Standard Article

J Vet Intern Med 2018;32:853–859

Zinc Deficiency-Like Syndrome in Fleckvieh Calves: Clinical and Pathological Findings and Differentiation from Bovine Hereditary Zinc Deficiency

M.C. Langenmayer, S. Jung, M. Majzoub-Altweck, F.M. Trefz, C. Seifert, G. Knubben-Schweizer, R. Fries, W. Hermanns, and N.S. Gollnick

Background: Zinc deficiency-like (ZDL) syndrome is an inherited defect of Fleckvieh calves, with striking similarity to bovine hereditary zinc deficiency (BHZD). However, the causative mutation in a phospholipase D4 encoding gene (PLD4) shows no connection to zinc metabolism.

Objectives: To describe clinical signs, laboratory variables, and pathological findings of ZDL syndrome and their utility to differentiate ZDL from BHZD and infectious diseases with similar phenotype.

Animals: Nine hospitalized calves with crusting dermatitis and confirmed mutation in PLD4 and medical records from 25 calves with crusting dermatitis or suspected zinc deficiency.

Methods: Prospective and retrospective case series.

Results: The 9 calves (age: 5–53 weeks) displayed a moderate to severe crusting dermatitis mainly on the head, ventrum, and joints. Respiratory and digestive tract inflammations were frequently observed. Zinc supplementation did not lead to remission of clinical signs in 4 calves. Laboratory variables revealed slight anemia in 8 calves, hypoalbuminemia in 6 calves, but reduced serum zinc concentrations in only 3 calves. Mucosal erosions/ulcerations were present in 7 calves and thymus atrophy or reduced thymic weights in 8 calves. Histologically, skin lesions were indistinguishable from BHZD. Retrospective analysis of medical records revealed the presence of this phenotype since 1988 and pedigree analysis revealed a common ancestor of several affected calves.

Conclusions and Clinical Importance: ZDL syndrome should be suspected in Fleckvieh calves with crusting dermatitis together with diarrhea or respiratory tract inflammations without response to oral zinc supplementation. Definite diagnosis requires molecular genetic confirmation of the PLD4 mutation.

Key words: BHZD; Bovine; Genetic defect; PLD4; ZDL syndrome.

In recent years, Bavarian Fleckvieh calves, displaying marked dyseratotic dermatitis in combination with a history of recurring episodes of respiratory and enteric diseases, attracted attention. These animals showed a striking clinical similarity to calves affected by bovine hereditary zinc deficiency (BHZD), a keratinization disorder caused by a mutation in the SLC39A4 gene. However, a molecular genetic study we conducted on 7 of the affected Fleckvieh calves excluded SLC39A4 as causative gene for the syndrome. Genome-wide resequencing of 1 affected calf identified a nonsense mutation (p.W215X) in a phospholipase D encoding gene (PLD4), located on bovine chromosome 21, as most likely polymorphism. The disease has an autosomal recessive mode of inheritance. A study on the progeny descending from carriers of the mutation showed a significant reduction of the survival rates ($P = 2.97 \times 10^{-5}$). Because of the phenotypic similarity to BHZD the condition was named zinc deficiency-like (ZDL) syndrome.

Knowledge about the PLD4 protein in mammal metabolic pathways is scarce. In mice, PLD4 was identified as an intracellular transmembrane glycoprotein with potential function in phagocytosis. In humans, PLD4 variants are associated with systemic sclerosis and rheumatoid arthritis.

Detailed description of clinical and pathological aspects of the ZDL syndrome in cattle might provide important guidance for future metabolomic studies connecting geno-
3: Sternal recumbency, head up. 854 Langenmayer et al for this study. institutional and national guidelines. No ethical approval was required for normal breeding routine on Bavarian Fleckvieh farms. Animal husbandry and care were carried out in accordance with institutional and national guidelines. No ethical approval was required for this study.

Materials and Methods
Clinical Case Series

A total of 9 Fleckvieh calves (8 purebred and 1 Fleckvieh calf with an Angus grand-sire in the ancestral father line [according to owner]) admitted to the Clinic for Ruminants with Ambulatory and Herd Health Services between October 2009 and April 2015 were included. Inclusion criteria were typical zinc-deficiency-like skin alterations and a molecular genetic validation of the PLD4-status by 5′-exonuclease assay. Two calves were monitored over several weeks, but 7 calves were euthanized within a few days after admission because of animal welfare reasons. A complete clinical examination was performed daily. Blood samples for laboratory analysis were routinely taken from the jugular vein for diagnostic purposes. Two calves were treated with oral zinc supplementation (Calf 2 = 480 mg and Calf 4 = 2,500 mg elementary zinc per day). These doses were within (Calf 2) and above (Calf 4) recommendations to treat BHZD. All affected animals descended from carrier matings, which happened by chance during normal breeding routine on Bavarian Fleckvieh farms. Animal husbandry and care were carried out in accordance with institutional and national guidelines. No ethical approval was required for this study. Laboratory variables were determined as previously described. In brief, serum chemistry variables and enzyme activities were determined with an automated analyzer system and a complete blood cell count was performed with a hematology analyzer (poch-H100iV DIFF, Sysmex).

Retrospective Analysis of Medical Records

Archive cases of Fleckvieh calves under the age of 8 months with a history of crusting dermatitis or suspected zinc deficiency were identified (1988–2008) and the histology reviewed. For genomic DNA isolation, 10 µm sections of these archival formalin-fixed paraffin-embedded (FFPE) tissue samples were deparaffinized 3 × 20 minutes in xylene and rehydrated in a graded ethanol series for 2 × 15 minutes. After, sections were incubated overnight in 1× PK-buffer (20 mM Tris-HCl, 4 mM EDTA, 100 mM NaCl), 10% SDS and 2 µL Proteinase K (20 mg/mL) at 56°C followed by phenol-chloroform-DNA-extraction. For up-concentration, DNA was alcohol-precipitated and redissolved and diluted in 1× TE-buffer. The published PLD4-variant was polymerase chain reaction (PCR)-amplified with 2 separated primer-systems resulting in a 244 and 426 bp fragment, respectively (1-for: CCTTGTTCCACCTTGAAT/rev: TTACAGACTGTCCACCCTTGG and 2-for: CTTTGTCAGGTGCCAGGT/rev: TTACAGACCTGCTC ATCC). Polymerase chain reaction products were enzymatically purified and sequencing reactions were done for both strands with BigDye Terminator v1.1 Cycle Sequencing Kit. Electrophoresis of purified sequencing reactions was performed on the ABI 3130xl Genetic Analyzer. The Phred/Phrap/Polyphred software suite was used for base calling, sequence alignment, and polymorphism detection. Sequences were viewed with Consed. To avoid false results because of DNA desamination, amplification reactions were performed repeatedly.

Pedigree Information

Pedigree information of archival and recent cases was provided by the Bavarian State Research Centre for Agriculture. Visual examination of the pedigrees gave the indication of a most recent common ancestor (MRCA) among the recent and former cases.

Results

Clinical Case Series

The study involved 2 female and 7 male calves. Calves were admitted at median 81 days of age (mean: 116 days; range: 37–372 days) and necropsied at median 90 days of age (mean: 122 days; range: 38–373 days). The owners observed first skin lesions at median age of 45 days (mean: 49 days; range: 21–95 days). Calves were presented with a history of recurrent diarrhea, fever, dyspnea, and runting. Calf 5 had a female twin, which deceased with 79 days (necropsy was not conducted). The twin’s first skin lesions were detected on the head, best obvious around the muzzle and peri-orbital regions and on the ventrum. Calves 2, 4, 6, and 9 were PO supplemented with zinc before admission to the Clinic for Ruminants (elementary zinc per day: Calf 2: 480 mg for 21 days; Calf 4: 333 mg for 24 days and 2500 mg for 11 days; Calf 6: dosage unknown, treatment for 24 days; Calf 9: 10–16 mg for 150 days). However, zinc supplementation did not lead to amelioration of the clinical signs.

Table 1. Summary of clinical findings/diagnoses of 9 calves affected by zinc deficiency-like syndrome.

| Finding/Diagnosis | Number of Calves |
|-------------------|------------------|
| Body condition    |                  |
| Poor              | 4/9              |
| Moderate          | 5/9              |
| Attitude          |                  |
| Pruritus          | 2/9              |
| Bruxism           | 3/9              |
| Behavior<sup>a</sup> |                  |
| Score 1: 6/8      | Score 2: 2/8     |
| Score 3: 3/8      |                  |
| Touch-sensitivity<sup>b</sup> |            |
| Score 1: 1/8      | Score 2: 4/8     |
| Score 3: 3/8      |                  |
| Posture<sup>c</sup> |                  |
| Score 1: 1/9      | Score 2: 7/9     |
| Score 3: 1/9      |                  |
| Enlargement of lymphonodi cervicales superficiales et subiliaci | 9/9 |
| Nasal discharge   | 5/9              |
| Stridor           | 8/9              |
| Coughing          | 3/9              |
| Diarrhea          | 7/9              |
| Fever (≥39.5°C)   | 5/8              |

<sup>a</sup>1: Approaches examiner voluntarily after some time, allows being gently touched and scratched. 2: Constantly moves away, seems frightened, avoids being touched.
<sup>b</sup>1: Approaches examiner voluntarily after some time, allows being gently touched and scratched. 2: Constantly moves away, seems frightened, avoids being touched.
<sup>c</sup>1: Normal. 2: Increased. 3: Highly sensitive to touch.
The results of the clinical examinations of Calves 1–9 are summarized in Table 1. A moderate to severe exudative and crusty generalized dermatitis was present in all calves (Fig 1a,b). Skin regions with mild lesions displayed loose flakes. In moderate to severely affected regions there were large plates with rhagades, consisting of crusts, keratin, and hair (Fig 1c). Removal of single plates was not possible without injuring the epidermis. Moderate to severe lesions were predominantly localized on the head (around the muzzle, eyes, and on the forehead, Fig 1b), ventrum (Regio sternalis, R. umbilicalis), joints (R. cubiti, R. carpi, R. genus/ plicae lateralis, R. tarsi), and inguinal and axillary region. The peripheral lymph nodes were enlarged in all calves (Fig 1d). Calves 1, 4, and 5 displayed interdigital small erosions. Several calves displayed mucosal lesions and excessive salivation, 5 calves displayed yellow plaques on their incisors. Calf 9 was infested by numerous Damalinia. Seven calves displayed signs of weakness: weight shifting from 1 hind leg to the other and sudden bending of knees (5 calves), constant muscle tremor (Calf 3), and inability to rise (Calf 1). Calves 3 and 5 were extremely exhausted, did not move after being placed in lateral recumbency and fell into deep sleep immediately. Additionally, Calf 3 was introverted and chewed idly during examination.

Laboratory variables are summarized in Table 2 (excluding Calf 7, where serum and blood were not investigated). Calves 1–8 presented slight anemia. Leukocytosis was present in 4 calves. Six calves displayed lowered serum albumin concentration and increased serum AST activity. Serum zinc concentrations were below the reference range in 3 calves, whereas the other 6 were within or above reference range.

**Table 2. Laboratory findings of 8 calves affected by zinc deficiency-like syndrome.**

| Variable (unit)         | Reference range | Median | Minimum–Maximum |
|-------------------------|-----------------|--------|-----------------|
| RBCs (×10^6/μL)         | 5–8             | 8.68   | 7.09–9.33       |
| Hemoglobin (g/dL)       | 10–14           | 7.5    | 4.9–11.4        |
| Hematocrit (%)          | 30–36           | 26.25  | 18.9–34         |
| MCV (FL)                | 40–60           | 29.9   | 26.7–38.3       |
| MCHC (g/dL)             | 25.8–33.8       | 29.6   | 26.3–33.0       |
| MCH (pg)                | 14.5–22.6       | 9.0    | 7.2–12.7        |
| WBCs (×10^3/μL)         | 4–10            | 12.7   | 4.3–44.6        |
| Urea (mmol/L)           | <5.5            | 4.4    | 1.6–7.9         |
| Creatinine (μmol/L)     | <110            | 40.3   | 25.7–69.6       |
| Total bilirubin (μmol/L)| <8.5            | 5.7    | 1.7–12.6        |
| Total protein (g/L)     | 55–70           | 57.6   | 51.2–69.9       |
| Albumin (g/L)           | 30–40           | 29.1   | 23.8–32.3       |
| Globulin (g/L)          | 10–40           | 29.9   | 21.8–45         |
| AST (U/L)               | <80             | 102.7  | 49.3–136.2      |
| GGT (U/L)               | <36             | 16.2   | 11.1–30.3       |
| GLDH (U/L)              | <16             | 8.2    | 2.9–37.5        |
| CK (U/L)                | <245            | 206    | 75–554          |
| Iron (μmol/L)           | 12–44           | 6.2    | 3.2–18.7        |
| Copper (μmol/L)         | 8–39            | 8.2    | 4.3–11.6        |
| Zinc (μmol/L)           | 10–20           | 11.7   | 8.3–36.8        |

Values outside the reference range are highlighted.

AST, aspartate transaminase; CK, creatine kinase; GGT, gamma-glutamyl transferase; GLDH, glutamate dehydrogenase; MCH, mean corpuscular hemoglobin; MCHC, mean corpuscular hemoglobin concentration; MCV, mean corpuscular volume; RBC, red blood cell; WBC, white blood cell.
range. High zinc concentrations in Calves 2, 4, and 6 correlated with the oral zinc substitution. Because of the advanced course of the disease and lack of improvement, all calves were euthanized and submitted for necropsy.

At necropsy, median body weight was 80 kg (mean: 86 kg; range: 55–120 kg). The major skin lesions, severe crusting with rhagades (Fig 2a), of all calves were distributed multifocally with emphasis on above-mentioned locations. The peripheral but also often multiple organ lymph nodes were massively enlarged. Seven calves had multiple erosions or ulcers of the oral mucosa with multifocal necrosis and blunting of buccal mucosal papillae (Fig 2b). Calves additionally displayed enteritis (9/9), bronchopneumonia (7/9), and thymic atrophy (5/8; mean weight 46.8 g).

Histologically, affected areas displayed diffuse severe serocellular crusting often with many colonies of cocci and Malassezia. There was diffuse severe laminated orthokeratotic hyperkeratosis and multifocal mild to moderate parakeratosis with many intracorneal serum lakes (Fig 2c). The epidermis was multifocally hyperplastic, ulcerated, and displayed subcorneal and suprabasilar vesiculation. In some locations, multiple epidermal keratinocytes were apoptotic and few epidermal and follicular keratinocytes displayed clear cytoplasmic vacuolation. The dermal ridges displayed superficial edema and multifocal interface vacuolation in 8/9 calves. The dermal blood vessels were congested and surrounded by mixed-cell inflammation, the lymphatics and sweat glands were dilated. The severity of skin lesions revealed by histological examination reflected the gross pattern with most severe lesions in skin sections of the head, ventrum, and joints.

Histologically, the mucosa of the oral cavity and muzzle was multifocally eroded or ulcerated with many infiltrating neutrophils and many suprabasilar apoptotic keratinocytes in the deep epithelial papillae (Fig 2d).

The lymph nodes displayed increased cortical and para-cortical areas, dilated sinuses, and increased sinusoidal neutrophils and macrophages. The thymus (examined in Calves 2–9) of 7 calves displayed a moderate to marked decrease in cortical size and cellularity resulting in a shifted cortico-medullary ratio.

Interstitial nephritis was detected in 3 calves and suppurative rhinitis, suppurative tracheitis and vasculitis in 2 calves, each.

BVDV- and OHV2-PCR of spleen was negative in 8/9 and 5/5 calves, respectively. Enteric Coronavirus was detected in 1/4 calves. Salmonella enrichment of intestines was negative in 5/5 calves. Bacterial lung culture revealed 1 calf with Mannheimia spp. and 1 with Pasteurella multocida.

Retrospective Analysis of Medical Records

Medical records of 25 Fleckvieh calves with crusting dermatitis or suspected zinc deficiency were identified in the archive of the Clinic for Ruminants. FFPE tissue samples from 13/25 calves were stored in the archive of the Institute of Veterinary Pathology. The 4 female and 9 male calves were between 5 weeks and 5 months old. Clinical signs/diagnoses in these 13 calves were remarkably similar to the recent cases and included poor to moderate body condition, similar skin lesions, lymph node enlargement, mucosal erosions, and inflammation and infections of the respiratory...
and digestive tract. Zinc deficiency was suspected in 9 calves and zinc has been supplemented in 5 calves but did not ameliorate the clinical signs. Twelve calves were BVDV negative, only 1 calf was antigen-positive. Anemia and leukocytosis were present in 10 calves.

The postulated W215X-variant in PLD4 was re-sequenced in all 13 FFPE-samples. Sample age varied between 9 and 26 years, the oldest sample was from 1988, the youngest from 2005. Twelve animals were homozygous carriers of the nonsense mutation and 1 male Fleckvieh calf carried the wild-type allele (sample from 1997, sample age 17 years). Retrospective histological evaluation of skin sections suspected ZDL-syndrome in 10 cases, 2 cases were ambiguous and the wild-type calf displayed sarcoptic mange in skin sections.

**Pedigree Information**

Pedigree analysis revealed familial relationships between the calves of the recent cases and those retrospectively analyzed. The MRCA of all calves with available pedigree information was identified and traced back until 1972 (Fig 3).

**Discussion**

Recently, German and international reports about ZDL syndrome increased the awareness of owners and veterinarians and led to an increased presentation of affected calves to veterinary clinics. The clinical presentation of ZDL syndrome is mainly characterized by crusting dermatitis on head, ventrum, and joints and is therefore strikingly similar to BHZD. Lesions further include mucosal ulcerations of upper digestive tract, inflammation of respiratory and digestive tracts, thymic atrophy, and lymphadenopathy. This study shows that this phenotype and the absence of treatment responses to zinc supplementation should alert the clinician about ZDL syndrome.

The age distribution and severity of clinical and pathological changes varied among these cases. The first detection of skin lesions was reported to be recognized by the farmers between the first and 19th week after birth. The time interval between recognition of clinical signs of skin disease and hospital admission ranged from 7 to 316 days. Calves were necropsied between 38 and 373 days after birth. As the affected calves carried the same genotype and showed a similar phenotype, other factors like individual predisposition, environmental or management factors might influence disease severity considerably and might provide an explanation for the marked age differences.

The predominately mild microcytic, normochromic anemia, and mild hypoalbuminemia of the calves is very likely because of the chronic inflammatory state. Leukocytosis reflects the severe inflammatory infiltration in skin, respiratory, or digestive tract. Decreased serum zinc concentrations can be associated with chronic inflammation as well. Additionally, the chronic enteritis and the catabolic state of the calves could be the cause of hypoalbuminemia. The concentrations of the trace elements copper and iron were below reference ranges in 3/7 and 4/5 calves, respectively. The low iron concentrations can be caused by the chronic inflammatory state or reduced alimentary uptake because of disease-associated anorexia. However, the causes and connections of the metabolic disturbances in these calves are not known. As copper, iron, and zinc concentrations also have synergistic and antagonistic properties in physiological states, the reason for the observed concentration changes in this disease deserves further study.
The similarity of the ZDL phenotype to BHZD is striking, even though the genetic cause and affected breeds are clearly distinct. Cl 12 Bovine hereditary zinc deficiency, first described in Holstein Friesian calves, affects a variety of cattle breeds including Fleckvieh, Holstein Friesian, Shorthorn, and Angus. Cl 2 3 8, 18, 19

Bovine hereditary zinc deficiency-affected calves show keratinization disorders, thymus hypoplasia, respiratory and digestive tract inflammations, oral mucosal ulcerations, impaired function of the immune system, and growth retardation as consequence of a malfunction of an intestinal zinc transporter. Cl 20–22 In zinc deficiency in Holstein Friesian cattle, a splice-site variant in the SLC39A4 gene is the underlying mutation. Cl 2 This gene encodes a zinc transporter which is crucial for intestinal zinc uptake. Cl 21 In humans, several additional mutations in this gene are also responsible for an inherited zinc deficiency syndrome known as acrodermatitis enteropathica. Cl 24–26

Because of the malfunction of the intestinal zinc transporter, BHZD calves do respond to oral zinc supplementation. Cl 20, 27 In one study, clinical signs developed when plasma zinc concentration dropped below 0.5 ppm and a plasma zinc concentration of ~1.0 ppm was maintained through oral supplementation to keep the animal without clinical signs. Cl 27 In the present study, the zinc doses were within (Calf 2) and above (Calf 4) the range of published values used to alleviate or resolve clinical disease of zinc deficient calves. Cl 8, 20, 27 In both calves, serum zinc concentrations responded to the treatment and were above the maintenance zinc concentrations of Machen et al. Cl 27 Unfortunately, the exact dose given to Calf 6 is not known, however, serum zinc concentration was at the upper limit of the reference range and was above the maintenance zinc concentration of Machen et al. Cl 27 Calf 9 was treated with 10–16 mg elemental zinc per day. This dose appears really low when compared to the other calves and its effects were probably minor (low serum zinc concentrations). Although serum zinc concentrations are not always useful to determine tissue zinc concentrations, Cl 28 elevation after treatment indicates that intestinal uptake took place. In zinc-treated BHZD calves, clinical signs improve after 1–3 weeks and completely resolve after 2–6 weeks. Cl 20, 22 However, the clinical signs observed in Calves 2, 4, and 6 did neither improve nor resolve, clearly separating this syndrome from BHZD clinically. Nonetheless, zinc supplementation might help the animal to survive longer, possibly because of positive effects of zinc on skin and immune system. Cl 29

Marked hypoplasia of the thymus with an average thymus weight of 18 g has been reported in 15 affected BHZD calves. Cl 30 Thymus weights in this study revealed thymic atrophy, yet the thymus was readily identifiable and weighed nearly 3 times more than described in BHZD. Although thymus weight/body weight ratio was lower than published ratios, Cl 2 gross and histologic examination could not support a finding of “marked hypoplasia.”

Bovine hereditary zinc deficiency occurs in various cattle breeds but has its primary incidence in Holstein Friesian calves Cl 20, 27, 32 unlike ZDL syndrome which has only been observed in Fleckvieh calves.

Differential diagnoses for ZDL syndrome include mange, acquired zinc deficiency, and ichthyosis. The mucosal lesions and lesions in the interdigital space should be differentiated from bovine viral diarrhea/mucosal disease (BVD/MD), malignant catarrhal fever (MCF), and blue tongue. In the medical records from 1988 to 2005, BVD/MD was the main differential diagnosis followed by zinc deficiency and to a small amount ichthyosis or MCF.

The gene PLD4 encodes Phospholipase D4, which belongs to the PLD superfamily, comprising 6 members (PLD1–6) widely distributed in tissues of humans and animals. Cl 33 PLD1 + 2 are well characterized phospholipid signaling enzymes, involved in cellular proliferation and vesicle trafficking. Cl 3 Enzymes with PLD activity or phospholipase domains are further involved in keratinocyte metabolism or the keratinization process. Cl 34–36 However, little is known about role and function of PLD4. Localized in the spleen and early microglia, a role in immunological pathways has been suggested. Cl 4 Its enzymatic activity is still enigmatic and possible nonenzymatic functions are discussed. Cl 35 There is considerable interspecies phenotypic variation between the few diseases and syndromes linked to PLD4 so far. Recent studies associated variants in PLD4 with systemic sclerosis and rheumatoid arthritis in the Japanese population. Cl 6, 7, 37 Mice carrying a nonsense mutation in PLD4 manifest a phenotype with thin hair and small size that seems quite distinct from ZDL syndrome in calves. Cl 28, 38, 39 These substantial differences between the species highlight the importance of different models, which allow investigation of PLD4 functions and its position within metabolic pathways.

Combining the available information and the present cases, the role of PLD4 in calves is very likely associated with immunological functions and keratinocyte differentiation. Infection of the respiratory and digestive tract and skin suggest some kind of immunosuppression. Regarding PLD4 and PLD function in microglia Cl 3 and neutrophils, Cl 3 respectively, the innate immune system is likely compromised. Hyper- and parakeratosis suggest a role of PLD4 in the keratinization process and differentiation of keratinocytes.

The gold standard to distinguish ZDL from BHZD is—of course—a genetic examination. Clinicians confronted with Fleckvieh calves displaying signs of zinc deficiency should be aware of ZDL and can differentiate ZDL from BHZD through unresponsiveness to oral zinc supplementation. However, lack of improvement or advanced cases should initiate a genetic examination to quickly identify ZDL calves. On pathological examination, dermal lesions together with thymic atrophy can indicate ZDL, but atrophy must be separated from disease- or age-associated involution. Dermal histopathology is not helpful to distinguish between these 2 syndromes.

Footnotes

a Hitachi 912, Boehringer Mannheim, Mannheim, Germany
b Applied Biosystems, Foster, CA
c Applied Biosystems
Acknowledgments

We acknowledge the excellent technical assistance of Ingrid Hartmann, Michaela Nützel, Doris Merl, and Heike Sperling. We thank the animal caretakers at the Clinic for Ruminants for their dedication and the owners of the calves and herd veterinarians for their support.

Conflict of Interest Declaration: Authors declare no conflict of interest.

Off-Label Antimicrobial Declaration: Authors declare no off-label use of antimicrobials.

Institutional Animal Care and Use Committee (IACUC) or Other Approval Declaration: Authors declare no IACUC or other approval was needed.

References

1. MacPherson EA, Beattie IS, Young GB. An inherited defect in Friesian calves. Nord Vet Med Suppl 1964;1:533–540.
2. Yuzbasyan-Gurkan V, Bartlett E. Identification of a unique splice site variant in SLC39A4 in bovine hereditary zinc deficiency, lethal trait A46: An animal model of acrodermatitis enteropathica. Genomics 2006;88:521–526.
3. Jung S, Pausch H, Langenmayer MC, et al. A nonsense mutation in PLD4 is associated with a zinc deficiency-like syndrome in Fleckvieh cattle. BMC Genomics 2014;15:623.
4. Otani Y, Yamaguchi Y, Sato Y, et al. PLD4 is involved in phagocytosis of microglia: Expression and localization changes of PLD4 are correlated with activation state of microglia. PLoS One 2011;6:e27544.
5. Yoshikawa F, Banno Y, Otani Y, et al. Phospholipase D family member 4, a transmembrane glycoprotein with no phospholipase D activity, expression in spleen and early postnatal microglia. PLoS One 2010;5:e13932.
6. Okada Y, Terao C, Ikari K, et al. Meta-analysis identifies nine new loci associated with rheumatoid arthritis in the Japanese population. Nat Genet 2012;44:511–516.
7. Terao C, Ohnura K, Kawaguchi Y, et al. PLD4 as a novel susceptibility gene for systemic sclerosis in a Japanese population. Arthritis Rheum 2013;65:472–480.
8. Dirksen G, Gründer H-D, Stöber M. Innere Medizin und Chirurgie des Rindes, 5th ed. Stuttgart, Germany: Parey; 2006.
9. Langenmayer MC, Scharr JC, Sauter-Louis C, et al. Natural Bessnoitia besnoiti infections in cattle: Hematological alterations and changes in serum chemistry and enzyme activities. BMC Vet Res 2015;11:32.
10. Langenmayer MC, Gollnick NS, Majzoub-Altwdeck M, et al. Naturally acquired bovine besnoitiosis: Histological and immunohistochemical findings in acute, subacute, and chronic disease. Vet Pathol 2015;52:476–488.
11. Ewing B, Green P. Base-calling of automated sequencer traces using phred. II. Error probabilities. Genome Res 1998;8:186–194.
12. Ewing B, Hillier L, Wendl MC, Green P. Base-calling of automated sequencer traces using phred. I. Accuracy assessment. Genome Res 1998;8:175–185.
13. Nickerson DA, Tobe VO, Taylor SL. PolyPhred: Automating the detection and genotyping of single nucleotide substitutions using fluorescence-based ressequencing. Nucleic Acids Res 1997;25:2745–2751.
14. Gordon D, Abajian C, Green P. Consed: A graphical tool for sequence finishing. Genome Res 1998;8:195–202.
15. Stockholm SL, Scott MA. Erythrocytes. In: Stockholm SL, Scott MA, eds. Fundamentals of Veterinary Clinical Pathology. Ames, IA: Blackwell Publishing; 2008:107–222.
16. Stockholm SL, Scott MA. Proteins. In: Stockholm SL, Scott MA, eds. Fundamentals of Veterinary Clinical Pathology. Ames, IA: Blackwell Publishing; 2008:369–414.
17. Thrall MA. Classification of and diagnostic approach to anemia. In: Thrall MA, ed. Veterinary Hematology and Clinical Chemistry. Philadelphia, PA: Lippincott Williams & Wilkins; 2003:83–88.
18. Schlerka G, Baumgartner W. Parakeratose bei einem Höhenfleckviehkalb. Wien Tierärztl Mschr 1976;63:19–22.
19. Vogt DW, Carlson CG, Miller RB. Hereditary parakeratosis in shorthorn beef calves. Am J Vet Res 1988;49:120–121.
20. Stöber M. [Parakeratosis in black pied calves. 1. Clinical findings and etiology]. Dtsch Tierärztl Wochenschr 1971;78:257–265.
21. Trautwein G. [Parakeratosis in black pied calves. 2. Pathological findings]. Dtsch Tierärztl Wochenschr 1971;78:265–270.
22. Weismann K, Flagstad T. Hereditary zinc deficiency (Adema disease) in cattle, an animal parallel to acrodermatitis enteropathica. Acta Dermatovener 1976;56:151–154.
23. Andrews GK. Regulation and function of ZIP4, the acrodermatitis enteropathica gene. Biochem Soc Trans 2008;36:1242–1246.
24. Park CH, Lee MJ, Kim HJ, et al. Congenital zinc deficiency from mutations of the SLC39A4 gene as the genetic background of acrodermatitis enteropathica. J Korean Med Sci 2010;25:1818–1820.
25. Wang F, Kim BE, Dufner-Beattie J, et al. Acrodermatitis enteropathica mutations affect transport activity, localization and zinc-responsive trafficking of the mouse ZIP4 zinc transporter. Hum Mol Genet 2004;13:563–571.
26. Wang K, Pugh EW, Griffen S, et al. Homozygosity mapping places the acrodermatitis enteropathica gene on chromosomal region 8q24.3. Am J Hum Genet 2001;68:1055–1060.
27. Machen M, Montgomery T, Holland R, et al. Bovine hereditary zinc deficiency: Lethal trait A 46. J Vet Diagn Invest 1996;8:219–227.
28. Jeejeebhoy K. Zinc: An essential trace element for parental nutrition. Gastroenterology 2009;137:S7–S12.
29. Prasad AS. Impact of the discovery of human zinc deficiency on health. J Am Coll Nutr 2009;28:257–265.
30. Brunnerstedt E. Animal model of human disease. Acrodermatitis enteropathica, zinc malabsorption. Am J Pathol 1977;87:725–728.
31. Lubis I, Ladds PW, Reilly LR. Age associated morphological changes in the lymphoid system of tropical cattle. Res Vet Sci 1982;32:270–277.
32. Andresen E, Flagstad T, Basse A, Brunnerstedt E. Evidence of a Lethal Trait, A 46, in black pied Danish cattle of Frisian descent. Nord Vet Med 1970;22:473–485.
33. Frohman MA. The phospholipase D superfamily as therapeutic targets. Trends Pharmacol Sci 2015;36:137–144.
34. Arun SN, Xie D, Howard AC, et al. Cell wounding activates phospholipase D in primary mouse keratinocytes. J Lipid Res 2013;54:581–591.
35. Grall A, Guaguere E, Plancais S, et al. PNPLA1 mutations cause autosomal recessive congenital ichthyosis in golden retriever dogs and humans. Nat Genet 2012;44:140–147.
36. Bollinger Bollag W, Bollag RJ. 1,25-Dihydroxyvitamin D3, phospholipase D and protein kinase C in keratinocyte differentiation. Mol Cell Endocrinol 2001;177:173–182.
37. Jin J, Chou C, Lima M, et al. Systemic sclerosis is a complex disease associated mainly with immune regulatory and inflammatory genes. Open Rheumatol J 2014;8:29–42.
38. Harris B, Ward-Bailey PF, Bergstrom DE, et al. Thin hair with small size (thss) is a new recessive hair mutation on Chromosome 12. Nat Genet 2001;27:432–434.
39. Harris BS, Fairfield HE, Gilbert GJ, et al. Thin hair with small size is the first Pkd4 mutant mouse. Available at: http://www.informatics.jax.org/reference/J:179503. Accessed November 30, 2017.
40. Harris BS, Fairfield HE, Gilbert GJ, et al. Thin hair with small size is the first Pkd4 mutant mouse. Available at: http://www.informatics.jax.org/reference/J:179503. Accessed November 30, 2017.