FETAL PROGRAMMING

Supplementing organic-complexed or inorganic Co, Cu, Mn, and Zn to beef cows during gestation: physiological and productive response of cows and their offspring until weaning

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Abstract

One hundred and ninety non-lactating, pregnant beef cows (three-fourth Bos taurus and one-fourth Bos indicus; 138 multiparous and 52 primiparous) were assigned to this experiment at 117 ± 2.2 d of gestation (day 0). Cows were ranked by parity, pregnancy type (artificial insemination = 102 and natural service = 88), body weight (BW), and body condition score (BCS) and assigned to receive a supplement containing: 1) sulfate sources of Cu, Co, Mn, and Zn (INR; n = 95) or 2) an organic-complexed source of Cu, Mn, Co, and Zn (AAC; Availa 4; Zinpro Corporation, Eden Prairie, MN; n = 95). The INR and AAC provided the same daily amount of Cu, Co, Mn, and Zn, based on 7 g of the AAC source. From day 0 to calving, cows were maintained in a single pasture and were segregated three times weekly into 1 of the 24 individual feeding pens to receive treatments. Cow BW and BCS were recorded on days −30, 97, upon calving, and at weaning (day 367). Milk production was estimated at 42 ± 0.5 d postpartum via weigh–suckle–weigh (WSW) method. Liver biopsies were performed in 30 cows per treatment on days −30, 97, upon calving, and the day after WSW. Calf BW was recorded at birth and weaning. Liver and longissimus muscle (LM) biopsies were performed in 30 calves per treatment upon calving and 24 h later, the day after WSW, and at weaning. No treatment effects were detected (P ≥ 0.49) for calf BW recorded at birth and weaning. Liver and longissimus muscle (LM) biopsies were performed in 30 calves per treatment upon calving and 24 h later, the day after WSW, and at weaning. No treatment effects were detected (P ≥ 0.49) for cow BCS during gestation, despite AAC cows having greater (P = 0.04) BW on day 97. Liver Co concentrations were greater (P < 0.01) for AAC compared with INR cows, and liver concentrations of Cu were greater (P = 0.02) for INR compared with AAC cows on day 97. Upon calving, INR cows had greater (P ≤ 0.01) liver Cu and Zn concentrations compared with AAC cows. No other treatment differences were noted (P ≥ 0.17) for cow and calf liver trace mineral concentrations. Cows receiving AAC had greater (P = 0.04) hepatic mRNA expression of metallothionein 1A at calving, and their calves had greater (P = 0.04) hepatic mRNA expression of superoxide dismutase at weaning. Milk production did not differ between AAC and INR cows (P = 0.70). No treatment effects were detected (P ≥ 0.29) for mRNA expression of LM genes associated with adipogenic or muscle development activities in calves at birth and weaning. Calf birth and weaning BW also did not differ (P ≥ 0.19) between treatments. In summary, supplementing AAC or INR to beef cows during the last 5 mo of gestation yielded similar cow–calf productive responses until weaning.

Key words: beef cows, gestation, offspring, physiology, production, trace minerals
Introduction

Maternal nutrition is a major extrinsic factor programming nutrient partitioning and development of fetal organ systems, leading to long-term effects on offspring production, health, and reproduction (Long et al., 2009, 2010). The majority of research conducted in this area has focused primarily on the energy and protein intake of gestating beef cows (Funston et al., 2010; Bohnert et al., 2013; Wilson et al., 2016), and limited knowledge exists on how trace mineral nutrition during gestation impacts offspring productivity. The fetus is completely dependent on the dam for its supply of trace minerals (Hidiroglou and Knipfel, 1981), which are essential for fetal developmental processes, such as protein synthesis, bone formation, and lipid metabolism (Hostetler et al., 2003). Inadequate maternal intake or transfer of trace minerals to the fetus can result in impaired fetal development and postnatal performance (Weiss et al., 1983).

One strategy to enhance trace mineral status in cattle is to feed organic sources (Spears, 1996). There has been considerable interest in the use of organic trace minerals in ruminant diets due to indications that these may be of greater bioavailability compared with their inorganic counterparts (Brown and Zeringue, 1994). Hostetler et al. (2003) reported that supplementing an organic Cu, Mn, and Zn to gestating sows increased concentrations of these trace elements in fetal tissues and reduced fetal loss by day 30 of gestation compared with sows supplemented with inorganic Cu, Mn, and Zn. Similarly, research from our group reported improved productivity responses in the offspring from beef cows receiving organic trace minerals during late gestation (Marques et al., 2016a). More specifically, supplementing beef cows with organic-complexed Co, Cu, Mn, and Zn enhanced the passage of Zn and Cu to fetal liver, resulting in increased growth rate until weaning and postweaning immunity to bovine respiratory disease.

The results from Marques et al. (2016a) were suggestive of programming effects of organic-complexed trace minerals on postnatal offspring development and health (Funston et al., 2010); however, the physiological mechanisms underlying these outcomes still warranted investigation. Nutritional management of beef cows impacts fetal development throughout the entirety of gestation (Wu et al., 2006), and Marques et al. (2016a) investigated the supplementation for approximately 90 d prior to calving. Hence, supplementing an organic trace mineral source may be even more beneficial to offspring development if offered to beef cows over a greater duration of gestation. To test this theory and expand on the novel results from Marques et al. (2016a), this experiment evaluated the effects of supplementing organic-complexed or sulfate sources of Co, Cu, Mn, and Zn to beef cows during the second and third trimesters of gestation on productive and physiological responses of the cow and their offspring.

Materials and Methods

This experiment was conducted at the Texas A&M – Beef Cattle Systems (College Station, TX, USA). All animals were cared for in accordance with acceptable practices and experimental protocols reviewed and approved by the Texas A&M – Institute of Animal Care of Use Committee (#2018/0093). This manuscript describes prepartum and postpartum responses of cows as well as offspring responses from birth through weaning. A companion manuscript (Harvey et al., 2021) describes the postweaning responses of the male and female offspring reared, respectively, as feedlot steers or replacement heifers.

Cow management and dietary treatments

One hundred and ninety non-lactating, pregnant beef cows (average three-fourth Bos taurus and one-fourth Bos indicus; 138 multiparous and 52 primiparous; initial body weight [BW] = 509 ± 5.7 kg; age = 4.6 ± 0.2 yr; initial body condition score [BCS] = 5.5 ± 0.1 according to Wagner et al., 1988) were assigned to this experiment at 117 ± 2.2 d of gestation (day 0 of the experiment). Cows were pregnant to artificial insemination (AI) using semen from a single Brangus sire (n = 83) or pregnant to six Brangus bulls according to the breeding management and pregnancy diagnosis described by Oosthuizen et al. (2020).

Prior to the beginning of the experiment (day 30), cows were ranked by pregnancy type (AI or natural service), parity, BW, and BCS and assigned to receive a supplement containing: 1) sulfate sources of Cu, Co, Mn, and Zn (INR; custom blend manufactured by Anipiro Xtraformance Feeds, Pratt, KS; n = 95) or 2) organic-complexed source of Cu, Co, Mn, and Zn (AAC; Availa 4; Zinpro Corporation, Eden Prairie, MN; n = 95). The AAC trace mineral source was based on a metal:amino acid complex ratio of 1:1 for Zn, Cu, and Mn in addition to cobalt glucoheptonate (Zinpro Corporation). The INR and AAC sources were mixed with dried distillers grain (Table 1) and formulated to provide the same daily amount of energy, protein, macro minerals, and trace minerals, including Cu, Co, Mn, and Zn (based on 7 g/cow daily of Availa 4; as in Marques et al., 2016a). From day 0 of the experiment until calving, cows were maintained in a single pasture dominated by bermudagrass (Cynodon spp.). All cows were gathered three times weekly (Tuesdays, Thursdays, and Saturdays) and individually sorted into 1 of the 24 feeding pens (one cow per pen; 6 × 9 m). Cows individually received their assigned supplement treatment (601 or 607 g of INR and AAC per cow each feeding; as-fed basis) and returned to pasture after their treatment was completely consumed. This process was repeated until all cows had been individually sorted into pens and consumed their treatments. From day 93 of the experiment until calving, cows also received daily 12 kg/cow of sorghum-sudangrass hay and 0.76 kg/cow of cubes containing no mineral additive (Table 2).

After calving, cow–calf pairs were moved to an adjacent pasture dominated by perennial ryegrass (Lolium perenne L.) and assigned to the general management of the research herd, which included free-choice inorganic trace mineral supplementation (Producers Special Pasture Mineral; Producers Cooperative Association, College Station, TX; containing 14% Ca, 7% P, 13% NaCl, 5% Mg, 9,900 mg/kg Zn, 2,500 mg/kg Cu, 100 mg/kg I, 4,000 mg/kg Mn, 26 mg/kg Se, 91 IU/g of vitamin A, 10 IU/g of vitamin D3, and 0.05 IU/g of vitamin E). This trace mineral supplement was the same fed to cows prior to the beginning of this experiment. Male calves were castrated using an elastic bander at approximately 30 d of age. Cows were assigned to the same reproductive management (day 214 to 287) and
pregnancy diagnosis (day 345 of the experiment) as described by Oosthuizen et al. (2020). Cows were inseminated by the same technician using semen from the same bull and batch and exposed to natural service (six mature bulls) for 60 d. All calves received vaccination against respiratory viruses (Triangle 5; Boehringer Ingelheim Animal Health USA Inc., Duluth, GA) and Clostridium (Covexin 8; Merck Animal Health, Omaha, NE) on day 345 of the experiment. Calves were weaned on day 367, when they were revaccinated against respiratory viruses (Titanium 5; Elanco Animal Health, Greenfield, IN) and Clostridium (Covexin 8; Merck Animal Health), and received an anthelmintic (Dectomax; Zoetis, Florham Park, NJ).

Sampling
Samples of all ingredients fed to gestating cows were collected monthly during the experiment, pooled across months, and analyzed for nutrient content by a commercial laboratory (Dairy One Forage Laboratory, Ithaca, NY). The nutritional profile of all ingredients is described in Tables 1 and 2. Individual BW and BCS (Wagner et al., 1988) were recorded from all cows prior to the beginning of the experiment (day –30) and during late gestation (day 97). Liver samples were collected via needle biopsy (TruCut biopsy needle; CareFusion Corporation, San Diego, CA) from 30 cows on days −30 and 97, whereas cows were randomly chosen from each treatment on each sampling day. The same biopsy and selection procedure was adopted for the liver and longissimus muscle (LM) samplings throughout the experimental period (Marques et al., 2016a; Schubach et al., 2019).

After calving, cow BW and BCS were recorded from all cows, while another subset of 30 cows were randomly selected from each treatment for colostrum sampling via hand milking (100 mL) and liver biopsy. Birth BW and sex were recorded from all calves, and calves from cows selected for colostrum sampling were assigned to liver and LM biopsies. Cows and calves were sampled immediately after calving was identified and completed, except for cows that calved at night whose calves

| Item | INR | AAC |
|------|-----|-----|
| Dried distillers grains | 193 | 193 |
| Macromineral mix | 60.0 | 60.0 |
| Inorganic trace mix | 4.67 | 0.00 |
| Organic trace mix | 0.00 | 7.00 |
| Nutrient profile (dry matter basis) | | |
| DM, % | 90.8 | 90.9 |
| Net energy for maintenance, Mcal/kg | 1.58 | 1.57 |
| Net energy for lactation, Mcal/kg | 1.50 | 1.49 |
| Crude protein, % | 26.0 | 25.7 |
| Ca, % | 4.52 | 4.53 |
| P, % | 2.91 | 2.88 |
| Mg, % | 0.64 | 0.63 |
| K, % | 1.13 | 1.13 |
| Na, % | 1.89 | 1.93 |
| Co, mg/kg | 62.0 | 65.0 |
| Cu, mg/kg | 605 | 600 |
| Fe, mg/kg | 1,883 | 1,931 |
| Mn, mg/kg | 983 | 967 |
| Se, mg/kg | 6.44 | 6.39 |
| Zn, mg/kg | 1,791 | 1,695 |

1Containing (DM basis) 339.7 g/kg CaHPO₄, 312.7 g/kg CaCO₃, 197.5 g/kg NaCl, 39.1 g/kg KCl, 18.9 g/kg MgO, 4.0 g/kg Fe₂O₃, 2.5 g/kg S, 0.9 g/kg Vit E, 0.8 g/kg Na₂O₃Se₃%, 0.7 g/kg Vit A, 0.1 g/kg C₂H₁₀I₂N₂, and 0.1 g/kg Vitamin D 400.
2Containing (DM basis) 420 g/kg ground corn, 213 g/kg ZnSO₄, 133 g/kg MnSO₄, 106 g/kg CuSO₄, and 8 g/kg CoSO₄.
3Available from Zinpro Corporation; Eden Prairie, MN, which contained (DM basis) 5.15% Zn from 1:1 Zn and AA complex, 2.86% Mn from 1:1 Mn and amino acid (AA) complex, 1.80% Cu from 1:1 Cu and AA complex, and 0.18% Co from cobalt glucoheptonate.
4Values obtained via wet chemistry analysis (Dairy One Forage Laboratory, Ithaca, NY).
5Calculated with equations described by the NRC (2000).

Table 2. Nutritional profile (dry matter basis) of feedstuffs offered to cows during the experiment

| Item | A | B | Supplemental cubes | Sorghum-sudangrass hay |
|------|---|---|-------------------|----------------------|
| Net energy for maintenance, Mcal/kg | 1.14 | 1.62 | 2.00 | 1.00 |
| Net energy for lactation, Mcal/kg | 1.18 | 1.02 | 1.33 | 0.44 |
| Crude protein, % | 10.7 | 20.5 | 48.0 | 11.8 |
| Neutral detergent fiber, % | 57.7 | 19.6 | 24.0 | 61.5 |
| Ca, % | 0.98 | 2.07 | 0.32 | 0.76 |
| P, % | 0.27 | 0.32 | 1.26 | 0.25 |
| Mg, % | 0.29 | 0.41 | 0.67 | 0.31 |
| K, % | 1.62 | 2.72 | 1.86 | 4.11 |
| Na, % | 0.02 | 0.05 | 0.28 | 0.02 |
| Co, mg/kg | 1.61 | 3.05 | 1.10 | 0.69 |
| Cu, mg/kg | 8 | 12 | 15 | 11 |
| Fe, mg/kg | 3,855 | 5,120 | 303 | 1,566 |
| Mn, mg/kg | 106 | 144 | 36 | 54 |
| Se, mg/kg | 0.40 | 0.20 | 0.19 | 0.12 |
| Zn, mg/kg | 42 | 35 | 66 | 57 |

1Values obtained from a commercial laboratory wet chemistry analysis (Dairy One Forage Laboratory, Ithaca, NY). Total digestible nutrients were calculated according to equations described by Weiss et al. (1992). Net energy for maintenance and lactation were calculated with equations described by the NRC (2000).
2Cows were maintained in a single pasture (A) dominated by bermudagrass (Cynodon spp.) and were supplemented with 12 kg of sorghum-sudangrass hay and 0.76 kg of cubes daily from day 93 until calving. Upon calving, cow–calf pairs were moved to an adjacent pasture (B) dominated by perennial ryegrass (Lolium perenne L.).
were sampled the next morning within 8 h of calving. Another liver biopsy was collected from the same set of calves 24 h after birth. When feasible, the expelled placenta was retrieved and immediately rinsed with nanopure water for 5 min, with 32 placentas retrieved (16 placentas/treatment). All placentas were expelled within 12 h after calving and not considered retained fetal membranes (Takagi et al., 2002). The five largest cotyledons were dissected from each placenta using curved scissors, given that the largest cotyledons are expected to be the most active regarding the nutrient transfer from the dam to the fetus (Senger, 2003).

Milk production was estimated from all cows, at 42 ± 0.5 d postpartum, via the weigh–suckle–weigh (WSW) method (Aguiar et al., 2015). More specifically, calves were separated from their dams for 8 h, weighed, allowed sucking for 30 min, and were weighed again. Milk yield was calculated as the difference in pre- and post-sucking calf BW and was adjusted to 24 h by multiplying by a factor of three. The day following WSW, another subset of 30 cow-calf pairs were randomly selected from each treatment for sampling of fresh milk via hand milking (100 mL) as well as liver biopsy from cows and calves. At weaning (day 367 of the experiment), cow BW and BCS were recorded at weaning, and another subset of 30 calves per treatment were randomly selected for liver and LM biopsy. Calf BW was recorded on days 367 and 368, and the average was considered as calf weaning BW.

Laboratorial analyses

All feed samples were analyzed by wet chemistry procedures for concentrations of crude protein (method 984.13; AOAC, 2006), acid detergent fiber (method 973.18 modified for use in an Ankrom 200 fiber analyzer, Ankrom Technology Corp., Fairport, NY; AOAC, 2006), and neutral detergent fiber (Van Soest et al., 1991; modified for use in an Ankrom 200 fiber analyzer, Ankrom Technology Corp.), macro- and trace minerals using inductively coupled plasma emission spectroscopy (Sirois et al., 1991), and Se according to method 996.16 of the AOAC (2006). Calculations for total digestible nutrients used the equation proposed by Weiss et al. (1992), and calculations for net energy for maintenance and lactation used the equations proposed by NRC (2000).

Liver, LM, cotyledon, colostrum, and milk samples were placed immediately on ice after collected and processed for storage within 8 h. Liver and LM samples were stored in duplicates at –80 °C, with or without 1 mL of RNA stabilization solution (RNAlater, Ambion, Inc., Austin, TX). Liver, cotyledon, colostrum, and milk samples were analyzed via inductively coupled plasma mass spectrometry for concentrations of Co, Cu, Mn, and Zn by the Michigan State University Diagnostic Center for Population and Animal Health (East Lansing, MI) according to Braselton et al. (1997). Total RNA was extracted from liver and LM samples stored in RNA stabilization solution using the TRIzol Plus RNA Purification Kit (Invitrogen, Carlsbad, CA). The number of isolated RNA were assessed via UV absorbance (NanoDrop Lite; Thermo Fisher Scientific, Wilmington, DE) at 260 nm and 260/280 nm ratio, respectively (Fleige and Pfaffl, 2006). Reverse transcription of extracted RNA and real-time reverse-transcription polymerase chain reaction (PCR) using gene-specific primers (20 pM each; Table 3) were completed as described by Rodrigues et al. (2015). Responses from the genes of interest were quantified based on the threshold cycle (CT), wherein the number of PCR cycles required for target amplification to reach a predetermined threshold. A portion of the amplified products was purified with the QIAquick PCR purification kit (Qiagen Inc., Valencia, CA) and sequenced at the Texas A&M AgriLife Genomics and Bioinformatics Service to verify the specificity of amplification. All amplified products represented only the genes of interest. The CT responses from the liver genes of interest were normalized to the geometrical mean of CT values of ribosomal protein L12 and cyclophilin, whereas the CT responses from the muscle genes of interest were normalized to the geometrical mean of CT values of ribosomal protein S9 and β-actin (Vandesompele et al., 2002). The coefficient of variation for the geometrical mean of reference genes across all liver and LM samples was 3.0% and 2.6%, respectively. Results are expressed as relative fold change (2^−ΔΔCT) as described by Ocón-Grove et al. (2008).

Statistical analysis

All cow and calf variables were analyzed with cow as the experimental unit and cow(treatment × parity) as the random variable. Quantitative data were analyzed using the MIXED procedure of SAS (SAS Inst. Inc., Cary, NC), and binary data were analyzed using the GLIMMIX procedure of SAS (SAS Inst. Inc.). All data collected beginning at calving were analyzed using gestation days receiving treatment as an independent covariate. Model statements for cow-related responses included the effects of treatment, parity, day for repeated measures, and all the resultant interactions. Model statements for calf-related responses and placental cotyledons analysis included the effects of treatment, parity, calf sex, day for repeated measures, and all resultant interactions. For repeated measures, the subject for the repeated statement was cow(treatment × parity) and the covariance structure utilized was autoregressive, which provided the best fit according to the lowest Akaike information criterion. Results are reported as least square means (covariately adjusted for post-calving responses) and separated using least square difference. Significance was set at P < 0.05, and tendencies were determined if P > 0.05 and ≤ 0.10. Results are reported according to main treatment effects if higher-order interactions containing treatments were nonsignificant or according to highest-order interaction detected.

Results and Discussion

Total feed intake of cows during gestation was not monitored individually as in Marques et al. (2016a) to calculate the daily consumption of Zn, Cu, Mn, and Co. Nonetheless, supplement treatments provided to each cow on a daily basis, approximately 400 mg of Zn, 140 mg of Cu, 230 mg of Mn, and 1.50 mg of Co (Table 1). Based on the BW of cows during gestation (~450 kg), composition of treatments and feedstuffs (Tables 1 and 2), and estimated daily feed intake of 9.0 kg/cow (dry matter basis; NASEM, 2016), diets fed during gestation contained at least 85 ppm of Zn, 26.6 ppm of Cu, 129 ppm of Mn, and 3.28 ppm of Co. As in Marques et al. (2016a), requirements for Cu, Mn, and Zn were exceeded by more than 200%, whereas the requirement for Co was exceeded by over 2,000% for both INR and AAC treatments (NASEM, 2016). A non-supplemented group was not included in this experiment to conform with current industry practices as the majority of U.S. cow–calf herds receive supplementation (USDA-APHIS, 2010), whereas basal dietary ingredients already provided Co, Cu, Mn, and Zn near or above adequate levels (Table 2; NASEM, 2016). Therefore, this experiment was designed to compare supplemental INR or AAC to gestating beef cows and not to investigate trace mineral deficiency or differences in trace mineral intake between experimental treatments.
Cow responses

Cow age, BW, and BCS on day −30 were similar (P ≥ 0.35) between treatments as designed (Table 4). Length of treatment administration also did not differ (P = 0.91) between INR and AAC cows (Table 4), and cows received treatments during the second and third trimesters of gestation. Cows receiving AAC had greater (P = 0.04; Table 4) BW at late gestation (day 97) compared with INR cows, although this difference was insufficient to impact concurrent BCS (P = 0.93; Table 4). No differences were detected (P ≥ 0.31) for cow calving BW and BCS between treatments (Table 4). These outcomes were expected given that AAC and INR cows were maintained in a single group under equivalent nutritional management until calving. Others have also reported that Cu, Co, Mn, and Zn supplementation as organic or inorganic sources resulted in similar cow BCS change (Stanton et al., 2000; Ahola et al., 2004; Marques et al., 2016a). It should be noted that both AAC and INR cows lost BCS during gestation (day effect; P < 0.01) due to inclement weather and excessive pasture flooding. Cows should gain BCS during gestation to optimize offspring productivity (Bohnert et al., 2013; Marques et al., 2016b). This BCS loss may have impacted the outcomes of this experiment but equivalently across treatments given the lack of BCS differences (P ≥ 0.49) during gestation between AAC and INR cows (Table 4).

No treatment differences were detected (P ≥ 0.39) for initial (day −30) liver Cu, Co, Mn, and Zn concentrations (Table 5), indicating similar trace mineral status between AAC and INR cows prior to treatment administration. In samples collected during late gestation (day 97), the liver Co concentrations were greater (P < 0.01) for AAC compared with INR cows, and the liver concentrations of Cu were greater (P = 0.02) for INR compared with AAC cows. No treatment differences were detected (P ≥ 0.27) for liver Mn or Co concentrations on day 97 (Table 5). Marques et al. (2016a) also reported greater liver Co, less liver Cu, and similar liver Zn and Mn between cows supplemented with sulfate or organic-complexed sources in samples collected during late gestation. Upon calving, INR cows had greater (P ≤ 0.01) liver Cu and Zn concentrations compared with AAC cows, and no treatment differences were detected (P ≥ 0.17) for liver Mn or Co concentrations (Table 5). Collectively, treatment effects on liver trace mineral status do not corroborate that organic trace mineral sources have increased absorption and retention compared with sulfate minerals (Spears, 1996; Hostetler et al., 2003). Hepatic Mn concentrations are not influenced by dietary Mn intake in ruminants (Underwood and Suttle, 1999; Ahola et al., 2004), and Marques et al. (2016a) reported that Mn supplementation as AAC or INR did not increase liver concentrations of this trace mineral in late-gestating cows. In

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### Table 3. Primer sequences, accession number, and reference for gene transcripts analyzed by real-time reverse transcription PCR

| Target                                     | Primer sequence          | Accession no.   | Source                  |
|--------------------------------------------|--------------------------|-----------------|-------------------------|
| **Liver samples**                          |                          |                 |                         |
| CUT                                        | Forward: GGGTACCTCTGCATTTGCTGT  | NM_001100381    | Han et al. (2009)       |
|                                           | Reverse: ATGCAATCTGCATTTGCTGT  |                 |                         |
| MT                                         | Forward: ACACACAGTCATTTGCTGT  | NM_001040492.2  | Gessner et al. (2013)   |
|                                           | Reverse: CGAAGCAGGATTTGCTGT  |                 |                         |
| SOD                                        | Forward: TGGTACATCTGATTTGCTGT  | NM_174615       | Gessner et al. (2013)   |
|                                           | Reverse: CAGCCTGACCATTTGCTGT  |                 |                         |
| Ribosomal protein L12                      | Forward: CACACAGTCATTTGCTGT  | NM_205797.1     | Gessner et al. (2013)   |
|                                           | Reverse: CGAAGCAGGATTTGCTGT  |                 |                         |
| Cyclophilin                                | Forward: GGGTACCTCTGCATTTGCTGT  | NM_178320.2     | Moriel et al. (2014)    |
|                                           | Reverse: ATGCAATCTGCATTTGCTGT  |                 |                         |
| **LM samples**                             |                          |                 |                         |
| FABP4                                      | Forward: AACTCTAGTAAGTGCTCTGAG  | AJ4160220       | Li et al. (2011)        |
|                                           | Reverse: CATAAACTCTGGCTGGAGTGA  |                 |                         |
| Myogenin                                   | Forward: GAGAACGGCAGTCAAGAGGAGGATTGA  | AF09174         | Muroya et al. (2002)    |
|                                           | Reverse: TCTGAGTGCTGCGGAGGATGTC  |                 |                         |
| PAX7                                       | Forward: GGGTACCTCTGCCGATGGCTCAG  | XM_616352.6     | Moriel et al. (2014)    |
|                                           | Reverse: GCTTCCTTGTCGATGTCAG  |                 |                         |
| PPAR-γ                                    | Forward: GCATTTCACCTGCCCATCTTGT  | AY137204        | Li et al. (2011)        |
|                                           | Reverse: GGGATACAGGCTCCACTTTG  |                 |                         |
| B-actin                                    | Forward: AGCAAGCAGAGTACAGATGACT  | NM_173979       | Bong et al. (2012)      |
|                                           | Reverse: ATCCAGCAGTCTGCATGCA    |                 |                         |
| Ribosomal protein L12                      | Forward: CCTCAGACCATCGTGTACAG  | AF479289        | Jeong et al. (2012)     |
|                                           | Reverse: CCTCCAGACCTCACCTCTTCTGTC  |                 |                         |
Weaning 5.26 5.02 0.06 <0.01
Late gestation 4.14 4.15 0.07 0.93
BCS
Weaning 463 469 6 0.48
Late gestation 428 443 5 0.04

(Sturtz et al., 2001; Miao and St. Clair, 2009). Hepatic oxidative damage by eliminating superoxide anion radicals

enzyme that uses Cu or Zn as cofactors and protects cells from
dictate be an indicator of Zn bioavailability, given that local Zn levels

hepatic cells (Coyle et al., 2002). Hepatic

liver, with Zn being the primary inductor of its synthesis in

intracellular metal-binding proteins abundantly found in the

expression was negatively associated with Cu intake and the

amounts of supplemental Cu during the experiment, and

highly speculatory. Both AAC and INR cows consumed similar

biological or experimental explanations would ultimately be

on each biopsy date; however, attempts to identify definitive biological or experimental explanations would ultimately be

highly speculative. Both AAC and INR cows consumed similar amounts of supplemental Cu during the experiment, and

marginal Cu levels noted in AAC cows should not be associated with inadequate intake of this trace mineral during gestation.

No treatment differences were detected (P ≥ 0.18; Table 6) for liver mRNA expression of Cu-transporter protein (CUT),

metallothionein 1A (MT), and copper-zinc-superoxide dismutase 1 (SOD) prior to treatment administration (day −30) or during

late gestation (day 97). At calving, MT expression was greater in AAC vs. INR cows (P = 0.04), but no treatment differences were detected for mRNA expression of CUT or SOD (P ≥ 0.19; Table 6). These genes are associated with Cu and Zn metabolism in the liver and provide additional insight into Zn and Cu status of AAC and INR cows during the experiment. The CUT is associated with Cu transport into hepatic cells and distribution of Cu into cellular organelles (Prohaska and Gybina, 2004; Han et al., 2009), and its mRNA expression partially modulated by liver Cu (Bauerly et al., 2005; Fry et al., 2013). The MT is a superfamily of intracellular metal-binding proteins abundantly found in the liver, with Zn being the primary inductor of its synthesis in hepatic cells (Coyle et al., 2002). Hepatic MT expression may be an indicator of Zn bioavailability, given that local Zn levels dictate MT mRNA levels (Wang et al., 2012). The SOD is an enzyme that uses Cu or Zn as cofactors and protects cells from oxidative damage by eliminating superoxide anion radicals (Sturtz et al., 2001; Miao and St. Clair, 2009). Hepatic SOD mRNA expression was negatively associated with Cu intake and the

Table 4. Performance responses of beef cows that received diets containing supplemental INR (n = 95) or AAC (n = 95) during gestation.

| Item                          | INR  | AAC  | SEM  | P-value |
|-------------------------------|------|------|------|---------|
| Cow age, yr                   | 4.47 | 4.60 | 0.25 | 0.71    |
| Days receiving treatments, d  | 167  | 166  | 3    | 0.91    |
| BW, kg                        | Initial | 485  | 496  | 8      | 0.35    |
|                              | Late gestation | 428  | 443  | 5      | 0.04    |
|                              | Calving        | 419  | 437  | 12     | 0.31    |
|                              | Weaning         | 463  | 469  | 6      | 0.48    |
| BCS                           | Initial | 5.44 | 5.51 | 0.07   | 0.49    |
|                              | Late gestation  | 4.14 | 4.15 | 0.07   | 0.93    |
|                              | Calving         | 4.50 | 4.54 | 0.07   | 0.73    |
|                              | Weaning         | 5.26 | 5.02 | 0.06   | <0.01   |

1INR and AAC cows received the same amount of supplemental Co, Cu, Mn, and Zn from sulfate sources or Availa 4 (Zinpro Corporation, Eden Prairie, MN). Cows were assigned to the experiment at 117 ± 2 d of gestation (day 0).

2BW and BCS (Wagner et al., 1988) were recorded before the beginning of the experiment (day −30; initial), on day 97 (late gestation), upon calving, and at weaning (day 367).

Table 5. Liver concentrations of Co, Cu, Mn, and Zn in beef cows that received diets containing supplemental INR (n = 95) or AAC (n = 95) during gestation.

| Item                          | INR  | AAC  | SEM  | P-value |
|-------------------------------|------|------|------|---------|
| Co, mg/kg                     | Initial | 0.165 | 0.155 | 0.009 | 0.44    |
|                              | Late gestation | 0.586 | 0.678 | 0.024 | 0.01    |
|                              | Calving       | 0.584 | 0.550 | 0.114 | 0.83    |
|                              | Early lactation | 0.140 | 0.154 | 0.009 | 0.31    |
| Cu, mg/kg                     | Initial | 7.40  | 6.69  | 11.0   | 0.65    |
|                              | Late gestation | 125  | 81.9  | 12.9   | 0.02    |
|                              | Calving       | 118  | 42.9  | 9.9    | <0.01   |
|                              | Early lactation | 136  | 154   | 15     | 0.39    |
| Mn, mg/kg                     | Initial | 8.94  | 8.58  | 0.29   | 0.39    |
|                              | Late gestation | 10.5 | 10.0  | 0.3    | 0.30    |
|                              | Calving       | 12.0 | 10.6  | 0.7    | 0.17    |
|                              | Early lactation | 8.40 | 7.48  | 0.47   | 0.18    |
| Zn, mg/kg                     | Initial | 148   | 139   | 7      | 0.40    |
|                              | Late gestation | 307  | 341   | 21     | 0.27    |
|                              | Calving       | 173  | 129   | 11     | 0.01    |
|                              | Early lactation | 139  | 155   | 9      | 0.21    |

1INR and AAC cows received the same amount of supplemental Co, Cu, Mn, and Zn from sulfate sources or Availa 4 (Zinpro Corporation, Eden Prairie, MN). Cows were assigned to the experiment at 117 ± 2 d of gestation (day 0).

2Liver samples were collected before the beginning of the experiment (day −30; initial), on day 97 (late gestation), upon calving when cows were 43 ± 0.5 d postpartum (early lactation), and at weaning (day 367) via needle biopsy (Arthington and Corah, 1995).

Concentrations of Co, Cu, Mn, and Zn were determined by the Michigan State University Diagnostic Center for Population and Animal Health (East Lansing, MI; Braselton et al., 1997). Samples were collected from 30 cows of each treatment randomly selected on each sampling day.

overall status in cattle (Hansen et al., 2009). In this experiment, AAC cows had greater mRNA expression of MT and similar mRNA expression of CUT and SOD compared with INR cows at calving, despite having less liver Cu and Zn concentration. These outcomes suggest that Cu and Zn status between AAC and INR cows may have not differed during gestation and upon calving as denoted by liver concentrations of these trace minerals.

Both AAC and INR cows were in early lactation and with similar (P = 0.69) days postpartum at the time of WSW (Table 7). No treatment differences were detected (P ≥ 0.19) for milk production as well as milk Co, Cu, Zn, and Mn concentrations (Table 7). Liver concentrations of these trace minerals the day after WSW also did not differ (P ≥ 0.18) between treatments (Table 5). Pregnancy rates to AI or overall (AI + bull breeding) did not differ (P ≥ 0.41) between INR and AAC cows (46.2% and 40.3% overall, SEM = 5.0; respectively). At weaning, no treatment differences were detected for cow BW (P = 0.31), while BCS was greater (P < 0.01) in INR compared with AAC cows (Table 4). Collectively, the lack of major treatment differences in measures obtained from WSW until weaning can be attributed to the similar nutritional management that AAC and INR cows received after calving, including inorganic trace mineral supplementation. Treatment differences in BCS at weaning do not follow this rationale, although both INR and AAC cows were within the threshold for adequate BCS in beef females (Richards et al., 1986). Nonetheless, similar milk production and
profile between AAC and INR cows indicate that any potential treatment effects on offspring responses were not caused by altered milk yield during early lactation. Hence, supplementing AAC or INR to gestating beef cows did not impact their post-calving lactation and reproductive performance, corroborating the results from previous research in this area (Stanton et al., 2000; Muehlenbein et al., 2001; Marques et al., 2016a).

Calf birth to weaning responses

No treatment differences were detected (P ≥ 0.51; Table 8) for calving rate and calf birth BW. No treatment differences were also detected (P ≥ 0.21) for the proportion of calves born to AI or proportion of male calves born (Table 8), whereas sex and genetic merit impact offspring developmental responses (Koger and Knox, 1945). Previous research also reported similar calf birth BW when supplementing gestating beef cows with inorganic or organic sources of trace minerals (Stanton et al., 2000; Sprinkle et al., 2006; Marques et al., 2016a). Hence, fetal weight was likely similar between AAC and INR cows despite differences detected for liver concentrations of Co, Zn, and Cu during gestation.

No treatment differences were detected (P ≥ 0.61) for Co, Cu, Mn, or Zn concentrations in the placental cotyledons (Table 9). No treatment differences were also detected (P ≥ 0.51) for liver Co, Cu, Mn, or Zn concentrations in calves at birth or 24 h after birth (Table 9) or concentrations of these trace minerals in the colostrum (P ≥ 0.48; Table 7). The fetus relies completely on liver concentrations of Co, Zn, and Cu as organic-complexed instead of sulfate sources. INR to gestating beef cows did not yield immediate effect on fetal trace mineral profile, but resulted in reduced Mn for 24 h later (Table 10), which suggest similar maternal Cu and Zn metabolism during early life (Coyle et al., 2002; Han et al., 2009; Hansen et al., 2009). Day effects were also observed (P < 0.01) for calf liver mRNA expression of CUT and MT, both of which increased 24 h after birth across treatments reflecting the greater activity in hepatic tissue and utilization of trace minerals in the neonate (Woodward and Suttle, 1999; López-Alonso et al., 2005). No treatment differences were detected (P ≥ 0.29) for mRNA expression of LM genes associated with adipogenic or muscle development activities at birth (Table 11). Myogenin is a regulatory factor that influences postnatal muscle growth through differentiation and fusion of satellite cells with existing fibers (Le Grand and Rudnicky, 2007; Du et al., 2010). These cells, both quiescent and activated, are marked by the expression of paired box gene 7 (PAX7), which is necessary for satellite cell specification and survival (Seale et al., 2000; Li et al., 2011). Trace minerals, such as Zn, are required for muscle differentiation through the activation and proliferation of satellite cells (Petrie et al., 1996; Ohashi et al., 2015). Peroxisome proliferator-activated receptor gamma (PPAR-γ) plays a pivotal role in adipocyte differentiation and associated gene expression (Housknecht et al., 2002), the process of which is initiated around mid-gestation in the ruminant (Du et al., 2010). Adipocyte fatty acid-binding protein (FABP4) is a target of PPAR-γ and highly involved in the differentiation of adipocytes by acting as an intracellular fatty acid chaperone (Michal et al., 2006). Several Zn-finger proteins participate in adipocyte determination and differentiation (Wei et al., 2013), while Zn has been shown to enhance adipogenesis in vitro (Tanaka et al., 2001). Despite the established role of trace minerals on muscle development or adipogenesis, supplementing gestating cows with Cu, Co, Mn, or Zn as organic-complexed instead of sulfate sources did not alter the mRNA expression of genes associated with these activities in the LM of neonatal calves.

No treatment differences were detected (P ≥ 0.44) for liver concentrations of Co, Cu, Mn, and Zn in calves the day after WSW (early life; Table 9). These results agree with the hepatic trace mineral profile of newborn calves, trace mineral profile of maternal milk at WSW, and similar nutritional management that cow–calf pairs received after calving. At weaning, no treatment differences were detected (P ≥ 0.18) for weaning rate, proportion of AI-sired and male calves weaned, weaning age, and calf weaning BW (Table 8). Liver concentrations of Co, Cu, Mn, and Zn also did not differ (P ≥ 0.17) between treatments (Table 9). Hepatic mRNA expression of SOD at weaning was greater (P = 0.04) in calves from AAC cows compared with INR cohorts, while no treatment differences (P ≥ 0.26) were detected in the hepatic mRNA expression of CUT and MT (Table 10) nor mRNA expression of FABP4, myogenin, PAX7, and PPAR-γ in the LM (Table 11). These outcomes suggest that supplementing AAC instead of INR to gestating beef cows did not yield immediate and longer-term effects in hepatic trace mineral profile and LM expression of adipogenic and muscle development genes.

Table 6. Expression of liver genes in beef cows that received diets containing supplemental INR (n = 95) or AAC (n = 95) during gestation.1,2

| Item          | INR   | AAC   | SEM  | P-value |
|---------------|-------|-------|------|---------|
| **CUT**       |       |       |      |         |
| Initial (−30) | 1.86  | 1.43  | 0.21 | 0.18    |
| Pre-calving (day 97) | 1.67  | 1.70  | 0.08 | 0.82    |
| Calving       | 1.95  | 1.75  | 0.11 | 0.19    |
| **MT**        |       |       |      |         |
| Initial (−30) | 3.39  | 2.38  | 0.73 | 0.36    |
| Pre-calving (day 97) | 26.1  | 22.8  | 3.8  | 0.54    |
| Calving       | 36.4  | 65.6  | 11.6 | 0.04    |
| **SOD**       |       |       |      |         |
| Initial (−30) | 2.29  | 2.76  | 0.27 | 0.24    |
| Pre-calving (day 97) | 1.99  | 2.11  | 0.13 | 0.51    |
| Calving       | 2.02  | 2.04  | 0.13 | 0.90    |

1INR and AAC cows received the same amount of supplemental Co, Cu, Mn, and Zn from sulfate sources or Avail 4 (Zinpro Corporation, Eden Prairie, MN). Cows were assigned to the experiment at 117 ± 2 d of gestation (day 0).
2Liver samples were collected via needle biopsy (Arthington and Corah, 1995) before the beginning of the experiment (day −30; initial), on day 97 (late gestation), and upon calving. Values are expressed as relative fold change compared within CT of reference genes analyzed within the same sample (Ocón-Grove et al., 2008).
resulting in similar calf growth until weaning. The upregulated hepatiic mRNA expression of SOD in calves from AAC may have impacted postweaning responses, which are discussed in the companion manuscript (Harvey et al., 2021).

Marques et al. (2016a) also did not report differences in weaning responses between cows supplemented with INR or AAC during late gestation, despite a 13-kg numerical increase in weaning BW from AAC cows. This experiment was designed with a greater number of cows and providing AAC and INR during a longer period during gestation to expand on the results reported by Marques et al. (2015a) and-wow. One can speculate that cow BCS loss during gestation observed herein may have limited the benefits of the AAC treatment, given the importance of cow nutritional status to postnatal offspring development (Bohnet et al., 2013; Marques et al., 2016b). As an attempt to address this question, AAC and INR cows were divided according to BCS change during gestation (day −30 to 97). Calf weaning BW also did not differ (P ≥ 0.51) between AAC and INR cows (n = 129) that lost a significant amount of BCS (−0.42 and −0.48 BCS change, SEM = 0.08; 176 vs. 178 kg of BW, SEM = 3; respectively) or cows (n = 43) whose BCS change was below the sensitivity of the BCS scale (−0.42 and −0.48 BCS change, SEM = 0.08; 176 vs. 178 kg of BW, SEM = 6; respectively). Therefore, the lack of differences in offspring productivity between AAC and INR cows is likely not associated with the unexpected cow BCS loss during gestation observed in this experiment.

### Overall conclusions

Supplementing beef cows with AAC during gestation did not improve liver concentrations of these trace minerals nor enhanced the transport of these trace minerals to the fetus or offspring. However, the results support the potential benefits of the AAC treatment, given the importance of cow nutritional status to postnatal offspring development.
Table 10. Expression of liver genes in calves born from beef cows that received diets containing supplemental INR (n = 95) or AAC (n = 95) during gestation1,2

| Item  | INR     | AAC     | SEM   | P-value |
|-------|---------|---------|-------|---------|
| CUT   |         |         |       |         |
| Birth | 2.19    | 2.23    | 0.11  | 0.78    |
| 24 h after birth | 2.51    | 2.53    | 0.11  | 0.91    |
| Weaning | 2.13    | 2.01    | 0.10  | 0.40    |
| MT    |         |         |       |         |
| Birth | 33.7    | 32.9    | 7.2   | 0.94    |
| 24 h after birth | 59.4    | 68.4    | 7.2   | 0.38    |
| Weaning | 39.9    | 54.1    | 8.8   | 0.26    |
| SOD   |         |         |       |         |
| Birth | 2.92    | 2.96    | 0.20  | 0.88    |
| 24 h after birth | 2.77    | 2.70    | 0.20  | 0.79    |
| Weaning | 1.78    | 1.98    | 0.07  | 0.04    |

1INR and AAC cows received the same amount of supplemental Cu, Mn, and Zn from sulfate sources or Availa 4 (Zinpro Corporation, Eden Prairie, MN). Cows were assigned to the experiment at 117 ± 2 d of gestation (day 0).

2Liver samples were collected via biopsy (Arthington and Corah, 1995) at birth, 24 h after birth, and at weaning (day 367). Values are expressed as relative fold change compared within CT of reference genes analyzed within the same sample (Ocón-Grove et al., 2008). Samples were collected from 30 cows of each treatment randomly selected on each sampling day.

Table 11. Expression of LM genes in calves born from beef cows that received diets containing supplemental INR (n = 95) or AAC (n = 95) during gestation1,2

| Item  | INR     | AAC     | SEM   | P-value |
|-------|---------|---------|-------|---------|
| PPAR4 |         |         |       |         |
| Birth | 155.7   | 108.0   | 31.4  | 0.29    |
| Weaning | 45.9    | 38.4    | 9.0   | 0.56    |
| Myogenin |         |         |       |         |
| Birth | 5.05    | 4.54    | 0.58  | 0.53    |
| Weaning | 4.20    | 4.42    | 0.51  | 0.77    |
| PAX7  |         |         |       |         |
| Birth | 3.42    | 3.42    | 0.25  | 0.99    |
| Weaning | 3.23    | 3.22    | 0.16  | 0.98    |
| PPAR-γ |         |         |       |         |
| Birth | 4.28    | 4.24    | 0.76  | 0.97    |
| Weaning | 2.14    | 2.03    | 0.21  | 0.71    |

1INR and AAC cows received the same amount of supplemental Cu, Mn, and Zn from sulfate sources or Availa 4 (Zinpro Corporation, Eden Prairie, MN). Cows were assigned to the experiment at 117 ± 2 d of gestation (day 0).

2Muscle samples were collected via needle biopsy (Schubach et al., 2019) at birth and at weaning (day 367). Values are expressed as relative fold change compared within CT of reference genes analyzed within the same sample (Ocón-Grove et al., 2008). Samples were collected from 30 cows of each treatment randomly selected on each sampling day.

colostrum compared with cohorts supplemented with sulfate sources. No major physiological and productive benefits were realized from cows receiving organic-complexed trace minerals, besides upregulated mRNA expression of hepatic genes associated with Cu and Zn metabolism at calving and in their offspring at weaning. The results from this manuscript indicate that supplementing Co, Cu, Zn, and Mn as organic-complexed or sulfate sources to beef cows during the last 5 mo of gestation yielded similar cow–calf productive responses until weaning. The companion manuscript (Harvey et al., 2021) describes the postweaning responses of the female and male offspring reared, respectively, as replacement heifers and feeder cattle.

Acknowledgments

Financial support for this research was provided by Zinpro Corporation (Eden Prairie, MN). K.M.H. was funded as Tom Slick Graduate Research Fellow at Texas A&M University. B.R. was supported by Coordenação de Aperfeiçoamento de Pessoal de Nível Superior, Brazil #88887.366406/2019-00. A.P.B. was supported by CAPES, Brazil (#88881.128327/2016-01).

Conflict of interest statement

J.R.R. is employed by the funder of this project (Zinpro Corporation, Eden Prairie, MN) and contributed to research design and data interpretation. However, the principal investigator (R.F.C.) and all other authors of this manuscript have no additional conflict of interest to report.

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