Genetic, management, and nutritional factors affecting intramuscular fat deposition in beef cattle — A review

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Abstract: Intramuscular fat (IMF) content in skeletal muscle including the *longissimus dorsi* muscle (LM), also known as marbling fat, is one of the most important factors determining beef quality in several countries including Korea, Japan, Australia, and the United States. Genetics and breed, management, and nutrition affect IMF deposition. Japanese Black cattle breed has the highest IMF content in the world, and Korean cattle (also called Hanwoo) is the second highest. Here, we review results of research on genetic factors (breed and sex differences and heritability) that affect IMF deposition. Cattle management factors are also important for IMF deposition. Castration of bulls increases IMF deposition in most cattle breeds. The effects of several management factors, including weaning age, castration, slaughter weight and age, and environmental conditions on IMF deposition are also reviewed. Nutritional factors, including fat metabolism, digestion and absorption of feed, glucose/starch availability, and vitamin A, D, and C levels are important for IMF deposition. Manipulating IMF deposition through developmental programming via metabolic imprinting is a recently proposed nutritional method to change potential IMF deposition during the fetal and neonatal periods in rodents and domestic animals. Application of fetal nutritional programming to increase IMF deposition of progeny in later life is reviewed. The coordination of several factors affects IMF deposition. Thus, a combination of several strategies may be needed to manipulate IMF deposition, depending on the consumer's beef preference. In particular, stage-specific feeding programs with concentrate-based diets developed by Japan and Korea are described in this article.

Keywords: Beef Cattle; Intramuscular Fat Deposition; Genetic Factors; Management; Nutrition

INTRODUCTION

Intramuscular fat (IMF) deposition or marbling in muscle including the *longissimus thoracis et lumborum*, *infraspinatus*, and *biceps femoris*, especially in Korean cattle (also called Hanwoo) and Japanese Black cattle, is one of the most important traits influencing beef quality and palatability attributes. Marbling is positively correlated with juiciness, tenderness, and palatability of beef [1]. For example, beef with higher IMF produces higher sensory traits, including tenderness, juiciness, flavor, and overall acceptability in Korean cattle [2] and in USA beef samples [3]. Most marbling fat is deposited between muscle fibre bundles in the perimysial connective tissue in beef cattle [4]. Marbling adipocytes are also found within muscle bundles in the *longissimus dorsi* muscle (LM) of high-quality grade Japanese Black cattle [5]. The production of highly marbled beef in Korea is essential to compete with cheap imported beef that has a relatively low marbling score (MS).

Several countries, including Korea, Japan, Australia and the USA, have considered IMF...
content as important to beef quality grade (QG), whereas European countries, including France and Germany, prefer leaner meat (less IMF content). Understanding the factors affecting IMF deposition and the mechanisms responsible for its deposition is important when manipulating IMF deposition. Our laboratory has studied IMF deposition for more than a decade to understand the mechanisms responsible for IMF deposition in beef cattle [6-12], and we have reviewed molecular mechanisms responsible for IMF deposition in cattle [13].

In this review, we first provide an overview of muscle and fat cell growth in ruminants. Factors affecting IMF deposition can be sub-categorized into genetic (breed and sex differences, heritability), management (weaning age, castration, slaughter weight and age, and environment), and nutritional factors (fat metabolism, fat digestion and absorption, glucose/starch availability, roughage vs concentrate ratio, dietary energy and protein levels, vitamins A, D, and C levels, fetal and neonatal nutritional programming, and stage-specific feeding systems). Here, we also review these factors affecting IMF deposition in cattle.

**OVERVIEW OF MUSCLE AND FAT CELL GROWTH IN RUMINANTS**

Skeletal muscle including the LM mainly consists of skeletal muscle fibres (myofibres), but it also contains fat cells, connective tissue or extracellular matrix, and various other cell types including satellite cells, immune cells and blood vessels. The ratio of muscle to fat cells varies depending on the amount of IMF deposited. Muscle and fat cell growth typically occur as mass increases due to both hyperplasia (increase in cell number) and hypertrophy (increase in cell size) [14,15]. It is important to identify the mechanisms responsible for initial muscle and fat cell growth during the developmental stage of cattle to better understand IMF deposition in later life. Muscle and fat cell development are dependent on myogenesis (muscle cell formation) and adipogenesis (fat cell formation), respectively [16]. The process of skeletal muscle and adipose tissue development during the fetal stage in laboratory animals and livestock including ruminants has been well described in reviews by Du et al [16-18]. Du et al [16] proposed that myogenic, adipogenic, and fibrogenic cells are derived from common mesenchymal progenitor cells. Briefly, they proposed that mesenchymal stem cells are first committed to the myogenic and adipogenic–fibrogenic cell lineages during fetal muscle development [16]. Then, myogenic progenitor cells further develop into muscle fibres and satellite cells, while adipogenic–fibrogenic progenitor cells develop into the stromal–vascular fraction of skeletal muscle, where adipocytes, fibroblasts, and resident adipogenic–fibrogenic progenitor cells reside [16]. Du et al [17] suggested that a shift in development of mesenchymal stem cells from myogenesis to adipogenesis may increase IMF deposition later in life.

Adipogenesis is a comprehensive term that includes commitment of mesenchymal stem cells to preadipocytes, determination and proliferation of preadipocytes, and differentiation of preadipocytes into mature adipocytes [19]. Adipogenesis normally begins around mid-gestation in ruminant animals [20]. Differentiation of adipogenic cells occurs during fetal and postnatal stages. In the fetal stage, the main process is increasing the number of adipocytes (hyperplasia) derived from adipogenic–fibrogenic progenitor cells [21]. It has been suggested that adipocyte formation in muscle begins approximately between late gestation and 250 d after birth in cattle [16]. Hyperplasia of fat cells mainly occurs from conception until birth, and hypertrophy is a greater contributor to IMF deposition than is hyperplasia from 250 d of age to slaughter [16,17]. Both hyperplasia and hypertrophy are involved in IMF deposition (Figure 1). Therefore, strategies that up-regulate both hyperplasia and hypertrophy of fat cells are effective ways to improve IMF deposition.

After birth, the adipogenic–fibrogenic progenitor cells remain on some parts of the stromal vascular cells in adipose tissue, later serving as precursors to preadipocytes and adipocytes [16,22]. The nutritional and physiological condition of the fetus and the early postnatal stage affects the number of adipocytes in adipose tissues of animals [16]. Thus, managing maternal nutrition is a potentially effective way to promote adipogenesis in fetal muscle [16]. Du et al [17] suggested that the order of effectiveness of nutritional management for manipulating IMF deposition is fetal stage>neonatal stage>weaning stage>older stages. Harper and Pethick [23] recommended that producers focus on individuals with high genetic capacity to create more preadipocytes within animal muscle. Then, preadipocytes can differentiate during the finishing stage when they fill with lipid to grow into visible marbling by hypertrophy. Albrecht et al [24] reported that hyperplasia of preadipocytes also plays an important role in marbling formation during postnatal growth of IMF because new marbling flecks appear after recruitment and filling with preadipocytes.

Four major adipose tissue depots exist in animals: abdominal, subcutaneous, intermuscular, and intramuscular. Adipogenesis in beef cattle is first initiated in abdominal (visceral) adipocytes around mid-gestation [16]. According to Du et al [16], this adipogenic phase extends to the neonatal period for abdominal adipocytes, to the early weaning stage for subcutaneous and intermuscular adipocytes, and to around 250 d of age for IMF. Thus, during the late fattening stage of beef cattle, marbling adipocytes may grow continuously even though development of abdominal, intermuscular, and subcutaneous fat may slow or cease.
GENETIC FACTORS

Breed and sex differences
Intramuscular fat content in the LM varies significantly depending on the cattle breed. The LM of Wagyu contains the highest fat content (36.5%; average value from two publications [25,26]), and the LM from Korean cattle contains the second highest fat content worldwide (13.7%; average value from four publications [10,27-29]), followed by Angus (9.3%; average value from three publications [30-32]), Hereford crossbred or Hereford (7.6%; average value from two publications [30,33]), and Brahman (2.8%; average value from four publications [32,34-36]) (Table 1). Bos taurus cattle generally have higher marbling and better tenderness than do Bos indicus animals.

Table 1. Intramuscular fat contents in several cattle breeds

| Breed               | No. of animals | Feeding system       | Sex      | Slaughter age±SE (mo) | Slaughter wt ±SE (kg) | Carcass wt ±SE (kg) | Intramuscular fat±SE (%) | Muscle type | Reference |
|---------------------|----------------|----------------------|----------|-----------------------|-----------------------|-----------------------|------------------------|-------------|-----------|
| Wagyu               | 6              | feedlot-concentrate  | Steer    | 26.0                  | 640 ± 19              | 393 ± 13              | 34.3 ± 1.7             | LT          | [25]      |
| Wagyu               | 10             | feedlot-concentrate  | Cow      | 22.0                  | 701 ± 17              | 444 ± 11.9            | 37.8 ± 1.00            | LT          | [26]      |
| KC                  | 41             | feedlot-concentrate  | Steer    | 28.7 ± 1.15           | 682 ± 71.6            | 397 ± 48.9            | 13.3 ± 4.40            | LD          | [10]      |
| KC                  | 10             | NP                   | Cow      | 22.8–44.4             | NP                    | NP                   | 14.3 ± 0.43            | LL          | [27]      |
| KC                  | 20             | feedlot-concentrate  | Steer    | 31.7                  | 752                   | 459                   | 15.3                   | LT          | [28]      |
| KC                  | 77             | feedlot-concentrate  | Steer    | 27.0–30.0             | NP                    | NP                   | 13.5                   | LD          | [29]      |
| Angus               | 10             | FS or SS             | Steer    | 27.5                  | NP                    | NP                   | 14.7                   | LL          | [30]      |
| Angus               | 450            | feedlot-concentrate  | Steer    | 27.5                  | 610                   | 330 ± 1.9             | 6.2 ± 0.21             | LT          | [31]      |
| Angus               | 9              | feedlot-concentrate  | Steer    | 27.5                  | NP                    | NP                   | 7.1                    | LD          | [32]      |
| Hereford            | 10             | FS or SS             | Steer    | 27.5                  | NP                    | NP                   | 8.3                    | LL          | [30]      |
| Hereford crossbred  | 228            | Pasture-grain        | Not specified | 30.0      | 678                   | 382                  | 6.9                    | LD          | [33]      |
| Brahman             | 7              | feedlot-concentrate  | Steer    | 27.5                  | NP                    | NP                   | 3.1                    | LD          | [32]      |
| Brahman             | 10             | feedlot-silage, forage | Bull and cow | 29.0      | 399                   | 234                  | 2.8                    | LT          | [34]      |
| Brahman             | 10             | feedlot-grass        | Bull and cow | 24.0–30.0 | 272                   | 148                  | 1.9                    | LD          | [35]      |
| Brahman             | 50             | Grazing              | Steer    | 31.0                  | 500                   | NP                   | 3.4                    | LD          | [36]      |

SE, standard errors; KC, Korean cattle; NP, not provided; LD, longissimus dorsi; LT, longissimus thoracis; LL, longissimus lumborum.
1 FS or SS = a forage-only system (FS) or a forage with high-energy supplemented system (SS).
Several management factors, including weaning age, castration, sex affects IMF content in the LM. Castrating bulls generally increases IMF content in several breeds, including Korean cattle and Holstein [9,38-40]. In Korean cattle, Park et al [39] compared marbling degree by sex (bulls, steers, and cows), and steers had the best MS, followed by cows.

Heritability
Heritability of the marbling trait has been reported in several cattle breeds. Utrera and Van Vleck [41] presented estimates of heritability for cattle carcass traits in 72 studies (1962 through 2004). They reported that the average breed type list included purebred or crossbred Angus, Brahman, Simmental, Charolais, Hereford, Jersey, Korean Brown cattle, Maine Anjou, Pinzgauer, Sahiwal, Salers, and South Devon with estimates MS heritability of 0.37 (range 0.30 to 0.57). MS heritability is relatively high in Korean cattle (0.57 in Yoon et al [42]; 0.64 in Park et al [43]). The heritability marbling phenotype in Japanese Wagyu is 0.40 to 0.55 [44-46], and MS heritabilities in Angus and Brahman are 0.48 [47] and 0.37 to 0.44 [48,49], respectively. The genetic potential of individual cattle can significantly affect IMF deposition. Greenwood et al [30] conducted a study to test effects of genotypes and nutrition during the immediate postweaning period on IMF contents in 5 muscles, marbling, and subcutaneous fat (rib fat or P8 rump fat) in Bos Taurus steers. The study was conducted with steer of three genotypes (Angus, Hereford and Wagyu×Angus): Angus was targeting high IMF and high subcutaneous fat, Hereford was targeting low IMF and high subcutaneous fat, and Wagyu sires Angus dams were targeting high IMF and lower subcutaneous fat. Following weaning, steers were grown with a forage-only system and forage with high-energy supplemented system to test effects of a high-energy grain-based substitution on fat deposition with 5 times serial slaughter. In their study, genotype differences were generally observed in IMF contents of 4 muscles (biceps femoris, supraspinatus, semitendinosus, longissimus lumborum, and infraspinatus), although semitendinosus muscle IMF did not differ among genotypes. The P8 rump fat depth was generally greater in Hereford than Angus, and Meat Standard Australia (MSA) MS was generally lower in Hereford than in Angus and Wagyu×Angus. However, substitution of equivalent amount of energy from grazed pasture with high-energy grain-based feed has not increased IMF contents. Selecting a sire is important for high-quality beef production; animals with high genetic potential will generally exhibit higher MS at slaughter age, whereas animals with low genetic potential will give low MS [50].

**MANAGEMENT FACTORS**

Several management factors, including weaning age, castration, slaughter age and weight, nutrition and other environmental conditions can affect IMF deposition.

**Weaning age**
Weaning age is known to affect IMF deposition and MS in beef. Several studies have reported that early weaning or early weaning combined with a high-grain diet increases IMF deposition in beef cattle. In two separate studies, early-weaned steers (weaned at 177 d [study 1] and 158 d [study 2] of age and placed on a finishing diet) increased MS and body weight gain, improved feed efficiency; and improved QG of the beef compared with normally weaned steers or normally weaned steers supplemented with grain [51]. In these studies, the normally weaned steers remained on pasture during nursing and weaning at 231 d (study 1) and 213 d of age (study 2), respectively; and they were then placed on a finishing diet. The steers of the normally weaned steers supplemented with grain were raised under the same conditions as the normally weaned animals, but they were supplemented with grain for 55 d on pasture (study 1, 177 to 231 d of age; study 2, 158 to 213 d of age). Effect of weaning age has been studied in Shorthorn calves [52]. In their study, early weaning calves (123 d of age; 145 kg of body weight) were compared with conventional weaning animals (259 d of age; 273 kg of body weight) in contemporary feeding management. Early weaned-animals showed lighter final body weight, and tended to eat less than conventionally weaned-animals. However, weaning treatment did not affect feed efficiency, post weaning growth, MS, and other carcass traits. Another study in Angus and Wagyu cross heifers raised under early weaning (142 days old) and accelerated finishing programs on a high concentrate diet showed increased IMF and greater weight gain compared with heifers weaned at 180 d of age and grown on pasture for 16 months before entering the feedlot [53]. In the Angus study, early weaning at 105 d of age and high concentrate feeding for 148 d followed by grazing and feedlot entry increased both the MS and hot carcass weight (HCW) compared with normal weaning at 253 d of age (normally weaned calves stayed with their dams to graze on pasture), although feedlot performance remained unchanged [54]. The authors suggested that metabolic imprinting by early weaning and a high concentrate feeding before grazing is a feasible method to improve MS and carcass weight.

A study was performed to understand the mechanism responsible for improved MS after early weaning and high-concentrate feeding in Angus and Angus × Simmental steers [55]. Early weaning (141 d of age) followed by a high-starch diet improved MS compared with normal weaning (222 d of age) followed by a starch creep-feed diet. A molecular study revealed that early weaning and a high-starch diet activated precocious pre-adipocyte differentiation and fat accumulation by activating adipogenic peroxisome proliferator-activated receptor-γ (PPAR-γ). The PPAR-γ signaling pathway regulates muscle growth, and weaning age increased muscle-specific PPAR-γ in early-weaned steers [56]. Therefore, the early-weaning protocol induced a metabolic switch from fat deposition to muscle growth, which could improve the IMF deposition in beef cattle.
receptor-gamma (PPARG) and CCAAT/enhancer-binding protein α pathways. In another study, Graugnard et al. [56] reported that a high-starch diet induced precocious upregulation of the adipogenic gene network in the \textit{longissimus lumborum} of early-weaned (average 155 d) Angus cattle. Smith and Johnson [57] suggested that the glucose contribution to fat synthesis decreases as cattle become fatter, whereas acetate use increases, especially in the IMF area. Those authors also suggested that providing dietary glucose sources (e.g., high-starch diet) at an early age can improve IMF deposition. Two-week-old Korean cattle calves were fed different diets, i.e., mother’s milk+roughage (control), milk replacement+concentrate, milk replacement+concentrate+roughage, and milk replacement+concentrate+30% starch [58]. All calves were then fed the same diet for 22 months, and muscle tissues were collected by biopsy. The RNA-Seq transcriptome and KEGG pathway analyses of differentially expressed genes showed that pathways for PPAR signaling, regulation of the actin cytoskeleton, biosynthesis of unsaturated fatty acids (FA), and Wnt signaling changed in the milk replacement+concentrate+roughage group compared with those in the control group, whereas the milk replacement+concentrate+30% starch diet group showed changes in insulin signaling and mTOR signaling pathways. The milk replacement+concentrate+roughage group and the milk replacement+concentrate group showed changes in the adipokine signaling pathway compared to control group. The authors suggested that feeding concentrate plus roughage or starch after early weaning at a young age could alter adipogenic and lipogenic potential in loin meat.

\section*{Castration}

Castration of bulls generally increases IMF content in several cattle breeds, although it reduces growth rate [40]. Castration of bulls profoundly increased MS, QG, and IMF contents in Korean cattle [9,39]. The castration rate of bulls has been increasing in Korea in the 21st century, and it was about 95% in 2013 [59]. Five QG levels are used in the Korean beef QG system [QG1++ (highest), QG1+, QG1, QG2, and QG3 (lowest)] [59]. A 2016 report of nationwide pooled data in Korean cattle revealed that the majority (76%) of bulls had a QG 3, whereas 85.5% of castrated cattle showed grades of QG 1 or higher (QG1, QG1+, and QG1++) [59; Figure 2]. Castration also increased MS in Canadian beef carcasses [60]. Japanese Black cattle bulls are normally castrated to get highly marbled beef in Japan [61]. Castration is also known to increase tenderness in several cattle breeds, including Korean cattle [62], Angus and Angus–Hereford cross breeds [63], and Brahman and Charolais cross breeds [64].

\section*{Slaughter age}

IMF content increases with increasing age in most cattle breeds. The IMF content in the LM of somatic-cell-cloned Japanese black steers increased with age at slaughter: 20 (23.7%), 25 (38.7%), and 30 months (41.1%) [65]. Purebred Japanese Black cattle show a linear increase in IMF content with increasing slaughter age from 16.7 months (500 d) to over 33.3 months (1,000 d) [66]. MS increased with increasing slaughter age from 16.2 months (485 d) to 20.3 months (610 d) in Holstein-Friesian and Norwegian dairy breeds [67]. In Angus, Hereford, and Wagyu×Angus, the IMF contents in the 5 muscles (\textit{biceps femoris}, \textit{supraspinatus}, \textit{semitendinosus}, \textit{longissimus lumborum}, and \textit{infraspinatus}) and MSA MS increased with the increasing slaughter stages (0, 168, 326, 431 and 585 d after weaning) [30]. Both MS and IMF contents increased with increasing

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure2.png}
\caption{Comparison of quality grade percentages among sexes of Korean cattle in 2016. Cow (n = 353,880), steers (n = 363,332), bulls (n = 20,264). Data source: Korea Institute for Animal Products Quality Evaluation [59].}
\end{figure}
slaughter age (2, 4, 6, 12, and 24 months) in bulls of German Angus, Galloway, and Holstein-Friesian breeds [24]. However, these changes with slaughter age were not observed in double-muscled Belgian Blue bulls [24].

The MS generally increases with slaughter age in Korean cattle. The MS standard ranges from 1 to 9 in Korea [59]. The MS of Korean cattle steers (81 animals) raised at the Livestock Improvement Main Center of the National Agricultural Cooperative Federation in Korea increased from 6.0 to 7.06 as slaughter age increased from 26 to 31 months [68]. The highest profit in this steer group was for a slaughter age of 31 months. In the same report [68], the MS in 5,826 Korean cattle steers from several farms in Gangwon Province was analyzed in 2010 to 2012; MS increased from 4.9 to 5.7 as slaughter age increased from 26 to 35 months. In this case, the optimum slaughter age for the highest profit was 28 months. This study suggested that changes in MS with slaughter age vary among farms. The difference may be due to genetic factors, management, or feed. Decreasing slaughter age may be better for reducing production costs when feed prices are high.

Pethick et al [69] suggested that a prolonged feeding period gives the cattle time to obtain high IMF levels because time is needed for muscle to reach maturity, followed by time for the muscle to fill up with IMF. However, feed efficiency decreases as animals age due to a decrease in the rate of muscle growth [70]. Thus, animals with low genetic potential for QG are not recommended for a long fattening stage, which would result in low feed efficiency. In addition, a long fattening period could result in low yield grade (YG) because long-term fattening increases backfat thickness and carcass weight, which negatively affect YG in Korean cattle [59]. Slaughter age needs to be optimised to get maximum profits for farmers by monitoring the QG and quantity grade of carcasses, feed costs, and carcass prices (auction prices).

Slaughter weight
IMF content or MS generally increases with slaughter weight in several cattle breeds. The National Beef Quality Audit of the USA (beef of steers and heifers in 28 packing plants during October–December 1991) found that MS increased with increasing slaughter weight from 173 to over 543 kg [71]. A linear increase in IMF content was observed in British and Japanese Black×Holstein-type cattle during a long fattening period in a feedlot, as carcass weight increased from 200 to 400 kg (reviewed in Pethick et al [69]). The MS and IMF content in Angus steers increases in a linear fashion with increasing HCW from 208 to 380 kg [72]. In the same study, LM area also increased linearly with increasing HCW. The IMF of Angus steers grown in South Dakota (USA) increased linearly with increasing HCW from 189 to 368 kg (live weight from 340 to 591 kg) [73]. In pooled data of different sexes (cows, bulls, and steers) of Korean cattle, MS and LM area increased with increasing carcass weight from 143 to 536 kg [39]. IMF contents in the 5 muscles (biceps femoris, supraspinatus, semitendinosus, longissimus lumborum, and infraspinatus) also increased along with the increase of HCW at 5 slaughter stages in Angus (from 114.8 to 459.1 kg), Hereford (from 114.0 to 452.2 kg), and Wagyu×Angus (from 116.8 to 459.2 kg) [30].

Feed efficiency decreases as animal age and slaughter weight exceed a certain level. Several factors, including MS or QG, feed costs, and carcass prices need to be considered to determine the optimum slaughter weight for the maximum profit.

Environments
Studies have demonstrated that environmental conditions, such as temperature stress, can negatively affect animal growth and beef quality in cattle [74]. In particular, exposing feedlot animals to hot or cold stress reduces growth and feed efficiency [75]. Cold stress during winter decreases animal performance and production efficiency [76,77]. Our recent study indicated that a lower ambient temperature decreased growth and feed efficiency in Korean cattle steers [78]. Heat stress can also negatively affect animal performance, including growth, milk production, reproduction, feed intake, overall health, and well-being in cattle [79,80].

British and British×Continental crossbred cattle grown with no shelter or windbreak had greater fat thickness and MS in winter than did animals fed in a protected area [74]. The authors suggested that mild heat stress in summer and mild cold stress in winter may be beneficial to fat deposition under ad libitum feeding conditions for cattle fed in an unprotected, unsheltered area. They also suggested that animals deposit fat in the winter to provide insulation for the body under cold stress. Animals may require more energy to meet the increased maintenance energy requirements. Young [81] suggested that energy requirements increase during winter when animals are under cold stress, as increased heat production is needed to maintain body temperature by shivering or other thermo-genic process. Angus crossbred cattle exposed to greater heat stress tend to have a greater MS [82]. The authors mentioned that slight heat stress might induce IMF deposition. Seasonal variations in carcass characteristics have been observed in Korean cattle steers by our group [2]. The YG was lowest in winter and highest in spring and summer. In contrast, backfat thickness, which is a negative YG factor, was highest in winter. Therefore, the poor YG in winter may be due, in part, to high backfat thickness. However, both MS and QG were best in autumn and worst in spring. In same study, neither MS nor QG was associated with air temperature. Our recent study also revealed that growth performance was not affected by mild heat stress in Korean cattle [83]. Overall, severe temperature stress may have significant negative effects on animal production, including growth performance and product quality, although mild temperature stress may differentially affect...
production.

**NUTRITIONAL FACTORS**

Several nutritional factors including fat metabolism, fat digestion and absorption, glucose/starch availability, roughage vs. concentrate ratio, dietary energy and protein levels, vitamins A, D, and C levels, fetal nutritional programming, and stage-specific feeding systems can affect IMF deposition.

**Fat synthesis and degradation metabolism**

IMF deposition results from a balance among uptake, synthesis, and degradation of triglycerides (TGs). TG synthesis is a key factor for IMF deposition [69]. Both non-esterified fatty acids (NEFAs) and a glycerol backbone are required for TG synthesis. The FA de novo synthesis occurs mainly in adipose tissue in ruminants [84], whereas it occurs in the liver in humans and rodents. FA for TG synthesis is derived from de novo synthesis or diet-originating FAs.

The de novo FA synthesis pathway of ruminants is summarized in Figure 3. Acetate or glucose is used as a substrate for FA synthesis [85]. The predominant substrate for FA synthesis in ruminants is acetate, which is mainly derived from ruminal fermentation. Glucose can also be used as a substrate for FA de novo synthesis in ruminants. Glucose is derived from either glucose synthesis via gluconeogenesis from propionate/lactate or glucose absorbed from the small intestinal digestion of the rumen bypass [86]. ATP citrate and NADP malate dehydrogenase are two key enzymes utilizing glucose in the FA de novo synthesis pathway. The activity of ATP citrate lyase is known to be low in ruminants [87]; thus, glucose is used as a lipogenic substrate less in ruminants than in monogastric animals [87]. However, glucose is believed to be preferred to acetate as a substrate for FA synthesis for IMF deposition [88]. These two enzymes are induced in adipose tissues of ruminants by increasing energy intake [89]. In addition, IMF has higher ATP-citrate lyase activity compared to subcutaneous fat [90]. Furthermore, IMF has a higher expression of glucose transporter 4, which is responsible for uptake of glucose in adipose tissues, compared to subcutaneous fat (reviewed in [90]). Using an in vitro system of adipose tissue from Angus and Wagyu steers, Rhoades et al [91] reported that the rate of glucose incorporation into FA was over two times higher than the rate of acetate incorporation into IMF, although they were similar in subcutaneous fat. The same authors suggested that feeding a corn-based diet increased glucose uptake in IMF, whereas feeding a hay-based diet may promote subcutaneous fat deposition via incorporation of acetate into FAs.

![Figure 3](https://www.ajas.info/1049)

**Figure 3.** Fatty acid (FA) synthesis pathway in ruminants. Microbes ferment polysaccharide into volatile fatty acids such as acetate, propionate, and butyrate in the rumen. The acetate and propionate can contribute for FA synthesis. Acetate is converted to acetyl-CoA in the cytoplasm of ruminant cells and is used to synthesize FA. Propionate is transported into mitochondria where it enters the tricarboxylic acid cycle via succinyl-CoA and can be utilized as a substrate for the production of glucose via gluconeogenesis. Glucose, which is synthesized from propionate or rumen-bypassed and absorbed in the small intestine, can generate citrate in the tricarboxylic acid cycle, where it can be transported into the cytosol. The citrate in the cytosol is degraded into oxaloacetate and acetyl-CoA by ATP-citrate lyase, and the acetyl-CoA can be used to synthesize FAs. Modified from Bauman et al [162].
Fatty acids of dietary origin are obtained from transportation of TG lipoprotein and subsequent uptake after lipoprotein lipase hydrolysis into tissues from the circulation [92,93]. Fat addition can affect IMF deposition. Feeding high-oil corn (1.3% more total lipid than typical corn) enhanced IMF deposition in predominantly Angus finishing crossbred steers [94].

In addition to FAs, a glycerol “backbone”, mainly supplied by glucose, is needed for TG synthesis. Therefore, increased glucose availability is important for providing glycerol for IMF deposition in ruminants [69], similar to monogastric animals such as pigs. Increasing propionate production for glucose synthesis is also important for IMF deposition, especially during the fattening period. For this reason, a large amount of concentrate is fed during the fattening period in Japanese Wagyu and Korean beef cattle [95,96].

High levels of IMF deposition requires excess energy consumption above the maintenance and normal production requirements [69]. Proliferative potential is lower in intramuscular preadipocytes than in subcutaneous preadipocytes [97]. IMF also has lower lipogenic activity than does subcutaneous fat [98]. Our research also showed that IMF had the lowest transcriptional activity for many adipose development and lipid metabolism genes, including adipogenesis, FA transport, lipogenesis, lipolysis, and FA oxidation among several fat depots (abdominal, perirenal, subcutaneous, and intramuscular) of Korean cattle [7]. These results suggest that higher energy levels are required for IMF deposition than for general body fat deposition, including for subcutaneous fat deposition.

TG hydrolysis by lipolysis generates NEFAs and glycerol, and it decreases IMF deposition, as lipolysis is a fat mobilization pathway. Stress increases circulating concentrations of cortisol, a typical stress marker [99]. Cortisol is also an important regulator of energy metabolism, including the regulation of lipolysis and lipogenesis, and increases plasma NEFA concentrations [100]. Cold stress stimulates lipolysis by increasing the secretion of stress hormones including epinephrine and norepinephrine in humans [101]. Cold stress causes white fat depot lipolysis in rats [102]. In addition, our recent research has shown that transport stress causes temporal increases in circulating cortisol and NEFA concentrations, along with a decrease in circulating TG concentration in pregnant Holstein heifers [103], consistent with transport stress also causing lipolysis. Thus, stressors, such as temperature and transportation, may have a negative effect on IMF deposition through lipolysis in beef cattle. Therefore, fat degradation by lipolysis should be minimized to prevent loss of IMF, and management and transport systems that minimize stress are needed. Fat supplementation for extra energy supply has been recommended during cold or heat stress to prevent fat degradation. Feeding medium-chain TGs, such as coconut oil increased the lipid content in the liver and carcasses of Holstein calves under cold stress [104]. However, some studies have revealed that mild temperature stress can increase body fat deposition and marbling for insulation and other purposes [74,82], as described above. Further study is warranted to clarify how stress intensity affects animal performance and fat deposition.

**Fat digestion and absorption**

Many of the FAs generated by digestion in the rumen pass into the small intestine, where they are absorbed into circulation via the lymph and blood. Most (~90%) dietary lipids reach the duodenum as NEFAs, unless they are protected [105]. Protected dietary fats are bypassed in the rumen without digestion, and they are digested and absorbed in the small intestine. Manipulating dietary fat digestion and absorption in the small intestine may enhance MS. Emulsifying fat may affect the digestion and absorption of fat. Emulsifier supplementation increased digestion and absorption of a high energy/fat diet in ruminants [106,107]. The emulsifier PROSOL (sodium stearoyl-2-lactylate) increases IMF deposition and QG in Korean cattle [108]. Bile salts are amphiphatic compounds that act as biological detergents, converting dietary fats into mixed micelles of bile salts and TGs [109]. Micelle formation profoundly increases the fraction of lipid molecules accessible to the action of intestinal lipase, aiding fat digestion in the intestine. Bile acid supplementation may enhance fat digestion. Ursodeoxycholic acid, which is a bile acid, is synthesized chemically from cholic acid. Ursodeoxycholic acid supplementation increased IMF and QG in Wagyu heifers [26], demonstrating that bile acid supplementation improves IMF deposition, probably by increasing intestinal fat digestion and absorption by increasing fat emulsification.

**Glucose/starch availability and roughage vs concentrate ratios**

Consuming surplus net energy is an important factor for deposition of IMF. Feeding cereal grains is one way to raise net energy supply because energy dense grains can be utilized in the rumen and the small intestine to produce volatile fatty acids and glucose for energy production. Maximum starch utilization can be achieved by optimizing fermentation of starch in the rumen to produce propionate (gluconeogenic precursor) and maximizing starch digestion and absorption in the small intestine. Increasing digestion of rumen-bypassed starch in the small intestine helps increase the glucose supply. A considerable amount of glucose is absorbed from the small intestine when high-starch diets are fed, especially during the beef cattle fattening period. Ørskov [110] reported that up to 42% of dietary starch may escape ruminal fermentation and reach the small intestine. Harmon and McLeod [111] reported that starch digestion is more efficient in the small intestine than during fermentation in the rumen: the total energetic efficiency of rumen-fermented starch was about 73% that of small in-
A meta-analysis of starch digestion in dairy cows revealed that the ruminal digestibility of corn was 57.4% [112], demonstrating that a substantial portion of corn can be digested in the small intestine. The same report estimated that up to 2 kg of starch/d can be digested in the small intestine of dairy cows; the average starch digestibility in the small intestine was 60.6%. Therefore, a better method to achieve high digestibility of starch in the small intestine is required to provide high post-absorptive glucose to the host animal. Pethick et al [69] reported that a high level of diet processing can increase the accessibility of dietary starch granules for both rumen and small intestine amylases, resulting in maximum glucose availability to the fattening animal [113]. The decreased flake thickness of steam-flaked sorghum increased starch digestion and liver gluconeogenesis [114]. Grain type and processing affect IMF content [115]: maize is more effective at increasing IMF content than is barley, followed by sorghum. Steam flaking further increases marbling in cases of maize and sorghum, and is more effective for sorghum. Bindon [115] suggested that maize produces increased glucose from digestion in the small intestine and that steam flaking increases digestion in both the rumen and small intestine. Pethick et al [69] suggested that feed that has a high glycemic index to allow rapid glucose absorption and concomitant high insulin levels through either grain feeding instead of grass finishing or grain processing provides increased net energy levels for FA synthesis.

Several studies have reported higher IMF content in high concentrate diets compared with low concentrate diets (reviewed in Wood et al [116]). Yearling Brangus×Hereford×Angus steers fed a high grain diet (79% corn) had more growth performance, greater MS, and higher grade meat than did steers fed a forage diet (winter wheat pasture with sorghum-Sudan and Bermuda grass pasture) [117]. Increasing the percentage of concentrate (60%, 75%, and 90% concentrate) in growing diets increased MS in steers [118]. Pooled results from Angus, Shorthorn, Murray Grey, and Hereford showed that the grain-finishing group had a greater IMF% than the grass-finishing group [115]. In bulls of two Portuguese breeds (Alentejana and Barrosã), the effects of high (70% concentrate 30% maize silage) and low isoenergetic and isonitrogenous (Alentejana and Barrosã), the effects of high (70% concentrate 30% maize silage) and low isoenergetic and isonitrogenous concentrates (30% concentrate 70% maize silage) on IMF contents were examined in the longissimus lumborum and the semitendinosus muscles [119]. The IMF contents in longissimus lumborum and semitendinosus muscle were higher in high-concentrate-fed than in low-concentrate-fed Barrosã bulls. However, this difference was not observed in Alentejana bulls, indicating a breed difference in IMF content in response to the roughage-to-concentrate ratio. American Wagyu steers fed a high-corn diet had greater fat accumulation than did animals fed a hay-based diet [120]. Similarly, dietary forage level affected MS. Jersey steers fed a low forage diet (6% Sudan grass, 6% alfalfa hay and 63.15% steam-flaked corn, dry matter [DM] basis) in a feedlot finishing diet had greater MS in the strip loin than did animals fed a high forage level diet (12% Sudan grass, 12% alfalfa hay and 51.20% steam-flaked corn) [121]. The effects of roughage/concentrate ratio on body fat deposition and the expression of angiogenic and adipogenic factors in adipose tissue have been examined in fattening Wagyu steers [122]. Steers were fed the same total digestible nutrient (TDN) but different dietary roughage/concentrate ratios (low-concentrate group: 35% roughage and 65% concentrate; high-concentrate group: 10% roughage and 90% concentrate). The IMF content in the LM of the high-concentrate group (31.7%±5.4%) was greater than that in the low-concentrate group (22.9%±5.3%). However, fat weights in the subcutaneous, mesenteric, and intermuscular regions were not different between the groups. The expression of adipogenic factors (CCAAT/enhancer binding protein beta, CCAAT/enhancer binding protein alpha, and PPARα) in the subcutaneous fat and IMF was higher in the high-concentrate group than that in the low-concentrate group, indicating that the dietary roughage/concentrate ratio affects adipogenic gene expression in fat tissues. The same authors suggested that total IMF within the LM in the high-concentrate group might be greater than that in the low-concentrate group by stimulating intramuscular preadipocyte differentiation. Concentrate diets are generally a major glucose source through either production of rumen propionate or glucose absorption from the small intestine from a rumen-bypassed concentrate, as described above, whereas roughage is a major acetate source that results from the rumen fermentation process. Higher IMF deposition with higher concentrate feeding may occur if IMF preferentially uses glucose as a substrate for FA synthesis, while subcutaneous fat uses acetate, as described by Smith and Crouse [88].

Dietary energy and protein levels
Studies in pigs indicate that a deficiency of essential amino acids such as lysine increases IMF content [123,124], and it has been suggested that an amino acid imbalance relative to energy levels may increase IMF content in cattle (reviewed in Pethick et al [69]). Limited information is available on the effects of the dietary energy-to-protein ratio on IMF content in cattle. Effects of dietary energy (TDN: 70%, 80% DM basis) and protein levels (crude protein [CP]: 11.9%, 14.3% DM basis) on growth performance, carcass characteristics and meat quality were examined in F1 Angus×Chinese Xiangxi yellow cattle [125]. Steers in the high-energy group had greater LM IMF content than those in the low-energy group, whereas dietary protein levels did not affect IMF content. Podolian bulls fed grazing pasture and fed a high protein (15% CP, DM basis) supplement had higher IMF content in the LM than did animals grazing pasture and fed a low protein (12% CP) supplement [126]. Japanese Black steers fed a 16% CP diet had...
greater backfat thickness than those fed a 12% CP diet [127]. However, dietary protein levels do not affect IMF deposition in Angus or Shorthorn steers (reviewed in Bindon [115]). Therefore, current studies are inconsistent in their results and conclusions on the effects of dietary protein levels affect IMF deposition.

**Vitamins**

Several vitamins, including vitamins A, D, and C, are implicated in regulating IMF deposition.

**Vitamin A**

Restricting vitamin A intake increases IMF deposition. Vitamin A (also known as retinol) inhibits adipocyte differentiation [128]. The activity of glycerol-3-phosphate dehydrogenase, a biomarker for adipogenesis, decreases in bovine preadipocytes cultured with retinol [129]. A negative correlation was observed between serum retinol concentrations and carcass MS values in Japanese Black cattle [130,131]. Several studies have shown that restricting vitamin A in the diet, or low beta-carotene (a precursor of vitamin A) intake, increases body fat, including IMF, deposition. A study of Angus-crossbred steers showed that animals not supplemented with vitamin A had higher MS and QG compared to vitamin A-supplemented animals [132]. The same authors reported that exclusion of a vitamin A supplement induced hyperplasia in IMF, but not in the subcutaneous fat depot. The lack of vitamin A supplementation also increased marbling compared with vitamin A supplementation in a study of Angus crossbred steers [133].

The appropriate age for, and duration of, vitamin A depletion or supplementation have been suggested as important factors that affect MS. Oka et al [134] reported that restricting vitamin A in 15-month-old cattle until the final finishing phase (29 months of age) increased MS, but not in 23-month-old cattle prior to finishing. They suggested that serum vitamin A level affects the LM MS during the early fattening stage and that restricting vitamin A during the later stages of growth affects growth rate. Oka et al [134] recommended selective restriction of dietary vitamin A from 14 to 22 or 25 months of age to improve MS and QG without adverse health effects (blindness, arthritis, and muscular edema), given the effect of vitamin A restriction from 23 months of age in improving MS is minimal. Continuous dietary vitamin A restriction during the late fattening phase is not recommended because prolonged vitamin A restriction may lead to health problems and retard growth rate [130]. Retinoic acid positively regulates growth hormone gene expression [135] and decreases fat deposition and IMF in steers [136]. A continuous supply of normal levels of dietary vitamin A during the entire growth period may also result in poor marbling and QG, even if the growth rate were normal and no health problems occurred. Kawachi [137] described that the number of adipocytes increases in muscle between 13 and 19 months of age, so feeding a vitamin A-deficient diet during this stage may stimulate marbling.

The appropriate concentration range for restriction of vitamin A has been suggested as an important factor for enhancing MS and preventing health-related problems, including the incidence of blindness and muscular edema. Kawachi [137] described that the plasma retinol level in Japanese fattening cattle is about 300 μg/L (100 IU/dL) and that a vitamin A-deficient diet decreases the plasma concentration to <90 μg/L (30 IU/dL). Oka [134] recommended that vitamin A-deficient levels not fall below 30 IU/dL to enhance MS and maintain health.

Alcohol dehydrogenase 1 (ADH) is the primary enzyme that oxidizes retinol to retinaldehyde [138], and retinaldehyde inhibits adipocyte differentiation [139]. Ward et al [140] examined nutrigenetic interactions between the alcohol dehydrogenase 1C (ADH1C) single nucleotide polymorphism (ADH1C c.-64T>C) and vitamin A levels and the impact on IMF deposition in Angus-cross steers. A significant interaction was detected between ADH1C c.-64T>C genotype and vitamin A supplementation for IMF content in the LM. The cattle with a TT genotype (approximately 6.15%) that were not supplemented with vitamin A had 22.9% greater IMF compared with those with the CC genotype (approximately 4.95%). Vitamin A-unsupplemented TT steers (approximately 6.15%) had 24.4% greater IMF content compared with supplemented TT steers (approximately 4.9%). They concluded that the TT genotype of ADH1C c.-64T>C is related to greater IMF than the CC genotype when vitamin A is restricted, and ADH1C c.-64T>C could be a genetic marker for enhancing marbling in feedlot steers.

**Vitamin D**

Vitamin D also inhibits adipogenesis in vitro. 1,25-Dihydroxyvitamin D$_3$ (active form of vitamin D) inhibits preadipocyte differentiation [141] and suppresses adipogenic PPARG gene expression in 3T3 L1 cells [142]. One study reported that 1,25-dihydroxyvitamin D$_3$ inhibited lipid accumulation by inhibiting key adipogenic factors such as PPARG [143]. Wang et al [144] suggested that the vitamin D receptor (VDR) competes with PPARG for binding to the retinoid X receptor (RXR). Higher VDR/RXR binding may decrease formation of the PPARG/RXR heterodimer, thus inhibiting adipogenesis (Figure 4). Few in vivo studies have been performed on the effect of vitamin D on beef marbling. While vitamin A restriction improves marbling, but vitamin D restriction did not significantly affect marbling in Angus crossbred steers [133]. Further study is needed to clarify the effects of dietary vitamin D on MS and QG in beef cattle.

Smith et al [145] suggested a possible connection between 1,25-dihydroxyvitamin D$_3$ and calcium homeostasis in the regulation of marbling. The increase in blood 1,25-dihydroxy-
Vitamin D, levels in response to decreased calcium intake may downregulate preadipocyte differentiation, thus decreasing beef marbling.

Vitamin C

Vitamin C can be synthesized from glucose in the liver of many mammalian species other than guinea pigs, humans, and other primates [146]. Thus, the dietary vitamin C requirement for beef cattle is not specified in the NRC [146]. Vitamin C or L-ascorbic acid has several roles including in oxidation–reduction reactions and collagen synthesis [147]. In addition, vitamin C may help to regulate adipogenesis. Vitamin C increased preadipocyte growth of 3T3 L1 cells [148] and also had a positive effect on differentiation of sheep preadipocytes in primary culture [149]. A rumen-protected form of vitamin C is preferred as a dietary supplement because vitamin C is most likely degraded in the rumen [146]. In vivo studies showed increased marbling in response to dietary vitamin C supplementation in beef cattle; dietary supplementation of rumen-protected L-ascorbic acid-2-phosphate during the final fattening period increased both MS and fat content in carcasses of Japanese Black cattle [150]. Supplementing rumen-protected vitamin C in a high sulfur diet increased the MS and the percentage of Choice grade beef in Angus cross steers [151]. Further study is necessary to clarify the effects of dietary vitamin C on MS and QG in other breeds of beef cattle.

Fetal nutritional programming

Fetal programming or prenatal programing is the concept that mammalian embryonic and fetal development are linked to environmental changes, thus affecting the entire life [152]. The development of adipose tissue coincides with myogenesis during a particular time in fetal development [18]. The gestation period is very important because it determines the fate of progenitor cells that will become muscle cells, fibroblasts, or adipocytes [16]. Nutritional manipulation of the fetus during gestation may affect animal performance and modify the quality and quantity of livestock products. Therefore, fetal nutritional programming via nutrition of dams during gestation may be one way to increase marbling in offspring later in life. Several studies have shown that fetal nutritional programming affects growth and/or beef quality, including marbling. For example, the effects of a 30% nutrient restriction in cows at early to mid-gestation on offspring (steers and heifers) growth and adiposity were tested in Angus×Gelbvieh [153]. Adipocyte diameter in subcutaneous, mesenteric, and omental fat was larger in offspring of dams nutritionally restricted than in animals from dams with normal dietary intake (100% of NRC recommendations), but marbling data were not reported. In their study, YG was reduced, and HCW tended to be reduced in offspring of dams nutritionally restricted. In addition, nutritional restriction during early and mid gestation reduced ovarian characteristics (wet ovary weight, luteal tissue weight) of heifers, indicating negative effects of fetal programming on productive efficiency of replacement heifers. Negative maternal energy status at mid-gestation (80% of the energy requirements for maintenance of body weight) did not affect MS, intramuscular fat percentage, and HCW, but increased the

Figure 4. Transcriptional regulation of adipogenesis gene expression by retinoic acid (RA) and vitamin D. RA (an active form of vitamin A) binds to the retinoic acid receptor (RAR), and vitamin D$_3$ (Vit D$_3$) binds to the vitamin D receptor (VDR). The RA and vitamin D$_3$ may inhibit adipogenic gene expression by competing for retinoid X receptor (RXR) binding with peroxisome proliferator-activated receptor γ (PPARγ), which is a key transcription factor for adipogenesis. Modified from Pyatt and Berger [163].
ratio of MS to 12th rib fat thickness and the ratio of percent intramuscular fat to 12th rib fat thickness in offspring compared with positive energy status [154]. This study indicates little or no effect of reduced maternal energy status during midgestation on offspring IMF deposition and muscle mass. Improving the nutritional condition of cows during pregnancy affects growth, fat deposition, and beef quality in steers; steers from cows grazing with better pasture (Thinopyrum–Elymus–Psathrystachus, CP 11.1% to 6.0%) during mid- to late gestation had greater live body weight at slaughter, HCW, and tenderness, and had an increased tendency for IMF deposition compared with offsprings of cows grazing on native range (Bouteloua–Hesperostipa–Pascopyron, CP 6.5% to 5.4%) [155]. Effects of nutritional restriction from early gestation to parturition and/or throughout lactation on offspring growth and production characteristics to 30 mo of age and health and reproductive performance of cows were studied in large maternal grazing systems with offsprings of Hereford cows mated to Piedmontese or Wagyu sires [33]. Fetal programming of nutritional restriction with pasture-based systems retarded offspring growth performance (body weight, feed intake, carcass weight) and production parameters including muscle weight, meat yield, and fat and bone weight at any given age. However, the fetal programming did not significantly affect beef quality characteristics at 30 mo of age. In addition, authors emphasize importance of survival and health of cows and calves and of subsequent reproductive performance of cows for relative contribution of fetal programming to the profitability of beef production systems when developmental programming is applied [33].

Radunz et al [156] tested the effects of three maternal late-gestational dietary energy sources (grass hay, corn, and dried corn distillers grains with solubles) on progeny postnatal growth and beef MS in an Angus cross-breed. The dams were limited in their feed to maintain similar energy intake among the groups. In the study, steers from dams fed a corn-based diet during late gestation had a lower MS compared with progeny from dams fed diets low in starch (grass hay or dried corn distillers grains with solubles). The authors suggested that the prepartum maternal dietary energy source during late gestation can alter the development of fetal adipose tissue, affecting IMF deposition in progeny. However, studies during postnatal periods (after weaning or finishing period) show that feeding a high starch diet along with early weaning in steers generally increased marbling, as described above [51,53,157]. Thus, the dietary energy source may differentially influence marbling and IMF deposition, depending on the feeding period.

Overall, nutritional manipulation during the early, mid-, or late gestation period may affect adiposity or marbling of beef, but these effects are generally small and often masked by postnatal nutritional status, as mentioned by Bell and Greenwood [158]. Further study is warranted to understand what mechanisms affect the adipogenic potential by nutritional modulation during the fetal period at specific gestation stages, thereafter affecting marbling and IMF deposition at the progeny slaughter stage. Proper nutritional levels and proper selection of the types of energy sources for dams during each period of gestation stages may improve future production efficiency of progeny. Some of the results presented above suggest restricted nutrition at certain stages of pregnancy has more favourable outcomes. Bell and Greenwood [158] also suggest that effects of fetal programming during early-mid-gestation on postnatal growth or composition are generally minor. One of the mechanisms responsible for the effect of fetal nutritional programming on growth and adiposity of progeny is epigenetic modification (DNA methylation, histone modification, and microRNA) of gene expression during early development and permanent changes in offspring phenotypes in later life. Epigenetic modifications have been suggested to occur during development (fetal programming), growth, and fat deposition in cattle [13,18,159]. Our previous study showed the possible involvement of DNA methylation in regulating tissue-specific expression of PPARG gene in IMF and muscle of beef cattle [6].

Stage-specific feeding systems

A significant proportion of consumers in Korea, Japan, and the USA prefer highly marbled beef to minimally marbled beef. Thus, Korean and Japanese beef production systems have focused on production of highly marbled beef to satisfy their customers and to protect their beef industry against imported beef that has relatively low MS. As described above, several nutritional factors affect IMF deposition, and these factors must be considered to prepare a proper diet to improve marbling at the proper time in the life cycle. Japan and Korea have developed stage-specific feeding systems to produce highly marbled beef. The feeding system for highly marbled beef from Japanese Black cattle has been described previously in a review by Gotoh et al [95]. Briefly, Japanese Black cattle are generally fed a high concentrate diet from 11 to 30 months of age to induce deposition of IMF. The concentrate constitutes 36.8% to 86.4% of feed from 11 to 18 months of age. During the final stage, from 18 months to slaughter (28 to 30 months of age), the concentrate is 84.2% to 86.4% of feed, and the rest is roughage (rice straw). In addition, Japanese farmers generally restrict vitamin A levels in the diet during the mid-fattening stage to produce highly marbled Japanese Black cattle beef.

Similarly, a growth stage-specific feeding system has generally been applied to Korean cattle to produce highly marbled beef. The Korean Feeding Standard for Hanwoo [160] describes how to feed Korean cattle at each stage (growth, early fattening, and late fattening stages). The feeding strategy for highly marbled Korean cattle beef has been also described by Jo et al [96]. Briefly, high-quality forage is fed ad libitum to
develop the rumen during the growing stage (7 to 12 months of age), and the concentrate (CP 15% to 16%, TDN 69% to 70%) is restricted to 1.5% to 1.6% of body weight at this time. The fattening period can be divided into early and late periods, during which the concentrate is mainly fed with a small portion of roughage. Feed comprising about 80% concentrate (1.6% to 1.9% of body weight; CP 12% to 13%, TDN 72% to 73%) is provided during the early fattening stage (13 to 21 months of age). During the late fattening phase, from 22 months to slaughter age (28 to 31 months of age), about 90% concentrate (CP 11% to 12%, TDN 73% to 74%) is fed, and roughage (rice straw and rye grass) is about 10% of the diet. Korean farmers often feed a total mixed ration during the growing and early finishing stages instead of feeding concentrate and roughage separately. The mixed ration is made by using agricultural by-products (soybean curd cake and pomace of citrus, apple, and carrot) [96].

**CONCLUSION**

In this article, we reviewed several factors (genetic, management, and nutritional factors) affecting IMF deposition in cattle (Table 2). IMF deposition can positively affect beef taste in Japan and Korea have developed stage-specific feeding systems to produce highly marbled beef. In this article, we reviewed several factors (genetic, management, and nutritional factors) affecting IMF deposition in cattle. The current stage-specific feeding program for producing Korean cattle and Japanese Wagyu is relatively high. Fetal nutritional programming at early, mid-, or late gestation period can affect adiposity of offspring in later life. Applying a single factor (e.g., nutritional or genetic) may not be adequate to increase IMF deposition. Development of new fusion technologies (e.g., nutrigenomics) with genetic, epigenetic, and nutritional factors may be more efficient to optimize IMF deposition and improve feed efficiency. Nutritional programming during the fetal and neonatal periods has been suggested as a promising method to manipulate IMF deposition. Further studies are warranted to test the feasibility of early nutritional programming for regulating IMF deposition in cattle.

There has been increasing concerns by consumers about the negative impact of extremely highly marbled beef on human health, such as cardiovascular diseases and arteriosclerosis.

**Table 2. Summary of factors affecting intramuscular fat (IMF) deposition in beef cattle**

| Factors                     | Description summary                                                                 |
|-----------------------------|-------------------------------------------------------------------------------------|
| **Genetic factors**         |                                                                                      |
| Breed differences           | IMF content varies with cattle breed                                                 |
| Sex                         | Sex affects IMF deposition                                                          |
| Heritability                | Heritabilities of marbling in Korean cattle and Japanese Wagyu are relatively high  |
| **Management factors**      |                                                                                      |
| Weaning age                 | Early weaning age generally increases IMF deposition                                |
| Castration                  | Castration generally increases IMF deposition                                        |
| Slaughter age               | IMF content increases with increasing age in most cattle breeds                     |
| Slaughter weight            | IMF content increases with increasing slaughter weight in several cattle breeds     |
| Environments                | Environmental conditions, such as ambient temperature, can affect IMF deposition     |
| **Nutritional factors**     |                                                                                      |
| Fat metabolism              | Triglyceride synthesis is a key factor for IMF deposition, whereas the TG hydrolysis decreases IMF deposition |
| Fat digestion and absorption| Manipulation of dietary fat digestion and absorption in small intestine may enhance IMF deposition |
| Glucose/starch availability | Maximum starch utilization is important for IMF deposition and can be achieved by both optimum rumen fermentation of starch and by maximum starch digestion and absorption in the small intestine |
| Roughage vs concentrate ratio| IMF content increases with increasing percentage of concentrate in the diet         |
| Vitamin A                   | Vitamin A restriction generally increases IMF deposition                              |
| Vitamin D                   | Vitamin D generally inhibits adipogenesis in *in vitro* studies                    |
| Vitamin C                   | Vitamin C has a positive effect on differentiation of sheep preadipocytes           |
| Fetal nutritional programming| Fetal nutritional manipulation at early, mid-, or late gestation period can affect adiposity of offspring in later life |
| Stage-specific feeding systems| Japan and Korea have developed stage-specific feeding systems to produce highly marbled beef |
Additional studies are warranted to produce beef with appropriate levels of IMF deposition to satisfy beef quality and taste, support human health, and sustain the animal industry, depending on the situation of each country.

CONFLICT OF INTEREST

We certify that there is no conflict of interest with any financial organization regarding the material discussed in the manuscript.

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