade, beef muscle and their interaction. Steak was  
205 the experimental unit. Panel session, serving round, and consumer were each treated as random

206 effects in the model. Differences were considered significant at  $P < 0.05$ . The CORR procedure  
207 of SAS was used to determine Pearson correlation coefficients. Principal component (PC)  
208 analysis was performed on volatile compounds using PROC FACTOR of SAS (v.9.3, Cary, NC).  
209 Three principal components, PC1, PC2 and PC3 were retained to determine treatment scores and  
210 correlation coefficients with consumer palatability scores and proximate data. The treatment PC  
211 scores and correlation coefficients were plotted together (x coordinate = PC1; y coordinate =  
212 PC2 or PC3 correlation coefficients) to evaluate relationships.

### 213 **3. Results and Discussion**

#### 214 3.1. *Chemical fat, collagen, moisture, and protein*

215 Proximate analysis was conducted for steaks from subprimals for which consumer and  
216 volatile flavor compound evaluations were obtained (Table 1). It is important to note that the  
217 samples for inclusion in this experiment were selected to give clear differences in the chemical  
218 fat content of the *Longissimus lumborum* between grades. Therefore, these data do not represent  
219 a random selection of samples from these USDA quality grades and are recorded to assist with  
220 the explanation of consumer and flavor analyses.

221 Percent chemical fat, collagen and moisture showed an interaction between USDA quality  
222 grade and muscle ( $P < 0.001, 0.01, 0.001$ , respectively; Table 1). In *Longissimus lumborum*  
223 steaks the chemical fat percentages of the various quality grades were similar to previous  
224 findings (Emerson, Woerner, Belk, & Tatum, 2013). As quality grade increased, fat content  
225 increased while moisture content decreased, as demonstrated in numerous previous studies (Hunt  
226 *et al.*, 2014; Von Seggern, Calkins, Johnson, Brickler, & Gwartney, 2005; Brackebush, McKeith,  
227 Carr, & McLaren, 1991; Romans, Tuma, & Tucker, 1965). The interaction between grade and  
228 muscle highlighted the fact that the relationship between grade and fat content differs markedly

229 among muscles. Intramuscular fat levels in *Psoas major*, *Gluteus medius* and *Semimembranosus*  
230 follow a similar pattern to the *Longissimus* samples, but the differences were much less distinct,  
231 with most difference occurring between Prime and Upper 2/3 Choice compared with the Low  
232 Choice, Select and Standard grades. As expected, an opposite pattern of effects was observed for  
233 percent moisture content, though the differences between muscles and grades were much  
234 smaller. Other researchers have also reported that moisture and fat content of beef muscles vary  
235 with quality grade (Hunt *et al.*, 2014; Von Seggern *et al.*, 2005).

236 There was no interaction for percent protein ( $P > 0.05$ ), but there were differences due to  
237 muscles ( $P < 0.01$ ) and grade ( $P < 0.05$ ), similar to Hunt *et al.* (2014). As expected, these  
238 differences, again small, follow the pattern for the percent moisture and mirror that for percent  
239 fat (Table 1). This trend reflects results reported by previous works (Hunt *et al.*, 2014;  
240 Brakebusch *et al.*, 1991; Romans *et al.*, 1965).

241 An interaction was present between grade and muscle for percent collagen ( $P < 0.01$ ), with  
242 levels increasing in higher grades of *Longissimus lumborum* but unaffected by grade in *Psoas*  
243 *major*. Prost *et al.* (1975) has previously reported that percent collagen of the *Psoas major* is  
244 unaffected by grade. Variation in percent collagen between muscles is well documented (Von  
245 Seggern *et al.*, 2005; McKeith, De Vol, Miles, Bechtel, & Carr, 1985; Prost, Pelczynska, &  
246 Kotolua, 1975). The effect of quality grade on percent collagen is less clear and often dependent  
247 on muscle (Von Seggern *et al.*, 2005), as found in this study.

### 248 3.2. Consumer palatability scores

249 Consumer evaluations of tenderness, juiciness, flavor liking, and overall liking of beef steaks  
250 from four muscles and five USDA quality grades are displayed in Table 2, along with the  
251 composite MQ4 value. The results show significant interactions between muscle and grade ( $P <$

252 0.05) for all attributes except tenderness. Surprisingly, USDA quality grade had no effect ( $P >$   
253 0.05) on consumer tenderness ratings and there was no interaction between muscle and grade ( $P$   
254  $> 0.05$ ). However, as expected from previous reports (Browning, Huffman, Egbert, & Jungst,  
255 1991; Christensen, Johnson, West, Marchall, & Hargrove, 1991; McKeith *et al.*, 1985),  
256 tenderness differed ( $P < 0.05$ ) between all the muscles (Table 2), with mean scores ranging from  
257 38 for *Semimembranosus* to 89 for *Psoas major*.

258 Juiciness was determined by consumers to be greatest among *Psoas major* steaks from Prime,  
259 Upper 2/3 Choice, Select, and Standard quality grades along with Prime *Longissimus lumborum*  
260 steaks ( $P < 0.05$ ; Table 2). Interestingly, Low Choice *Psoas major* and Low Choice  
261 *Semimembranosus* steaks received lower scores than the rest of the quality grades for these  
262 muscles, but the same effect was not observed for *Gluteus medius* and *Longissimus lumborum*  
263 muscles. Thus, juiciness scores differed between muscles and were generally greater in Prime  
264 and Upper 2/3 Choice grades. These are the same grades that had greater percent fat supporting  
265 the documented belief that percent fat is related to juiciness (Lorenzen *et al.*, 1999; Lorenzen *et*  
266 *al.*, 2003; Savell, Cross, & Smith, 1986; Smith *et al.* 1984). Flavor liking scores followed similar  
267 trends (Table 2) to juiciness where an interaction ( $P < 0.05$ ) for flavor liking was due to lower  
268 flavor liking scores within *Psoas major* and *Semimembranosus* Low Choice grade receiving  
269 lower scores than expected.

270 The MSA MQ4 value, as previously described, assessed meat eating quality based on  
271 weighted calculations. This value has been shown to predict consumer satisfaction and avoids the  
272 difficulty consumers have in distinguishing between attributes (Watson *et al.*, 2008). In this data  
273 the MQ4 values followed similar trends as overall liking and flavor liking (Table 2).

274 Generally, the effect of USDA quality grade on juiciness, flavor liking, overall liking, and  
275 MQ4 was found to be dependent on muscle (Table 2). For most muscles, these attributes did not  
276 show consistent increases in consumer score with increasing quality grade. Specifically, the  
277 *Longissimus lumborum* muscle was the only muscle possessing a linear ranking with quality  
278 grade for juiciness, flavor liking, overall liking, and MQ4. This is likely the effect of fat level  
279 within the different muscles. The maximum difference in fat content between USDA Prime  
280 *Longissimus lumborum* and USDA Standard *Longissimus lumborum* was close to 12% (where  
281 samples were selected on percent fat), whereas the range in percent fat was only 5.2% in the  
282 *Psoas major*, 5.5% in the *Gluteus medius*, and 4.9% in the *Semimembranosus* (Table 1).  
283 Additionally, USDA quality grade did not have an effect ( $P > 0.05$ ) on fat content for muscles  
284 other than the *Longissimus lumborum*, especially for the lowest three quality grade treatments  
285 (Table 2).

### 286 3.3. Volatile compounds

287 A total of 26 volatile compounds representing pathways of cooked beef flavor development  
288 (e.g., thermal oxidation of lipids, Maillard reaction) were selected and quantified. Table 3 shows  
289 the mean quantities of volatiles collected from different muscles while Table 4 presents the  
290 quantities for those volatile compounds which showed a significant interaction ( $P < 0.05$ ). None  
291 of the compounds differed ( $P > 0.05$ ) due to quality grade as a first order effect. Some of the  
292 interactions were influenced by particularly low quantities detected for one muscle/grade  
293 interaction, especially for some *Psoas major* samples.

294 Five compounds (2,3-butanedione, heptane, 3-hydroxy-2-butanone, octane, and methyl  
295 pyrazine) differed ( $P < 0.05$ ) between muscles independent of quality grade (Table 3). The  
296 alkanes, heptane and octane, were found in greatest ( $P < 0.05$ ) quantities from *Psoas major*

297 steaks while being similar ( $P > 0.05$ ) to *Gluteus medius* and *Semimembranosus* steaks but  
298 differing ( $P < 0.05$ ) from *Longissimus lumborum* steaks (Table 3). Alkanes are formed from the  
299 oxidation of long-chain fatty acids (Mottram, 1991). In this study, alkanes did not appear to be  
300 related to percent fat.

301 The ketones, 3-hydroxy-2-butanone and 2,3-butanedione were both present in greatest ( $P <$   
302 0.05) abundance in the headspace of *Gluteus medius* and *Semimembranosus* steaks compared  
303 with *Longissimus lumborum* and *Psoas major* steaks (Table 3). These compounds can arise from  
304 the 2,3-enolisation pathways which form part of the Maillard reaction (Hurrell, 1982). This could  
305 arise from elevated levels of reducing sugars and amino acids or from a higher pH, which favors  
306 2,3-enolisation. Other Maillard products are not similarly affected (Table 3) so the role of pH  
307 within muscles may be worthy of further investigation.

308 Methyl pyrazine was found in the greatest ( $P < 0.05$ ) abundance among *Longissimus*  
309 *lumborum* steaks compared with *Psoas major* and *Semimembranosus*, while *Gluteus medius*  
310 steaks were intermediate and similar ( $P > 0.05$ ) to all other muscles (Table 3). Similar trends for  
311 other pyrazines were not significant ( $P > 0.05$ ; Table 4). Nitrogen-containing pyrazines are  
312 known to be some of the final products of the Maillard reaction (Back, 2007). Although they  
313 occur at lower abundances, compared with lipid degradation volatile compounds, these  
314 compounds have low odor thresholds which contribute roasted flavors (Buttery & Ling, 1997).

315 Certain aldehydes have been shown to be the result of Strecker degradation of amino acids.  
316 Degradation of alanine, isoleucine, leucine, methionine, phenylalanine, and valine leads to the  
317 development of acetaldehyde, 2-methylbutanal, 3-methylbutanal, methional, and  
318 phenylacetaldehyde (Cerny, 2007). Benzaldehyde, is another volatile compound potentially  
319 resulting from the Strecker degradation of the amino acid phenylglycine (MacLeod, & Ames,

320 1987; Mottram, & Edwards, 1983). However, as phenylglycine is not an amino acid which  
321 occurs in muscle, a different mechanism of formation must be responsible in this case. In our  
322 study, benzaldehyde was found to be greater ( $P < 0.05$ ) in *Psoas major*, *Gluteus medius*, and  
323 *Semimembranosus* steaks.

324 Interactions ( $P < 0.05$ ) were found between muscle and USDA grade for seven compounds  
325 (acetaldehyde, 2-propanone, dimethyl sulfide, hexanal, benzaldehyde, octanal, and nonanal;  
326 Table 4). The effect of quality grade on the *n*-aldehydes, octanal, and nonanal, depended on  
327 muscle (Table 4). In the case of *Longissimus lumborum* and *Psoas major*, there was a clear and  
328 significant increase in quantities detected with a decrease in grade. Interestingly the fat content  
329 of these muscles decreased with quality grade (Table 1). Formation of aldehydes occurs in  
330 cooked meat through the thermal oxidation of fatty acids such as oleic, linoleic, and linolenic  
331 acid (Cerny, 2007). Each of these aldehydes have previously been identified in beef odor  
332 (Mottram, 1991).

333 Among volatile compounds found to have interactions between USDA quality grade and  
334 muscle (Table 4), acetaldehyde, 2-propanone and dimethyl sulfide were all found to be greatest  
335 among Upper 2/3 Choice *Psoas major* steaks ( $P < 0.05$ ). Interestingly, Upper 2/3 Choice *Psoas*  
336 *major* steaks received the greatest score for flavor liking by consumers (Table 2). Sulfur-  
337 containing compounds, including dimethyl sulfide, contribute to meaty flavor notes (Gasser &  
338 Grosch, 1990). The sum of sulfur-containing compounds (dimethyl sulfide, dimethyl disulfide,  
339 methanethiol, and methional) were collectively found to be greatest ( $P < 0.05$ ) among *Psoas*  
340 *major* steaks.

341 Overall, these data indicate that the pattern of volatile compounds differs between muscles.  
342 *Psoas major* was characterized by greater levels of the sulfur-containing thiols and sulfides;



343 these and other sulfur-containing compounds are known to contribute to the meaty and roasted  
344 characteristics of beef flavor (Mottram, 1991). It is of interest that *Psoas major* steaks  
345 consistently received the greatest scores for flavor liking though relationships between the  
346 attributes may mean that this score was influenced by tenderness (Table 2). This phenomenon  
347 has been described as a halo-effect where one favorable attribute influences consumer's  
348 perception of other attributes (Roerber, *et al.*, 2000). As previously described tenderness is often  
349 considered to be the most influential beef palatability attribute and this may have some impact on  
350 flavor liking in this study within the notoriously tender *Psoas major* muscle. *Longissimus*  
351 *lumborum* steaks tended to give greater amounts of pyrazines (Table 3), known to contribute to  
352 roasted and nutty characteristics (Mottram, 1991), but lower concentrations of benzaldehyde and  
353 short chain ketones. *Gluteus medius* and *Semimembranosus* steaks gave high levels of some short  
354 chain ketones known to participate in a range of flavor forming reactions and tended to give  
355 more *n*-aldehydes, though there was considerable variability between USDA grades (Table 4).  
356 These differences would be expected to influence and explain differences in perceived flavor  
357 quality between the different muscles.

#### 358 3.4. Correlations

359 Pearson correlations between proximate data and consumer palatability scores are  
360 displayed in Table 5. As expected, moisture was inversely related with chemical fat ( $r = -0.97$ ;  $P$   
361  $< 0.001$ ). This inverse relationship between moisture and fat content in multiple beef muscles is  
362 very similar to previous work, where a similarly highly significant correlation ( $r = -0.92$ ) was  
363 found (Jeremiah, Dugan, Aalhus, & Gibson, 2002).

364 There is an apparent correlation between increased chemical fat and increased collagen  
365 ( $P < 0.001$ ). Previously, accumulation of collagen during animal physiological maturation was

366 documented to impact palatability, specifically tenderness (Berry, Smith, & Carpenter, 1974;  
367 Breidenstein, Cooper, Cassens, Evans, & Bray, 1968; Romans *et al.*, 1965). However, in this  
368 study similarly young 'A' maturity carcasses were selected for all grades. A weak positive  
369 correlation was observed between collagen and juiciness, flavor liking, and overall liking ( $P <$   
370  $0.05$ ), but not tenderness (Table 5). It is difficult to propose any direct causative link between  
371 more collagen and higher consumer scores.

372 Overall liking was greatly correlated with flavor liking, juiciness and tenderness ( $P < 0.001$ ;  
373 Table 5) indicating that consumers find it difficult to differentiate fully between attributes.  
374 Percent fat was correlated with overall liking, tenderness, juiciness, and flavor liking ( $P < 0.001$ ),  
375 as expected from previous work (McKeith *et al.*, 1985; Tatum, Smith, Berry, Murphey,  
376 Williams, & Carpenter, 1980). There was also a tendency for negative correlations of *n*-  
377 aldehydes with flavor liking, overall liking, and percent fat (Table 6).

378 Negative correlations of long chain *n*-aldehydes (octanal and decanal) with percent fat (Table  
379 6) may be due to the retention of volatile compounds by fat, delaying flavor release as described  
380 previously (Farmer, Hagan, Oltra, Devlin, & Gordon, 2013; Chevance *et al.*, 2000; Chevance &  
381 Farmer, 1999). However, this effect was not apparent for other compounds or compound groups,  
382 which showed no significant correlations with percent fat ( $P > 0.05$ ; data not tabulated). Instead,  
383 these results may indicate a greater potential for oxidation of unsaturated fatty acids of the polar  
384 lipid fraction within beef steaks having low total percent fat. Within beef with a lower total fat  
385 content, a greater proportion of the fat includes polar lipids (Wood *et al.*, 2008). Polar lipids are  
386 known to be more susceptible to oxidation (Mottram, 1998). Previously, volatile compounds  
387 associated with lipid oxidation were increased up to 4-fold in response to increased proportions  
388 of polyunsaturated fatty acids (Elmore, Mottram, Enser, & Wood, 1999).

### 389 3.5. Principal component analysis

390 Principal component analysis (PCA) was conducted in order to explore relationships between  
391 multiple volatile compounds and muscles of different quality grades. Volatile compounds were  
392 used to determine principal components (PCs). When PCA was conducted for all grade and  
393 muscle treatments PC1 explained 39.8%, PC2 explained 29.4%, and PC3 explained 20.8% of the  
394 variation associated with volatile compounds (Figures 1 and 2). Plots revealed that PC1  
395 separated Upper 2/3 Choice *Psoas major* from most of the samples on the basis of increased  
396 quantities of many of the Maillard products and reduced quantities of lipid oxidation products.  
397 Secondly, PC2 tended to separate *Longissimus lumborum* steaks from many of the other muscles  
398 and was associated with an overall lack of volatiles. Principle Component 3 separated *Psoas*  
399 *major* steaks of all grades from many of the remaining samples, with the *Psoas major* being  
400 associated with greater quantities of sulfur-containing Maillard products.

401 Volatile compounds segregated into clusters of similar compound classes (Figures 1 and 2).  
402 Pyrazines, Strecker aldehydes, and sulfur compounds were found to be positively related with  
403 PC1, while lipid oxidation products, aldehydes, ketones, and alkanes were clustered together and  
404 negatively related with PC1. Figure 2 revealed that PC3 separated the treatments on the basis of  
405 different groups of Maillard products. This collinear divergence of compound groups may make  
406 it possible to use related compounds as “markers” for flavor compounds of greater odor  
407 significance which are difficult to detect. Most volatile compounds were located on the positive  
408 side of PC2 while percent fat was on the negative side, a similar finding was reported in a recent  
409 work (Farmer *et al.*, 2013) where lower fat content beef was related with greater quantities of  
410 volatile compounds. It was suggested by Farmer *et al.*, (2013) that lower intramuscular fat  
411 content leads to increases in volatile compounds, due to the solubility of volatile aroma

412 compounds in lipids, as previously observed in frankfurters (Chevance & Farmer, 1999;  
413 Chevance *et al.*, 2000).

414 *Longissimus lumborum* showed an association with chemical fat content and an absence of  
415 volatile compounds compared with other muscles regardless of quality grade (Figure 1). Upper  
416 2/3 Choice *Psoas major*, which diverted from the remaining treatments was associated with  
417 groupings of sulfur-containing compounds and Maillard products and was greatly separated from  
418 *n*-aldehydes. The data in Table 4 show that this treatment gave unusually (and consistently) high  
419 levels of acetaldehyde, 2-propanone and sulfur-containing compounds.

420 Figure 2 confirms that Maillard products are closely associated with flavor development  
421 (Mottram, 1998) and in this study flavor liking. More specifically, sulfur-compounds were  
422 greatly associated with flavor liking. This may reflect the importance of these and other sulfur-  
423 containing compounds for aspects of beef flavor.

#### 424 **4. Conclusions**

425 The results of this study indicate that there is potential to gain understanding of flavor  
426 differences between beef muscles through the analysis of volatile flavor compounds in  
427 association with palatability and chemical measurements. Similar to previous studies USDA  
428 quality grade affected consumer flavor and overall liking dependent on muscle. Beef muscle type  
429 greatly influenced volatile compounds. Some volatile compounds were negatively correlated  
430 with percent fat, while others were not related to fat content. Volatile compounds from similar  
431 compound classes and from the same pathways of formation behaved, similarly, with Maillard  
432 products being most closely related with flavor liking. This clear relationship between  
433 palatability scores and volatile compound classes suggests that differences in the pattern of  
434 volatile compounds between muscles may play a valuable role in explaining consumer liking.

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595

**Table 1.** Proximate Data<sup>1</sup> of raw beef steaks from five USDA Quality Grades and four muscles

USDA Quality Grade	Muscle	%			
		Fat	Collagen	Moisture	Protein
Prime	<i>Psoas major</i>	8.1 <sup>b</sup>	1.8 <sup>cde</sup>	69.5 <sup>de</sup>	21.2
Upper 2/3 Choice	<i>Psoas major</i>	6.9 <sup>bcd</sup>	1.9 <sup>bcde</sup>	70.3 <sup>cde</sup>	21.4
Low Choice	<i>Psoas major</i>	3.8 <sup>defghi</sup>	1.7 <sup>de</sup>	73.1 <sup>abc</sup>	21.7
Select	<i>Psoas major</i>	3.5 <sup>efghi</sup>	1.9 <sup>bcd</sup>	72.5 <sup>abc</sup>	22.5
Standard	<i>Psoas major</i>	2.9 <sup>fghij</sup>	1.8 <sup>de</sup>	73.1 <sup>abc</sup>	22.1
Prime	<i>Longissimus lumborum</i>	13.1 <sup>a</sup>	2.1 <sup>ab</sup>	64.0 <sup>f</sup>	21.7
Upper 2/3 Choice	<i>Longissimus lumborum</i>	7.9 <sup>b</sup>	2.0 <sup>abcd</sup>	68.7 <sup>e</sup>	21.9
Low Choice	<i>Longissimus lumborum</i>	4.5 <sup>defg</sup>	1.7 <sup>de</sup>	70.4 <sup>cde</sup>	23.2
Select	<i>Longissimus lumborum</i>	2.9 <sup>fghi</sup>	1.7 <sup>de</sup>	71.3 <sup>cd</sup>	23.1
Standard	<i>Longissimus lumborum</i>	1.3 <sup>ij</sup>	1.6 <sup>e</sup>	73.5 <sup>ab</sup>	23.3
Prime	<i>Gluteus medius</i>	7.1 <sup>bc</sup>	2.3 <sup>a</sup>	69.0 <sup>e</sup>	21.7
Upper 2/3 Choice	<i>Gluteus medius</i>	4.3 <sup>defgh</sup>	1.7 <sup>de</sup>	71.8 <sup>bc</sup>	21.8
Low Choice	<i>Gluteus medius</i>	1.6 <sup>ij</sup>	1.6 <sup>e</sup>	72.4 <sup>abc</sup>	23.3
Select	<i>Gluteus medius</i>	2.9 <sup>fghij</sup>	1.9 <sup>bcd</sup>	71.8 <sup>bc</sup>	22.9
Standard	<i>Gluteus medius</i>	2.6 <sup>fghij</sup>	1.9 <sup>bcd</sup>	72.3 <sup>abc</sup>	22.9
Prime	<i>Semimembranosus</i>	5.6 <sup>cde</sup>	1.9 <sup>bcd</sup>	70.6 <sup>cde</sup>	22.5
Upper 2/3 Choice	<i>Semimembranosus</i>	5.0 <sup>cdef</sup>	2.1 <sup>abc</sup>	71.4 <sup>cd</sup>	21.8
Low Choice	<i>Semimembranosus</i>	2.0 <sup>hij</sup>	1.8 <sup>cde</sup>	72.8 <sup>abc</sup>	23.2
Select	<i>Semimembranosus</i>	2.5 <sup>ghij</sup>	1.9 <sup>bcd</sup>	72.3 <sup>abc</sup>	23.2
Standard	<i>Semimembranosus</i>	0.7 <sup>j</sup>	1.6 <sup>e</sup>	74.1 <sup>a</sup>	23.1
Std. Error		1.5	0.2	1.5	0.8
<i>P</i> value		<0.001	0.004	<0.001	0.809
	<i>Psoas Major</i>	5.0	1.8	71.7	21.8 <sup>b</sup>
	<i>Longissimus lumborum</i>	5.9	1.8	69.6	22.7 <sup>a</sup>
	<i>Gluteus medius</i>	3.7	1.9	71.5	22.5 <sup>a</sup>
	<i>Semimembranosus</i>	3.2	1.9	72.2	22.8 <sup>a</sup>
	Std. Error	0.4	0.1	0.4	0.2
	<i>P</i> value	<0.001	0.766	<0.001	0.006
Prime		8.5	2.0	68.3	21.8 <sup>b</sup>
Upper 2/3 Choice		6.0	1.9	70.5	21.7 <sup>b</sup>
Low Choice		2.9	1.7	72.1	22.9 <sup>a</sup>
Select		2.9	1.9	71.9	22.9 <sup>a</sup>
Standard		1.9	1.7	73.2	22.9 <sup>a</sup>
Std. Error		0.5	0.1	0.5	0.2
<i>P</i> value		<0.001	0.019	<0.001	0.028

596 <sup>abcde fghij</sup> Means within a column lacking a common superscript differ ( $P < 0.05$ ).

597 <sup>1</sup> Chemical percentages of fat, moisture, protein, and collagen determined of raw steaks by  
598 AOAC official method (2007.04; Anderson, 2007)

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600

**Table 2.** Consumer palatability scores<sup>1</sup> of grilled beef steaks from five USDA Quality Grades and four muscles

USDA Quality Grade	Muscle	Tenderness	Juiciness	Flavor Liking	Overall Liking	MQ4
Prime	<i>Psoas major</i>	94.1	85.9 <sup>a</sup>	84.8 <sup>a</sup>	89.1 <sup>a</sup>	89.7 <sup>a</sup>
Upper 2/3 Choice	<i>Psoas major</i>	90.2	86.3 <sup>a</sup>	86.1 <sup>a</sup>	88.1 <sup>ab</sup>	84.9 <sup>abc</sup>
Low Choice	<i>Psoas major</i>	81.4	55.5 <sup>efg</sup>	67.9 <sup>abcde</sup>	67.1 <sup>bcd</sup>	71.5 <sup>bcde</sup>
Select	<i>Psoas major</i>	94.1	73.7 <sup>abcd</sup>	84.9 <sup>a</sup>	86.3 <sup>ab</sup>	87.4 <sup>ab</sup>
Standard	<i>Psoas major</i>	90.1	81.4 <sup>ab</sup>	75.7 <sup>ab</sup>	82.7 <sup>ab</sup>	82.6 <sup>abcd</sup>
Prime	<i>Longissimus lumborum</i>	76.6	75.7 <sup>abc</sup>	78.4 <sup>a</sup>	78.1 <sup>ab</sup>	77.9 <sup>bcd</sup>
Upper 2/3 Choice	<i>Longissimus lumborum</i>	67.9	69.9 <sup>bcde</sup>	68.8 <sup>abcd</sup>	69.2 <sup>bc</sup>	69.8 <sup>de</sup>
Low Choice	<i>Longissimus lumborum</i>	71.3	67.8 <sup>cde</sup>	73.6 <sup>abc</sup>	68.4 <sup>bc</sup>	70.6 <sup>cde</sup>
Select	<i>Longissimus lumborum</i>	60.4	59.3 <sup>ef</sup>	64.6 <sup>bcde</sup>	61.9 <sup>cd</sup>	62.3 <sup>efg</sup>
Standard	<i>Longissimus lumborum</i>	68.2	59.2 <sup>ef</sup>	56.4 <sup>ef</sup>	58.7 <sup>cd</sup>	60.8 <sup>efg</sup>
Prime	<i>Gluteus medius</i>	54.9	62.5 <sup>def</sup>	65.2 <sup>bcde</sup>	63.4 <sup>cd</sup>	62.7 <sup>efg</sup>
Upper 2/3 Choice	<i>Gluteus medius</i>	61.2	69.2 <sup>bcde</sup>	72.9 <sup>abc</sup>	69.5 <sup>bc</sup>	67.8 <sup>def</sup>
Low Choice	<i>Gluteus medius</i>	47.6	60.2 <sup>ef</sup>	61.3 <sup>bcdef</sup>	58.0 <sup>cde</sup>	55.6 <sup>efgh</sup>
Select	<i>Gluteus medius</i>	51.9	50.4 <sup>fgh</sup>	57.2 <sup>def</sup>	54.9 <sup>cde</sup>	55.4 <sup>efgh</sup>
Standard	<i>Gluteus medius</i>	48.1	50.8 <sup>fgh</sup>	56.6 <sup>def</sup>	51.6 <sup>def</sup>	52.2 <sup>ghi</sup>
Prime	<i>Semimembranosus</i>	36.6	62.7 <sup>def</sup>	59.7 <sup>cdef</sup>	52.1 <sup>def</sup>	52.9 <sup>fghi</sup>
Upper 2/3 Choice	<i>Semimembranosus</i>	33.9	61.6 <sup>def</sup>	56.8 <sup>def</sup>	41.6 <sup>ef</sup>	44.8 <sup>hi</sup>
Low Choice	<i>Semimembranosus</i>	32.2	38.6 <sup>h</sup>	49.4 <sup>f</sup>	37.2 <sup>f</sup>	39.1 <sup>i</sup>
Select	<i>Semimembranosus</i>	39.4	55.1 <sup>fg</sup>	64.9 <sup>bcde</sup>	57.5 <sup>cde</sup>	55.6 <sup>efgh</sup>
Standard	<i>Semimembranosus</i>	42.3	44.3 <sup>gh</sup>	52.5 <sup>ef</sup>	44.4 <sup>ef</sup>	46.2 <sup>hi</sup>
Std. Error		7.5	7.1	7.3	7.8	7.0
<i>P</i> value		0.107	0.024	0.032	0.019	0.033
	<i>Psoas major</i>	89.4 <sup>a</sup>	76.6	79.9	82.7	83.2
	<i>Longissimus lumborum</i>	69.4 <sup>b</sup>	66.4	68.3	67.3	68.3
	<i>Gluteus medius</i>	54.1 <sup>c</sup>	58.6	62.6	59.5	58.7
	<i>Semimembranosus</i>	38.4 <sup>d</sup>	52.5	56.6	46.6	47.7
	Std. Error	3.7	3.4	3.0	3.1	2.9
	<i>P</i> value	<0.001	<0.001	<0.001	<0.001	<0.001
Prime		68.8	71.7	72.0	70.7	70.8
Upper 2/3 Choice		61.1	71.8	71.2	67.1	66.8
Low Choice		57.8	55.5	63.0	57.7	59.2
Select		63.3	59.6	67.9	65.2	65.2
Standard		63.0	58.9	60.3	59.4	60.4
Std. Error		5.9	3.2	3.8	4.2	4.1
<i>P</i> value		0.735	<0.001	0.135	0.174	0.268

602 <sup>abcdefghi</sup> Means within a column lacking a common superscript differ ( $P < 0.05$ ).

603 <sup>1</sup> Consumer rated each steak on a 100-mm continuous line scale for flavor, tenderness, juiciness, and overall liking.  
604 On the scale, 0 was verbally anchored as not tender, not juicy, dislike flavor extremely, and dislike overall  
605 extremely. Similarly, 100 was verbally anchored as very tender, very juicy, like flavor extremely, and like overall  
606 extremely. Meat quality, 4 variables score (MQ4) reflecting a weighted consumer score between 1 and 100 was  
607 calculated using standard Meat Standard Australia weightings of 30% for tenderness, flavor and overall liking and  
608 10% for juiciness.

**Table 3.** Least-squares means of volatile flavor compounds (ng) from grilled beef steaks of four muscles

Volatile compound	Linear Retention Indices	Beef Muscles				Std. Error	P value
		<i>Longissimus lumborum</i>	<i>Psoas major</i>	<i>Gluteus medius</i>	<i>Semi-membranosus</i>		
<i>n-Aldehydes</i>							
Acetaldehyde	412	2.52 <sup>b</sup>	6.77 <sup>a</sup>	2.05 <sup>b</sup>	1.59 <sup>b</sup>	0.81	<0.001
Pentanal	697	28.65	33.39	34.84	38.29	9.99	0.859
Hexanal	795	12.24	10.01	13.72	15.18	4.68	0.779
Heptanal	898	0.83	1.09	1.28	1.27	0.16	0.051
Octanal	1002	0.79	1.17	1.11	1.18	0.18	0.188
Nonanal	1107	1.36	1.96	1.94	1.89	0.24	0.103
Decanal	1205	0.22	0.18	0.28	0.23	0.04	0.219
Sum n-Aldehydes		44.16	47.55	53.08	57.48	15.16	0.858
<i>Strecker Aldehydes</i>							
3-Methyl butanal	652	52.43	39.74	41.75	50.49	9.21	0.467
2-Methyl butanal	659	87.38	49.28	71.21	84.45	15.03	0.139
Benzaldehyde	960	0.36 <sup>b</sup>	0.58 <sup>a</sup>	0.54 <sup>a</sup>	0.48 <sup>a</sup>	0.04	<0.001
Phenylacetaldehyde	1045	0.06	0.07	0.07	0.07	0.01	0.711
Sum Strecker aldehydes		139.99	90.44	114.53	136.55	24.81	0.277
<i>Ketones</i>							
2-Propanone	496	2.85 <sup>b</sup>	13.97 <sup>a</sup>	3.78 <sup>b</sup>	4.55 <sup>b</sup>	1.49	<0.001
2,3-Butanedione	560	6.87 <sup>bc</sup>	6.34 <sup>c</sup>	9.45 <sup>ab</sup>	10.53 <sup>a</sup>	1.39	0.033
2-Butanone	597	1.94	2.84	1.92	1.99	0.43	0.235
3-Hydroxy-2-butanone	705	61.44 <sup>b</sup>	65.59 <sup>b</sup>	135.28 <sup>a</sup>	123.33 <sup>a</sup>	13.11	<0.001
<i>Sulfides</i>							
Dimethyl sulfide	519	0.41 <sup>c</sup>	3.03 <sup>a</sup>	1.03 <sup>bc</sup>	1.38 <sup>b</sup>	0.42	<0.001
Dimethyl disulfide	744	0.35	0.52	0.32	0.28	0.07	0.065
<i>Thiols</i>							
Methanethiol	423	0.02	0.04	0.02	0.03	0.01	0.134
Methional	911	0.23	0.28	0.28	0.25	0.04	0.504
Sum Sulfur containing		1.02 <sup>b</sup>	3.81 <sup>a</sup>	1.63 <sup>b</sup>	1.95 <sup>b</sup>	0.44	<0.001
<i>Furans</i>							
2-Pentyl furan	994	0.03	0.06	0.05	0.06	0.02	0.359
<i>Pyrazines</i>							
Methyl pyrazine	833	0.24 <sup>a</sup>	0.12 <sup>b</sup>	0.16 <sup>ab</sup>	0.08 <sup>b</sup>	0.05	0.029
2-5/6-Dimethyl pyrazine	925	0.73	0.35	0.56	0.29	0.18	0.100
Trimethyl pyrazine	1000	0.19	0.91	0.17	0.73	0.05	0.172
2-Ethyl-3,5/6-dimethyl pyrazine	1086	0.09	0.07	0.12	0.06	0.02	0.184
Sum pyrazines		1.25	0.64	1.01	0.52	0.29	0.079
<i>Alkanes</i>							
Heptane	700						
Heptane	800	30.83 <sup>b</sup>	57.63 <sup>a</sup>	40.83 <sup>ab</sup>	42.35 <sup>ab</sup>	7.89	0.034
Octane		1.36 <sup>b</sup>	2.15 <sup>a</sup>	1.77 <sup>ab</sup>	1.71 <sup>ab</sup>	0.23	0.014

**Table 4.** Least-squares means of volatile flavor compounds (ng) from grilled beef steaks of five USDA quality grades and four muscles with significant interactions ( $P < 0.05$ )

USDA Quality Grade	Muscle	Acetaldehyde	2-Propanone	Dimethyl sulfide	Hexanal	Benzaldehyde	Octanal	Nonanal	Sum Sulfur containing
Prime	<i>Psoas major</i>	1.46 <sup>c</sup>	5.81 <sup>cd</sup>	0.91 <sup>cd</sup>	5.23 <sup>bcd</sup>	0.26 <sup>e</sup>	0.69 <sup>cd</sup>	1.47 <sup>bcd</sup>	1.04 <sup>ed</sup>
Upper 2/3 Choice	<i>Psoas major</i>	18.39 <sup>a</sup>	35.55 <sup>a</sup>	9.44 <sup>a</sup>	4.12 <sup>bcd</sup>	0.91 <sup>a</sup>	0.51 <sup>cd</sup>	1.26 <sup>bcd</sup>	10.59 <sup>a</sup>
Low Choice	<i>Psoas major</i>	2.28 <sup>c</sup>	1.92 <sup>cd</sup>	0.58 <sup>cd</sup>	3.91 <sup>cd</sup>	0.55 <sup>bcd</sup>	0.57 <sup>cd</sup>	1.10 <sup>cd</sup>	1.57 <sup>cde</sup>
Select	<i>Psoas major</i>	9.43 <sup>b</sup>	15.72 <sup>b</sup>	3.54 <sup>b</sup>	9.66 <sup>bcd</sup>	0.72 <sup>ab</sup>	1.98 <sup>ab</sup>	2.29 <sup>bc</sup>	4.44 <sup>b</sup>
Standard	<i>Psoas major</i>	2.28 <sup>c</sup>	10.83 <sup>bc</sup>	0.97 <sup>cd</sup>	27.12 <sup>ab</sup>	0.47 <sup>cde</sup>	2.11 <sup>a</sup>	3.69 <sup>a</sup>	1.39 <sup>de</sup>
Prime	<i>Longissimus lumborum</i>	2.90 <sup>c</sup>	4.13 <sup>cd</sup>	0.49 <sup>d</sup>	9.00 <sup>bcd</sup>	0.28 <sup>e</sup>	0.41 <sup>d</sup>	0.72 <sup>d</sup>	1.01 <sup>e</sup>
Upper 2/3 Choice	<i>Longissimus lumborum</i>	3.15 <sup>c</sup>	2.32 <sup>cd</sup>	0.19 <sup>d</sup>	10.83 <sup>bcd</sup>	0.33 <sup>de</sup>	0.72 <sup>cd</sup>	1.21 <sup>cd</sup>	0.75 <sup>e</sup>
Low Choice	<i>Longissimus lumborum</i>	1.64 <sup>c</sup>	3.00 <sup>cd</sup>	0.38 <sup>d</sup>	10.87 <sup>bcd</sup>	0.34 <sup>de</sup>	0.81 <sup>cd</sup>	1.43 <sup>bcd</sup>	1.01 <sup>e</sup>
Select	<i>Longissimus lumborum</i>	2.35 <sup>c</sup>	3.29 <sup>cd</sup>	0.69 <sup>cd</sup>	12.22 <sup>bcd</sup>	0.40 <sup>de</sup>	0.78 <sup>cd</sup>	1.32 <sup>bcd</sup>	1.36 <sup>ed</sup>
Standard	<i>Longissimus lumborum</i>	2.56 <sup>c</sup>	1.53 <sup>d</sup>	0.30 <sup>d</sup>	18.28 <sup>abcd</sup>	0.46 <sup>cde</sup>	1.23 <sup>bc</sup>	2.14 <sup>bc</sup>	0.98 <sup>e</sup>
Prime	<i>Gluteus medius</i>	1.53 <sup>c</sup>	3.11 <sup>cd</sup>	0.68 <sup>cd</sup>	7.55 <sup>bcd</sup>	0.42 <sup>de</sup>	0.76 <sup>cd</sup>	1.30 <sup>bcd</sup>	0.15 <sup>ed</sup>
Upper 2/3 Choice	<i>Gluteus medius</i>	2.29 <sup>c</sup>	9.62 <sup>bc</sup>	2.55 <sup>bc</sup>	13.79 <sup>bcd</sup>	0.39 <sup>de</sup>	0.88 <sup>cd</sup>	1.64 <sup>bcd</sup>	3.00 <sup>bcd</sup>
Low Choice	<i>Gluteus medius</i>	3.33 <sup>c</sup>	2.66 <sup>cd</sup>	0.92 <sup>cd</sup>	21.13 <sup>abc</sup>	0.52 <sup>bcd</sup>	1.37 <sup>abc</sup>	2.25 <sup>bc</sup>	1.57 <sup>de</sup>
Select	<i>Gluteus medius</i>	1.63 <sup>c</sup>	1.37 <sup>d</sup>	0.45 <sup>d</sup>	16.48 <sup>bcd</sup>	0.82 <sup>a</sup>	1.57 <sup>abc</sup>	2.44 <sup>b</sup>	1.29 <sup>ed</sup>
Standard	<i>Gluteus medius</i>	1.46 <sup>c</sup>	2.15 <sup>cd</sup>	0.56 <sup>cd</sup>	9.64 <sup>bcd</sup>	0.58 <sup>bcd</sup>	0.99 <sup>cd</sup>	2.05 <sup>bc</sup>	1.24 <sup>ed</sup>
Prime	<i>Semimembranosus</i>	1.77 <sup>c</sup>	3.82 <sup>cd</sup>	0.91 <sup>cd</sup>	25.81 <sup>abc</sup>	0.66 <sup>abc</sup>	1.57 <sup>abc</sup>	2.18 <sup>bc</sup>	1.56 <sup>ed</sup>
Upper 2/3 Choice	<i>Semimembranosus</i>	1.81 <sup>c</sup>	1.21 <sup>d</sup>	0.28 <sup>d</sup>	4.64 <sup>bcd</sup>	0.42 <sup>de</sup>	0.73 <sup>cd</sup>	1.33 <sup>bccd</sup>	0.77 <sup>e</sup>
Low Choice	<i>Semimembranosus</i>	3.08 <sup>c</sup>	9.73 <sup>bc</sup>	3.49 <sup>b</sup>	7.55 <sup>bcd</sup>	0.39 <sup>de</sup>	1.21 <sup>bc</sup>	2.14 <sup>bc</sup>	4.01 <sup>bc</sup>
Select	<i>Semimembranosus</i>	0.27 <sup>c</sup>	2.75 <sup>cd</sup>	0.61 <sup>cd</sup>	35.74 <sup>a</sup>	0.40 <sup>de</sup>	1.55 <sup>abc</sup>	2.37 <sup>bc</sup>	1.17 <sup>ed</sup>
Standard	<i>Semimembranosus</i>	1.37 <sup>c</sup>	5.26 <sup>cd</sup>	1.63 <sup>bcd</sup>	2.18 <sup>d</sup>	0.52 <sup>bcd</sup>	0.84 <sup>cd</sup>	1.40 <sup>bcd</sup>	2.22 <sup>bcd</sup>
	Std. Error	1.29	2.75	0.74	7.29	0.08	0.31	0.41	0.72
	<i>P</i> value	<0.001	<0.001	<0.001	0.017	<0.001	0.028	0.037	<0.001

610 <sup>abcde</sup> Means within a column lacking a common superscript differ ( $P < 0.05$ ).



**Table 5.** Pearson correlation coefficients (r) of consumer palatability scores<sup>1</sup> and proximate data<sup>2</sup> of grilled beef steaks from five USDA Quality Grades<sup>3</sup> and four muscles<sup>4</sup>

	Overall Liking	Tenderness	Juiciness	Flavor Liking	% Collagen	% Fat	% Moisture
Tenderness	0.79***						
Juiciness	0.75***	0.65***					
Flavor	0.85***	0.61***	0.65***				
% Collagen	0.10*	0.01	0.14**	0.13*			
% Fat	0.27***	0.22***	0.29***	0.27***	0.70***		
% Moisture	-0.23***	-0.16***	-0.24***	-0.23***	-0.68***	-0.97***	
% Protein	-0.28***	-0.25***	-0.29***	-0.26***	-0.57***	-0.64***	0.50***

611 <sup>1</sup> Consumer rated each steak on a 100-mm continuous line scale for flavor, tenderness, juiciness, and overall liking. On the scale, 0  
612 was verbally anchored as not tender, not juicy, dislike flavor extremely, and dislike overall extremely. Similarly, 100 was verbally  
613 anchored as very tender, very juicy, like flavor extremely, and like overall extremely.

614 <sup>2</sup> Chemical percentages of fat, moisture, protein, and collagen determined of raw steaks by AOAC official method (2007.04;  
615 Anderson, 2007).

616 <sup>3</sup> Beef quality grades included: Prime, Upper 2/3 Choice, Low Choice, Select, and Standard.

617 <sup>4</sup> Beef muscles included: *Psoas major*, *Longissimus lumborum*, *Gluteus medius*, and *Semimembranosus*.

618 \* Significant correlation ( $P < 0.05$ )

619 \*\* Significant correlation ( $P < 0.01$ )

620 \*\*\* Significant correlation ( $P < 0.001$ )

**Table 6.** Pearson correlation coefficients (r) between *n*-aldehydes, flavor liking<sup>1</sup>, overall liking<sup>1</sup> and % fat<sup>2</sup> for grilled beef steaks from five USDA Quality Grades<sup>3</sup> and four muscles<sup>4</sup>

	Flavor liking	Overall liking	% Fat
<i>n</i> -Aldehydes			
Pentanal	-0.15	-0.13	-0.16
Hexanal	-0.17	-0.14	-0.16
Heptanal	-0.18	-0.16	-0.28**
Octanal	-0.19	-0.15	-0.39***
Nonanal	-0.24*	-0.17	-0.41***
Decanal	-0.25*	-0.22*	-0.19
Sum C <sub>5</sub> -C <sub>10</sub> n-Aldehydes	-0.18	-0.15	-0.17

621 <sup>1</sup> Consumer rated each steak on a 100-mm continuous line scale for flavor liking and overall  
 622 liking. On the scale, 0 was verbally anchored as dislike flavor extremely, and dislike overall  
 623 extremely. Similarly, 100 was verbally anchored as like flavor extremely, and like overall  
 624 extremely.

625 <sup>2</sup> Chemical percentages of fat, moisture, protein, and collagen determined of raw steaks by  
 626 AOAC official method (2007.04; Anderson, 2007).

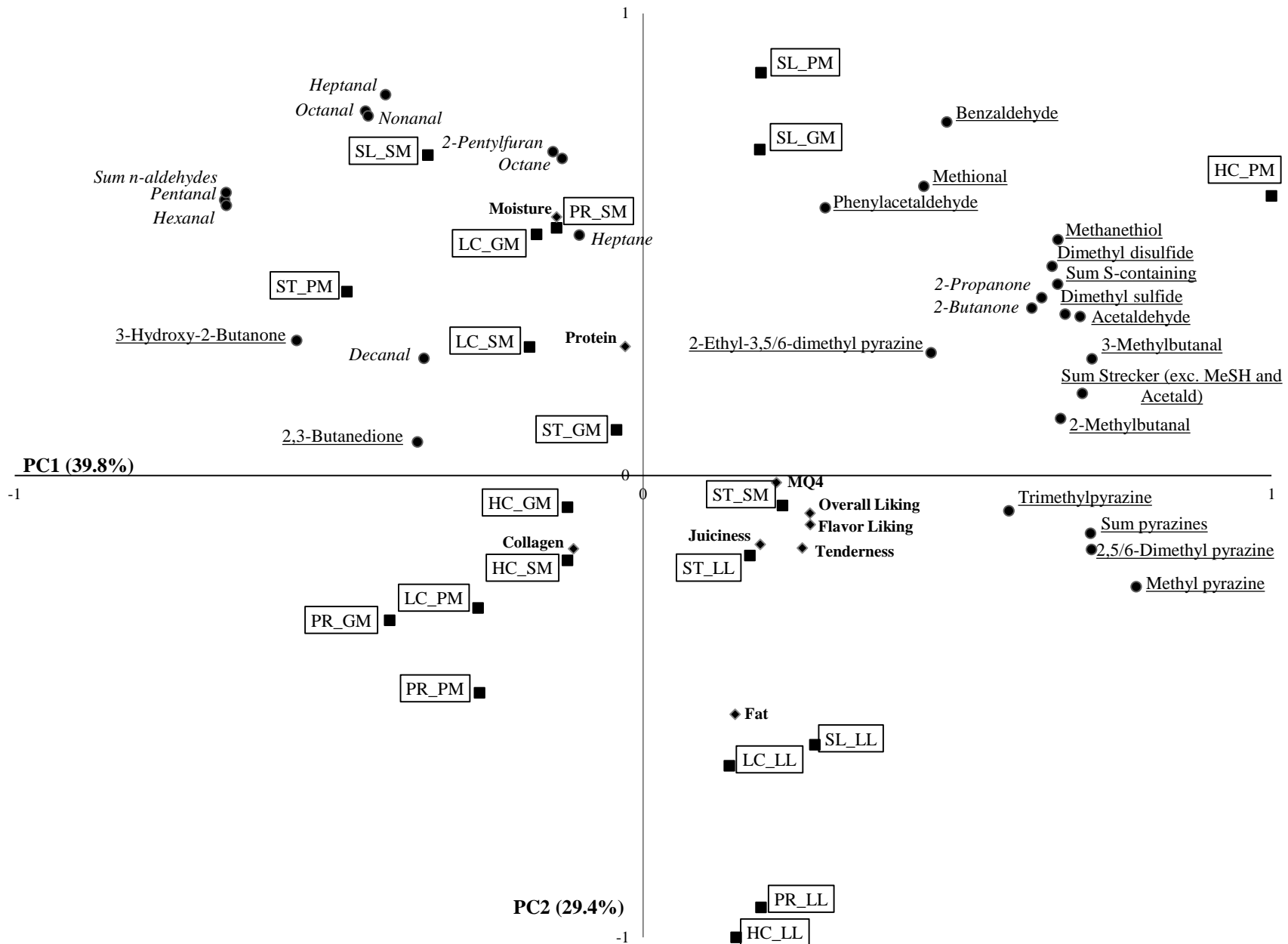
627 <sup>3</sup> Beef quality grades included: Prime, Upper 2/3 Choice, Low Choice, Select, and Standard.

628 <sup>4</sup> Beef muscles included: *Psoas major*, *Longissimus lumborum*, *Gluteus medius*, and  
 629 *Semimembranosus*.

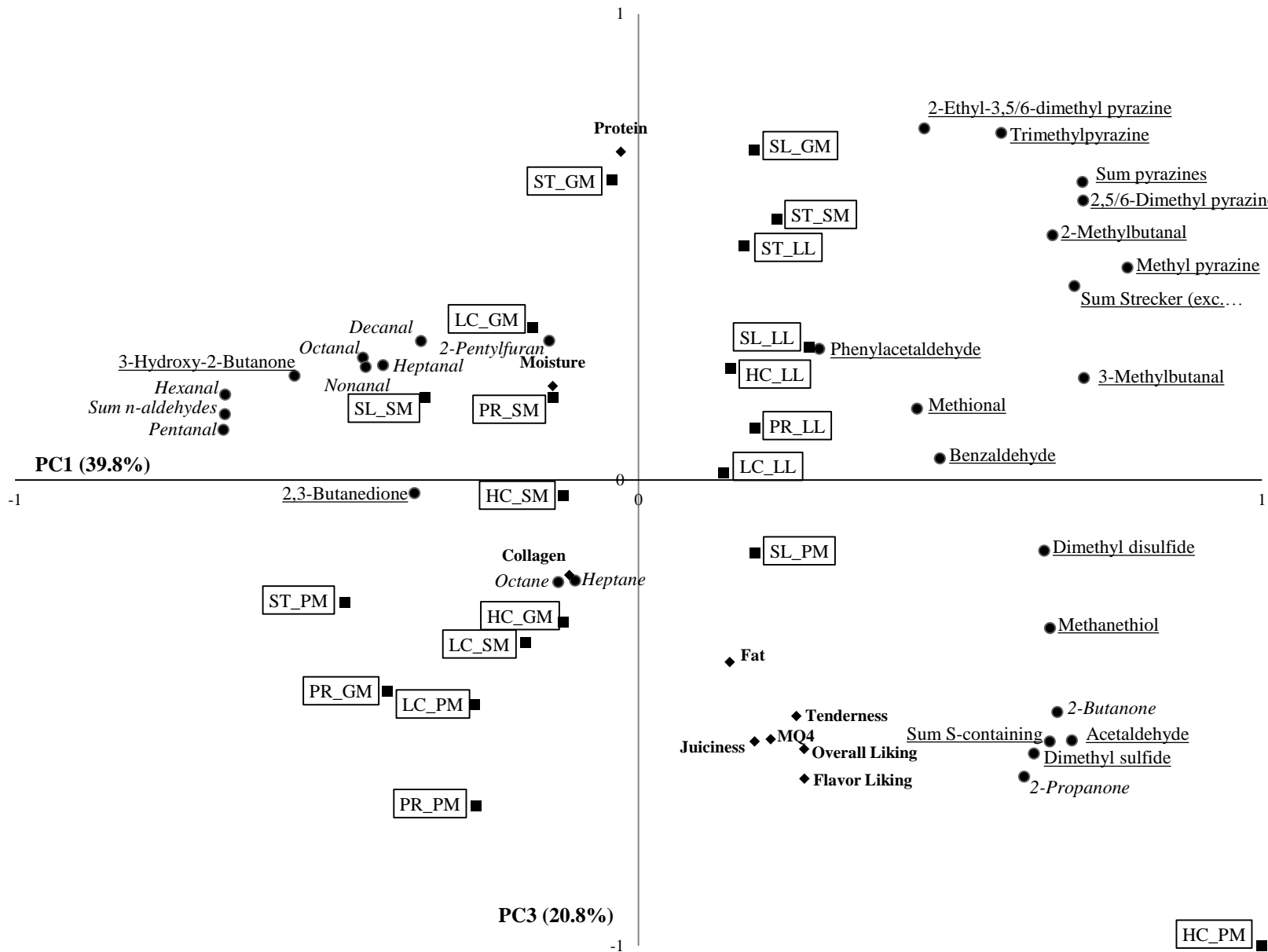
630 \* Significant correlation ( $P < 0.05$ )

631 \*\* Significant correlation ( $P < 0.01$ )

632 \*\*\* Significant correlation ( $P < 0.001$ )



633 **Figure 1.** Principal component (PC) analysis for volatile compounds, of five USDA quality grades (Prime = PR, Upper 2/3 Choice = HC, Low Choice = LC, Select = SL, Standard = ST) and four  
634 muscles (*Psoas major* = PM, *Longissimus lumborum* = LL, *Gluteus medius* = GM, *Semimembranosus* = SM). Volatile compound groups shown with different formatting: Maillard products and lipid  
635 oxidation products. **Consumer palatability traits and proximate data (%)** were correlated on the same axes.  
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**Figure 2.** Principal component (PC) analysis for volatile compounds, of five USDA quality grades (Prime = PR, Upper 2/3 Choice = HC, Low Choice = LC, Select = SL, Standard = ST) and four muscles (*Psoas major* = PM, *Longissimus lumborum* = LL, *Gluteus medius* = GM, *Semimembranosus* = SM). Volatile compound groups shown with different formatting: Maillard products and lipid oxidation products. **Consumer palatability traits and proximate data (%) were correlated on the same axes.**