Continuous milking of dairy cows disrupts timing of peak IgG concentration appearance in mammary secretions

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The length of the dry period in commercial dairy production is under close scrutiny. While the main concern is the composition and volume of milk produced, the evaluation of colostrum quality under these new paradigms has suggested a decline in IgG concentrations, while some reports indicate no change. Colostrum quality has been defined as an adequate concentration (> 50 mg/ml) of immunoglobulin in the secretions to provide the newborn with maximal disease resistance. We investigated the appearance of IgG in mammary pre- and post partum secretions in cows without a dry period (continuously milked, Dry0) and compared the secretions with cows that experienced a dry period of 60 d (Dry60). Blood was collected during the experimental period and plasma analysed for progesterone (P4) and prolactin (Prl). Approximately ~6 d relative to parturition, the Dry0 animals exhibited increased concentration of IgG in their secretions to an average of ~35 mg/ml that remained rather constant through subsequent pregnancy and following parturition. Dry0 cows were producing an average IgG concentration in parturition colostrum of 44·2 ±17·6 mg/ml that was not different than that of controls (66·86 ±16·8 mg/ml). However, Dry0 cows exhibited high variation, different peak times (day) of IgG concentration including times that occurred both pre and post parturition. IgG mass of the Dry0 cows remained rather constant pre- and post partum and did not show the same declining mass following parturition that was shown for the Dry60 cows. The change in plasma P4 and Prl were shown to have no timing effect on colostrum IgG concentration.

Keywords: Continuous milking, colostrum, IgG, progesterone, prolactin.

Dairy production of milk is affected by a number of factors (Wiggins et al. 2002; Mansfeld et al. 2012). The length of the non-lactating period between successive lactations is one of the factors. The non-lactating period involves involution, a process of mammary gland remodelling, and is followed by a period of mammary epithelial cell proliferation and differentiation (Capuco et al. 1997). The industry adopted the period of 50–60 d as the optimum for most cows to maximise their lactational performance based upon research findings (O’Connor & Oltenacu, 1988).

During the later stages of pregnancy and generally overlapping the traditional dry period, the mammary gland epithelial cells of the dairy cow first differentiate to conduct colostrogenesis that results in the secretion of colostrum (Baumrucker & Bruckmaier, 2014). Colostrum immunoglobulin content is critical for the newborn because the transfer of these proteins in utero does not occur in ruminant species.

Colostrum feeding and timing of the feeding is critical for lowering the risk of disease (Donovan et al. 1998; Quigley & Drewry, 1998; Weaver et al. 2000). Colostrum quality has been traditionally determined by IgG content, but both protein concentration (11·8–74·2 mg/ml; Kehoe et al. 2007) and mass (30 g to >2 kg; Baumrucker et al. 2010) is highly variable among animals. Poor colostrum quality is one of the contributors to the risk of intestinal disease (McGuire et al. 1976; Quigley & Drewry, 1998). Industry standard guidelines describe the feeding of high-quality colostrums (>50 mg/ml IgG) within the first few hours of birth (Tyler et al. 1996, 1999).
Recent studies have explored the possibility of shortening the dry period (Annen et al. 2004; Klusmeyer et al. 2009) or eliminating the dry period between lactations resulting in continuous milking (Fitzgerald et al. 2007; Madsen et al. 2008; Schlamberger et al. 2010; Collier et al. 2012). Colostral IgG concentration has been reported to be similar for cows with short dry periods compared with cows with conventional dry periods (Annen et al. 2004; Rastani et al. 2005; Watters et al. 2008; Klusmeyer et al. 2009). Cows with no dry period, however, were reported to have decreased IgG concentration in colostrum compared with cows with a conventional dry period (Annen et al. 2004; Rastani et al. 2005; Klusmeyer et al. 2009). The variance in IgG concentrations have been attributed to many factors (Baumrucker & Bruckmaier, 2014). To our knowledge, studies which describe the IgG content in late gestation of continuously milked cows are absent. The objective of this study was to determine the appearance of IgG in mammary secretions pre- and post partum in dairy cows that were continuously milked and compare them with those in animals experiencing a 60-d dry period.

Material and methods

Animals and design

The Institutional Animal Care and Use Committee of Wageningen University and Research centres approved the experimental protocol. Sixteen Holstein-Friesian dairy cows from Wageningen University were used in the study. Two groups of cows with comparable milk production in previous lactation (10 057 ± 285 kg/305 d) were composed of: continuously milked cows (Dry0; n = 8) and cows that experienced a 60-d dry period (Dry60; n = 8). Cows were housed in a freestall and during lactation, cows were milked twice daily (5:00 and 16:30). The drying-off protocol for cows with the Day60 dry period consisted of a transition to the far-off ration at day 7 before drying-off, and milking once daily at day 4 before drying-off cows. At drying-off, Day60 cows were treated with an intramammary antibiotic (Supermastidol; Virbac Animal Health, Barneveld, The Netherlands). Overall milk yield on day 1 following parturition was 21.7 ± 1.7 kg/d for the Dry60 and 13.1 ± 8.0 kg/d for the Dry0 group. During the experimental period of ~day 7 pre-partum to day 3 post partum a daily morning secretion sample was collected from machine-milked animals. Ration composition was described earlier (Van Knegsel et al. 2013). In short, pre-partum feeding for dry cows consisted of a dry cow ration; lactating cows received a lactating cow ration supporting a milk yield of 25 kg/d. From 10 d before the expected calving date, all cows were fed a glucogenic concentrate and, when lactating, cows received in the milking parlour 1 kg/d of standard lactation concentrate. Forage composition consisted pre-partum of grass silage, corn silage, wheat straw and a protein source (rapeseed meal or soyabean meal) in a ratio of 39:25:25:11 (dry matter basis). Post partum, forage consisted of grass silage, corn silage, straw, and a protein source in a ratio 51:34:2:13 (dry matter basis).

Blood sampling

Tail vein blood samples were taken in 9-ml vacuum tubes coated with EDTA and kept on wet ice and centrifuged at 2500 g at 4 °C for 15 min to harvest plasma. Plasma was subsequently stored at −20 °C until analysis. Mammary secretions were sampled once a day (5 ml) and stored at −20 °C until analysis.

Plasma analyses

Plasma prolactin (Prl) was determined by radioimmunoassay as described previously by Bruckmaier et al. (1992). For the analysis of plasma progesterone (P4), a radioimmunoassay kit (no. IM1188, Beckman Coulter GmbH, Krefeld, Germany) was used.

Colostrum analysis

Colostral IgG concentration was determined with a modified enzyme-linked immunosorbent assay (Bovine IgG ELISA QuantiTitation Set; Cat. No. E10-118; Bethyl Laboratories Inc., Montgomery TX, USA) as described by Baumrucker et al. (2014). Results were expressed as IgG concentration in mg/ml. Mass of IgG secreted by the mammary gland was calculated by multiplying respective concentrations by the corresponding milk yields.

Statistical analysis

Data were analysed using SAS Proc Mixed procedure (SAS, 2007) that included treatment (dry period length), sampling time-point and their interaction as fixed effects. Sampling points were treated as repeated factor within animals. Analysis of peak IgG concentrations between treatment groups was conducted with Proc GLM with IgG=Treatment.

Results

Secretions

Figure 1 shows the milk secretion occurring in the two treatment groups. Cows that were continuously milked were producing 8.28 ± 5.52 kg/d before parturition. At the first milking following parturition, the secretion weight increased to 10.4 ± 6.0 kg/d. The Dry60 group produced less secretion weight at the day 0 (6.07 ± 3.7 kg/d) when compared with the Dry0 group. While the Dry60 group increased secretion dramatically in the day 2 milking (29.4 ± 1.17 kg/d) the Dry0 cows increased at a lower rate reaching 13.95 ± 8.5 kg/d at day 2 milking.

Blood hormones

Blood plasma P4 and Prl concentrations occurring during the experimental period are shown in Fig. 2. Plasma P4 was
different between Dry0 and Dry60 during the pre-partum period with the Dry60 group exhibiting higher concentration than the Dry0 group during the period day −4 to day 0 (Trt; \( P = 0.01 \)). While both treatments declined in P4 around the same day to parturition, that was different at day 0 when compared with previous days (\( P < 0.01 \)) and there was no treatment*time to parturition interaction (Trt*TP; \( P = 0.34 \)). While the Dry0 cow plasma was different than that of the control group (Dry60) during the period from day −4 to day −1, at parturition P4 concentrations for the two treatment groups were not different. Analysis of area under the curve of progesterone indicated a treatment effect (\( P < 0.01 \)) with Dry0 showing greater amount of P4 in the plasma.

Blood plasma Prl concentrations are also shown in Fig. 2. At no time was the prolactin concentration different between the two treatment groups (Trt; \( P = 0.60 \)). Increase in Prl was observed at day −1 (TP; \( P < 0.01 \)), peaked at parturition (day 0) and declined rapidly in the days following parturition. There was no Trt*TP interaction (\( P = 0.90 \)). Analysis of area under the curve for Prl indicated a difference between the two treatment groups with Dry60 showing greater amount of Prl in the plasma (\( P = 0.01 \)).

Figure 3 shows the concentration of IgG occurring in the secretions between the two groups. The Dry0 group had low IgG concentration of \( \approx 9 \) mg/ml at day −7 but then significantly increased in the secretions to a mean of 35·4 ± 6·3 mg/ml across the period from day −5 to day −1. The first milking after parturition with the Dry0 group showed a concentration of 44·2 ± 17·6 mg/ml with a concentration decline following parturition. The Dry60 group had a concentration of 66·86 ± 16·8 mg/ml at the first milking after parturition (day 0) and then showed a concentration decline to 45·12 ± 7·6 mg/ml at day 1 and continued to decline. Treatment groups were not different in concentration after parturition (\( P = 0.36 \)), but the high variation in the Dry0 group shown in Fig. 3 contributed to this finding. While time to parturition changes in IgG concentration was significant (\( P < 0.001 \)), there was no treatment*time to parturition interaction (\( P = 0.34 \)). Interestingly, the Dry0 group treatment cows expressed high variation in the IgG concentration at any one time point around parturition after the initial increase.

**Timing of peak IgG concentrations**

Table 1 shows the timing of IgG peak concentrations between the two groups. While the Dry60 group all exhibited the highest IgG concentration at day 0, the Dry0 group had variable highest concentration days that ranged from −4 d to +3 d relative to parturition. The average of the peak concentrations of the Dry0 group was 63·73 ± 42·77 and was not different than the Dry60 group peak concentration average of 66·86 ± 16·8 (\( P = 0.83 \)). However, this is an artificial comparison in that it compares peak IgG concentrations that for the Dry0 cows appear on different days. Furthermore, two cows within the Dry0 group (4341 and 4978) were clearly deficient in IgG concentration relative to calf feeding and reduction of infection risks (Quigley & Drewry, 1998). Conversely, only one cow within the Dry60 group (2338) had below the IgG concentration recommended for optimal calf health (Tyler et al. 1999).

**Secretions of the Dry0 cows**

Figure 4 shows the variance in milk secretion among the Dry0 cows from day −5 to day 3. The asterisk (*) in Fig. 4 indicates the day of maximal IgG concentration that corresponds with Table 1 data. Most cows maintained a rather constant secretion rate until around parturition when milk secretion increased. Only one cow (4978) had a decline in milk secretion that plummeted after parturition and then began a gradual increase.

**IgG mass in pre-partum secretions**

Figure 5 shows the mass recovered in the secretions following parturition. The Dry0 group showed an average mass of 255 ± 101 g during the four collection days and was not different between the days sampled (days 0−3). The Dry60 group was significantly greater in IgG mass at days 0 and 1 (\( P < 0.01 \)) when compared with the Dry0 group, but the two were not different in mass at days 2 and 3 of milking. There was a significant time to parturition effect (TP, \( P = 0.01 \)) due to the drastic decline in the Dry60 treatment group. Interestingly, the sum of IgG mass in the four milkings after parturition was 1028 ± 104 g compared with 2693 ± 557 g for Dry0 and Dry60, respectively. Thus, less mass of IgG was produced after parturition in the Dry0 cows. However, if the pre-parturition mass is taken into consideration for the Dry0 cows (35·4 mg/ml × 6070 l/d × 6 pre-partum days), then an additional 1289 g IgG was produced by the Dry0 group. Adding the known post parturition 1028 g to the theoretical 1289 g from the pre-partum calculation suggests that these
cows produced an average of 2317 g of IgG during their colostrogenesis period. This appears to be similar to the 2793 g produced for the Dry60 cows.

Discussion

Colostrum quality is important for the newborn dairy calf. Provision of adequate mass of IgG gives the newborn the least risk of disease (Quigley & Drewry, 1998). Shortened dry periods or continuous milking with alterations in the time of the dry period are under investigation (Swanson, 1965; Annen et al. 2004; Kuhn et al. 2006; Fitzgerald et al. 2007; Madsen et al. 2008; Klusmeyer et al. 2009; Schlamberger et al. 2010) but the effect of a shortened dry period on the production of colostrum is controversial with some reports of lower colostrum IgG concentrations and some reports of no change.

It is surprising that although milked twice daily, the Dry0 cows recovered approximately the same concentration of IgG in their subsequent secretions after the initial increase occurring at day -5. Gross et al. (2014) showed that the transcytosis of IgG into mammary secretions can be very rapid when the gland is emptied during the pre-partum period.

The continuously milked cows were producing ~8.28 kg/d of secretions prior to parturition. Considering that a newborn calf can consume approximately 2 kg of colostrum in the first drinking, sufficient is present pre and post parturition in the continuously milked cows (Dry0) to provide the calf with an adequate volume. However, the IgG concentration of the Dry0 cows averaged below the recommended values (50 mg/ml; Tyler et al. 1999) and two of the Dry0 cows had very poor IgG concentrations.
The initiation and termination of colostrum formation has been thought to occur with the decline in P4 concentration in the blood and to be further inhibited by the increased appearance of Prl (Guy et al. 1994; Barrington et al. 1999, 2000). Our finding that pre-partum P4 is at a lower concentration in the blood of the continuously milked cows (Dry0) may be explained by the appearance of P4 in pre-partum mammary secretions. Keller et al. (1976) showed that pre-partum mammary secretions contain *1 μg/ml of P4* and this could account for the lower concentrations occurring in the blood pre-partum. Because milk removal was not occurring in the Dry60 cows, the P4 was at a higher concentration (~7 ng/ml vs. ~5·8 ng/ml). This also could be an explanation for the higher area under the curve for Prl in the Dry60 cows.

### Table 1. Timing of peak IgG concentration and mass occurring in the continuously milked cows (Dry0) compared with cows dry for 60 d (Dry60)

<table>
<thead>
<tr>
<th>Dry60 cows</th>
<th>Day</th>
<th>IgG, mg/ml</th>
<th>Milk, litre</th>
<th>IgG, g</th>
<th>Dry0 Cows</th>
<th>Day</th>
<th>IgG, mg/ml</th>
<th>Milk, litre</th>
<th>IgG, g</th>
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<tr>
<td>5279</td>
<td>0</td>
<td>60·43</td>
<td>15·6</td>
<td>945·7</td>
<td>5158</td>
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<tr>
<td>5198</td>
<td>0</td>
<td>68·68</td>
<td>20·76</td>
<td>1426·0</td>
<td>4918</td>
<td>−4</td>
<td>77·50</td>
<td>32·65</td>
<td>2530·7</td>
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<tr>
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<td>0</td>
<td>77·26</td>
<td>8·61</td>
<td>664·8</td>
<td>4376</td>
<td>2</td>
<td>118·26</td>
<td>11·67</td>
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</tr>
<tr>
<td>5200</td>
<td>0</td>
<td>68·31</td>
<td>20·20</td>
<td>1380·2</td>
<td>4978</td>
<td>3</td>
<td>15·66</td>
<td>5·62</td>
<td>88·0</td>
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<td>0</td>
<td>65·61</td>
<td>10·50</td>
<td>802·3</td>
<td>4439</td>
<td>0</td>
<td>89·34</td>
<td>6·89</td>
<td>616·1</td>
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<tr>
<td>2338</td>
<td>0</td>
<td>25·51</td>
<td>22·98</td>
<td>563·4</td>
<td>4801</td>
<td>0</td>
<td>33·57</td>
<td>4·18</td>
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<tr>
<td>Mean</td>
<td></td>
<td>66·40</td>
<td>19·59</td>
<td>1299·4</td>
<td>63·73</td>
<td>11·32</td>
<td>744·7</td>
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<td>2·98</td>
<td>255·1</td>
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</tr>
</tbody>
</table>

We showed that the neither the timing of the decline in P4 nor the timing of the increase in Prl has any effect upon the appearance of IgG concentration or mass in the secretions of continuously milked cows. While the concentration of IgG in the post-partum secretions of Dry0 and Dry60 cows declines, the mass continues to appear at a constant rate in Dry0 cows that either hormone seems to affect.

The variation in the appearance of peak of IgG concentration among the continuously milked cows is interesting. On average, somewhere around 6 d pre-partum, colostrum concentration appears to increase in the Dry0 cows. As a consequence of continuous milking, the range in peak IgG concentration for each cow was large (days *t = C0 to +3*) indicating that each animal seemed to reach its peak concentration at different times during the experiment, one reaching that peak on day 3 following parturition. This suggests that each animal initiated maximal colostrum transfer rates at different time independent of parturition and P4 and Prl changes. This finding also suggests that a...
standard collection time either pre- or post partum for continuously milked animals would be difficult to establish. Nevertheless, because these cows are continuously milked and the IgG is actively removed, perhaps providing a day or two of non-milking prior to parturition would allow for the IgG to accumulate so that the IgG concentration and mass would be higher following parturition. Gross et al. (2014) showed rapid recovery of IgG concentrations and mass within a short period prior to parturition.

When considering IgG mass, we showed that the sum of post-partum mass and pre-partum mass from the Dry0 cows was 2317 g. This sum appears to be equal to that of the Dry60 group production of IgG (2793 g). However, the mass of IgG secreted by Dry0 cows following parturition is less than that of the Dry60 cows on days 0 and 1. Based upon this finding, it appears that the colostrogenesis capacity of continuously milked cows is not mass transfer impaired; rather the timing of the mass transfer is altered.

Conclusions

The lack of a dry period (continuous milking) resulted in the first detected increase occurring around 6 d before parturition. Furthermore, the first milked colostrum following parturition of the Dry0 cows averaged a lower IgG concentration than that of the traditional Dry60 cows, but owing to high variation among Dry0 cows, this was not statistically different in concentration. However IgG was different in total mass between Dry0 and Dry60 cows. We attribute the Dry0 cow colostrum lower concentration at parturition to continuous removal of IgG, the variance of peak IgG concentration (day ~ 4 to +3), and the variance of IgG concentrations among the Day0 animals. We conclude that the typical timing of the peak IgG concentration in Dry0 cows is disrupted by the milking process (removal and replacement of the secretions), but not linked to changes in P4 or PRL concentration changes in the blood.

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