Effects of stage of lactation and time of year on plasmin-derived proteolytic activity in bovine milk in New Zealand

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Summary. The objective of this study was to determine the effects of stage of lactation (SOL) and time of year on plasmin-derived proteolytic activity in the milk of pasture-fed dairy cows in New Zealand. Four herds of 20 Friesian cows were used, one herd calving in each of January, April, July and October. Cows grazed ryegrass/white clover pasture only, except during June (winter) when all cows received supplementary pasture silage. Milk samples were collected on four occasions during the year (spring, summer, autumn and winter) from each cow in milk, to give a total of three samples per cow (early, mid and late lactation; c. 30, 120 and 220 days after calving, respectively). Milk samples were analysed for plasmin-derived proteolytic activity. There was no effect of either SOL or time of year on plasmin activity and therefore yields of plasmin followed patterns in milk yield (highest in early lactation and in summer). There were effects of both SOL and time of year on plasminogen-derived and total plasmin plus plasminogen-derived activity, both of which were highest in late lactation and in spring. Changes in plasminogen-derived activity and total plasmin plus plasminogen-derived activity due to SOL were not only due to the decrease in milk yield associated with advancing lactation, because enzyme yields were also increased with advancing lactation. Similarly, effects of time of year on plasminogen-derived activity and total plasmin plus plasminogen-derived activity could not be attributed solely to concomitant changes in milk yield, and may be influenced by the variation in the quality and quantity of feed during the year inherent in a pasture-based dairy system. Effects of SOL on proteolytic activity were greater than, and independent of, effects of time of year.

Keywords: Milk protein, plasmin, proteolysis, lactation, seasonality.

The New Zealand dairy industry is based around the use of pasture as a low-cost feed source, which has led to widespread adoption of seasonal calving in order to
maximize pasture utilization. Most cows calve just before spring, and thus nearly all
cows are at a similar stage of lactation (SOL) at any time during the year. This
practice has created irregularities in the supply of milk to processors in terms of both
quantity and composition, and is accompanied by seasonal variations in the
manufacturing properties of the milk.

While a number of changes in milk occur during late lactation, the deterioration
in cheesemaking properties is often attributed to increased activity of the blood-
derived serine proteinase plasmin (EC 3.4.21.7) at that time (reviewed by Fox, 1992;
Lucey, 1996). Such an increase can occur through an increased presence in milk of
plasmin from the blood, or from the increased activation of the inactive zymogen
plasminogen (Korycka-Dahl et al. 1983; Politis, 1996). Plasmin hydrolyzes β-casein,
generating several γ-caseins and proteose peptones (Fox, 1992), while \( \alpha_{51}\)-casein
(Eigel, 1977; Andrews & Alichanidis, 1983) and \( \alpha_{52}\)-casein (Le Bars & Gripon, 1989;
Visser et al. 1989) are also susceptible to degradation. In turn, casein proteolysis has
a detrimental effect on the coagulation properties of milk (Grufferty & Fox, 1988).

The precise cause of changes in plasmin-derived proteolytic activity in milk
throughout the year is difficult to determine in a pasture-based, seasonally calving
dairying system. The levels of milk plasmin and plasminogen increase with
advancing lactation (Donnelly & Barry, 1983; Schaar, 1985; Politis & Ng Kwai
Hang, 1988; Politis et al. 1989a). In New Zealand, however, the effects of SOL are
confounded by the concomitant changes in the nutritional status of cows that results
from variations in the availability and quality of pasture as the dairying season
progresses. Environmental changes in temperature and day length may also impact
on milk composition and the plasmin-plasminogen system. This study was designed
to quantify the respective effects of SOL and time of year, and their potential
interaction, on proteolytic activity in milk.

**MATERIALS AND METHODS**

**Cows and design**

Details of animal management, cow age, genetic merit, live weight, condition
score, estimated dry matter intake, and composition of pasture and silage offered to
cows at time of sampling were reported by Auldist et al. (1998). Briefly, 80 mixed-age
Friesian cows were divided into four herds of approximately 20 cows each.
Insemination of each herd was staggered so that calving occurred at intervals of 3
months. Thus, one herd calved during each of January, April, July and October. This
meant that during any season there were three herds in milk, each at a different SOL
(and one herd dry), grazing similar pasture and exposed to similar management
practices and environmental factors.

Cows were managed as separate herds on the same farm and milked twice daily
through a common dairy. Each herd was offered a daily allowance of ryegrass/white
clover pasture sufficient to meet metabolizable energy requirements deemed
appropriate for the level of production, based on the UK dairy cow feeding standards
(Agricultural Research Council, 1984). In winter (June sampling) when insufficient
pasture was available to meet requirements, each herd received approximately 20% of
their dry matter intake as silage. On four occasions during the year (September: spring;
December: summer; March: autumn; June: winter) milk yield was recorded
and a milk sample collected from each cow at morning milking. Each cow was
therefore sampled during early lactation (approximately 30 d in milk), mid lactation
(approximately 120 d in milk) and late lactation (approximately 220 d in milk).
Milk analyses

Immediately following milking, an aliquot of fresh milk was centrifuged at 490 g at room temperature (25 °C) for 15 min and the fat fraction discarded. A portion (2–5 ml) of the skim milk was incubated with 50 mM 6-amino-n-hexanoic acid (Sigma Chemical Co., St. Louis, MO, USA) for 2 h at room temperature to dissociate plasmin and plasminogen from casein micelles (Politis et al. 1993). Treated skim milk was then centrifuged at 100000 g at 4 °C for 1 h and the supernatant (milk serum fraction) was stored at −20 °C until analysis for plasmin and plasminogen-derived activity using a modification of the method of Richardson & Pearce (1981).

To measure total enzyme (plasmin plus plasminogen-derived) activity, 30 µl milk serum, 25 µl 0.26 Sigma units/ml human urine urokinase (EC 3.4.21.73; Sigma) and 50 mM-Tris buffer (pH 7.5 to a total volume of 160 µl) was placed into each well of a 96-well microtitre plate. The microplate was incubated at 37 °C for 1 h to allow activation of plasminogen to plasmin. Then, after equilibration to 20 °C, 40 µl 1 mM-coumarin peptide (N-succinyl-Ala-Phe-Lys 7-amido-4-methylcoumarin; Sigma) was added and the fluorescence was measured every 30 s for 20 min at 20 °C using a Bio-Tek FL500 fluorescence microplate reader (Bio-Tek Instruments Inc., Winooski, VT, USA). The excitation and emission wavelengths were 360 ± 40 and 460 ± 40 nm, respectively.

Plasmin activity was measured without the addition of urokinase (25 µl water was added in its place). Plasminogen-derived activity was calculated by difference. Under the conditions described above, a change in fluorescence intensity of 375 in 1 min in the reaction mixture was defined as one unit of activity. Enzyme activities are reported as units/ml and yields (yield = activity x milk yield).

Somatic cell count (SCC) was measured using an automated cell counter (Fossomatic 215, Foss Electric, Hillerød, Denmark; Auldist et al. 1998).

Statistical analyses

Time of year, SOL and their interaction were analysed using the restricted maximum likelihood method of the mixed model procedure in SAS (1997). Cow was specified as a random effect and SOL and time of year were specified as fixed effects. Graphs of residuals versus fitted showed variation increasing with plasmin activity and yield and thus square root transformations were applied to these results before analysis.

On any sampling occasion, cows with SCC > 400000 cells/ml (between 1 and 6 cows per group) were excluded from the analyses to avoid the confounding effect of SCC on plasmin and plasminogen-derived activity in milk.

RESULTS

As expected, milk yields were highest in summer and lowest in winter, and consistently highest during early lactation and lowest during late lactation (Table 1). There was a significant interaction between SOL and time of year, with the effects of SOL being greatest in winter.

Neither SOL nor time of year significantly affected plasmin activity and the pattern of results for yield of plasmin were similar to those for milk yield (Table 1). Thus, there was a significant effect of SOL and time of year, with plasmin yield highest for cows in early lactation during summer.

Both SOL and time of year significantly affected plasminogen-derived activity
Table 1. Milk yields, and yields and activities of plasmin, plasminogen-derived enzyme and total plasmin plus plasminogen-derived enzyme in milk from cows in early (E), mid (M) and late (L) lactation during spring (September), summer (December), autumn (March) and winter (June)

(Values are means for all cows in each herd, n = 20)

<table>
<thead>
<tr>
<th></th>
<th>Spring</th>
<th></th>
<th></th>
<th>Autumn</th>
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<th>Winter</th>
<th>s.e.d. within herds</th>
<th>s.e.d. across herds</th>
<th>Main effects</th>
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<tbody>
<tr>
<td></td>
<td>E</td>
<td>M</td>
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<td>E</td>
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<td>L</td>
<td>E</td>
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<tr>
<td>Milk yield</td>
<td>13.2</td>
<td>9.6</td>
<td>8.9</td>
<td>14.8</td>
<td>11.8</td>
<td>8.8</td>
<td>13.6</td>
<td>9.9</td>
<td>8.5</td>
<td>11.6</td>
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<tr>
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<td>134</td>
<td>129</td>
<td>144</td>
<td>132</td>
<td>126</td>
<td>130</td>
<td>134</td>
<td>132</td>
<td>132</td>
<td>129</td>
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<tr>
<td>(× 10⁴), units/ml</td>
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<tr>
<td>Plasminogen</td>
<td>16.5</td>
<td>28.4</td>
<td>35.5</td>
<td>19.1</td>
<td>21.8</td>
<td>25.7</td>
<td>11.3</td>
<td>25.8</td>
<td>26.2</td>
<td>9.3</td>
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<td>(× 10⁴), units/ml</td>
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<tr>
<td>Total enzyme</td>
<td>18.3</td>
<td>30.0</td>
<td>37.7</td>
<td>11.8</td>
<td>23.4</td>
<td>27.4</td>
<td>13.4</td>
<td>25.6</td>
<td>27.9</td>
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<tr>
<td>Ratio (plasminogen:plasmin)</td>
<td>10</td>
<td>18</td>
<td>18</td>
<td>6</td>
<td>14</td>
<td>16</td>
<td>7</td>
<td>14</td>
<td>16</td>
<td>5</td>
</tr>
<tr>
<td>Square root plasmin yield (units)</td>
<td>1.54</td>
<td>1.25</td>
<td>1.36</td>
<td>1.61</td>
<td>1.37</td>
<td>1.21</td>
<td>1.57</td>
<td>1.31</td>
<td>1.21</td>
<td>1.38</td>
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<td>Plasminogen yield (units)</td>
<td>22</td>
<td>28</td>
<td>31</td>
<td>15</td>
<td>26</td>
<td>24</td>
<td>15</td>
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<td>Total enzyme yield (units)</td>
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<td>30</td>
<td>33</td>
<td>18</td>
<td>28</td>
<td>25</td>
<td>17</td>
<td>25</td>
<td>24</td>
<td>13</td>
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* P < 0.05, ** P < 0.01, NS not significant.
† SOL, stage of lactation.
‡ A change in fluorescence intensity in the reaction mixture of 375 in 1 min at 20 °C was defined as one unit of activity.
but there was no interaction (Table 1). Mean plasminogen-derived activity increased with advancing lactation, more than doubling from early to late lactation. Highest plasminogen-derived activities were found in spring and lowest in winter. In contrast to milk yield, which was highest in summer and early lactation, yield of plasminogen followed the overall pattern (averaged across all cows) spring > summer > autumn > winter and early < mid > late lactation (Table 1). Activity and yield of total enzyme were significantly influenced by SOL and time of year, and followed the same pattern as plasminogen.

There were effects of both SOL and time of year on the ratio of plasminogen to plasmin, although there was no interaction between these effects (Table 1). The ratio increased with advancing lactation, more than doubling between early and mid lactation.

The relationship between milk yield and the yields of plasmin, plasminogen and total plasmin plus plasminogen during each SOL and at each time of year are shown in Fig. 1. Milk yield decreased with advancing lactation, by 24% from early to mid lactation, and by a further 20% from mid to late lactation. Yield of plasmin essentially mirrored this decline with advancing lactation. In contrast, there was a
marked increase (45%) in plasminogen and total enzyme yields from early to mid lactation then a decrease (8%) from mid to late lactation.

The pattern in rise and fall of plasmin yield across seasons was essentially the same as that of milk yield: both increased from spring to peak in summer and then declined in autumn and further in winter. Yields of plasminogen and total enzyme differed from this general pattern by decreasing steadily from spring to winter.

**DISCUSSION**

Increases in plasminogen-derived enzyme activity associated with advancing lactation observed in the current study agree with previous reports (Politis *et al.* 1989a; Bastian *et al.* 1991). Reports of the response of plasmin activity to SOL, however, are inconsistent. In the current study there was no effect of SOL on plasmin activity. In contrast, previous reports have shown variously a gradual increase in plasmin activity with advancing lactation (Schaar, 1985), a peak at 5 months lactation (Baldi *et al.* 1996), a dramatic peak during late lactation (Davies & Law, 1977; Donnelly & Barry, 1983; Politis *et al.* 1989a) and differing results depending upon parity (Bastian *et al.* 1991).

The varying response of plasmin activity to SOL may simply represent inherent variation in cows, since wide ranges of plasmin activities have been reported within a SOL (Richardson, 1983b). Management practices such as level of feeding and milking frequency (Lacy-Hulbert *et al.* 1999), and cow factors such as mastitis (Auldist & Hubble, 1998), are known to contribute to this variation. Alternatively, the variable SOL effects reported could be due to differences in the onset of involution (Politis *et al.* 1989a).

Increases in plasminogen-derived and total plasmin plus plasminogen-derived activities observed in this study were not only due to the concentrating effects of decreased milk yield with advancing lactation, because there were considerable increases in yields of plasminogen and total enzyme from early lactation. These findings are consistent with a substantial increase in the rate of transport of total enzyme into the gland from early to mid lactation, and then a small decrease from mid to late lactation. This contrasts with another blood protein, serum albumin, also measured in these cows, for which the increase in concentration in milk with advancing lactation was due largely to the concomitant decrease in milk yield (Auldist *et al.* 1998).

Owing to the increase in plasminogen-derived activity with advancing lactation, coupled with relatively constant levels of plasmin, the ratio of plasminogen:plasmin increased during mid and late lactation. This ratio can be used to represent the degree of activation of the enzyme, or the relative rates of transport into milk (Politis *et al.* 1989a). These results therefore are consistent with less activation of the enzyme during mid and late lactation than early lactation. This is contrary to observations by Politis *et al.* (1989a) that at the end of lactation this ratio was half that in early lactation. Gilmore *et al.* (1995) who reported that plasminogen activator activity in the casein fraction of late lactation milk was approximately two-fold higher than in early or mid lactation, and Bastian *et al.* (1991) who reported that plasmin as a percentage of total plasmin plus plasminogen-derived activity increased dramatically in the last 3 months of lactation. Instead, the present study corroborates the proposition of Richardson (1983b) that increased proteolase activity occurs because more plasmin and plasminogen enter milk, rather than solely because of increased plasminogen activation. This notion is consistent with the fact that a loosening of
mammary tight junctions, which also occurs during advancing lactation, is positively correlated with plasmin and plasminogen-derived activity in milk (Stelwagen et al. 1994).

There was a strong influence of time of year on activities and yields of plasminogen and total enzyme, with activities highest in spring followed by summer. An opposite finding was observed by Bastian et al. (1991) that activities were greatest during autumn and winter, but these authors did not include cow management details and it is difficult to rationalize these contrasting findings. Level of feeding can, however, have an influence on proteolytic activity (Nicholas, 1998), and the quantity and quality of feed in the pasture-based systems of New Zealand varies considerably, especially between seasons (Auldist et al. 1998; McCall & Smith, 1998).

Overall, the effects of SOL on proteolytic activity in milk from grazing cows were greater than, and independent of, any influences of time of year. While the adoption of all-year-round calving may help smooth out changes in bulk milk proteolysis due to SOL, the effects of SOL and time of year appear to be additive and so seasonal calving could, for example, result in late lactation cows in spring having significantly higher levels of enzyme than they would normally have had in autumn. Thus all-year-round calving may result in an undesirable reduction in milk quality for manufacturing at certain times of the year. The effects of time of year need to be examined further, given that the present results under New Zealand conditions are the opposite of previous studies from the Northern Hemisphere, which have significantly different feeding and housing conditions. Importantly, the mechanisms driving this time of year effect, particularly the higher levels of plasmin and plasminogen-derived activity during spring, need to be examined.

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