Brief communication

Dietary supplementation with safflower seeds differing in fatty acid composition differentially influences serum concentrations of prostaglandin F metabolite in postpartum beef cows

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Abstract – Synthesis and secretion of prostaglandin F2α (PGF2α) is elevated following parturition and exerts divergent effects on the re-establishment of fertile estrous cycles in cows. The objective of these experiments was to determine if oil seed supplements differing in fatty acid composition differentially influence serum concentrations of the specific PGF2α metabolite, PGFM. Safflower seed supplements were formulated to provide 5% of dry-matter intake as fat. In Trial 1, 24 multiparous beef cows were individually fed control (beet pulp-soybean meal) or cracked high-linoleate safflower seed (78% 18:2n-6) supplements for 80 d postpartum. Linoleate supplemented cows had greater (P < 0.001) serum concentrations of PGFM than control cows. In Trial 2, primiparous beef cows (n = 36) were individually fed control (cracked corn-soybean meal), cracked high-linoleate (76% 18:2n-6) or -oleate (72% 18:1n-9) safflower seed supplements for 92 d postpartum. As in Trial 1, serum concentrations of PGFM were greater (P ≤ 0.04) in linoleate than control or oleate supplemented cows. Serum concentrations of PGFM, however, did not differ (P = 0.40) among oleate and control supplemented cows. Although potential impacts on reproductive performance remain to be proven, dietary oil supplements high in linoleate, but not oleate, increased serum concentrations of PGFM compared to control supplements.

beef cows / lipid supplementation / prostaglandin

1. INTRODUCTION

Feeding oil supplements to cattle is an approach used to enhance the amount of energy contained in diets. In addition to helping meet energy demands, supplemental lipids influence reproductive performance in bovine females. Supplementing cows with dietary lipid increased concentrations of serum lipids [1, 2], altered ovarian follicular development [2, 3], and prostaglandin (PG) synthesis [4–6]. In late–pregnant ewes, diets high in linoleic acid increased plasma concentrations of arachidonic acid [7] and production of PG from endometrial tissue [7, 8].
Specific fatty acid content of lipid supplements affects ruminal biohydrogenation and duodenal flow of fatty acids [9]. Thus, the apparent availability of fatty acids for metabolism is affected by the source of the dietary fat. Linoleic acid that escapes ruminal degradation and is subsequently absorbed from the digestive tract can be desaturated and elongated to form arachidonic acid, a direct precursor of PGF$_{2\alpha}$ [10]. The objectives of these experiments were to determine if supplemental safflower-seeds high in linoleic acid affect serum concentrations of the PGF$_{2\alpha}$ metabolite, PGFM, and if safflower seed supplements differing in fatty acid content differentially influence serum concentrations of PGFM in beef cows. It was hypothesized that the fatty acid content of oil seed supplementation during the postpartum period may differentially affect prostaglandin synthesis.

2. MATERIALS AND METHODS

Animal care and treatments were approved by the University of Wyoming Animal Care and Use Committee.

2.1. Trial 1: animals

Twenty-four spring-calving multiparous Angus × Gelbvieh rotationally-crossed beef cows (473.9 ± 9.2 kg) were blocked by weight and body condition score (4.5; 1 = emaciated, 9 = obese) [11] and randomly allocated to one of two dietary treatments within 24 h of parturition. The cows were maintained in pens (6 m × 20 m) and provided rations in individual feeding stations twice daily, at 0700 and 1600, for 80 d postpartum. The diets were formulated to provide equal amounts of protein and energy using nutrient requirements for 36-mo, 499 kg lactating cows producing 9.1 kg of milk per d during peak lactation [12]. Supplements consisted of liquid molasses with the addition of beet pulp-soybean meal (control) or cracked high-linoleate safflower seeds (77.7% 18:2n-6, Tab. I; n = 12 per group). The quantity of feedstuffs provided was based on initial cow body weight, calculated as the average of weights obtained on the first and second days of the experiment. Nutrient analysis of the supplements and hay is reported in Table II. In order to provide an equal quantity of energy, control and linoleate supplemented cows were fed the supplement at 0.5% and 0.3% of their body weight, respectively. Both groups were fed chopped (Haybuster, Model 2620, Duratech Industries, Jamestown, ND) native grass hay at 2.1% of their body weight. The supplement was fed first with hay provided after the supplement was completely consumed. Cows remained in their stations for approximately 2 h, or until consumption ceased. All animals had ad libitum access to trace mineralized salt and water.

2.2. Trial 2: animals

Thirty-six spring-calving primiparous Angus × Gelbvieh rotationally-crossed beef cows (487.9 ± 10.5 kg; 5.5 body condition score) [11] were blocked by d of parturition and randomly assigned to one of three dietary supplement treatments beginning 3 d postpartum. Primiparous cows were assigned to 1 of 6 pens (6 m × 20 m) equipped with individual feeding stations, large round bale

Table I. Fatty acid content (% total fatty acids-as fed) of supplemental L-Trial 1 (Linoleic, Trial 1) C (Control, Trial 2), L (Linoleic, Trial 2), O (Oleic, Trial 2), diets$^a$.

<table>
<thead>
<tr>
<th>Fatty Acid</th>
<th>L-Trial 1</th>
<th>C</th>
<th>L</th>
<th>O</th>
</tr>
</thead>
<tbody>
<tr>
<td>16:0</td>
<td>6.7</td>
<td>13.0</td>
<td>6.4</td>
<td>5.2</td>
</tr>
<tr>
<td>18:0</td>
<td>2.1</td>
<td>2.6</td>
<td>2.6</td>
<td>1.7</td>
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<tr>
<td>18:1n-9</td>
<td>6.3</td>
<td>25.9</td>
<td>11.2</td>
<td>72.2</td>
</tr>
<tr>
<td>18:2n-6</td>
<td>77.7</td>
<td>48.5</td>
<td>75.7</td>
<td>18.5</td>
</tr>
<tr>
<td>18:3</td>
<td>0.3</td>
<td>2.4</td>
<td>0.2</td>
<td>0.2</td>
</tr>
<tr>
<td>Others</td>
<td>6.9</td>
<td>7.6</td>
<td>3.9</td>
<td>2.2</td>
</tr>
</tbody>
</table>

$^a$ Control diet for Trial 1 consisted of Beet Pulp with < 1% ether extract and therefore not a significant source of fatty acids.
self-feeders, and heated watering troughs. Supplements were fed once daily from d 3 through d 92 postpartum. At 0630, the dams were separated from their calves and locked into feeding stations. The supplements consisted of dried molasses with the addition of cracked corn and soybean meal (control), cracked high-linoleate (76% 18:2n-6, Tab. I), or cracked high-oleate (72% 18:1n-9, Tab. I) safflower seeds (n = 12 per group). Safflower seed supplements were formulated to provide 5% of dry matter intake as fat. Estimates for hay intake and cow performance were based on Alderton et al. [13]. By using estimated hay intake, the supplements were formulated to contain equal amounts of protein and energy and meet NRC [12] nutritional requirements for a 24-mo, 499 kg lactating cow (565 kg mature BW) producing 9.1 kg milk·d⁻¹ during peak lactation. Tabular values for TDN were utilized to formulate supplements [14]. Nutrient analysis of supplements and hay are reported in Table II. To ensure an equal quantity of energy, the amounts of supplement fed were 1.46 kg to control, 1.62 kg to linoleate, and 1.43 kg to oleate supplemented cows. All animals had ad libitum access to hay, trace mineralized salt, and water. Forage intake did not differ among supplemental treatments [15].

### Table II. Nutrient analysis (% DM) of control (C), linoleate (L), and oleate (O) supplements and hay fed during Trial 1 and Trial 2.

<table>
<thead>
<tr>
<th></th>
<th>Trial 1</th>
<th></th>
<th></th>
<th>Trial 2</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C</td>
<td>L</td>
<td>O</td>
<td>Hay</td>
<td>C</td>
<td>L</td>
</tr>
<tr>
<td>ADF</td>
<td>9.76</td>
<td>31.20</td>
<td>34.73</td>
<td>47.17</td>
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<td>27.14</td>
</tr>
<tr>
<td>NDF</td>
<td>22.39</td>
<td>42.82</td>
<td>45.06</td>
<td>75.71</td>
<td>38.55</td>
<td>42.68</td>
</tr>
<tr>
<td>CP</td>
<td>18.51</td>
<td>20.14</td>
<td>17.78</td>
<td>7.75</td>
<td>12.63</td>
<td>16.55</td>
</tr>
<tr>
<td>Fatb</td>
<td>3.80</td>
<td>31.36</td>
<td>29.32</td>
<td>1.11</td>
<td>0.55</td>
<td>27.14</td>
</tr>
<tr>
<td>IVOMDC</td>
<td>91.53</td>
<td>48.12</td>
<td>40.22</td>
<td>42.20</td>
<td>74.71</td>
<td>56.42</td>
</tr>
</tbody>
</table>

a Control supplement was corn-soybean (Trial 1) and soybean-beet pulp (Trial 2).
b Fat content of supplement is reported as Total Fatty Acids (Trial 1) and Ether Extract (Trial 2).
c IVOMD = in vitro organic matter digestibility. Fatty acids are largely undigested by ruminal microbes but highly digested postruminally; therefore, diets were formulated to have equal total quantities of TDN (NRC, 1982).

#### 2.3. Collections

Twice weekly blood samples for the analysis of serum concentrations of PGFM were collected via venipuncture of the coccygeal vein into vacuum tubes (Becton Dickinson Company, Franklin Lakes, NJ) beginning at d 25 through the completion of each feeding trial. Blood was allowed to clot overnight at 4 °C. Serum was separated by centrifugation (700 × g), and stored at −20 °C until analysis.

Additional blood samples were collected over a 4-d period starting at d 30, d 60, and d 90 postpartum for determination of plasma fatty acids in Trial 2. In order to partially account for diurnal variation, blood samples were collected four times during the AM and PM hours at each collection period. Collection frequencies in relation to feeding were 0, 3, 6, 9, 12, 15, 18, and 21 h with 0 representing immediately prior to supplementation. All blood samples were collected from the coccygeal vein via venipuncture using 10 mL EDTA coated Vacutainer (Becton Dickinson Company, Franklin Lakes, NJ) tubes. Blood was centrifuged at 4 °C at 700 × g within 30 min of collection. Plasma was pooled by sampling period, immediately frozen at −20 °C, freeze dried, and stored in plastic vials under nitrogen gas at −80 °C until analysis. Total lipids were
extracted from duplicate 500 mg samples of freeze-dried blood plasma. Fatty acid methyl esters were prepared using 14% BF₃ in methanol to insure methyl esterification of esterified and non-esterified fatty acids [16].

2.4. Laboratory analysis

Serum concentrations of PGFM were quantified in extracts of serum by RIA [17] and inferred concentrations of PGF₂α [18]. In brief, 2 mL aliquots of serum were acidified with 150 µL of 1.0 M formic acid and extracted twice with 5 mL of ethyl acetate. The combined extracts were dried under N₂ and reconstituted in 1 mL of gel-PBS. When necessary, the samples were further diluted with gel-PBS before they were assayed. Recovery of [³H]-PGFM was 94.5% [17]. Antigen (PGFM-tyrosine methyl ester) was labeled with I¹²⁵ using chloramine T and separated on Sephadex G-25. Rabbit anti-PGFM at a working dilution of 1:24 000 was utilized with a cross-reactivity of ~1% to PGE₂. No other significant cross-reactivity of PG are noted with this antibody [17]. Samples from Trial 1 were analyzed in a single assay with an intra-assay CV of 8.0%. Samples from Trial 2 were analyzed in three assays with intra- and inter-assay CV of 10.8% and 16.5%, respectively.

Isolated plasma fatty acid methyl esters (Trial 2) were separated using a Hewlett-Packard 5890 GLC (Hewlett-Packard, Avondale, PA) equipped with a flame ionization detector and a 100 m × 0.25 mm (i.d.) fused silica capillary column (SP-2560, 0.2 µm film thickness, Supelco, Bellefonte, PA). Oven temperature was maintained at 175 °C for 40 min, and then increased to 240 °C at 10 °C per min. Injector and detector temperatures were 275 °C. Helium was the carrier gas with a split ratio of 50:1 and 0.8 mL·min⁻¹ column flow. Fatty acid peaks were recorded and integrated using a Hewlett-Packard 3396 integrator. Fatty acids were identified by comparing retention times of peaks with fatty acid methyl ester standard (Nu-Chek Prep, Inc., Elysian, MN and Matreya, Inc., Pleasant Gap, PA).

2.5. Statistical analysis

Concentrations of PGFM and relative weight percentages of fatty acids were analyzed as a split-plot using GLM procedures of SAS (Ver. 8.0, SAS Inst. Inc., Cary, NC) with treatment as the main effect, and day and treatment by day interactions tested as sub-plot effects. Animal within treatment was the error term for treatment effects. Differences among means were separated using Fisher protected LSD [19]. Type III sums of squares were used, and least squares means and associated standard errors are reported.

3. RESULTS

Serum concentrations of PGFM in the multiparous cows used in Trial 1 were greater (P < 0.001) in cows consuming high-linoleate safflower seed supplement (469 ± 24 pg·mL⁻¹) than in cows consuming control supplement (328 ± 23 pg·mL⁻¹, Fig. 1). Significant day (P = 0.12) or treatment by day interactions (P = 0.83) were not detected.

In contrast, serum concentrations of PGFM in the primiparous cows used in
Trial 2 were influenced by day ($P < 0.001$) and declined from 600 ± 33 pg·mL$^{-1}$ at d 25 to 363 ± 42 pg·mL$^{-1}$ by 92 d postpartum in all cows (Fig. 2). As in Trial 1, the concentrations of PGFM were influenced by treatment ($P = 0.01$) and were greater ($P \leq 0.04$) in cows fed high-linoleate safflower seeds (647 ± 62 pg·mL$^{-1}$) than cows fed either high-oleate safflower seeds (371 ± 68 pg·mL$^{-1}$) or control supplement (452 ± 68 pg·mL$^{-1}$). Treatment by day interactions were not detected ($P = 0.13$).

In Trial 2, relative weight percentages of plasma linoleic ($P < 0.001$), oleic ($P < 0.001$) and arachidonic ($P = 0.04$) acid differed by treatment. Relative plasma concentration of linoleic acid was the greatest ($P < 0.001$) in linoleic-supplemented cows (29.2 ± 0.6%) compared with cows fed oleic (18.6 ± 0.6%) or control (21.2 ± 0.6%) supplements. Cows supplemented with high-oleic safflower seeds had the highest ($P < 0.001$) relative concentrations of plasma oleic acid (21.5 ± 0.4%) compared with linoleic- (10.5 ± 0.4%) or control- (13.4 ± 0.4%) supplemented cows. The relative weight percentage of plasma arachidonic acid was low in all cows; however, differences ($P = 0.04$) were noted among treatments. Relative plasma concentrations of arachidonic acid were greater ($P = 0.01$) in linoleic-supplemented cows (1.5 ± 0.1%) than in control cows (1.2 ± 0.1%). Cows supplemented with high-oleic safflower seeds had intermediate relative plasma concentrations (weight %) of arachidonic acid (1.4 ± 0.1%) that did not differ ($P \geq 0.1$) from either control- or linoleic-supplemented cows.

4. DISCUSSION

Uterine production of PGF$_{2\alpha}$ is important during the early postpartum period. Synthesis and secretion of PGF$_{2\alpha}$ can decrease the number of days to complete uterine involution and hence, the length of postpartum infertility [20]. Early pregnancy is associated with an attenuation of PG synthesis and secretion that may be partially caused by limited substrate availability [21]. Increases in substrate availability from diets high in PG precursors during early pregnancy could increase production of PGF$_{2\alpha}$ and contribute to early luteolysis and embryo mortality [10]. Increased plasma concentrations of PGFM were noted in cows supplemented with high-linoleic safflower seeds in Trial 1. In Trial 2, fatty acid composition of fat supplements differentially influenced plasma concentrations of fatty acids and serum concentrations of PGFM. Although the relative weight percentage of arachidonic acid was low in all cows, cows supplemented with high-linoleate safflower seeds had increased plasma concentrations of arachidonic acid over cows fed the control supplement. Increases in plasma arachidonic acid were also noted in ewes fed a diet high in linoleic acid [7]. Relative plasma concentrations of arachidonic acid were similar in oleic and linoleate supplemented cows possibly due to the amount of linoleic acid present in the high-oleic safflower seed supplement. Although relative weight % of plasma arachidonic acid was low compared to plasma concentrations of oleic and linoleic acid, these levels may be representative.
Plasma concentrations of PGFM were also greater in linoleate-supplemented cows than in cows fed oleate- or control-supplements. Similarly, Brahman cows supplemented with 530 g·d⁻¹ lipid from rice bran containing approximately equal proportions of oleic and linoleic acid tended to have increased concentrations of serum PGFM [4]. Likewise, beef cows fed 230 g·d⁻¹ calcium soaps of fatty acids containing 40% oleic and 9% linoleic acid also had increased concentrations of PGFM early in the postpartum period [5].

Total flow and postruminal disappearance of unsaturated fatty-acids, especially linoleic acid, were the greatest in cattle fed high-linoleate safflower seeds [9] presumably leading to an increase in milk linoleic acid [22]. Cis-linoleic acid (18:2n-6) can be desaturated and elongated to form dihomo-γ-linolenic acid which serves as an immediate precursor for series one PG or can be desaturated further to arachidonic acid (20:4), an immediate precursor for series two PG [23]. Although not apparent in the present study, linoleic acid may also exert inhibitory effects on PG synthesis by competing with arachidonic acid for binding to cyclooxygenase [23]. Staples et al. [23] suggested the amount of particular fatty acids reaching the target tissues likely influence whether PG synthesis is stimulated or inhibited. Increased concentrations of PGFM in plasma from lipid supplemented cows could be related to increased substrate availability. Scholljegerdes et al. [9] reported an approximate three fold increased availability of intestinal 18:2n-6 for linoleate- compared to control- or oleate-supplemented cows. Moreover, plasma 18:2n-6 greatly increased and 20:4 was the greatest for the linoleate treatment.

Serum concentrations of PGFM decreased with time postpartum in Trial 2 but not in Trial 1. The reasons for this difference are not apparent, but could be related to differences in cow parity [24]. Elevated circulating concentrations of PGFM during the postpartum interval can exert divergent effects on subsequent fertility. High serum concentrations of PGFM early postpartum decreases the time required for uterine involution [20]. Filley et al. [5] reported a negative correlation between the duration of increased PGF₂α production and the duration of the postpartum anestrous period. Conversely, high serum concentrations of PGF₂α later in the postpartum interval may reduce subsequent fertility by increasing the incidence of estrous cycles with abbreviated luteal phases resulting in early embryonic mortality [10, 21, 25].

In conclusion, serum concentrations of PGFM were differentially influenced by dietary oil supplements unique in fatty acid composition. Although potential effects of fat supplementation-induced changes in PGFM on postpartum reproduction have not been proven [26], appropriate supplementation with lipids may provide an alternative method to manipulate PG production during the postpartum interval in order to optimize reproductive performance.

REFERENCES


