ABSTRACT

Twelve dairy heifers were used to examine the clinical response of an alimentary oligofructose overload. Six animals were divided into 3 subgroups, and each was given a bolus dose of 13, 17, or 21 g/kg of oligofructose orally. The control group (n = 6) was sham-treated with tap water. Signs of lameness, cardiovascular function, and gastrointestinal function were monitored every 6 h during development of rumen acidosis. The heifers were euthanized 48 and 72 h after administration of oligofructose. All animals given oligofructose developed depression, anorexia, and diarrhea 9 to 39 h after receiving oligofructose. By 33 to 45 h after treatment, the feces returned to normal consistency and the heifers began eating again. Animals given oligofructose developed transient fever, severe metabolic acidosis, and moderate dehydration, which were alleviated by supportive therapy. Four of 6 animals given oligofructose displayed clinical signs of laminitis starting 39 to 45 h after receiving oligofructose and lasting until euthanasia. The lameness was obvious, but could easily be overlooked by the untrained eye, because the heifers continued to stand and walk, and did not interrupt their eating behavior. No positive pain reactions or lameness were seen in control animals. Based on these results, we conclude that an alimentary oligofructose overload is able to induce signs of acute laminitis in cattle. This model offers a new method, which can be used in further investigation of the pathogenesis and pathophysiology of bovine laminitis.

(Key words: bovine, laminitis, lameness, oligofructose)

Abbreviation key: PAO = postadministration of oligofructose, PCV = packed cell volume.

INTRODUCTION

Lameness and its welfare implication have become more widely recognized problems in recent years particularly in intensive dairy farming (Hoblet et al., 2000; Nelson and Cattell, 2001). Many of the major claw horn lesions causing lameness are believed to result from so-called laminitis. However, in many cases the various authors were referring to pododermatitis aseptica diffusa (sole hemorrhage) and to associated severe lesions such as pododermatitis circumspecta (sole ulceration) and had no direct evidence that the lamellae themselves were directly affected. Nevertheless, experimental studies and epidemiological surveys have identified risk factors that are associated with an increased prevalence of laminitis-associated lesions: housing (Vermunt and Greenough, 1996; Webster, 2001), management (Colam-Ainsworth et al., 1989; Bergsten, 1994), nutrition (Manson and Leaver, 1989), and calving (Offer et al., 2000). However, despite considerable research over recent years, the causal agents and the pathophysiological mechanisms of bovine laminitis remain unclear. A plausible and reproducible experimental model for the study of bovine laminitis does not currently exist.

Analysis of clinical cases precipitated by feeding a large amount of readily fermentable carbohydrate shows that signs of laminitis are often preceded by ruminal acidosis or gastrointestinal disease (Maclean, 1966; Yeruham et al., 1999). Different experimental models have tried to mimic this situation, by the administration of substances into the rumen to cause changes in the gastrointestinal environment. Attempts to induce acute laminitis with intraruminal infusion of lactic acid were reported to be partially successful in sheep (Morrow et al., 1973) but unrewarding in cattle (Anderson, 1981). Several studies have unsuccessfully tried to induce laminitis using an alimentary carbohydrate overload model (Hyldgaard-Jensen and Simesen, 1966; Boosman et al., 1990; Momcilovic et al., 2000). Although 3 of 6 steers showed very early signs of laminitis 12 to
16 h after starch overload (Suber et al., 1979), the clinical signs defining acute laminitis were not specified and it was unclear how comparison between animals was carried out. Christmann et al. (2002) evaluated hemodynamics in the digits of anesthetized steers given a grain overload. A successful induction of acute laminitis was reported without explaining how the criteria used for a positive diagnosis relate to the clinical syndrome.

A relationship between rumen acidosis and detection of endotoxin in the rumen generated the hypothesis of endotoxin being the trigger factor for laminitis. However, endotoxin injections have not been shown to induce clinical signs of acute laminitis in cattle (Boosman et al., 1991; Ohtsuka et al., 1997). Histamine and other vasoactive amines have been suggested as causal factors of laminitis. Nilsson (1963) reported that subcutaneous injections of histamine induced acute laminitis in cattle. Takahashi and Young (1981) were partially successful in their attempts to induce clinical laminitis when grain overload was combined with histamine administration. Unsuccessful attempts to reproduce the model were later reported by Boosman (1990), leaving the role of histamine debatable.

In summary, the reports on the potential of starch to induce clinical laminitis seem conflicting, and it becomes tempting to hypothesize the involvement of other carbohydrates. The results from a recent study undertaken by Pollitt and van Eps (accepted) demonstrated that alimentary oligofructose overload consistently resulted in the development of acute laminitis in horses. Oligofructose, as fructan, is one of the most abundant nonstructural carbohydrates in several plant species including many grasses (Cairns and Longland, 1998). Plants accumulate fructan depending on the prevailing growth conditions. Stressful conditions such as high light intensity and low temperature (comparable to spring and autumn in temperate climates) can result in very high fructan concentrations (Longland and Cairns, 2000).

Therefore, the purpose of the present study was to examine the clinical response of cattle dosed with an alimentary overload of oligofructose, with particular emphasis on development of laminitis.

MATERIALS AND METHODS

Animals

Twelve nonpregnant dairy heifers (3 Jersey, 6 Holstein-Friesian, 2 Ayrshire, and 1 Guernsey) between 325 and 505 kg (mean = 408 kg) were subjected to a 4-wk period of close handling and acclimatization. The animals originated from 3 dairy farms (all animals could not be purchased from same source), all of which kept heifers at grass from an early age. During acclimatization, the animals were kept in a paddock with a soft surface and were fed mixed grass-lucerne (alfalfa) hay ad libitum to ensure good ruminal function. At the end of this period, all animals tolerated clinical examination without any excitement (no changes in heart or respiratory rates), they could be walked by hand, accepted lifting of the distal front limb, foot palpation, and hoof testing (hind limbs were not examined for safety reasons). At the beginning of the trial, the heifers were assigned to 2 groups of 6 animals: the treatment group was challenged with oligofructose (Raftilose P95; Orafti Group, Tienen, Belgium), and the control group was given tap water (6 L per 100 kg). The oligofructose group was further divided in subgroups of 2 animals, each given a bolus dose of 13, 17, or 21 g/kg of oligofructose. Oligofructose was dissolved in tap water (solubility = 80 g/100 mL of water; 6 L per 100 kg heifer was used) and administered into the rumen by stomach tube. Five percent of the main dose was given as a priming dose twice a day for 3 d before the experiment, to gradually allow adjustment of the forestomach microbiota to the new source of carbohydrate.

Data

Baseline information on the cardiovascular (heart rate, packed cell volume, and standard base excess) and gastrointestinal status (rumen and feces pH, rumen contractions, and rectal temperature) as well as signs of lameness (hoof testing, digital pulse strength, and lameness examination) were recorded before the first priming dose (at ~72 h) and during the priming period (at ~48 and ~24 h). The clinical response was monitored at the time of administration of the main dose, and then at 6-h intervals, starting at 9 h postadministration of oligofructose (PAO), until the heifers were euthanized. Time points for observation are shown in Table 1.

At the beginning of the study, the response to oligofructose was unknown and a careful approach was taken to avoid severe distress in a large number of animals. Initially, the response was observed in only 2 animals given the lowest dose of oligofructose. No unexpected or uncontrollable side effects were noticed. These 2 animals were euthanized at 72 h PAO. The decision to shorten the study period and euthanize the remaining 10 animals at 48 h PAO was made because lameness was observed as an early sign.

All animals had catheters placed in the jugular vein before the main dose of oligofructose was given. A handheld blood-gas analyzer (i-STAT; Sensor Devices Incorporated, Waukesha, WI) was used to monitor the resulting metabolic acidosis and degree of dehydration to enable early alleviation with supportive therapy. A pilot study showed that the i-STAT resulted in a slight un-
Table 1. Results of hoof testing in 6 animals given oligofructose versus 6 control animals. Oligofructose was administered at 0 h. Only pain reaction in the front claws was examined. Independent monitors without prior knowledge of previous recordings monitored animals in a rotation system.

<table>
<thead>
<tr>
<th>Time PAO² (h)</th>
<th>Monitor²</th>
<th>Animals given oligofructose¹</th>
<th>Control animals¹</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>−72</td>
<td>A</td>
<td>−</td>
<td>−</td>
</tr>
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<td>−48</td>
<td>A</td>
<td>−</td>
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<td>−24</td>
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<tr>
<td>9</td>
<td>B</td>
<td>−</td>
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<tr>
<td>15</td>
<td>C</td>
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<tr>
<td>21</td>
<td>A</td>
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<td>27</td>
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<td>−</td>
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</tr>
<tr>
<td>33</td>
<td>B</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>39</td>
<td>C</td>
<td>LFL, LFM</td>
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<tr>
<td>45</td>
<td>A</td>
<td>LFM</td>
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<td>51</td>
<td>A</td>
<td>LFL, LFM</td>
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<tr>
<td>57</td>
<td>B</td>
<td>All</td>
<td>−</td>
</tr>
<tr>
<td>63</td>
<td>C</td>
<td>All</td>
<td>−</td>
</tr>
<tr>
<td>69</td>
<td>A</td>
<td>LFL, LFM</td>
<td>−</td>
</tr>
</tbody>
</table>

1°Pain reactions: − = no pain reaction; LFL = pain reaction in left front lateral claw; LFM = pain reaction in left front medial claw; RFL = pain reaction in right front lateral claw; RFM = pain reaction in right front medial claw; All = pain reaction in all 4 front claws.

2PAO = Postadministration of oligofructose.

3A,B,C = 3 veterinarians who monitored the animals.

derestimation of packed cell volume (PCV), and an equation (actual PCV = 0.99 × i-STAT PCV + 3.04, n = 16) was therefore used to adjust the readings. Intravenous fluid therapy was instituted if PCV increased to more than 42%. Sodium bicarbonate was administered orally or i.v. if standard base excess dropped below −8 mmol/L. Signs of lameness were identified by observing the animal standing in the box, walking or trotting in a straight line, and while the animal was turned in small circles on a hard surface. The 3 monitors (all veterinarians), blinded to previous results, recorded the clinical responses of the animals in a nonconsecutive rotation. Lameness was classified as none, mild, moderate, or severe. ‘Mild lameness’ was recorded if slight lameness was seen when turning the heifers in circles or if the lameness was detectable only in the trot (e.g., head nodding or other uneven or asymmetric movement of hind or front). ‘Moderate lameness’ was recorded if lameness was obvious when walking and if the animal was reluctant to walk from soft bedding to a hard concrete surface (tender feet). ‘Severe lameness’ was recorded if lifting of feet was impossible or if animals only could stand for a few minutes.

In the present paper, acute laminitis is defined as a clinical disease with a rapid onset of foot pain and detectable lameness, which is seen shortly after alimentary overload of readily fermentable carbohydrate. Signs of claw inflammation (e.g., warmth and increased pulsation) may be present but no claw lesions can be identified visually (Ossent et al., 1997).

Statistical Analyses

An animal was regarded as having developed clinical laminitis if 2 consecutive positive hoof tests (positive pain reaction) were obtained 6 h apart in the same claw, and lameness was observed independently by at least 2 of the 3 monitors. The null hypothesis of no association between exposure to oligofructose and development of laminitis was examined using one-sided Fisher’s exact test at a 5% level of statistical significance. The effect of oligofructose on other clinical variables was compared graphically; however, observations obtained between 45 and 72 h PAO for the 2 animals euthanized at 72 h PAO were not used. Results from the 3 groups given different oligofructose dosages were not pooled so that trends in the dose-response relationship could be examined. The mean value of observations in the control group was compared with the mean value in the groups given the 3 doses of oligofructose (GraphPad Prism, version 4.00 for Windows, GraphPad Software, San Diego, CA).

The protocol was approved by the Animal Ethics Committee, The University of Queensland, and animals were inspected by the Animal Welfare Officer during the study (animal ethics approval certificate SVS/125/...
ACUTE BOVINE LAMINITIS: A NEW INDUCTION MODEL

Figure 1. Clinical response to oligofructose. Number of rumen contractions is depicted as mean values for 6 control heifers (◆) and heifers administered 3 different doses of oligofructose (n = 2 per dosage): 13 g/kg (○), 17 g/kg (▲), and 21 g/kg (◆). The standard error of the mean is not shown for groups given oligofructose due to low numbers in each group.

Figure 2. Clinical response to oligofructose. Changes in rumen pH are depicted as mean values for 6 control heifers (◆) and heifers administered 3 different doses of oligofructose (n = 2 per dosage): 13 g/kg (○), 17 g/kg (▲), and 21 g/kg (◆). The standard error of the mean is not shown for groups given oligofructose due to low numbers in each group.

Figure 3. Response to oligofructose. Changes in packed cell volume (PCV) are depicted as mean values for 6 control heifers (◆) and heifers administered 3 different doses of oligofructose (n = 2 per dosage): 13 g/kg (○), 17 g/kg (▲), and 21 g/kg (◆). The standard error of the mean is not shown for groups given oligofructose due to low numbers in each group.

02/RVA/UC). The heifers were euthanized by captive bolt and exsanguination at the end of the study. A post-mortem examination of the gastrointestinal tract was carried out and claw biopsies were processed for histology. Preliminary findings from histopathological examination of claw biopsies are presented in this paper. A detailed characterization will be provided in a separate publication.

RESULTS

All heifers given oligofructose drank large amounts of water (approximately 50 to 80 L/head) during the first 12 h PAO. In 3 animals, fluid distension of the rumen was palpated at 9 h PAO, but thereafter, neither gas nor fluid distension of the rumen was observed. All animals given oligofructose developed profuse, watery diarrhea, which began at 9 h and continued until 33 h in 3 animals, until 39 h in 2, and until 45 h PAO in 1 animal. During this period, animals were depressed and stopped eating. The 2 animals given the highest oligofructose dose (21 g/kg) stopped eating at 9 h, the remaining 4 stopped by 15 h PAO. Rumen contractions ceased or decreased from 9 to 15 h in animals given oligofructose, and increased again slowly from 27 h PAO (Figure 1). The change in rumen contractions coincided with change in rumen pH and diarrhea (Figure 2). Changes in fecal pH appeared to follow the changes in rumen pH (data not shown). All animals started to eat and ruminate without rumen fluid replacement being necessary. No changes in the above variables were recorded in the control group.

All animals given oligofructose had a short-term increase in PCV that peaked around 21 h PAO (mean
Figure 4. Response to oligofructose. Changes in standard base excess are depicted as mean values for 6 control heifers (◆) and heifers administered 3 different doses of oligofructose (n = 2 per dosage): 13 g/kg (○), 17 g/kg (▲), and 21 g/kg (◆). The standard error of the mean is not shown for groups given oligofructose due to low numbers in each group.

Figure 5. Clinical response to oligofructose. Changes in heart rate are depicted as mean values for 6 control heifers (◆) and heifers administered 3 different doses of oligofructose (n = 2 per dosage): 13 g/kg (○), 17 g/kg (▲), and 21 g/kg (◆). The standard error of the mean is not shown for groups given oligofructose due to low numbers in each group.

Figure 6. Clinical response to oligofructose. Changes in rectal temperatures are depicted as mean values for 6 control heifers (◆) and heifers administered 3 different doses of oligofructose (n = 2 per dosage): 13 g/kg (○), 17 g/kg (▲), and 21 g/kg (◆). The standard error of the mean is not shown for groups given oligofructose due to low numbers in each group.

PCV ≥38% in all groups. The increase tended to be more pronounced and prolonged in the 2 animals receiving the high dose (Figure 3). From 9 h PAO, all animals given oligofructose developed a profound metabolic acidosis (mean standard base excess below −12 mmol/L in all groups) (Figure 4). Dehydration and metabolic acidosis was further reflected in heart rates, which appeared to increase more in animals given the highest dose of oligofructose (>90 beats/min at 15 to 39 h PAO) (Figure 5). Five animals of 6 exposed to oligofructose developed a transient episode of fever, which coincided with the presence of diarrhea. It appeared that higher doses of oligofructose correlated with increases in rectal temperature (Figure 6). Observations for control animals were within normal reference intervals throughout the study.

Intravenous infusions of 5% sodium bicarbonate were necessary to alleviate metabolic acidosis in all animals given oligofructose. The 2 animals given 13 g/kg of oligofructose received 500 mL of bicarbonate i.v. at 15 and 21 h PAO, and received an oral dose of bicarbonate (300 g at 21 h PAO), which tended to overcorrect the acidosis (Figure 4). Both heifers given 17 g/kg of oligofructose received 1000 mL of bicarbonate i.v. at 21 and 27 h PAO and an additional 500 mL of bicarbonate was given i.v. at 27 h PAO. Animals given 21 g/kg of oligofructose received bicarbonate i.v. at 15 (500 mL), 21 (1000 mL), and 27 h PAO (1000 mL). One of the heifers given the highest dose of oligofructose also needed 500 mL i.v. at 39 and 45 h PAO.

Four animals with high PCV received i.v. fluids, and all animals given oligofructose received 300 to 650 mL of calcium borogluconate (400 g/L, Unical C.B.G., Mav-
The results of hoof testing are shown in Table 1. Five of 6 animals given oligofructose had at least 2 consecutive positive pain reactions in the same claw. All positive pain reactions were strong and were observed at 33 h PAO or later. Positive pain reactions were not observed in control animals. Palpation of digital arteries, in an attempt to detect increased pulse amplitudes during development of laminitis, appeared to be highly variable between observers and animals, and without any obvious relationship to oligofructose exposure (data not shown). Digital pulse amplitude was therefore discarded in further analyses.

Lameness was observed in animals 1, 3, 4, and 6 from 39 h PAO or later. Lame animals were distributed almost equally between the 3 dose groups: 1 of 2 heifers that received 13 g/kg, 2 of 2 that received 17 g/kg, and 1 of 2 that received 21 g/kg. Lameness was obvious on walking in animals 1, 3, and 6, whereas it was only detectable in the trot in animal 4. All lame heifers continued to display lameness at a constant level for the duration of the study. Two heifers given oligofructose showed no signs of lameness. None of the control animals displayed signs of lameness during the study.

Four of the 6 animals given oligofructose had 2 consecutive positive pain reactions in the same claw and displayed signs of lameness. These animals were therefore classified as ‘laminitis positive’. Two animals exposed to oligofructose and all 6 animals in the control group were classified as ‘laminitis negative’. Fisher’s exact test (one-sided) showed a significant association between oligofructose exposure and development of laminitis ($P = 0.03$).

The postmortem examination of the rumen showed that animals given oligofructose had a mild and diffuse red discoloration of the epithelium, confined to the ventral and cranial parts of the rumen. Mild signs of inflammation were also seen in the cecum and upper colon where lesions were more localized and appeared in stripes. No increase in intestinal wall thickness was observed. Histopathological examination of claw biopsies was carried out and preliminary results are presented in Figures 7 and 8. In summary, clear differences were observed between the lamellar regions of treated and control animals.

**DISCUSSION**

Oligofructose is a carbohydrate belonging to the fructan group. It consists of fructose units linked to each other by $\beta(2$ to $1)$ bonds. Fructans are important storage polysaccharides found in common fruits, vegetables, and grasses (Gupta and Kaur, 1997; Longland and Cairns, 2000). The fructan content of some grass types can, under certain growth conditions, reach very high concentrations and a possible role in the pathogenesis of grass-induced laminitis in horses has been proposed (Cairns and Longland, 1998; Longland and Cairns,
This hypothesis has recently been reinforced by an experimental study. It was shown that oral administration of oligofructose did induce acute laminitis in a high percentage of horses (Pollitt and van Eps, accepted). In cattle, the occurrence of lameness in grass-fed animals has recently been reviewed (Westwood et al., 2003). It is suggested that ruminal acidosis and laminitis should be considered in the etiology of lameness in pasture-fed dairy herds. The purpose of the present investigation was therefore to examine the clinical response to an alimentary oligofructose overload in dairy heifers.

A direct association between oral oligofructose administration and clinical signs of acute laminitis was demonstrated in 4 of 6 animals, whereas no animals in the control group could be classified as laminitis positive. To the best of our knowledge, this is the first time it has been possible to induce acute bovine laminitis in a high percentage of animals, using an alimentary carbohydrate overload model that parallels the clinical engorgement situation. Additionally, the clinical findings present some interesting similarities to the oligofructose model reported to cause acute laminitis in horses (Pollitt and van Eps, accepted). Oligofructose given at doses of 7.5 to 12.5 g/kg caused a transient, watery diarrhea beginning 12 to 16 h PAO that ceased by 36 to 44 h PAO, and which coincided with depression and inappetence. No gas distension of colon or cecum was reported. All horses developed metabolic acidosis, mild dehydration, and a transient fever. At 28 to 32 h PAO, all horses started to show signs of acute laminitis. Thus, despite the differences in hoof anatomy and gastrointestinal function between these 2 species, the similarities in the clinical course of the disease are striking. It therefore seems reasonable to hypothesize that pathogenesis and pathophysiology of oligofructose-induced acute laminitis is similar in both species.

The lameness displayed by 4 of 6 heifers given oligofructose was obvious at the lameness examination, but could easily be overlooked, because animals continued to stand and walk and did not interrupt their eating behavior despite the lameness. In a study of lameness perception, Whay et al. (2002) reported that farmers identified fewer than 25% of lame cows in their herd. Previous attempts to induce laminitis in cattle did not objectively define how acute laminitis was evaluated clinically (Suber et al., 1979; Andersson, 1981; Boosman et al., 1991). Whay (2002) states that in recognizing lameness it is important not to confine identification and examination only to those individuals showing severe signs. An effort to increase the sensitivity of the lameness examination was therefore sought in the present study. In preparation for objective clinical examination, animals were trained 4 wk before experimentation to accept handling without excitement. Hoof testing was carried out to detect low-grade pain in individual claws, and time intervals between clinical examinations were short. The likelihood of false-positive reactions was decreased by the use of 3 observers alternating in a rotation system, and by strictly defining the signs required for a positive classification. The successful detection of low-grade lameness in the present study suggests that the concept of ‘subclinical’ laminitis (Hoblett et al., 2000), especially when used in a broader scientific context, should be used with caution. ‘Subclinical laminitis’ implies that signs are below the threshold of clinical detection. Use of the more diligent examination protocol described here puts in doubt the concept of subclinical laminitis and suggests that most subclinical laminitis is in fact clinical.

Mammals do not possess enzymes to digest fructan directly, but rely on the activity of the intestinal microflora to degrade the fructan polymers (Longland and Cairns, 2000). Fructanolytic activity has been demonstrated in bacteria isolated from the rumen as well as the equine hindgut (Kasperowicz and Michalowski, 2002; Bailey et al., 2003). The sudden availability of excess fructan, acting as a specific substrate to microorganisms, induces a selective and explosive proliferation of gram-positive bacteria. Recent results of in vitro studies, using equine cecal contents and starch or fructan as the carbohydrate source, have shown that Strep. toccoci and Lactobacilli are the main species involved in bacterial overgrowth. Streptococcus bovis and 5 Lactobacilli spp. were identified as having the capacity to decarboxylate certain amino acids to produce vasoactive amines (Bailey et al., 2003). These results support a theory put forward by Baxter et al. (1989) that substances resulting in vasoconstriction could cause lamellar hypoxia and thereby initiate the development of laminitis. However, a second hypothesis on the pathophysiology of equine laminitis suggests that Strep. bovis exotoxins cause the uncontrolled upregulation of matrix metallo-proteinase activity, which results in enzymatic degradation of basement membrane components (Pollitt and Daradka, 1998; Mungall et al., 2001). The loss of basement membrane anchoring filaments and hemidesmosomes leads to a mechanically unstable dermo-epidermal junction. In the horse, detachment of basement membrane is thus followed by displacement of the distal phalanx in the hoof capsule.

A dramatic increase of Strep. bovis (in rumen quantities) was also noted in cattle fed large amounts of concentrates (Tajima et al., 2001). Proliferation of Strep. bovis has been recognized as responsible for excessive production of bacterial mucopolysaccharides, which increases the viscosity of rumen fluid and leads to bloat in feedlot animals (Cheng et al., 1998). In the present
study, severe abdominal distension due to bloat was not observed. A few animals experienced a brief, transient, and moderate rumen distension at 9 h PAO, but this coincided with a significant intake of water. Thus, the pathophysiology of the different theories, in relation to bovine and equine laminitis, remains to be investigated. In particular, the extreme complexity of the rumen microbiota and the changes that occur during carbohydrate overload must be addressed.

**CONCLUSIONS**

The present article describes the development of a new experimental model for the study of acute laminitis in cattle. An alimentary overload of oligofructose induced a transient period of gastrointestinal disease, which was followed by signs of mild to moderate degrees of lameness in 4 of 6 animals. Training of animals to accept close handling, and a strict definition for the positive classification of laminitis was used to detect clinical cases, which could have been overlooked in a field situation. The model offers a new method that could be used in further investigation of the pathogenesis and pathophysiology of bovine laminitis.

**ACKNOWLEDGMENTS**

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