INTRODUCTION

Managing nutrient supply to match demand is critical for sustainable and efficient livestock production. The California Net Energy System (CNES) was developed during the late 1950s and 1960s, as documented in the classic publication of Lofgreen and Garrett (1968). This system was a significant step forward and has served the beef cattle industry well during the past 50 yr. Energy requirements for maintenance (NEm) and gain (NEg), described by Lofgreen and Garrett (1968) as modified by expanding databases (Garrett 1980) and for specific situations [National Research Council (NRC), 1984, 1996, 2000; The National Academies of Sciences, Engineering, and Medicine (NASEM), 2016], are used worldwide as a basis for feeding beef cattle and often other ruminant species as well.

Concepts of developmental programming, or the idea that stressors during critical windows of development can have both short- and long-term consequences in offspring, began to emerge about three decades ago based on human epidemiological studies (Barker, 1992, 2004). Research with animal models, including livestock, has since demonstrated that developmental programming is probably universal and that consequences on offspring growth, development, and health are likely much larger than previously thought (Wu et al., 2006; Caton and Hess, 2010; Funston et al., 2012; Reynolds and Caton, 2012; Greenwood et al., 2017; Hoffman et al., 2017; Reynolds et al., 2017). Maternal stress, particularly nutritional stress, is one of the major drivers of negative consequences of developmental programming in offspring. Livestock can often experience a poor or compromised nutritional environment during gestation. For example, extensive livestock production systems such as those experienced by grazing animals in the Intermountain Region of the Western United States and similar environments throughout the world often result in compromised nutrient supply during gestation. In the United States, livestock can experience a poor nutritional environment during pregnancy as a result of 1) breeding of young dams, which increases competition for nutrients among the maternal and fetal systems; 2) presence of multiple fetuses; 3) selection for increased milk production, which results in competition for nutrients between the mammary gland and gravid uterus; and 4) naturally occurring environmental temperature stress (which complicates nutrient supply and fetal growth) or conditions that compromise feed quality or quantity that coincide with breeding and gestation of livestock
Compromised offspring may have altered metabolic and body composition outcomes (Du et al., 2017; Greenwood et al., 2017; Reed and Govoni, 2017; Reynolds and Vonnahme, 2017) at various points during their postnatal growth curves. Metabolic and body composition changes could influence NEm or NEg requirements. The goal of this review was to examine the potential impacts of maternal nutrition and developmental programming on offspring energy requirements.

MATERNAL NUTRITION AND DEVELOPMENTAL PROGRAMMING

Developmental Programming

Growth-restricted or developmentally impaired newborns have an increased risk of health complications throughout life, including metabolic, growth-related, and reproductive complications. Originally referred to as “the Barker hypothesis,” or “fetal programming,” the concept is that poor maternal nutrition (or other types of stress such as young or old maternal age, environmental heat stress, etc.) during critical windows of in utero or early postnatal development can have long-term effects on offspring health and well-being (Barker, 1992; Paneth and Susser, 1995; Armitage et al., 2004; Barker et al., 2004; Wu et al., 2006; Caton and Hess, 2010; Reynolds et al., 2010, 2017; Reynolds and Caton, 2012; Meyer and Caton, 2016). The concept of developmental programming was originally based on epidemiological studies in humans, but evidence of developmental programming of growth and well-being in livestock is found in older published literature and is often referred to as the maternal effect. For example, the crossbreeding experiments of Walton and Hammond with large (Shire) and small (Shetland) horses demonstrated that uterine environment impacts both birth weight and adult size (Hammond, 1927; Walton and Hammond, 1938).

In many species, including livestock, compromised fetal and/or neonatal growth can result in 1) increased neonatal morbidity and mortality; 2) altered postnatal growth; 3) changed body composition (e.g., increased fat, reduced muscle growth); 4) metabolic disorders (e.g., poor glucose tolerance and insulin resistance); 5) cardiovascular disease; and 6) dysfunction of organs and/or organ systems, (including adipose, brain, cardiovascular, endocrine, gastrointestinal, immune, kidney, liver, mammary gland, muscle, pancreas, placenta, and reproductive; Rhind et al., 2001; Sheldon and West, 2004; Wu et al., 2006; Anway et al., 2008; Caton and Hess, 2010; Du et al., 2010; Reynolds et al., 2010; Long et al., 2011, 2012; Shankar et al., 2011; Bartol and Bagnell, 2012; Connor et al., 2012; Meyer et al., 2012a; Reynolds and Caton, 2012; Spencer et al., 2012; Symonds et al., 2012; Jackson et al., 2013; Xiong and Zhang, 2013; Cardoso et al., 2014; Kilcoyne et al., 2014; Schmidt et al., 2014; Zambrano et al., 2014; Meyer and Caton, 2016; Reynolds and Vonnahme, 2016). Clearly, developmental programming can affect multiple organs and organ systems when assessed in various animal models, including livestock (Caton and Hess, 2010; Reynolds et al., 2010, 2017; Reynolds and Caton, 2012).

Impacts of Maternal Nutrition

Maternal nutrient supply is a major driver of developmental programming events and consequently, offspring outcomes (Wallace, 1948; Wallace et al., 1999; Wu et al., 2006; Caton et al., 2007; Caton and Hess, 2010; Funston et al., 2012; Reynolds and Caton, 2012; Robinson et al., 2013; Vonnahme et al., 2015, Meyer and Caton, 2016; Reynolds et al., 2017; Reynolds and Vonnahme, 2017). Fetal growth trajectory is affected by maternal nutrient intake even from very early stages of embryonic development, when nutrient requirements for conceptus growth are negligible in proportion to maternal needs (NRC, 1996, 2007; Robinson et al., 1999; NASEM, 2016).

Maternal nutrient restriction includes any event that decreases fetal nutrient supply during critical developmental windows (Caton and Hess, 2010; Reynolds and Caton, 2012). Restriction of fetal nutrient supply can result from many things, including compromised maternal nutrient supply, placental dysfunction, deranged maternal metabolism, physiological or environmental extremes, or combinations of the aforementioned or other scenarios. Effects of fetal nutrient restriction during gestation depend on timing, level, and/or duration of compromised nutrition (Reynolds and Caton, 2012; Reynolds et al., 2013; Vonnahme et al., 2015; Zhang et al., 2015). A majority of the data (Reed et al., 2007; Swanson et al., 2008; Vonnahme et al., 2015) demonstrate that maternal nutrient restriction during the last two-thirds of gestation can decrease fetal growth and offspring birth weights in sheep.

Nutrient restriction in beef cattle can decrease birth weights and result in slower postnatal growth (Robinson et al., 2013). Likewise, relative to controls
fed at requirements, feeding of low or high levels of metabolizable protein to mature beef cows resulted in decreased birth weights (Sletmoen-Olsen et al., 2000). Others (Martin et al., 2007; Larson et al., 2009) reported that protein supplementation of cows during the last third of pregnancy had little influence on calf birth weights. Conversely, Spitzer et al. (1995) and Stalker et al. (2007) reported that greater body condition during late gestation (a proxy for greater nutrient intake) can increase calf birth weights. Available data are taken to imply that birth weights in sheep, when compared with beef cattle, are more susceptible to maternal nutrient restriction, which could reflect differing placental growth patterns between sheep and cattle (Reynolds et al., 2005; Vonnahme and Lemley, 2012).

Neonates that are growth restricted in utero are at risk of postnatal complications, which may result in poor growth and development and concomitant negative consequences that include poor productivity and reduced longevity later in life (Wu et al., 2006; Caton and Hess, 2010; Funston et al., 2012; Reynolds and Caton, 2012; Reynolds and Vonnahme, 2017). Maternal undernutrition and restricted fetal growth are associated with decreased growth efficiency and altered body composition (Greenwood et al., 1998, 2000; Wu et al., 2006; Caton et al., 2007; Larson et al., 2009; Robinson et al., 2013). Birth weights in cattle are related to postnatal growth performance (Robinson et al., 2013; NASEM, 2016); however, nutrient restriction of dams can alter composition of offspring growth in the absence of birth weight differences (Reynolds and Caton, 2012). Altered postnatal metabolism or growth resulting from perturbed maternal nutrition can result in management challenges for livestock producers because nutritional management decisions are often based on the averages of groups of animals. Therefore, management approaches that mitigate negative effects of developmental programming have the potential to improve efficiency of ruminant livestock production, which will help address the grand challenge of doubling livestock production to feed the projected global population of 9.6 billion by the year 2050 (Reynolds et al., 2015; United Nations News Centre, 2015).

Much of the aforementioned discussion focused on altered total nutritional supply, which is most often achieved via differential intakes; however, a substantial body of literature focuses on supply of specific nutrients and developmental programming outcomes. Across species, the major classes of nutrients (carbohydrates, protein, lipids, vitamins, and minerals) have been investigated in models of developmental programming, and in each case, examples exist where maternal supply of a major nutrient class affected offspring outcomes. In other cases, realimentation or biological plasticity allows for compensation or protection from adverse outcomes.

Research with beef cattle in Nebraska (Funston et al., 2012) demonstrated that protein supplementation during gestation can have long-term positive effects on the offspring, including changes in weaning weight, carcass characteristics, and reproductive traits, when compared with non-supplemented controls. Funston et al. (2012) also showed that such long-term effects are not necessarily foreshadowed by differences in birth weight. When investigating prepartum maternal dietary energy source in beef cows, Radunz et al. (2012) reported that differing maternal dietary energy sources altered offspring adipose tissue development, glucose metabolism, insulin sensitivity, and long-term intramuscular fat deposition. Lan et al. (2013) reported that maternal dietary starch levels in pregnant sheep affected fetal DNA methylation and gene expression. Wang et al. (2015) demonstrated that imprinted gene expression and DNA methyltransferase in calves are influenced by maternal dietary starch. The work of Wang et al. (2015) indicated that epigenetic mechanisms can play a major role in the regulation of offspring responses to altered maternal nutrition, which is supported by recent reviews by Meyer et al. (2012a) and Reynolds et al. (2017). In a review of developmental programming in cattle, Robinson et al. (2013) concluded that fetal programming was pronounced and might explain considerable variation in growth and production traits including body weight, intake, carcass and muscle weights, and lean, fat, and bone weights.

Although global nutrient restriction and excess (driven by intake changes) and altered supply of major nutrient classes can clearly have effects on offspring outcomes, supply of specific nutrients in the maternal diet can also result in changes in offspring outcomes and have long-term consequences. Research from our laboratories investigating supranutritional selenium supply in maternal diets (Caton et al., 2007; Ward et al., 2008; Vonnahme et al., 2010; Camacho et al., 2012; Meyer et al., 2013, Yunusova et al., 2013; Caton et al., 2014a, 2014b) has demonstrated changes in lamb birth weight, growth, nutrient digestion, glucose metabolism and insulin sensitivity, visceral fat content, intestinal vascularity, and endocrine profiles in some but not all studies.

Supplementing ruminally protected arginine to pregnant ewes fed either adequate or nutrient-restricted diets increased the growth trajectory
of lambs (Peine et al., 2013, 2018). Lassala et al. (2010, 2011) reported that intravenous administration of L-arginine in nutrient-restricted ewes from day 60 until parturition, or in triplet- and quadruplet-bearing ewes from days 100 to 121 of gestation increased birth weights of lambs. McCoad et al. (2013) demonstrated that intravenous L-arginine in twin-bearing ewes from day 100 of pregnancy until term increased birth weight of females but not males and increased brown adipose stores in both males and females. Supplemental methionine in Holstein cows (Penagaricano et al., 2013) resulted in changes in the transcriptome of flushed embryos, including genes involved in embryonic development and immune responses.

The aforementioned discussion clearly demonstrates that altered maternal nutrition, from global nutrient supply (changing intake) to supply of specific nutrients, can have both short- and long-term consequences on offspring development. Some of these consequences, such as growth and reproductive rates, contribute to variation observed in livestock herds and likely present both recognized and unrecognized management challenges. Matching energy supply to demand in livestock is key to efficient production systems. This is particularly the case in beef cattle where maintenance energy expenditure for the cow herd is one of the priority demands, which he articulated as 1) basal metabolism; 2) food-gathering activities; 3) growth; 4) basic energy reserves; 5) maintenance of pregnancy; 6) lactation; 7) additional energy reserves; 8) estrous cycles and pregnancy establishment; and 9) accumulation of excess energy reserves. Competition for nutrients, particularly in first- and second-calf growing heifers, is a serious management concern in terms of conception and maintenance of pregnancy, cow longevity in the herd, and overall beef system production efficiencies.

Approximately 75% of fetal growth occurs during the last 2 to 3 mo of pregnancy in ruminants (Robinson et al., 1977; NASEM, 2016; Reynolds et al., 2018). The CNES partitions energy demand into NEm and NEg, with energy being used more efficiently for maintenance than for gain. Because efficiency of energy use for maintenance and pregnancy in beef cattle varies similarly with metabolizable energy (ME) concentration in the diet, and for convenience, NASEM (2016) expressed the net energy requirements for pregnancy in terms of NEm. Gravid uterine demand for energy is greater during the last third of pregnancy because energy retention is greater during this time as gravid uterine mass is increasing rapidly. Most energy systems report energy demands of pregnancy to be minimal during the first half of pregnancy because energy retention in the gravid uterus is minimal. Energy requirements associated with advancing pregnancy as reported by the NASEM (2016), NRC (1984, 1996, 2000), and Commonwealth Scientific and Industrial Research Organisation (CSIRO; 1990) are shown in Table 1.

The data in Table 1 demonstrate that energy requirements increase with advancing stage of pregnancy and are clearly increased during the last half of pregnancy. Under the current paradigm of expressing energy requirements, energy demand is low and practically insignificant during the first half of pregnancy; however, energy demand during the earliest stages of pregnancy must be important to embryonic survival and growth. In fact, a recent review (Bridges et al., 2013) indicated that pregnancy is not limited by fertilization rates but is greatly affected by early embryonic loss, which is largely, but not exclusively, affected by nutritional, environmental, and disease-related stress. Stress can drive nutrient partitioning toward immunological responses that likely create transitory, immunologically driven nutrient supply restriction for growth-related functions, which affect embryonic growth and survival.

**ENERGY DEMANDS DURING PREGNANCY**

A primary driver of whole-herd beef cattle production efficiencies is reproduction (Dickerson, 1970; Dziuk and Bellows, 1983; Koch and Algeo, 1983). The NASEM (2016) stated:

Undernutrition is a major factor affecting reproductive performance, resulting in delayed attainment of puberty, extended periods of postpartum anestrus, as well as compromised conception rates, embryonic survival, sexual behavior, and, as emerging documentation suggests, compromised developmental programming of offspring.

Beef cow nutrient requirements vary across the normal annual production cycle, with energy requirements being least immediately after weaning a suckling calf, increasing during pregnancy, and being greatest at peak lactation. Lactational demands peak early in beef cows and usually coincide with the annual breeding season, which further compounds nutrient competition between successful breeding and lactation. According to Short et al. (1990), cows partition nutrients to meet priority demands, which he articulated as 1) basal metabolism; 2) food-gathering activities; 3) growth; 4) basic energy reserves; 5) maintenance of pregnancy; 6) lactation; 7) additional energy reserves; 8) estrous cycles and pregnancy establishment; and 9) accumulation of excess energy reserves. Competition for nutrients, particularly in first- and second-calf growing heifers, is a serious management concern in terms of conception and maintenance of pregnancy, cow longevity in the herd, and overall beef system production efficiencies.
The Importance of Early Pregnancy

As stated previously, total energy requirements for pregnancy increase as gestation advances; however, nutrient supply to the developing conceptus is critical for survival and growth starting very early in pregnancy. Caton and Hess (2010) and Meyer and Caton (2016), building on earlier work by Fowden et al. (2006), discussed critical developmental windows and potential effects of maternal nutrient restriction on fetal and postnatal developmental outcomes. Most large-animal models of developmental programming focus on perturbations to the maternal system during mid-to-late gestation and resulting effects on the offspring. During early gestation (days 0 to 50), the conceptus grows from one cell to a fully formed embryo with fully recognizable organ systems and a functional placenta. Tissue-doubling time during this stage of gestation is exceptionally high. Consequently, nutrient supply needs to support rapid growth and development to ensure survival of the embryo and establishment of a viable pregnancy. During the early phase of fetal development, differentiation and vascularization of uteroplacental tissues as well as fetal organogenesis occur, all of which are critical events for normal fetal development (Funston et al., 2010). In addition, dams that undergo stress (nutritional, environmental, etc.) during early, but not late gestation, are likely to produce a normal birth weight offspring that still suffers from poor growth and metabolic problems because of the stress early in pregnancy (Ford et al., 2007; Vonnahme et al., 2007; Reynolds and Caton, 2012).

Recently, our laboratory developed an ovariohysterectomy technique (McLean et al., 2016) designed to investigate developmental programming responses to moderate nutrient restriction during the first 50 d of pregnancy. In these studies, postpubertal heifers were fed to gain 0.5 kg/d (control) or −0.08 kg/d (moderate restriction) for the first 50-d postbreeding. At various times during early pregnancy, ovariohysterectomies were conducted and tissues harvested. Results from these studies demonstrated nutrient and metabolite changes in fetal fluids (Figure 1). For example, at day 50 of gestation, glucose, methionine, and glutamine were decreased in allantoic fluids in moderately restricted heifers. Amniotic glucose was also decreased, whereas amniotic glutamine was increased in moderately restricted heifers. Maternal serum homocysteine also was increased in the moderately restricted heifers, suggesting compromised one-carbon metabolism.

Data generated from these studies on fetal muscle from the hind limb and fetal liver at day 50 of gestation revealed that in fetal liver and muscle, a total of 548 and 317 genes, respectively, were differentially expressed as a result of moderate nutrient restriction, of which 201 and 144 genes, respectively, were false-discovery-rate protected. Pathway analysis was performed on the differentially expressed genes to determine the functional categories of pathways or ontologies associated with factors known to affect production efficiencies. In fetal liver, five functional categories of interest were affected by moderate nutrient restriction during the first 50 d of gestation (Crouse et al., 2017b): metabolic pathways (n = 43 genes), protein kinases (n = 47 genes), nucleosome core proteins (n = 22 genes), mRNA splicing (n = 7 genes), and complement/coagulation cascades (n = 6 genes). In fetal muscle, three functional categories of interest were affected by moderate nutrient restriction: skeletal muscle (n = 74 genes), embryogenesis (n = 14 genes), and signaling cascades (n = 18 genes).
Further analyses of our data demonstrated that in fetal liver, nine histone genes were upregulated in moderately nutrient restricted (RES) compared with control (CON) heifers including members of the histone H1, H2A, H2B, and H4 families (Crouse et al., 2017a). The 13 differentially expressed histone-modifying transcripts included genes associated with acetylation and deacetylation, methylation, phosphorylation, and ubiquitination. Of particular note, HDAC10 was 2.67-fold greater (q < 0.05) in liver of RES fetuses. In addition, the histone deacetylase complex gene, CIRI, was 2.22-fold greater (q < 0.05) in liver of RES fetuses. Only one gene associated with histone modifications, SET, was lower (1.77-fold, \(P = 0.006; q = 0.16\)) in liver of RES compared with CON fetuses. The SET gene is involved in preventing H4 lysine acetylation. These data imply that maternal nutrient restriction very early in pregnancy initiates developmental programming through epigenetic remodeling of the fetal genome in beef cattle (Crouse et al., 2017a).

In fetal muscle from the hind limb (data from our laboratory), differentially expressed genes include the myogenic genes MYOG and MYOD1 (1.49- and 1.39-fold greater in RES than CON fetuses, respectively), both of which play important roles in skeletal muscle cell differentiation and fiber development. Four members of the Wnt signaling pathway, namely WNT5A, FZD1, APC2, and FZD10, were upregulated in RES fetuses (1.32- to 2.11-fold greater than CON). The Wnt pathway is critical in promoting the differentiation of myocytes from progenitor stem cells. Additional genes upregulated in fetal hind limb muscle of RES compared with CON fetuses included members of the troponin (TNNC1, TNNC2, TNNH1, TNNI2, TNNT1, TNNT2, TPM2), myosin (MYL1, MLY2, MLY4, MLY7, MYL6B, MLY9, MYH8, MLYPF), and actin (ACTA1, ACTA2, ACTG2) families. Therefore, we conclude that early gestation is an important period of myogenic developmental programming, and is sensitive to maternal nutrition in cattle (Ward et al., 2017).

In conclusion, although most investigations of developmental programming events in cattle focus on mid-to-late gestation, recent data from our laboratory indicate that moderate changes in maternal nutrition during the first 50 d of pregnancy can alter nutrient and metabolite concentrations in fetal fluids and gene expression in fetal liver and muscle. Whether these changes alter short- or long-term NEm or NEg requirements remains to be determined.

### OFFSPRING MAINTENANCE REQUIREMENTS

#### Maintenance Requirements

The CNES, as established by Lofgreen and Garrett (1968), used a comparative slaughter
technique to measure retained energy (RE) and regressed daily ME intake on daily heat production (HE) in Mcal/d, which was calculated by difference (ME – RE). Estimates of HE were expressed per unit of body weight (BW) raised to the ¼ power as a standard metabolic scaling approach (i.e., metabolic body size). Because of the techniques used in their studies, W more closely represented shrunk body weight (SBW) than live body weight. Lofgreen and Garrett (1968) then regressed calculated daily metabolism on daily heat production (adapted from Prezotto et al., 2014) indicate that low-birth-weight lambs had slower growth rates, differing body compositions at a given empty body weight (EBW), and lower RE. In addition, Greenwood et al. (1998) indicated the differences in observed fat and energy content of lambs at 17.5 kg of EBW were attributed to an approximately 30% decrease in maintenance energy requirements for low- compared with high-birth-weight lambs. Robinson et al. (2013) indicated that maintenance energy requirements are less in growth-retarded calves, particularly during the early postnatal phase, and that at any given age, growth-compromised offspring could have different nutrient requirements than their normally growing counterparts.

Whole-animal or specific tissue oxygen consumption reflects energy use. Tissue oxygen consumption reflects energy use and mitochondrial function. Hepatic and small intestinal tissues are major consumers of whole-body energy supply (Koong et al., 1985; Reynolds et al., 1991; Caton et al., 2000). Prezotto et al. (2014) investigated fetal hepatic and small intestinal oxygen consumption at 130 d of gestation from control- and restricted-fed ewes. First-parity ewes were fed a complete pelleted diet at either control (requirements for 140 g of daily growth) or restricted (60% of controls) levels of dietary intake from days 50 to 130 of gestation. At day 130, tissues were harvested and fetal hepatic and small intestinal oxygen consumption measurements were obtained. Data shown in Table 2 (adapted from Prezotto et al., 2014) indicate that both hepatic and small intestinal oxygen consumption in vitro were decreased in fetuses from restricted compared with control fed ewes at 130 d of gestation. Decreases in hepatic oxygen consumption in this study likely resulted from changes in liver mass and not because of changes in oxygen use per unit of tissue; however, small intestinal in vitro oxygen consumption was increased per unit of tissue, suggesting altered tissue energy use. In a follow-up study, with mature ewes fed control-intake, restricted-intake (60% of controls), or restricted-intake plus rumen-protected arginine, in vitro oxygen consumption of hepatic and small intestinal tissues were investigated in 54-d-old offspring (Prezotto et al., 2014).

Developmental Programming of Offspring Maintenance Requirements

Classically designed, definitive studies assessing the effects of maternal plane of nutrition on offspring NEm requirements are not available in the literature. Nonetheless, some of the early investigations into animal energetics and livestock performance provide evidence that compromised offspring at birth underperforms when compared with more normal offspring. For example, Armsby and Fries (1911) reported that “scrub” steers used energy less efficiently than “good” steers. More recently Greenwood et al. (1998) indicated that low-birth-weight lambs had slower growth rates, differing body compositions at a given empty body weight (EBW), and lower RE. In addition, Greenwood et al. (1998) indicated the differences in observed fat and energy content of lambs at 17.5 kg of EBW were attributed to an approximately 30% decrease in maintenance energy requirements for low- compared with high-birth-weight lambs. Robinson et al. (2013) indicated that maintenance energy requirements are less in growth-retarded calves, particularly during the early postnatal phase, and that at any given age, growth-compromised offspring could have different nutrient requirements than their normally growing counterparts.

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Data shown in Table 3 (adapted from Prezotto et al., 2018) showed that hepatic but not jejunal oxygen consumption was decreased in lambs from nutrient-restricted dams. In fact, hepatic oxygen consumption when expressed per gram of tissue, per whole tissue, or per unit of body weight was decreased in lambs from nutrient-restricted ewes. These data indicate that liver energy use was less in offspring from nutrient-restricted dams. Data from Prezotto et al. (2014, 2018) described earlier are supported by in vitro oxygen consumption data in fetal calves from cows fed control or restricted and then realimented diets (Prezotto et al., 2016). These authors suggested that both cows and fetal calves during gestation can modulate maintenance energy requirements in response to nutrient restriction and realimentation. Their conclusions seem reasonable given known modulations in maintenance energy requirements in nutrient-restricted and then realimented growing cattle, which results in compensatory growth (NASEM, 2016). Existing data indirectly suggest that maintenance requirements of offspring might indeed be programmed by maternal nutritional inadequacies, with a greater likelihood of programming occurring at nutritional extremes and early in the postnatal period. Additional research directed toward assessing the effects of maternal nutrition and developmental programming on offspring maintenance energy requirements will help to delineate these responses and their effects on offspring production outcomes.

**Table 2.** Hepatic and small intestinal oxygen consumption in fetal lambs at 130 d of gestation from ewes feed control (CON) or restricted (RES) diets (adapted from Prezotto et al., 2014)

| Item                                      | Treatments† | CON       | RES       | SEM       | P-value |
|-------------------------------------------|-------------|-----------|-----------|-----------|---------|
| Liver oxygen consumption                  |             |           |           |           |         |
| µmol/min per mg fresh tissue              |             | 60.3      | 60.5      | 2.24      | 0.76    |
| mol/min per liver                         |             | 5.79      | 4.67      | 0.38      | 0.01    |
| Small intestine oxygen consumption        |             |           |           |           |         |
| µmol/min per mg fresh tissue              |             | 67.1      | 53.3      | 4.2       | 0.009   |
| µmol/min per mg protein                   |             | 586       | 780       | 57        | 0.004   |

†Treatments were control (CON; fed at requirements for minimal gain and fetal growth) and restricted (RES) diets fed at 60% of CON.

**Table 3.** Hepatic and small intestinal oxygen consumption in 54-d-old lambs from mature ewes fed a control diet (CON) or a restricted (RES) diet with or without supplemental rumen-protected arginine (adapted from Prezotto et al., 2018)

| Item                                      | Treatments† | CON       | RES       | RES + ARG | SEM       | P-value¹ |
|-------------------------------------------|-------------|-----------|-----------|-----------|-----------|---------|
| Liver oxygen consumption                  |             |           |           |           |           |         |
| mol/min per g fresh tissue                |             | 0.39      | 0.35      | 0.34      | 0.02      | 0.09    |
| mol/min per liver                         |             | 181       | 146       | 155       | 10        | 0.04    |
| mol/min per kg BW                         |             | 8.0       | 6.7       | 6.8       | 0.4       | 0.02    |
| Small intestine oxygen consumption        |             |           |           |           |           |         |
| mol/min per g fresh tissue                |             | 0.35      | 0.38      | 0.35      | 0.02      | 0.60    |
| mol/min per jejunum                       |             | 58        | 71        | 60        | 12        | 0.62    |
| mol/min per kg BW                         |             | 2.4       | 3.2       | 2.5       | 0.5       | 0.50    |

†Treatments were control (CON; fed at requirements for fetal growth), restricted (RES) diets fed at 60% of CON, and restricted plus supplemental rumen-protected arginine (RES + ARG).

¹P-value is for the contrast of CON vs. RES plus RES + ARG.

translate basic science to industry innovation
RE vs. observed RE, in Mcal/d. When adjusting cattle to fit the Garrett (1980) database, the weight equivalent of the NRC (1984) medium-framed steer (EQSBW) is calculated as:

$$\text{EQSBW} = \text{SBW} \times \left( \frac{\text{SRW}}{\text{FSBW}} \right)$$

where

- EQSBW is the BW equivalent to the NRC (1984) medium-frame steer;
- SBW is the shrunk BW being evaluated;
- SRW is the standard reference weight for the expected final body fat; and
- FSBW is the final shrunk BW at the expected final body fat.

Within the NASEM system, there are various adjustments made to FSBW, which include decreasing FSBW by 25 to 45 kg for nonuse of an estrogenic implant; increasing FSBW by 25 to 45 kg for use of an implant containing trenbolone acetate plus estrogen; increasing FSBW by 6 to 36 kg for use of a β-adrenergic agonist; increasing FSBW by 25 to 45 kg for extended periods at slow rates of gain; and decreasing FSBW by 25 to 45 kg for continuous use of a high-energy diet from weaning. When problems arise in predicting NEg, they could be related to choosing the wrong FSBW, transitory effects of previous plane of nutrition, gut fill, anabolic implants, variations in NEm requirement, or the ME value assigned to the feed and the dietary NEm and NEg derived from the ME (NASEM, 2016).

Unfortunately, definitive studies that assess NEg requirements of cattle from dams with compromised maternal nutrition are not available in the literature. Considerable data exist, however, regarding the effects of maternal nutrition and developmental programming of muscle and fat accretion. Because NEg is estimated from RE, which depends on the composition of gain, it would seem reasonable that altered body composition at a given age could alter NEg requirements; however, this remains to be determined.

**Maternal Undernutrition and Muscle Development**

Decreased growth rate and feed efficiency pose a significant economic impact to the beef industry. Clearly, maternal nutritional status is one of the factors programming nutrient partitioning and ultimately growth and development of fetal skeletal muscle (Wallace, 1948; Wallace et al., 1999; Godfrey and Barker, 2000; Rehfeldt et al., 2004; Stickland et al., 2004). Growth-restricted neonates are not only at risk of immediate postnatal complications, but also might be “programmed” to exhibit poor growth and productivity, as well as diseases, later in life (Barker et al., 1993; Godfrey and Barker, 2000). This growth restriction seems to be especially important when fetal muscle development (myogenesis) is adversely affected (Handel and Stickland, 1987a, 1987b; Dwyer et al., 1993). Fetal skeletal muscle has a lower priority for nutrient partitioning compared with the brain and heart in response to challenges during fetal development, rendering fetal muscle particularly vulnerable to nutrient deficiency (Bauman et al., 1982; Close and Pettigrew, 1990). The fetal period is crucial for lifetime skeletal muscle development because no net increase in the number of muscle fibers occurs after birth (Gl ore and Layman, 1983; Greenwood et al., 2000; Nissen et al., 2003).

Several studies in a range of mammalian species have shown that maternal undernutrition during gestation can significantly decrease the number of muscle fibers and myocyte nuclei in the offspring (Bedi et al., 1982; Wilson et al., 1988; Ward and Stickland, 1991). For example, a lower ratio of secondary to primary myofibers and decreased sizes of the muscle fascicles were observed in muscle of fetuses gestated in nutrient-restricted ewes (Zhu et al., 2004). Therefore, muscle fiber type development can be influenced by maternal nutritional status depending on the energy needs of the muscle and species observed.

Nutrient restriction of heifers during the first two-thirds of gestation decreased fetal growth and calf birth weight (Micke et al., 2010). Nutrient restriction to 85% of ME compared with 140% in multiparous Angus-Simmental cows resulted in increased expression of myogenic genes MYOG and MYOD1 in offspring of restricted vs. control cows at day 247 of pregnancy (Paradis et al., 2017). Early prenatal nutritional restriction of ewes resulted in a decrease in the number of myofibers but an increased diameter of muscle fibers in offspring at 8 mo of age (Zhu et al., 2006). The finding of enlarged muscle fibers has been confirmed in other muscles in both bovine and ovine fetuses and at 8 mo of age in lambs when nutrient intake of dams was restricted during early gestation (Du et al., 2005, 2010). In addition, lambs born from ewes that were fed restricted diets during early- and mid-gestation had increased subcutaneous fat depots, reduced muscle size, and dysregulated glucose uptake compared with lambs from control ewes (Ford et al., 2007). Ewes that were nutrient-restricted to day 31 of gestation had lambs with decreased muscle fiber density in the triceps brachii compared with lambs of control-fed dams (McCoard et al., 1997).
In the bovine, primary muscle fibers develop during the first 2 mo after conception (Russell and Oteruelo, 1981). Secondary muscle fibers, which make up the majority of muscle fibers, form between 2 and 7 to 8 mo of gestation (Russell and Oteruelo, 1981). The formation of secondary myofibers partially overlaps with the formation of intramuscular adipocytes and fibroblasts (Du et al., 2010). The three cell types, myocytes, adipocytes, and fibroblasts, produce the basic structure of skeletal muscle and form at different time points in gestation. These data clearly show that maternal nutrient restriction during pregnancy can affect muscle development in offspring and that timing of restriction can have differential effects on muscle fiber development and growth as seen in postnatal phenotypic responses of offspring. Moreover, these data are consistent with our recent studies, described in The Importance of Early Pregnancy section, showing altered gene expression in hind limb muscle of fetuses from heifers that were nutrient-restricted during the first 50 d of gestation.

**Composition of Gain**

Robinson et al. (2013) concluded in their review that:

Fetal programming and related maternal effects are most pronounced and explain substantial amounts of variation for growth-related production characteristics such as BW, feed intake, carcass weight, muscle weights, meat yield, and fat and bone weights at any given age but are less evident when assessed at the same BW and carcass weight.

Body compositional changes resulting from compromised maternal nutrition seem to be more pronounced early in the postnatal and growing phases and less pronounced as offspring approach finished market weight, which is likely a result the high degree of plasticity of cattle body tissues and their ability to recover from early nutritional insults. Nonetheless, examples of differences in carcass composition are prevalent in the literature and are most often reflected by increased body fatness. Growth-restricted offspring may take more time to reach market weight (Funston et al., 2012), which could directly impact total energy required to finish cattle. Unfortunately, many published reports do not contain estimates of days on feed, particularly in response to birth weight or previous plane of maternal nutrition. While investigating effects of maternal nutrition on steer offspring, Underwood et al. (2010) reported maternal nutritional management could alter average daily gain, hot carcass weight, and 12th rib fat thickness at slaughter. Likewise, Radunz et al. (2012) reported that prepartum energy source of cows altered marbling score and intramuscular fat content in offspring at finish. Data from Nebraska (Stalker et al., 2006, 2007; Larson et al., 2009) indicate that steers born to non-protein-supplemented dams had lower dry matter intake and hot carcass weight, and decreased marbling score in some but not all studies.

Clearly, maternal nutritional plane can alter muscle development, even at the very early stages of growth. Tissue plasticity seems to compensate for some of these effects during steer growth and finishing; however, differences often persist until slaughter. Reduction of follicle numbers in the ovaries of offspring from restricted dams observed during gestation in beef heifers was still present at 86 wk of age (Mossa et al., 2013), indicating potential negative effects on fertility. Furthermore, heifers born to non-protein-supplemented dams had lower adjusted 205-d weaning weights, a lower percentage pregnancy after breeding (Martin et al., 2007), and decreased age at puberty (Funston et al., 2010). The degree to which NEg requirements of growing cattle are altered by maternal nutrient supply is not directly assessed in the literature. Additional research in this direction is needed to determine whether NEg requirements are changed in offspring from dams in nutritionally compromised environments.

**CONCLUSIONS**

Developmental programming is convincingly documented in the literature, can be driven by suboptimal maternal nutrition, and occurs in major livestock species, including beef cattle. Definitive studies that assess the effects of maternal nutrition and the resulting developmental programming events on NEm and NEg requirements in beef cattle have not been conducted. Indirect evidence included within this review suggests it is likely that energy requirements of offspring are affected by maternal nutrition in beef cattle, and that these events are at least partially controlled by epigenetic events during development that persist postnatally. Timing of the maternal nutritional insult(s) is important, and emerging data suggest that early pregnancy is likely much more important than previously thought. Additional research in the area of maternal nutrition and offspring energetics will provide insight that could lead to altered management practices and increased efficiencies of beef cattle production.

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LITERATURE CITED

Anway, M.D., S.S. Rekow, and M.K. Skinner. 2008. Transgenerational epigenetic programming of the embryonic testis transcriptome. Genomics 91:30–40. doi:10.1016/j.ygeno.2007.10.002

Armitage, J.A., I.Y. Khan, P.D. Taylor, P.W. Nathanielsz, and L. Poston. 2004. Developmental programming of the metabolic syndrome by maternal nutritional imbalance: how strong is the evidence from experimental models in mammals? J. Physiol. 561(Pt 2):355–377. doi:10.1113/jphysiol.2004.072009

Armsby, H.P., and J.A. Fries. 1911. The influence of type and of age upon the utilization of feed by cattle. Bulletin No. 128. Washington (DC): U.S. Department of Agriculture, Bureau of Animal Industry.

Barker, D.J.P. Editor. 1992. Fetal and infant origins of adult disease. London: BMJ Publishing Group.

Barker, D.J.P. 2004. Developmental origins of well being. In Banuelos, G., Z. Lin, and X. Yin, editors. Selenium in the environment and human health. London: CRC Press, Taylor & Francis Group. p. 159–161.

Barker, J.S., M.L. Bauer, and H. Hidari. 2000. Metabolic components of energy expenditure in growing beef cattle. Asian-Australas. J. Anim. Sci. 13:701–710. doi:10.5713/ajas.2000.702.

Caton, J.S., J.B. Taylor. 2014b. Moderating the effects of maternal nutrition on birth weight and postnatal nutritional metabolism. Proceedings of International Symposium on Energy and Protein Metabolism 9–13 September 2007. Vichy (France): EAAP Publication No. 124. p. 101–102. doi:10.3920/978-90-8686-613-7

Close, W.H., and J.E. Pettigrew. 1990. Mathematical models of sow reproduction. J. Reprod. Fertil. Suppl. 40:83–88.

Crouse, M.S., J.S. Caton, R.A. Cushman, K.J. McLean, C.R. Dahlen, P.P. Borowicz, L.P. Reynolds, and A.K. Ward. 2017a. Maternal nutrition during the first 50 days of gestation alters expression of histone and histone modifying genes in bovine fetal liver. 36th International Society of Animal Genetics Conference; July 16 to 21, 2017; Dublin, Ireland. p 32.

Crouse, M.S., J.S. Caton, R.A. Cushman, K.J. McLean, C.R. Dahlen, P.P. Borowicz, L.P. Reynolds, and A.K. Ward. 2017b. Moderate nutrient restriction influences transcript abundance of genes impacting production efficiencies of beef cattle in fetal liver, muscle, and cerebrum by d 50 of gestation. Proc. West. Sec. Amer. Soc. Anim. Sci. 68:42–47. doi:10.2527/asasws.2017.0014

Caton, J. S., M. L. Bauer, and H. Hidari. 2000. Metabolic components of energy expenditure in growing beef cattle. Asian-Australas. J. Anim. Sci. 13:701–710. doi:10.5713/ajas.2000.702.

Caton, J.S., and B.W. Hess. 2010. Maternal plane of nutrition: impacts on fetal outcomes and postnatal offspring responses. In: Hess, B.W., T. Delcurto, J.G.P. Bowman, and R.C. Waterman, editors. Proceedings of the 4th Grazing Livestock Nutrition Conference 9–10 July, 2010; Estes Park, CO. Champaign (IL): Western Section: American Society of Animal Science. p. 104–122.

Caton, J.S., A.M. Meyer, R.D. Yunusova, P.P. Borowicz, L.P. Reynolds, D.A. Redmer, C.J. Hammer, and K.A. Vonnahme. 2014a. Effects of maternal selenium supply and nutritional plane on offspring intestinal biology. Trace Elements in Man and Animals (TEMA-15) Meeting Abstr. 22–26 June 2014; Orlando, FL.

Caton, J.S., T.L. Neville, L.P. Reynolds, C.J. Hammer, K.A. Vonnahme, A.M. Meyer, and J.B. Taylor. 2014b. Biofortification of maternal diets with selenium: postnatal growth outcomes. In Banuelos, G., Z. Lin, and X. Yin, editors. Selenium in the environment and human health. London: CRC Press, Taylor & Francis Group. p. 159–161.

Caton, J., K. Vonnahme, J. Reed, T. Neville, C. Effertz, C. Hammer, J. Luther, D. Redmer, and L. Reynolds. 2007. Effects of maternal nutrition on birth weight and postnatal nutrient metabolism. Proceedings of International Symposium on Energy and Protein Metabolism 9–13 September 2007. Vichy (France): EAAP Publication No. 124. p. 101–102. doi:10.3920/978-90-8686-613-7

Dickerson, G. 1970. Efficiency of animal production mold-
Dwyer, C.M., J.M. Fletcher, and N.C. Stickland. 1993. Muscle cellularity and postnatal growth in the pig. J. Anim. Sci. 71:3339–3343. doi:10.2527/1993.71123339x

Dziuk, P.J., and R.A. Bellows. 1983. Management of reproduction in beef cattle, sheep and pigs. J. Anim. Sci. 57 (Suppl. 2):355–379. doi:10.2527/animalsci1983.57supplement_2355

Ferrell, C.L., W.N. Garrett, N. Himman, and G. Griecht. 1976. Energy utilization by pregnant and non-pregnant heifers. J. Anim. Sci. 42:937–950. doi:10.2527/jas1976.424937x

Ferrell, C.L., and L.P. Reynolds. 1987. Oxidative metabolism of gravid uterine tissues of the cow. In: Airle, V.A., P.W. Moe, H.F. Tyrell, and P.J. Reynolds, editors. Energy Metabolism of Farm Animals: Proceedings of the 10th Symposium; September 1985; EAAP Publication No. 32. New York (NY): Rowman & Littlefield. p. 298–301.

Ford, S.P., B.W. Hess, M.M. Schwope, M.J. Nijland, J.S. Gilbert, K.A. Vonnahme, W.J. Means, H. Han, and P.W. Nathanielsz. 2007. Maternal undernutrition during early to mid-gestation in the ewe results in altered growth, adiposity, and glucose tolerance in male offspring. J. Anim. Sci. 85:1285–1294. doi:10.2527/jas.2005-624

Fowden, A.L., D.A. Giussani, and A.J. Forhead. 2006. Intraterine programming of physiological systems: causes and consequences. Physiology (Bethesda). 21:29–37. doi:10.1152/physiol00050.2005

Funston, R.N., J.L. Martin, D.C. Adams, and D.M. Larson. 2010. Winter grazing system and supplementation of beef cows during late gestation influence heifer performance. J. Anim. Sci. 88:4094–4101. doi:10.2527/jas.2010-3039

Funston, R.N., A.F. Summers, and A.J. Roberts. 2012. Alpharma Beef Cattle Nutrition Symposium: implications of nutritional management for beef cow-calf systems. J. Anim. Sci. 90:2301–2307. doi:10.2527/jas.2011-4568

Garrett, W.N. 1980. Energy utilization by growing cattle as determined in 72 comparative slaughter experiments. In: Mount, L. E., editor, Energy Metabolism: Proceedings, 8th Symposium on Energy Metabolism; September 1979; Cambridge, England. EAAP Publication No. 26. London (England): Butterworths. p. 3–8.

Glore, S.R., and D.K. Layman. 1983. Cellular growth of skeletal muscle in weanling rats during dietary restrictions. Growth 47:403–410.

Godfrey, K.M., and D.J. Barker. 2000. Fetal nutrition and adult disease. Am. J. Clin. Nutr. 71 (5 Suppl):1344S–1352S. doi:10.1093/ajcn/71.5.1344s

Greenwood, P., E. Clayton, and A. Bell. 2017. Developmental programming and beef production. Anim. Front. 7:38–47. doi:10.2527/af2017-0127

Greenwood, P.L., A.S. Hunt, J.W. Hermanson, and A.W. Bell. 1998. Effects of birth weight and postnatal nutrition on neonatal sheep: I. Body growth and composition, and some aspects of energetic efficiency. J. Anim. Sci. 76:2354–2367. doi:10.2527/1998.7692354x

Greenwood, P.L., A.S. Hunt, J.W. Hermanson, and A.W. Bell. 2000. Effects of birth weight and postnatal nutrition on neonatal sheep: II. Skeletal muscle growth and development. J. Anim. Sci. 78:50–61. doi:10.2527/2000.78150x

Hammond, J. 1927. The physiology of reproduction in the cow. Cambridge (UK): Cambridge University Press.

Handel, S.E., and N.C. Stickland. 1987a. Muscle cellularity and birth weight. Anim. Sci. 44:311–317. doi:10.1017/S0003356100018687

Handel, S.E., and N.C. Stickland. 1987b. The growth and differentiation of porcine skeletal muscle fibre types and the influence of birthweight. J. Anat. 152:107–119.

Hoffman, M.L., S.A. Reed, S.M. Pillai, A.K. Jones, K.K. McFadden, S.A. Zinn, and K.E. Govoni. 2017. Physiology and endocrinology symposium: the effects of poor maternal nutrition during gestation on offspring postnatal growth and metabolism. J. Anim. Sci. 95:2222–2232. doi:10.2527/jas.2016.1229

Jackson, L.M., A. Mytinger, E.K. Roberts, T.M. Lee, D.L. Foster, V. Padmanabhan, and H.T. Jansen. 2013. Developmental programming: postnatal steroids complete prenatal steroid actions to differentially organize the GnRH surge mechanism and reproductive behavior in female sheep. Endocrinology 154:1612–1623. doi:10.1210/en.2012-1613

Kilcoyne, K.R., L.B. Smith, N. Atanassova, S. Macpherson, C. McKinnell, S. van den Driesche, M.S. Jobling, T.J. Chambers, K. De Gendt, G. Verhoeven, et al. 2014. Fetal programming of adult Leydig cell function by androgenic effects on stem/progenitor cells. Proc. Natl. Acad. Sci. USA. 111:E1924–E1932. doi:10.1073/pnas.1320735111

Kleiber, M. 1961. The fire of life: an introduction to animal energetics. New York (NY): Wiley & Sons.

Koch, R.M., and J.W. Algeo. 1983. The beef cattle industry: changes and challenges. J. Anim. Sci. 57 (Suppl. 2):28–43. doi:10.2527/animalsci1983.57supplement_228x

Koong, L.J., C.L. Ferrell, and J.A. Nienaber. 1985. Assessment of interrelationships among levels of intake and production, organ size and fasting heat production in growing animals. J. Nutr. 115:1383–1390. doi:10.1093/jn/115.10.1383

Lan, X., E.C. Cretney, J. Kropp, K. Khateeb, M.A. Berg, F. Peñagaricano, R. Magnness, A.E. Radunz, and H. Khatib. 2013. Maternal diet during pregnancy induces gene expression and DNA methylation changes in fetal tissues in sheep. Front. Genet. 4:49. doi:10.3389/fgene.2013.00049

Larson, D.M., J.L. Martin, D.C. Adams, and R.N. Funston. 2009. Winter grazing system and supplementation during late gestation influence performance of beef cows and steer progeny. J. Anim. Sci. 87:1147–1155. doi:10.2527/jas.2008-1323

Lassala, A., F.W. Bazer, T.A. Cudd, S. Datta, D.H. Keisler, M.C. Satterfield, T.E. Spencer, and G. Wu. 2010. Parenteral administration of L-arginine prevents fetal growth restriction in undernourished ewes. J. Nutr. 140:1242–1248. doi:10.3945/jn.110.125658

Lassala, A., F.W. Bazer, T.A. Cudd, S. Datta, D.H. Keisler, M.C. Satterfield, T.E. Spencer, and G. Wu. 2011. Parenteral administration of L-arginine enhances fetal survival and growth in sheep carrying multiple fetuses. J. Nutr. 141:849–855. doi:10.3945/jn.111.138172

Lofgreen, G.P., and W.N. Garrett. 1968. A system for expression of net energy requirements and feed values for growing and finishing cattle. J. Anim. Sci. 8:793–806. doi:10.2527/jas1968.273793x

Long, N.M., M.J. Prado-Coooper, C.R. Krehbiel, U. DeSilva, and R.P. Wettemann. 2011. Effects of nutrient restriction of bovine dams during early gestation on postnatal growth, carcass and organ characteristics, and gene expression in adipose tissue and muscle. J. Anim. Sci. 88:3251–3261. doi:10.2527/jas.2009-2512
Long, N.M., C.B. Tousley, K.R. Underwood, S.I. Paisley, W.J. Means, B.W. Hess, M. Du, and S.P. Ford. 2012. Effects of early- to mid-gestational undernutrition with or without protein supplementation on offspring growth, carcass characteristics, and adipocyte size in beef cattle. J. Anim. Sci. 90:197–206. doi:10.2527/jas.2011–4237

Martin, J.L., K.A. Vonnahme, D.C. Adams, G.P. Lardy, and R.N. Funston. 2007. Effects of dam nutrition on growth and reproductive performance of heifers calves. J. Anim. Sci. 85:841–847. doi:10.2527/jas.2006–337

McCoard, S.A., S.W. Peterson, W.C. McNabb, P.M. Harris, and S.N. McCutcheon. 1997. Maternal constraint influences muscle fibre development in fetal lambs. Reprod. Fertil. Dev. 9:675–681. doi:10.1071/R97061

McCoard, S., F. Sales, N. Wards, Q. Sciascia, M. Oliver, J. Koolaard, and D. van der Linden. 2013. Parenteral administration of twin-bearing ewes with L-arginine enhances the birth weight and brown fat stores in sheep. Springerplus 2:684. doi:10.1186/2193-1801-2-684

McLean, K.J., C.R. Dahlen, P.P. Borowicz, L.P. Reynolds, M.R. Crosswhite, B.W. Neville, S.D. Walden, and J.S. Caton. 2016. Technical note: a new surgical technique for ovariohysterectomy during early pregnancy in beef heifers. J. Anim. Sci. 94:5089–5096. doi:10.2527/jas.2016-0761

Meyer, A.M., and J.S. Caton. 2016. Role of the small intestine in developmental programming: impact of maternal nutrition on the dam and offspring. Adv. Nutr. 7:169–178. doi:10.3945/an.115.010405

Meyer, A.M., J.S. Caton, B.W. Hess, S.P. Ford, and L.P. Reynolds. 2012a. Epigenetics and effects on the neonate that may impact feed efficiency. In: Hill, R., editor, Feed efficiency in the beef industry. Wiley-Blackwell. p. 195–224.

Meyer, A.M., T.L. Neville, J.J. Reed, J.B. Taylor, L.P. Reynolds, D.A. Redmer, C.J. Hammer, K.A. Vonnahme, and J.S. Caton. 2013. Maternal nutritional plane and selenium supply during gestation impact visceral organ mass and intestinal growth and vasculature of neonatal lamb offspring. J. Anim. Sci. 91:2628–2639. doi:10.2527/jas.2012–5953

Meyer, A.M., J.J. Reed, T.L. Neville, J.B. Taylor, L.P. Reynolds, D.A. Redmer, K.A. Vonnahme, and J.S. Caton. 2012b. Effects of nutritional plane and selenium supply during gestation on visceral organ mass and indices of intestinal growth and vasculature in primiparous ewes at parturition and during early lactation. J. Anim. Sci. 90:2733–2749. doi:10.2527/jas.2011–4524

Mick, G.C., T.M. Sullivan, R.J. Soares Magalhaes, P.J. Rolls, S.T. Norman, and V.E. Perry. 2010. Heifer nutrition during early- and mid-pregnancy alters fetal growth trajectory and birth weight. Anim. Reprod. Sci. 117:1–10. doi:10.1016/j.anireprosci.2009.03.010

Mossa, F., F. Carter, S.W. Walsh, D.A. Kenny, G.W. Smith, J.L. Ireland, T.B. Hildebrandt, P. Lonergan, J.J. Ireland, and A.C. Evans. 2013. Maternal undernutrition in cows impairs ovarian and cardiovascular systems in their offspring. Biol. Reprod. 88:92. doi:10.1095/biolreprod.112.107235

NASEM. 2016. Nutrient requirements of beef cattle, 8th rev. ed. Washington (DC): National Academic Press.

Nissen, P.M., V.O. Danielsen, P.F. Jørgensen, and N. Oksbjerg. 2003. Increased maternal nutrition of sows has no beneficial effects on muscle fiber number or postnatal growth and has no impact on the meat quality of the offspring. J. Anim. Sci. 81:3018–3027. doi:10.2527/2003.81123018x

NRC. 1984. Nutrient requirements of beef cattle, 6th rev. ed. Washington (DC): National Academic Press.

NRC. 1996. Nutrient requirements of beef cattle, 7th rev. ed. Washington (DC): National Academic Press.

NRC. 2000. Nutrient requirements of beef cattle: update 2000, 7th rev. ed. Washington (DC): National Academic Press.

NRC. 2007. Nutrients requirements of small ruminants, sheep, goats, cervids and new world camels. Washington (DC): National Academic Press.

Paneth, N., and M. Susser. 1995. Early origin of coronary heart disease (the “barker hypothesis”). BMJ 310:411–412. doi:10.1136/bmj.310.6977.411

Paradis, F., K.M. Wood, K.C. Swanson, S.P. Miller, B.W. McBride, and C. Fitzsimmons. 2017. Maternal nutrient restriction in mid-to-late gestation influences fetal mRNA expression in muscle tissues in beef cattle. BMC Genomics 18:632. doi:10.1186/s12864-017-4051-5

Peine, J.L., G.Q. Jia, M.L. Van Emon, T.L. Neville, J.D. Kirsch, C.J. Hammer, S.T. O’Rourke, L.P. Reynolds, and J.S. Caton. 2013. Effects of maternal nutrition and rumen-protected arginine supplementation on ewe and postnatal lamb performance. Proc. West. Sec. Amer. Soc. Anim. Sci. 64:80–83.

Peine, J.L., G. Jia, M.L. Van Emon, T.L. Neville, J.D. Kirsch, C.J. Hammer, S.T. O’Rourke, L.P. Reynolds, and J.S. Caton. 2018. Effects of maternal nutrition and rumen-protected arginine supplementation on ewe performance and postnatal lamb growth and internal organ mass. J. Anim. Sci. 96:3471–3481. doi:10.1093/jas/sky221

Penagaricano, F., A.H. Souza, P.D. Carvalho, A.M. Driver, R. Gamba, J. Kropp, K.S. Hackbart, D. Luchini, R.D. Shaver, M.C. Wiltbank, et al. 2013. Effect of maternal methionine supplementation on the transcriptome of bovine preimplantation embryos. PLoS One 8:e72302. doi:10.1371/journal.pone.0072302

Prezotto, L.D., L.E. Camacho, C.O. Lemley, F.E. Keomanivong, J.S. Caton, K.A. Vonnahme, and K.C. Swanson. 2016. Nutrient restriction and reactualization in beef cows during early and mid-gestation and maternal and fetal hepatic and small intestinal in vitro oxygen consumption. Animal 10:829–837. doi:10.1017/S1751731115002645

Prezotto, L.D., C.O. Lemley, L.E. Camacho, F.E. Doscher, A.M. Meyer, J.S. Caton, B.J. Awda, K.A. Vonnahme, and K. C. Swanson. 2014. Effects of nutrient restriction and melatonin supplementation on maternal and foetal hepatic and small intestinal energy utilization. J. Anim. Physiol. Anim. Nutr. (Berl). 98:797–807. doi:10.1111/jpn.12142

Prezotto, L.D., J.F. Thorson, P.P. Borowicz, J.L. Peine, M. Bedenbaugh, S.M. Hileman, C.A. Lents, J.S. Caton, and K.C. Swanson. 2018. Influences of maternal nutrient restriction and arginine supplementation on visceral metabolic and hypothalamic circuitry of offspring. Domest. Anim. Endocrinol. 65:71–79. doi:10.1016/j.domaniend.2018.06.001

Radunz, A.E., F.L. Fluharty, A.E. Relling, T.L. Felix, L.M. Shoup, H.N. Zerby, and S.C. Loerch. 2012. Prepartum dietary energy source fed to beef cows: II. Effects on progeny postnatal growth, glucose tolerance, and
and carcass composition. J. Anim. Sci. 90:4962–4974. doi:10.2527/jas.2012-5098
Reed, S.A., and K.E. Govoni. 2017. How mom’s diet affects offspring growth and health through modified stem cell function. Anim. Front. 7:25–31. doi:10.2527/af.2017-0125
Reed, J.J., M.A. Ward, K.A. Vonnahme, T.L. Neville, S.L. Julius, P.P. Borowicz, J.B. Taylor, D.A. Redmer, A.T. Grazul-Bilska, L.P. Reynolds, et al. 2007. Effects of selenium supply and dietary restriction on maternal and fetal body weight, visceral organ mass and cellularity estimates, and jejunal vascularity in pregnant ewe lambs. J. Anim. Sci. 85:2721–2733. doi:10.2527/jas.2006-785
Rehfeldt, C., I. Fiedler, and N.C. Stickland. 2004. Number and size of muscle fibers in relation to meat production. In: te Pas, M.F.W., M.E. Everts, and H.P. Haagsman, editors. Muscle development of livestock animals: physiology, genetics, and meat quality. Cambridge (MA): CABI Publishing. p. 1–38.
Reynolds, L.P., P.P. Borowicz, J.S. Caton, K.A. Vonnahme, J.S. Luther, C.J. Hammer, K.R. Maddock Carlin, A.T. Grazul-Bilska, and D.A. Redmer. 2010. Developmental programming: the concept, large animal models, and the key role of utero-placental vascular development. J. Anim. Sci. 88 (13 Suppl):E61–E72. doi:10.2527/jas.2009-2359
Reynolds, L.P., P.P. Borowicz, K.A. Vonnahme, M.L. Johnson, A.T. Grazul-Bilska, J.M. Wallace, J.S. Caton, and D.A. Redmer. 2005. Animal models of placental angiogenesis. Placenta 26:689–708. doi:10.1016/j.placenta.2004.11.010
Reynolds, L.P., and J.S. Caton. 2012. Role of the pre- and postnatal environment in development of offspring energy requirements. Mol. Cell. Endocrinol. 354:54–59. doi:10.1016/j.mce.2011.11.013
Reynolds, L.P., Grazul-Bilska, A.T., & Borowicz, P.P. 2018. Placental angiogenesis. In: Skinner, M. K., editor. Developmental biology of the uterus: insights into mechanisms and developmental disruption. Mol. Cell. Endocrinol. 354:34–53. doi:10.1016/j.mce.2011.09.035
Reynolds, C.K., H.F. Tyrrill, and P.J. Reynolds. 1991. Effects of diet forage-to-concentrate ratio and intake on energy metabolism in growing beef heifers: whole body energy and nitrogen balance and visceral heat production. J. Nutr. 121:994–1003. doi:10.1093/jjn/121.7.994
Reynolds, L.P., and K.A. Vonnahme. 2016. Triennial reproduction symposium: developmental programming of fertility. J. Anim. Sci. 94:2699–2704. doi:10.2527/jas.2015-0131
Reynolds, L.P., and K.A. Vonnahme. 2017. Livestock as models for developmental programming. Anim. Front. 7:12–17. doi:10.2527/af.2017-0123
Reynolds, L.P., K.A. Vonnahme, C.O. Lemley, D.A. Redmer, A.T. Grazul-Bilska, P.P. Borowicz, and J.S. Caton. 2013. Maternal stress and placental vascular function and remodeling. Curr. Vasc. Pharmacol. 11:564–593. doi:10.2174/1570161111311050003
Reynolds, L.P., A.K. Ward, and J.S. Caton. 2017. Epigenetics and developmental programming in ruminants – long-term impacts on growth and development. In: Scanes, C.F., and R. Hill, editors. Biology of domestic animals. Milton Park (UK): CRC Press/Taylor & Francis Group.
Reynolds, L.P., M.C. Wulster-Radcliffe, D.K. Aaron, and T.A. Davis. 2015. Importance of animals in agricultural sustainability and food security. J. Nutr. 145:1377–1379. doi:10.3945/jn.115.212217
Rhind, S.M., M.T. Rae, and A.N. Brooks. 2001. Effects of nutrition and environmental factors on the fetal programming of the reproductive axis. Reproduction 122:205–214. doi:10.1530/rep.0.1220205
Robinson, J.J. 1977. The influence of maternal nutrition on ovine foetal growth. Proc. Nutr. Soc. 36:9–16. doi:10.1079/PNS19770003
Robinson, D.L., L.M. Cafe, and P.L. Greenwood. 2013. Meat science and muscle biology symposium: developmental programming in cattle: consequences for growth, efficiency, carcass, muscle, and beef quality characteristics. J. Anim. Sci. 91:1428–1442. doi:10.2527/jas.2012-5799
Robinson, J.J., K.D. Sinclair, and T.G. McEvoy. 1999. Nutritional effects on foetal growth. Anim. Sci. 68:315–331. doi:10.1017/S1357729800050323
Russell, R.G., and F.T. Oteruelo. 1981. An ultrastructural study of the differentiation of skeletal muscle in the bovine fetus. Anat. Embryol. (Berl). 162:403–417. doi:10.1007/BF00301866
Schmidt, K.L., E.A. Macdougall-Shackleton, K.K. Soma, and S.A. Macdougall-Shackleton. 2014. Developmental programming of the HPA and HPG axes by early-life stress in male and female song sparrows. Gen. Comp. Endocrinol. 196:72–80. doi:10.1016/j.ygcen.2013.11.014
Shankar, K., Y. Zhong, F. Lau, M.L. Blackburn, J.R. Chen, S.J. Borengasser, M.J. Ronis, and T.M. Badger. 2011. Maternal obesity promotes a proinflammatory signature in rat uterus and blastocyst. Endocrinology 152:4158–4170. doi:10.1210/en.2010-1078
Sheldon, B.C., and S.A. West. 2004. Maternal dominance, maternal condition, and offspring sex ratio in ungulates. Am. Nat. 163:40–54. doi:10.1086/381003
Short, R.E., R.A. Bellows, R.B. Staigmiller, J.G. Berardinelli, S.L. Julius, P.P. Borowicz, J.B. Taylor, D.A. Redmer, and E.E. Custer. 1990. Physiological mechanisms controlling anestrus and infertility in postpartum beef cattle. J. Anim. Sci. 68:799–816. doi:10.2527/1990.683799x
Sletmoen- Olson, K.E., J.S. Caton, L.P. Reynolds, and K.C. Olson. 2000. Undergraded intake protein supplementation: I. Effects on forage utilization and performance of perennial ryegrass beef cows fed low-quality hay during gestation and lactation. J. Anim. Sci. 78:449–455. doi:10.2527/2000.782449x
Spencer, T.E., K.A. Dunlap, and J. Filant. 2012. Comparative developmental biology of the uterus: insights into mechanisms and developmental disruption. Mol. Cell. Endocrinol. 354:34–53. doi:10.1016/j.mce.2011.09.035
Spitzer, J.C., D.G. Morrison, R.P. Wettemann, and L.C. Faulkner. 1995. Reproductive responses and calf birth and weaning weights as affected by body condition at parturition and postpartum weight gain in primiparous beef cows. J. Anim. Sci. 73:1251–1257. doi:10.2527/1995.7351251x
Stalker, L.A., D.C. Adams, T.J. Klopenstein, D.M. Feuz, and R.N. Funston. 2006. Effects of pre- and postpartum nutrition on reproduction in spring calving cows and calf feedlot performance. J. Anim. Sci. 84:2582–2589. doi:10.2527/jas.2005-5098
Stalker, L.A., D.C. Adams, T.J. Klopenstein, and R.N. Funston. 2006. Effects of pre- and postpartum nutrition on reproduction in spring calving cows and calf feedlot performance. J. Anim. Sci. 84:2582–2589. doi:10.2527/jas.2005-5098

Stickland, N.C., S. Bayol, C. Ashton, and C. Rehfeldt. 2004. Manipulation of muscle fiber number during prenatal development. In: te Pas, M.F.W., M.E. Everts, and H.P. Haagsman, editors. Muscle development of livestock animals: physiology, genetics, and meat quality. Cambridge (MA): CABI Publishing. p. 69–82.

Swanson, T.J., C.J. Hammer, J.S. Luther, D.B. Carlson, J.B. Taylor, D.A. Redmer, T.L. Neville, J.J. Reed, L.P. Reynolds, J.S. Caton, et al. 2008. Effects of gestational plane of nutrition and selenium supplementation on mammary development andcolostrum quality in pregnant ewe lambs. J. Anim. Sci. 86:2415–2423. doi:10.2527/jas.2008-0996

Symonds, M.E., M. Pope, D. Sharkey, and H. Budge. 2012. Adipose tissue and fetal programming. Diabetologia 55:1597–1606. doi:10.1007/s00125-012-2505-5

Underwood, K.R., J.F. Tong, P.L. Price, A.J. Roberts, E.E. Grings, B.W. Hess, W.J. Means, and M. Du. 2010. Nutrition during mid to late gestation affects growth, adipose tissue deposition, and tenderness in cross-bred beef steers. Meat Sci. 86:588–593. doi:10.1016/j.meatsci.2010.04.008

United Nations News Centre. 2015. World population projected to reach 9.6 billion by 2050. Available from http://www.un.org/en/development/desa/news/population/un-report-world-population-projected-to-reach-9-6-billion-by-2050.html. Accessed December 28, 2015.

Vonnahme, K.A., and C.O. Lemley. 2012. Programming the offspring through altered urogenital hemodynamics: how maternal environment impacts uterine and umbilical blood flow in cattle, sheep and pigs. Reprod. Fertil. Develop. 24:97–104. doi:10.1071/RD11910

Vonnahme, K.A., C.O. Lemley, J.S. Caton, and A.M. Meyer. 2015. Impacts of maternal nutrition on vasculature of nutrient transferring tissues during gestation and lactation. Nutrients 7:3497–3523. doi:10.3390/nu7053497

Vonnahme, K.A., J.S. Luther, L.P. Reynolds, C.J. Hammer, D.B. Carlson, D.A. Redmer, and J.S. Caton. 2010. Impacts of maternal selenium and nutritional level on growth, adiposity, and glucose tolerance in female offspring in sheep. Domest. Anim. Endocrinol. 39:240–248. doi:10.1016/j.domaniend.2010.06.005

Wang, X., X. Lan, A.E. Radunz, and H. Khatib. 2015. Maternal nutrition during pregnancy is associated with differential expression of imprinted genes and DNA methyltransferases in muscle of beef cattle offspring. J. Anim. Sci. 93:35–40. doi:10.2527/jas.2014-8148

Ward, A.K., M.S. Crouse, R.A. Cushman, K.J. McLean, C.R. Dahlen, P.P. Borowicz, L.P. Reynolds, and J.S. Caton. 2017. Maternal nutrient restriction in early gestation upregulates myogenic genes in cattle fetal muscle tissue. 36th International Society of Animal Genetics Conference; July 16 to 21, 2017; Dublin, Ireland. p. 177.

Ward, M.A., T.L. Neville, J.J. Reed, J.B. Taylor, D.M. Hallford, S.A. Soto-Navarro, K.A. Vonnahme, D.A. Redmer, L.P. Reynolds, and J.S. Caton. 2008. Effects of selenium supply and dietary restriction on maternal and fetal metabolic hormones in pregnant ewe lambs. J. Anim. Sci. 86:1254–1262. doi:10.2527/jas.2007-0509

Ward, S.S., and N.C. Stickland. 1991. Why are slow and fast muscles differentially affected during prenatal undernutrition? Muscle Nerve 14:259–267. doi:10.1002/mus.88104301

Wilson, S.J., J.J. Ross, and A.J. Harris. 1988. A critical period for formation of secondary myotubes defined by prenatal undernourishment in rats. Development 102:815–821.

Wu, G., F.W. Bazer, J.M. Wallace, and T.E. Spencer. 2006. Intraterine growth retardation: implications for the animal sciences. J. Anim. Sci. 84:2316–2337. doi:10.2527/jas.2006-156

Xiong, F., and L. Zhang. 2013. Role of the hypothalamic-pituitary-adrenal axis in developmental programming of health and disease. Front. Neuroendocrinol. 34:27–46. doi:10.1016/j.yfrne.2012.11.002

Yunusova, R., T.L. Neville, K.A. Vonnahme, C.J. Hammer, J.J. Reed, J.B. Taylor, D.A. Redmer, L.P. Reynolds, and J.S. Caton. 2013. Impacts of maternal selenium supply and nutritional plane on visceral tissues and intestinal biology in offspring. J. Anim. Sci. 91:2229–2242. doi:10.2527/jas.2012-5134

Zambrano, E., C. Guzmán, G.L. Rodríguez-González, M. Durand-Carbajal, and P.W. Nathanielsz. 2014. Fetal programming of sexual development and reproductivity. Mol. Cell. Endocrinol. 382:538–549. doi:10.1016/j.mce.2013.09.008

Zhang, S.T., R.H. Reguault, P.L. Barker, K.J. Botting, I.C. McMillen, C.M. McMillan, C.T. Roberts, and J.L. Morrison. 2015. Placental adaptations in growth restriction. Nutrients. 7:360–389. doi:10.3390/nu7010360

Zhu, M.J., S.P. Ford, W.J. Means, B.W. Hess, P.W. Nathanielsz, and M. Du. 2006. Maternal nutrient restriction affects properties of skeletal muscle in offspring. J. Physiol. 575(Pt 1):241–250. doi:10.1113/jphysiol.2006.112110

Zhu, M.J., S.P. Ford, P.W. Nathanielsz, and M. Du. 2004. Effect of maternal nutrient restriction in sheep on the development of fetal skeletal muscle. Biol. Reprod. 71:1968–1973. doi:10.1095/biolreprod.104.034561

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