Clinical Responses to Intramammary Endotoxin Infusion in Dairy Cows Subjected to Feed Restriction<sup>1</sup>

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**ABSTRACT**

Nonpregnant, midlactation primiparous Holstein cows were fed ad libitum (n = 12) or at 80% of maintenance energy requirements (n = 12) to determine whether feed restriction influences clinical response to endotoxin-induced mastitis. After 2 wk of ad libitum or restricted feeding, one mammary quarter per cow was infused with 100 µg of endotoxin. Within 3 to 6 h of intramammary infusion, endotoxin increased mean rectal temperature, heart rate, and milk somatic cell count and immunoglobulin (IgG) concentration; and decreased blood leukocyte count and rumen motility in both restricted and ad libitum-fed cows. Mean serum and milk tumor necrosis factor-alpha (TNF-α) concentrations showed only modest increases following endotoxin infusion. Restricted fed cows had slightly different acute fever responses and significantly increased heart and respiration rates than ad libitum fed cows. However, feed restriction did not influence mean total leukocyte count, rumen motility, serum TNF-α concentrations or milk IgG and TNF-α concentrations. Thus, results of this study suggest that energy balance does not significantly alter clinical symptoms following acute endotoxin-induced mastitis, at least in midlactation cows. As such, negative energy balance may not underlie the increases in severe coliform mastitis commonly observed in periparturient dairy cows.

(**Key words:** energy balance, mastitis, acute phase response, dairy cow)

**Abbreviation key:** EB = energy balance; LPS = lipopolysaccharide; NEB = negative energy balance; PEB = positive energy balance; TNF-α = tumor necrosis factor-alpha.

**INTRODUCTION**

In dairy cows, bacterial invasion and growth in the udder is most often the cause of mammary inflammation or mastitis. Intramammary infusion of endotoxin can be used as a noninfectious provocative tool to study the inflammatory response in coliform mastitis because many of the clinical signs of intramammary coliform infections arise from the release of endotoxin from the infecting microbes (Jain et al., 1978; Hill et al., 1981; Guidry et al., 1983; Erskine et al., 1989). These clinical signs include increases in body temperature, heart, and respiration rates and reductions in rumen motility, appetite, and milk yield (Lohuis et al., 1988; Eades, 1993). Therefore, clinical coliform mastitis and endotoxin-induced mastitis are as much systemic diseases as they are inflammations of the mammary gland.

Several factors influence the duration and severity of systemic disease caused by IMI. These include the causative pathogen, stage of lactation, age, immune status of the cow, genetics, and nutritional status (Harmon, 1994). Nutritional status may be of particular importance in dairy cows around the time of parturition (Burvenich et al., 1999). Periparturient dairy cows experience multiple changes in physiology and metabolism associated with expulsion of the fetus and lactogenesis. The tremendous increase in energy demand for these processes, in combination with a steady decline in daily feed intake, induces moderate to severe negative energy balance (NEB) in most periparturient dairy cows. Noteworthy, however, is the significant increase in the incidence of severe coliform mastitis at this time. Therefore, a causal link between energy partitioning among tissues and physiological processes, energy intake, and altered disease susceptibility around parturition is likely more than coincidence.

In a previous study, we showed that dietary-induced NEB did not significantly influence general immune
status in otherwise healthy dairy cattle (Perkins et al., 2001). However, Holstein steers were used as the animal model in those studies, and no attempt was made to determine the relevance of imposed NEB to clinical inflammatory diseases in those steers. The objective of the present study was to determine if NEB impacts the acute clinical response to endotoxin-induced mastitis in dairy cows. Nonpregnant, midlactation primiparous Holstein cows fed either ad libitum or at 80% of maintenance requirements and challenged intramammarily with endotoxin were used for this purpose.

MATERIALS AND METHODS

Animals and Diets

Twenty-four nonpregnant, primiparous Holstein cows between 77 and 118 DIM were used as the experimental animals. To be eligible for use in the study, each cow had to exhibit positive energy balance for >2 consecutive weeks and no signs of clinical mastitis or other diseases. Once these facts were established, cows were blocked into groups of two or four based on calving date and availability. Cows within a block were randomly assigned to receive one of two dietary treatments, a positive energy balance diet (PEB; n = 12) or a NEB; (n = 12). Both groups were fed a TMR balanced for 40 kg of daily milk production. On a DM basis, the diet contained 35% corn silage, 14% alfalfa, 14% corn grain, 6% cottonseeds, 7% corn distillers grains, 17% protein supplements, and 7% mineral/vitamin supplement. The diet was 32% NDF and 19% CP. The calculated energy density was 1.74 Mcal of NE/kg, and 35% of the CP was estimated to be undegraded in the rumen. All minerals were fed to meet or exceed requirements of cows fed ad libitum, and trace minerals and vitamins A, D, and E were fed to meet requirements of NEB cows (NRC, 1989). The diet was formulated to contain Se at 0.43 mg/kg and vitamin E at 10 IU/kg. However, the PEB group was fed ad libitum, while the NEB group was fed at a restricted level to ~80% of energy requirements for maintenance and milk production. Cows were fed their respective treatment diets for 14 d before intramammary endotoxin challenge. Calculated energy balance was monitored throughout this 14-d period so the amount of feed offered to the NEB cows could be adjusted twice weekly to maintain feed energy intake at 80% of requirements as milk yield decreased. All cows were implanted with 6 mg of norgestomet (Synchro-Mate B, Sanofi Animal Health, Overland Park, KS) on d 9 of this dietary treatment period to ensure that animals had similar profiles of ovarian hormones during subsequent intramammary endotoxin challenges. Dietary treatments were continued through an additional 2 d after endotoxin challenge, after which the study was concluded.

Energy requirement for maintenance was calculated (NRC, 1989), where requirement = [body weight (kg)^0.75] × [0.08 Mcal/kg]. Energy in milk was determined as in Tyrrell and Reid (1965), where milk energy output (Mcal/day) = [milk weight (lb)] × [41.63 × (% fat) + 24.13 × (% protein) + 21.6 × (% lactose) − 11.72]/1000. The total energy requirement was then calculated as [energy for maintenance + milk energy output]. Calculated energy balance (EB) = [NEE consumed in feed] − [NEE requirement for maintenance and milk]. If EB < 0, then EB = EB/0.82 to account for inefficiencies of using body tissues to meet metabolic energy needs. To make these various calculations, feed intake and milk yield were measured daily; concentrations of milk fat, protein, lactose, and SCC were determined 3 times per week by Michigan DHIA (East Lansing, MI) with a Bentley 2000 (Bentley Instrument, Chaska, MN); cows were weighed on 2 consecutive days per week, and DM content of the diet 2 times per week.

Indicators of Energy Balance

Several indicators of EB were monitored during the 14-d dietary treatment period to insure that PEB was maintained in the ad libitum fed cows and that NEB was established in the feed-restricted cows. Energy balance indicators included DMI (kg/d), 3.5% FCM yield (kg/d), biweekly serum NEFA concentration (µEq/L), and calculated energy balance (Mcal/d; see above). Before feeding, blood for serum NEFA analysis was collected via tail venipuncture into 10-ml Vacutainer tubes containing no anticoagulant 14, 11, 7, 4, and 0.5 d before endotoxin infusion, as well as immediately before endotoxin infusions. Blood was allowed to clot overnight at 4°C, serum was harvested following centrifugation of the blood tubes (1550 × g for 25 min at 4°C), and samples were stored at −20°C until assayed. Serum NEFA concentrations were determined enzymatically (NEFA-C Kit; Wako Pure Chemicals, Inc., Osaka, Japan) with an assay adapted for microtiter plates (Johnson and Peters, 1993).

Intramammary Endotoxin Challenge

Milk samples from each quarter of all cows were aseptically collected 12 h before endotoxin infusions and cultured on sheep blood agar plates to ensure that single quarters selected for endotoxin challenge were not infected with common mastitis-causing pathogens. Endotoxin was prepared by vortexing 100 µg of Escherichia coli O111:B4 lipopolysaccharide (LPS; Sigma Chemical Co., St. Louis, MO) in 5 ml of pyrogen-free
Figure 1. Daily means and SEM for DMI (a), 3.5% milk FCM production (b), serum NEFA concentration (c), and calculated energy balance (d) for cows fed ad libitum (control, ■) or restricted to 80% of energy requirements (feed-restricted, ○) for 14 d before intramammary endotoxin infusions (on d 0). All cows were in positive energy balance up to d −13, at which time dietary treatments were commenced. The restriction fed cows were in clear negative energy balance when endotoxin was administered on d 0.

Indicators of Clinical Disease

A variety of systemic and local inflammatory indicators were used to monitor the clinical response to intramammary endotoxin challenge. Monitoring was initiated at 12 h before infusion (to establish baseline values), and then intensively at 0, 2, 3, 4, 6, 9, 12, 24, and 36 h postendotoxin infusion. Systemic indicators included fever (°C; determined by rectal temperature); heart rate (beats/min), respiration rate (breaths/min), and rumen contraction rate (contractions/min), each determined by auscultation; total leukocyte count (millions of cells/L of blood) was determined by electronic cell counting [Coulter Particle Counter (Beckman Coulter, Fullerton, CA)]; and serum tumor necrosis factor-alpha (TNF-α) concentration (ng/ml; determined by radioimmunoassay as in Kahl et al., 1997). Acute changes in milk concentrations of serum proteins, such as IgG, have been used as indicators of capillary permeability that accompanies the local inflammatory response in the mammary gland (Schalm, 1977; Guidry et al., 1983). Therefore, we also monitored milk IgG from the mammary quarter infused (mg/dl; determined
by radial immunodiffusion using a RID kit from VMRD, Pullman, WA) at 0, 3, 6, 9, 12, and 24 h postinfusion. Milk TNF-α concentrations (ng/ml; determined by radiomunnoassay as in Kahl et al., 1997) were monitored in a subset of samples (0, 3, and 6 h postinfusion). Milk samples for the IgG and TNF-α assays were processed into whey (Harmon, 1994) and the samples were stored at −20°C until assays were performed at the end of the animal work. Also within 24 h of endotoxin infusion, milk samples from the quarter infused were analyzed for SCC (Michigan DHIA, East Lansing) using a Somaticount 500 (Bentley Instrument). Maximum sensitivity for the Somaticount 500 is 9.999 million cells per milliliter. Somatic cell counts for 11 samples were greater than the maximum sensitivity and were deleted before statistical analysis of this dataset. Somatic cell counts were determined in milk from all quarters before infusion and at 36 h and 1 wk postinfusion.

Statistical Analyses

The various datasets of this study were analyzed using repeated measures analysis (SAS PROC MIXED, SAS Institute Inc., Cary, NC) modeling correlated residuals within cow (Littell et al., 1996). Erratic heterogeneous variance was observed on initial examination of empirical distributions of residuals for each dataset, so all datasets were ln-transformed before statistical analyses. This transformation resolved the unequal variance distributions in each dataset. The linear models used to statistically analyze the transformed data included fixed effects of treatment (PEB or NEB), time (relative to start of dietary treatment or relative to endotoxin infusion), and the treatment by time interaction. Models also included a random block effect and error term. Covariance between residuals within cow was modeled either as compound symmetry or as heterogeneous compound symmetry (Littell et al., 1996). Analyses relative to endotoxin infusion used observations from the first 24 h after infusion with first observation made before infusion as a covariate. Significant differences were declared when \( P \leq 0.05 \). Trends toward significance were declared when \( P \leq 0.10 \).

RESULTS

Indicators of Energy Balance

Four indicators of EB were used in this study to insure that cows fed ad libitum maintained PEB and those restricted to 80% of maintenance requirements achieved NEB. Data presented in Figure 1 summarize results of these dietary treatments on EB over the 14 d prior to endotoxin challenge. Figure 1a shows that the NEB group consumed approximately 40% less feed per day than the PEB group \( (P < 0.0001) \). This resulted in decreased FCM production for the NEB group that was −10% below that of PEB cows \( (P < 0.0001; \text{Figure } 1b) \). Also, serum NEFA concentrations rose dramatically to >500 µEq/L by d 7 in the NEB group, compared with the maintained concentration of −105 µEq/L in the PEB group \( (P < 0.0001; \text{Figure } 1c) \). Finally, mean calculated EB in the NEB group dropped to approximately −6.2 ± 1.1 Mcal/d, a value that was significantly lower \( (P < 0.0001) \) than the mean EB of PEB cows \( (5.8 ± 1.1 \text{ Mcal/d; Figure } 1d) \). These four EB indicators demonstrated our success in inducing and maintaining chronic NEB by restriction feeding. Similarly, these indicators showed that ad libitum feeding was successful in maintaining PEB in the control group of cows.

Mean Clinical Response to Intramammary Endotoxin Infusion

We intensively monitored a variety of systemic and local inflammatory indicators following intramammary infusion of endotoxin into one mammary quarter per cow. Endotoxin infusion caused acute (within 6 h) fever and increases in heart and respiration rates \( (P < 0.0001; \text{Figure } 2) \). Each of these clinical indicators of the systemic inflammatory response to endotoxin returned to baseline values by 24 h postinfusion. As expected (Jain et al., 1978; Guidry et al., 1983; Kahl et al., 1997), endotoxin infusion also acutely reduced (−60%) mean blood leukocyte counts (Figure 3) and increased milk SCC \( (P < 0.0001; \text{Figure } 4) \) and IgG concentration \( (P < 0.0001; \text{Figure } 5) \), and caused modest elevations in milk TNF-α concentrations \( (P < 0.0001; \text{Figure } 6b) \) that were not so pronounced in serum (Figure 6a). Mean blood leukocyte counts and milk IgG concentrations returned to baseline levels by 24 and 36 h, postinfusion, respectively, but milk SCC was still elevated 36 h after infusion. Milk SCC returned to baseline within 1 wk of infusion. TNF-α concentrations in blood and milk stayed modestly elevated over the 6 h postinfusion period for which they were monitored. In combination with the indicators of EB in Figure 1, the mean clinical response to intramammary endotoxin infusion was repeatable enough to achieve the main objective of the study, which was to determine whether NEB impacts the acute clinical response to endotoxin-induced mastitis in dairy cows.

Effects of Energy Balance on the Clinical Response to Endotoxin Infusion

The effects of dietary treatment and treatment × time interaction were tested for each indicator of clinical
response to endotoxin infusion. First, there was a significant treatment × time interaction for the fever response to endotoxin infusion ($P < 0.0001$; Figure 2a). NEB cows had lower rectal temperatures 3 and 4 h postendotoxin infusion than did PEB cows ($P < 0.05$) and maintained higher rectal temperatures between 9 and 12 h postinfusion ($P < 0.05$). However, peak fever response (∼40.8°C at 6 h) and time to full recovery from fever (at 24 h) were not influenced by NEB.

Negative energy balance influenced the tachycardia response following intramammary infusion of endotoxin in a similar manner as the fever response with a significant treatment by time interaction ($P < 0.0001$). For example, NEB cows exhibited slower heart rates than PEB cows 2, 3, and 4 h postendotoxin infusion ($P < 0.05$) but had higher heart rates 9 h postendotoxin ($P < 0.05$; Figure 2b). Peak heart rate response (∼110 beats/min at 6 h) and time to full recovery of heart rate (at 24 h) were not influenced by NEB (Figure 2b).

Mean respiration rate was generally lower in NEB cows than PEB cows, even before the infusion of endotoxin ($P < 0.01$; Figure 2c), likely as a result of reduced DMI (Figure 1a) and milk yield (Figure 1b). Mean respiration rates began to rise ∼4 h postendotoxin infusion. In the PEB cows, this response was modest at best, peaking at 6 h postinfusion (∼20% higher than at 3 h) and returning to normal by 9 h. However, mean respiration rates in the NEB cows continued to increase between 4 and 12 h postendotoxin infusion, reaching a peak 12-h rate that was almost 40% greater than the 3-h rate (Figure 2c). Respiration rate for NEB cows returned to preendotoxin infusion rates by 24 h postinfusion. Data in Figure 2c thus suggest that the respiratory response to intramammary endotoxin may be slightly more intense and prolonged when cows are in NEB than when they are in PEB.

The acute response of the rumen to endotoxin infusion was not affected by EB, both dietary treatment
groups exhibiting similar decreases in rumen motility between 2 and 24 h postinfusion (Figure 2d). Also, EB did not influence total blood leukocyte (Figure 3; $P > 0.10$), milk IgG (Figure 5; $P > 0.10$), or serum and milk TNF-α (Figure 6; $P > 0.10$) responses to intramammary endotoxin infusion.

**DISCUSSION**

As evidenced by daily DMI, milk yield, calculated energy balance, and biweekly serum NEFA concentrations (Figure 1), feed restriction caused acute and sustained NEB in the lactating dairy cows of this study. Our intramammary endotoxin infusions caused the expected clinical signs of systemic and local disease, evidenced by fever response, increased respiration and heart rates, decreased rumen motility (Figure 2) and blood leukocyte counts (Figure 3), slightly increased serum TNF-α concentrations (Figure 6a), and more pronounced increases in milk SCC (Figure 4), IgG (Figure 5) and TNF-α concentrations (Figure 6b).

The main question addressed by this study was whether dietary-imposed NEB alters responsiveness of lactating dairy cows to intramammary endotoxin challenge. The only notable effects of imposed dietary energy deficiency in this study were on fever, tachycardia, and respiration rates, which were slightly lower in NEB cows than PEB cows up to 6 h postendotoxin infusion (Figure 2). Fever and tachycardia are caused by interactions between various cytokine networks and the neuroendocrine system (reviewed in Elsasser et al., 2000; Kelley, 1988). Although NEB had minimal effects on blood and milk TNF-α concentrations in this study (Figure 6), it is possible that nutrient deficit altered the sensitivity of cytokine-neuroendocrine networks in the feed restricted cows (Elsasser et al., 2000). In support of this possibility, Shuster et al. (1996) reported that early-lactation dairy cows (probably in NEB) with coliform mastitis had significantly higher rectal temperatures than mastitic midlactation cows (probably in PEB), as well as significantly higher TNF and interleukin-8 responses. Therefore, it is possible that the differences in cytokine, fever, and tachycardia responses ob-
Figure 6. Hourly means and SEM for tumor necrosis factor-alpha (TNF-α) concentrations in serum (a) and milk (b) following intramammary infusion of endotoxin in cows fed ad libitum (control, ■) or restricted to 80% of maintenance requirements (feed-restricted, ○).

served by Shuster et al. (1996) were related to energy balance. It might be interesting to substantiate this possibility through more holistic experiments designed to explore multiple cytokine-neuroendocrine gene expression cascades in relevant tissues of endotoxin-challenged cows with imposed NEB versus PEB. That said, whole animal clinical responses such as peak fever and tachycardia and return of these to baseline were not influenced by EB in the present study. Therefore, it appears that the impact of NEB on acute clinical responses during experimental endotoxin-induced mastitis is minimal. This was further supported by our data showing no effects of NEB on total blood leukocyte counts, serum TNF-α concentrations, or milk SCC, TNF-α, and IgG concentrations. However, these rather gross indicators of response to experimentally induced endotoxin mastitis may not tell the full story about potential roles for NEB in more subtle modulation of potent cytokine-neuroendocrine networks (Elsasser et al., 2000), or about possible influences of nutrient deficiency on the severity of acute mastitis from natural coliform infections.

Certainly, other studies have been published that support the notion of altered immune function in cattle during nutrient deficiency. For example, surgical removal of the mammary gland before calving to remove the demands for energy at the onset of lactation accelerates recovery of circulating leukocyte numbers after calving (Kimura et al., 1999). In addition, deficiencies in specific nutrients such as selenium and vitamin E that accompany NEB often result in increased incidence and severity of mastitis, while supplementation of diets with Se and vitamin E significantly reduces frequency and duration of clinical mastitis (Smith et al., 1984; Erskine et al., 1989). The NEB cows of the present study were feed-restricted and thus may have had concurrent deficiencies in some or all vitamins and minerals, as well as in protein. It is possible that these single-nutrient deficiencies played a role in altering the fever and tachycardia responses to intramammary endotoxin infusion in the current group of cows, perhaps through effects on the cytokine-neuroendocrine axis. Indeed, other researchers have shown that the nutritional influence on cytokine responses to endotoxin is an intricate interplay between protein and energy status, low-protein diets resulting in a TNF-α response to endotoxin greater in magnitude than high-protein diets (Kahl et al., 1997; Elsasser et al., 2000). Where periparturient cows are concerned, NEB and accompanying single nutrient deficiencies may be quite different than in midlactation cows because immunosuppression exists as a separate but interconnected contributor to altered mammary inflammatory responses around parturition. Therefore, it would be interesting to repeat the current study on midlactation cows with imposed immunosuppression, to determine whether EB impacts clinical responses in this scenario. Furthermore, endotoxin, while a convenient inducer of mammary inflammation, is noninfectious and may be inappropriate for studies designed to assess host defenses related to acute coliform mastitis. Therefore, future NEB studies of this nature may benefit from using mastitis-causing coliforms to induce mammary inflammation.

In summary, results of the current study suggest that NEB has a minimal impact on acute clinical responses to intramammary endotoxin challenge. It is thus tempting to conclude that NEB is not an underlying susceptibility factor for severe coliform mastitis. However, additional NEB studies with immunosuppressed midlactation cows challenged intramammarily with mastitis-causing coliforms are required to substantiate this con-
clusion and extend its implications to the early-lactation cow.

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**REFERENCES**


