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Effects of Intensive Alfalfa Feeding on Meat Quality and Fatty Acid Profile of Korean Native Black Goats

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Abstract The aim of this study was to determine meat quality characteristics and fatty acid composition of Korean native black goats (KNBG) finished on intensive feeding of alfalfa (ALF) and conventional feeding of commercial concentrate pellets (CCP) with low-energy common grasses. Ten KNBG (12 months old) were divided into two groups and subjected to either ALF or CCP treatments. The goats were slaughtered after 6 months of feeding with experimental diets to investigate meat quality characteristics and fatty acid compositions of *longissimus lumborum* muscle. There were no significant differences in proximate chemical composition, collagen, or myoglobin content between ALF and CCP groups of goats. Meat color, water-holding capacity, or tenderness was not significantly different between the two groups either. However, proportions of monounsaturated fatty acids and polyunsaturated fatty acids were significantly ($p < 0.05$) different between the two groups. The proportion of oleic acid was significantly ($p < 0.05$) higher in ALF goat whereas proportion of linoleic and arachidonic acids were significantly ($p < 0.05$) higher in CCP goat. Results suggest that KNBG finished with intensive feeding of alfalfa could produce goat meat with desirable fatty acids for human diets.

Keywords Korean native black goat, goat meat quality, fatty acid profile, alfalfa, goat feed

Introduction

Goats are appropriate under semi-arid and punitive environmental conditions because of their browsing behaviors and adaptation to severe climates. For these reasons, in many Asian countries, goat meat is the main source of animal protein. Korean native black goat (KNBG; *Capra hircuscoreanae*) has been domesticated about 2000 years back in Korea. Consumption of KNBG has been improved significantly with increasing consumption of meat products in last decades. KNBG has conventionally been known as healthy food, with nutritional features superior in other meat sources generally

consumed. According to Kim et al. (1993), KNBG has lower cholesterol and saturated fat contents compared to beef and pork. In general, as compared to other ruminants, goat meat has desirable fatty acids with moderately higher proportion of polyunsaturated fatty acids (Mushi et al., 2009).

It is well known that diet leads to alterations in meat quality traits and nutrients, because biological characteristics of muscular tissue can be affected by diet changes (Renand et al., 2001). It has been shown that diet is one of the most important factor influencing carcass yield, cutability and meat qualities of many species (Wood et al., 2008), especially goats (Warmington and Kirton, 1990). Alfalfa (*Medicago sativa* L.) is valuable grass due to its uniform distribution and high growth rate during the growing season. Alfalfa also has an outstanding nutritive value, leading to tremendous outcomes for feeding lambs (Fraser et al., 2004). Alfalfa is the main pasture legume for numerous species of livestock (Van Keuren and Matches, 1988) and alfalfa hay is often used to supply crude protein in finishing diets for ruminants. It is cultivated in more than 80 countries in an area exceeding 35 million ha (Radović et al., 2009). World production of alfalfa was around 436 million tons in 2006 (FAO, 2006). The cost of high quality of alfalfa bales is US \$80–100/Metric Tons.

In general, various kinds of forages are available for small ruminants in relation to finishing system such as energy and protein. Because growing animals require more nutrients than mature animals, the selection of forage species is much important. However, the effect of intensive alfalfa feeding on meat quality and fatty acid profile of KNBG is currently unclear. Therefore, the objective of this study was to assess meat quality characteristics and fatty acid compositions of KNBG finished on intensively-managed alfalfa or conventional feeding of commercial concentrate pellets (CCP) with *ad libitum* low-energy common grasses in Korea.

Material and Methods

Animals and sampling

Ten KNBG (approximately 12 month of age) were allocated into two groups and subjected to either intensive feeding of alfalfa grasses (ALF: crude protein, 15.3%; fiber, 28.6%) or conventional feeding of CCP (crude protein, 15.15%; crude fat, 3.88%; crude ash, 6.75%; moisture 11.5%; TDN, 68%) with *ad libitum* low-energy common grasses including rice straw, Italian ryegrass, and Timothy grass. Because the most abundant roughage for ruminants is rice straw in Korea, it was used as major source of low-energy common grass in this experiment. Animals were stratified by two dietary treatments and assigned into five pens for each dietary group. They were slaughtered at 6 months after feeding experiment diets. Feed were provided *ad libitum* twice daily. Animals were allowed free access to freshwater placed in water buckets.

Goats were slaughtered according to standard protocol at an average live body weight of 48.3±3.8 kg. They were subjected to electrical stunning (about 220 voltages) followed by exsanguination, skinning, evisceration, and washing procedures. The chilling process of carcasses were carried out in chilling room at 2°C for 24 h. Left and right *longissimus lumborum* (LL) muscles were cut to measure meat quality traits.

Meat quality traits

Meat color was determined by measuring L* (lightness), a* (redness/greenness) and b* (yellowness/blueness) values using a Minolta Chromameter (Minolta CR-300, Japan) that was calibrated with a white plate (Y=93.5, X=0.3132, y=0.3198). An average value of three different locations on the meat surface was used for statistical analysis.

Water-holding capacity (WHC) was evaluated by three methods including the released water (RW) %, drip loss (DL) %, and cooking loss (CL) %.

and cooking loss (CL) %, DL % and CL % were determined as described previously (Hwang et al., 2010). RW % was determined by the method of Joo (2018). A meat sample (3.0 ± 0.05 g) was placed on a previously dried and weighed filter paper (Whatman No. 1 of 11 cm of diameter) with two thin plastic films. After weighing them with electrical balance, the filter-paper and plastic film with meat sample were positioned between Plexiglas plates. A load of 2.5 kg and free mechanical force was applied for 5 min. The wet filter paper and plastic films were quickly weighed after precisely removing the compressed meat. The percentage of RW was calculated as follows: $RW \% = [(Damp\ filter-paper\ and\ plastic\ films\ weight) - (filter-paper\ and\ plastic\ films\ weight) / Meat\ sample\ weight] \times 100$.

Warner-Bratzler shear force (WBSF) was carried out for the cooked sample based on AMSA (1995) guideline. Toward parallel into myofiber direction sample core (1.0 cm-diameter) were inserted. The cores were sheared perpendicular to myofiber's direction using an Instron tensile testing machine (Model 4443, Instron Corp., USA) with a V-shaped shear blade. Ultimate force was obtained using 100 N load cell tension applied at a crosshead speed of 250 mm/min. The full-scale load was 50 kg. Maximum peak force recorded during the test was stated as shear force. The average of five readings was recorded for each sample group.

Chemical analysis

Moisture content and crude ash content were determined by applying the oven-drying method (AOAC, 1995). Crude protein was determined by the Kjeldahl method (AOAC, 1995). Fat content was determined using the Folch method (Folch et al., 1957) as described previously (Hwang et al., 2010).

Collagen content was measured by the method of AOAC (1995). Briefly, a 4 g of meat sample was taken and 30 mL of H_2SO_4 was added to the sample for hydrolysis at $105^\circ C$ for 16 h. Water was poured into funnel to reach up to total volume of 500 mL after hydrolysis. A filtered solution of 5 mL was diluted to 100 mL. For measurement of hydroxyproline content, 2 mL diluted solution was taken after adding 1 mL of oxidation solution (50 mM of chloramine-T hydrate, 156 mM citric acid, 375 mM NaOH, 661 mM sodium acetate trihydrate, 29% v/v 1-propanol, pH 6.0). Vortexing of samples were carried out and then samples were kept at room temperature for 20 min by adding 1 mL of color reagent. The vortexed samples were covered with aluminum foil and placed in water bath at $60^\circ C$ for 15 min. Before measuring the absorbance level, the samples were cooled down by holding it to running water for 3 min. The absorbance level was carried out by using UV spectrometer at wavelength of 558 nm. From standard curve (1.2, 2.4, 3.6, and 4.8 μg hydroxyproline per mL of H_2O), Hydroxyproline content was calculated. Results were computed by following formula: Total collagen = hydroxyproline $\times 8$.

Myoglobin (Mb) concentration was determined by the method of Warriss (1979). Fatty acids composition was determined using a gas chromatograph (HP6890N, Hewlett-Packard, USA) equipped with a HP7683 (Hewlett-Packard) automatic sampler as described previously (Hwang et al., 2010). Atherogenic index was calculated as content ratio of saturated fatty acid (SFA)/unsaturated fatty acid (UFA) using the following formula suggested by Ulbricht and Southgate (1991): AI (atherogenic index) = $[C12:0 + 4(C14:0) + C16:0] / [MUFA + PUFA]$.

Statistical Analysis

Statistical analysis was performed with one-way analysis of variance (one-way ANOVA) by applying SPSS (SPSS 16.0, Chicago, IL, USA). Differences between mean values of two dietary treatments were obtained by Duncan's multiple range test. Significance was defined at $p < 0.05$. Results were expressed as the least square mean values of three independent replications except that WBSF was recorded as the average of five measurements.

Results and Discussion

Changes in meat quality traits

Meat color measurements of LL muscles from KNBG finished on ALF and CCP are shown in Table 1. There were no significant differences in lightness (L^*), redness (a^*), or yellowness (b^*) between ALF and CCP goats. These results support previous reports that dietary treatments have no effect on meat color (Dransfield et al., 1990). Instead, meat color is more affected by breed (Kadim et al., 2003), slaughter weight (Beraiain et al., 2000), and aging (Kadim et al. 2003; Kannan et al., 2006). According to Priolo et al. (2002), feeding of low-energy diets to ruminants causes darker color (lower L^* values) of *Longissimus* muscle as compared to finishing them with high energy diets. Finishing lambs with grass had darker color meat as compared to concentrate fed lambs (Priolo et al., 2002). Realini et al. (2004) have also reported that pasture-fed steers had darker color on the surface of *Longissimus* muscle in contrast to concentrate-fed steers. However, steers that fed on several combinations of forage-concentrate diets for a short period had similar instrumental color values of *Longissimus* muscle (French et al., 2001). It is recommended that feeding period should be extended to improve the meat color by diet. In current study, there was no significant difference in myoglobin content in loins between the two groups of goats either. This implies that intensive feeding of alfalfa does not influence myoglobin concentration or meat color of KNBG.

As shown in Fig. 1, there was no significant difference in WHC of goat meats between the two groups (ALF and CCP). There were no significant differences in RW % or DL % between the two groups either. Beraiain et al. (2000) have reported a higher range of WHC in *longissimus dorsi* (LD) muscles of lambs by slaughter weight. Kadim et al. (2003) have also reported differences in WHC of different muscles among goat breeds (*Batina*, *Dhofari*, and *Jabal Akdhar*) in Oman. Although several studies reported that WHC of goat meat was influenced by breeds, slaughter weight, animal age, and muscle differences, our data clearly showed that the WHC of KNBG loin was not significantly affected by intensive feeding (alfalfa or CCP for goat).

Results of WBSF and collagen content of ALF and CCP goats are presented in Table 2. There were no significant differences in WBSF or collagen content between ALF and CCP goats. This might be due to the lack of significant difference in meat color and WHC parameters between the two treatments. Safari et al. (2009) have reported no changes in CL or shear force among goat fed on different nutritional regimes. Similar results have been reported by Kannan et al. (2006) and Smith et al. (1979). Connective tissue is a crucial element of organisms. It applies structural function as an aggregator to support cells. Meat tenderness is strongly related to collagen content and solubility. Our results clearly showed no difference in WBSF due to no difference in collagen content between ALF and CCP goats. This implies that tenderness or collagen content is not

Table 1. Color measurements and myoglobin % of *longissimus lumborum* muscle from goat finished on intensive feeding of alfalfa (ALF) or conventional feeding of commercial concentrate pellets (CCP) with low-energy common grasses

| Color traits | ALF | CCP |
|----------------------|------------|------------|
| Lightness (L^*) | 33.37±0.58 | 33.11±0.54 |
| Redness (a^*) | 17.41±0.72 | 17.50±0.86 |
| Yellowness (b^*) | 1.87±0.23 | 1.77±0.19 |
| Myoglobin (mg/g) | 7.34 ±0.25 | 6.93±0.59 |

Data are presented as least square mean±SE.

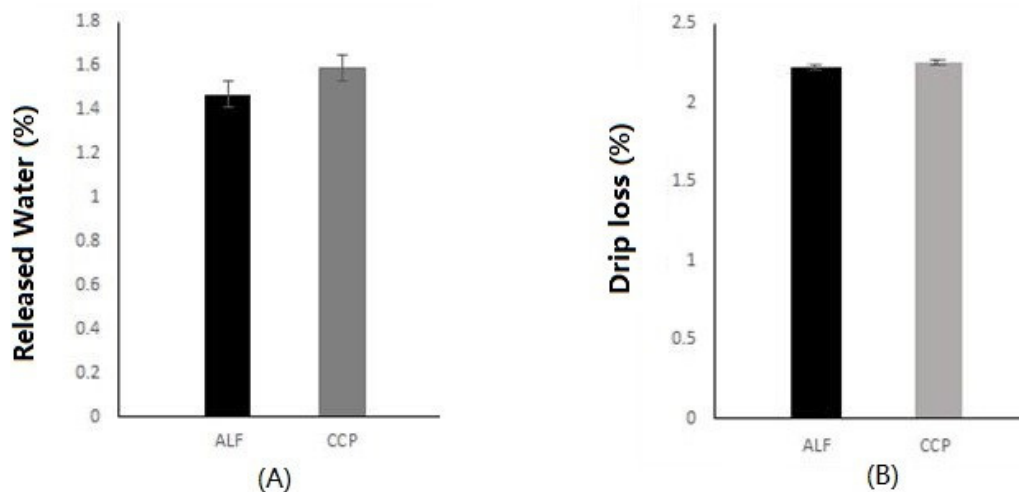


Fig. 1. Water holding capacity (A: released water %, B: drip loss %) of *longissimus lumborum* from goat finished on intensive feeding of alfalfa (ALF) or conventional feeding of commercial concentrate pellets (CCP) with low-energy common grasses. Data are presented as least square mean \pm SE.

Table 2. Shear force and collagen content of *longissimus lumborum* muscle from goat finished on intensive feeding of alfalfa (ALF) or conventional feeding of commercial concentrate pellets (CCP) with low-energy common grasses

| Parameters | ALF | CCP |
|---------------------------|-----------------|-----------------|
| WBSF(kg/cm ²) | 3.92 \pm 0.14 | 4.05 \pm 0.16 |
| Collagen (%) | 0.67 \pm 0.18 | 0.90 \pm 0.16 |

Data are presented as least square mean \pm SE.

influenced by diet of KNBG. This is similar to results of Van Niekerk and Casey (1988) on Boer goats. Our results showed that meat quality traits including color, WHC, and tenderness were not changed by intensive feeding of alfalfa compared to conventional feeding CCP for KNBG.

Proximate chemical and fatty acid composition

Effects of intensive feeding of alfalfa on proximate chemical composition of KNBG are shown in Table 3. Goat meat did not significantly differ in moisture, crude protein, fat, or ash content between ALF and CCP groups. Generally, the important factors which effect the chemical composition of meat are breed, age, sex, anatomical location of muscle and nutrition (Lawrie, 1998). However, in the current study, the chemical composition of KNBG meat was not influenced by different feeding of diet for 6 months before slaughter. It is normally recognized that alfalfa contains higher crude protein than common grasses (Turner et al., 2014). This designates that intensive feeding of alfalfa could substitute conventional feeding of CCP for KNBG because of its higher protein content. Schmidt et al. (2013) have stated that total ash, protein, or lipid concentration in longissimus muscle does not vary when beef steers are finished on monocultures of alfalfa. Results of proximate chemical analysis of KNBG were similar to those reported by Hopkins-Shoemaker (2006). Animals fed on higher concentrate diet had higher fat content in muscle (French et al., 2001; Geay et al., 2001). Marino et al. (2006) have also reported that the fat content in LD muscle of concentrate fed steers were two times higher compared to pasture-fed steers.

Fatty acid compositions of KNBG meat finished on ALF and CCP are summarized in Table 4. Proportions of

Table 3. Proximate chemical analysis of *longissimus lumborum* muscle from goat finished on intensive feeding of alfalfa (ALF) or conventional feeding of commercial concentrate pellets (CCP) with low-energy common grasses

| Parameters | ALF | CCP |
|------------|------------|------------|
| Moisture | 75.82±0.09 | 76.00±0.15 |
| Protein | 21.14±0.07 | 21.06±0.17 |
| Fat | 1.91±0.06 | 1.64±0.08 |
| Ash | 1.13±0.04 | 1.30±0.07 |

Data are presented as least square mean±SE.

Table 4. Fatty acid profiles of *longissimus lumborum* muscle from goat finished on intensive feeding of alfalfa (ALF) or conventional feeding of commercial concentrate pellets (CCP) with low-energy common grasses

| Fatty acids | ALF | CCP | SEM | p-level |
|-------------|--------------------|--------------------|-------|---------|
| C12:0 | 0.08 | 0.08 | 0.006 | 0.3466 |
| C14:0 | 2.00 ^A | 1.78 ^B | 0.09 | 0.0061 |
| C14:1 | 0.13 | 0.10 | 0.02 | 0.1281 |
| C16:0 | 23.89 | 22.83 | 2.07 | 0.4398 |
| C16:1 | 3.26 ^A | 2.89 ^B | 0.192 | 0.01 |
| C18:0 | 17.12 | 18.16 | 1.59 | 0.3773 |
| C18:1 n9 | 44.58 ^A | 41.00 ^B | 1.240 | 0.001 |
| C18:2 n6 | 4.56 ^B | 6.00 ^A | 0.360 | 0.002 |
| C18:3 n3 | 1.71 | 2.05 | 0.51 | 0.3227 |
| C20:0 | 0.08 | 0.08 | 0.01 | 0.7422 |
| C20:4 n6 | 1.61 ^B | 3.35 ^A | 0.522 | 0.0008 |
| C20:5 n3 | 0.73 ^B | 1.47 ^A | 0.371 | 0.0137 |
| C22:6 n3 | 0.10 ^B | 0.21 ^A | 0.061 | 0.0186 |
| SFA | 43.17 | 42.92 | 1.129 | 0.7314 |
| MUFA | 47.97 ^A | 44.00 ^B | 1.30 | 0.0014 |
| PUFA | 8.86 ^B | 13.08 ^A | 1.346 | 0.001 |
| PUFA/SFA | 0.21 ^B | 0.30 ^A | 0.035 | 0.002 |
| n6 | 6.17 ^B | 8.12 ^A | 1.780 | 0.1222 |
| n3 | 2.53 ^B | 3.73 ^A | 0.805 | 0.0463 |
| AI | 0.57 | 0.53 | 0.092 | 0.142 |

^{A,B} Treatment means within a row with different letters are different at the p level indicated. SFA, saturated fatty acids; MUFA, monounsaturated fatty acids; PUFA, polyunsaturated fatty; AI, atherogenic index.

monounsaturated fatty acids (MUFA) and polyunsaturated fatty acids (PUFA) were significantly ($p < 0.05$) different between ALF and CCP goats. However, proportions of SFA were not significantly different between ALF (43.17%) and CCP (42.92%) in goats. ALF goat had significantly ($p < 0.05$) higher proportion of MUFA than CCP goat mainly due to higher proportion of oleic acid (C18:1). The significantly lower proportion of PUFA in ALF goat was due to lower proportions of linoleic acid (C18:2) and arachidonic acid (C20:4). However, proportion of palmitic acid (C16:0) or stearic acid (C18:0) in ALF goat was not significantly different from that in CCP goat, resulting in no difference in SFA between ALF and CCP goats. All these

results suggest that fatty acid composition of KNBG meat could be strongly influenced by intensive feeding of alfalfa.

According to Webb et al. (2005) and Marino et al. (2006), the meat obtained from pasture-fed animals consist equivalent quantity of SFA, a lower amount of MUFA and higher content of PUFA in comparison to concentrate-fed animals. Fraser et al. (2004) reported the similar amount of fatty acid in comparison to present study, showing that C18:1, C16:0, and C18:0 were the majority of fatty acids in *longissimus* muscle from lambs finished on various kinds of grasses including alfalfa. Solaiman et al. (2011) have also reported that C18:1, C16:0, and C18:0 are the major fatty acids in *longissimus* muscle of Boer goat kids fed on annual ryegrass. In the present study, proportions of C18:1, C16:0, and C18:0 in LL muscles from ALF goat were 44.58%, 23.89%, and 17.12%, respectively. These results suggest that prominent fatty acids in KNBG meat might not change with intensive feeding of alfalfa. However, our data showed that intensive feeding of alfalfa might improve the palatability of KNBG because oleic acid (C18:1 cis-9) and stearic acid (C18:0) were the major constituents of intramuscular fat in goats (Rhee et al., 2000).

Proportion of most PUFA in ALF goat were significantly ($p < 0.05$) lower than those in CCP goat. Particularly, proportions of linoleic acid (C18:2) and arachidonic acid (C20:4) as prominent n-6 PUFA were significantly ($p < 0.05$) lower in ALF goat. However, there was no significant ($p > 0.05$) difference in proportion of linolenic acid (C18:3), the prominent n-3 PUFA, between ALF and CCP goats. These differences in fatty acid compositions might be due to different crude protein and TDN contents in goat diet. Alteration in fatty acid compositions could be attributed to changes in microbial distribution in the rumen due to increased crude protein and TDN level (Majdoub-Mathlouthi et al., 2013). Our data suggest that desirable composition of fatty acids for human health could be obtained by intensive feeding with alfalfa to decrease n-3 and n-6 PUFA simultaneously. In addition, our results clearly showed that AI was not significantly changed by intensive feeding of alfalfa.

Conclusions

KNBG finished on ALF and CCP had similar meat quality traits including color measurements, myoglobin content, WHC measurements, WBSF, and collagen content. Proximate chemical compositions of ALF goat were not significantly different from that those of CCP goat either. However, proportions of MUFA and PUFA were significantly different between ALF and CCP goats. Especially, the proportion of oleic acid was higher in ALF goat whereas proportions of linoleic and arachidonic acids were higher in CCP goat. These results suggest that KNBG finished with intensive feeding of alfalfa could produce goat meat with desirable fatty acids for human diets.

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