tetraacetic acid (EGTA). During infusion, blood samples were collected every 15 min until 60% of pre-infusion iCa concentrations were achieved. Samples were collected post-infusion at 0, 2.5, 5, 10, 15, 30, and every 30 min thereafter until 90% of pre-infusion iCa was reached.

**Results**

We utilized a likelihood ratio to determine whether the variances of the 2 meters were equal when measuring iCa concentrations during EGTA challenge and recovery period. Variance for the iSTAT was 0.02745, while variance for the Horiba meter was 0.14447. The chi-square value was significant ($P<0.0001$). These results indicate that the Horiba meter is more variable when measuring blood iCa concentration and produces values that are significantly higher than values obtained from the iSTAT ($P<0.001$).

**Significance**

The Horiba LAQUAtwin meter is less likely to accurately identify a cow with clinical or subclinical hypocalcemia than the Abaxis Vetscan iSTAT.

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**Effect of prepartum energy balance on neutrophil function following pegbovigrastim treatment in periparturient cows**

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**Introduction**

Treatment with granulocyte colony stimulating factor (G-CSF) increases neutrophil (PMN) count and enhances PMN function in the periparturient cow. It was hypothesized that prepartum undernutrition might reduce the effect of a commercial recombinant bovine G-CSF product (pegbovigrastim; Imrestor, Elanco) on PMN count and function. Hence the objective of this study was to test the effect of undernutrition for 1 month prior to calving on the response to IMR.

**Materials and Methods**

Cows (n=99) on pasture in a research herd in New Zealand were blocked by expected calving date and BCS and randomly assigned in a 2 by 2 factorial design to be fed to exceed energy requirements prepartum (FULL), or restricted to approximately 85% of prepartum energy requirements (RES). At approximately 7 d before expected calving and on the day of calving, half the cows in each feed group were injected with the labelled dose of IMR while the remaining half were injected with saline. Blood samples were collected pre-and post-calving for complete blood count, biochemistry and in vitro assessment of PMN function including phagocytosis, myeloperoxidase (MPO) release, and oxidative burst.

**Results**

Energy restriction prepartum resulted in lower body weight (96 ± 0.4% vs 101 ± 0.5% of initial body weight for RES vs FULL cows at calving; $P < 0.001$), and a higher proportion of cows with elevated concentrations (>0.4 mmol/L) of fatty acids (NEFA) in blood (35/41 (85%) vs 23/41 (56%) elevated for RES vs FULL cows at 7 d before calving; $P < 0.001$).

Treatment with IMR increased PMN count from 6 days before to 21 days after calving (9.8 ± 0.2 vs. 3.9 ± 0.2 x 10⁹/mL; $P < 0.001$). There was a time by IMR interaction ($P < 0.001$) for proportional release of MPO by PMN, with higher release at 4 d post-calving in IMR cows (0.80 (95% CI = 0.72 to 0.88) vs 0.59 (95% CI = 0.53 to 0.64), $P < 0.05$). There was no effect of prepartum energy restriction, nor energy restriction by IMR interactions for any of the white blood cell counts or functional tests.

**Significance**

It is concluded that IMR treatment results in significant increases in PMN count, and enhances PMN function as indicated by increased MPO release. The response to IMR was not affected by restricted pre-partum energy intake.