Selenium Status in Heifers, Late Pregnancy Cows and Their Calves in the Šumava Region, Czech Republic

P. SLAVÍK¹, J. ILLEK², T. ZELENÝ³

¹Department of Animal Nutrition, ²Clinic of Ruminant Diseases, University of Veterinary and Pharmaceutical Sciences, Brno, Czech Republic ³Veterinary Centre, Sušice, Czech Republic

Received February 7, 2006
Accepted November 15, 2007

Abstract


The objective of this study was to ascertain selenium status in beef cattle in different stages of production in the Šumava region. In the region, blood collections and analyses for selected metabolic variables were performed in 54 animals in different production stages (18 heifers, 18 cows in late pregnancy and 18 calves aged 3 weeks on the average). Three herds were studied. The selenium status was determined both directly by measuring serum selenium (Se) contents and indirectly by measuring glutathione peroxidase (GSH-Px) activity in whole blood. The mean serum selenium concentration in all the animals under study (n = 54) was 30.6 µg/l +/- 2.91, and mean GSH-Px activity was 167.01 µkat/l +/- 92.39. In heifers, mean serum selenium concentration was 34.81 µg/l +/- 13.84; mean GSH-Px activity was 186.96 µkat/l +/- 112.15. In late pregnancy cows, mean serum selenium concentration was 26.58 µg/l +/- 8.01, mean GSH-Px activity was 94.55 +/- 35.72 µkat/l. In calves, mean serum selenium concentration and GSH-Px activity were 30.41 µg/l +/- 12 and 219.54 µkat/l +/- 64.41, respectively. There was a statistically significant difference between the heifers and late pregnancy cows in both variables under study. However, between the late pregnancy cows and the calves, only the difference in GSH-Px activity was significant. The results indicate severe Se deficiency in the animals under study. It means apart from other things that mineral licks used did not provide enough minerals to meet the basic requirements of the animals.

Cattle, calves, selenium, beef cattle, glutathione peroxidase

Based on the socioeconomic changes that have led to fundamental agricultural reforms, extensive beef cattle rearing systems have started to develop in marginal agricultural regions as one of the ways of environmentally friendly landscape management. Although this type of production is not in the forefront of scientific interest, there are many serious problems of a factor nature, mainly in calf rearing. This study focused on the problems related to selenium. The objective of this study was to ascertain the selenium status in cattle of different production stages in the region under observation. First reports on the occurrence of selenium deficiency in this region and its impact on the health status were published more than 30 years ago (Kursa 1969).

Selenium performs many functions in the body. These include an effect on the development and motility of sperm, influence on immunity indicators (improvement of bactericidal activity of neutrophile granulocytes, increase in antibody production), effect on fertility (fewer cases of retained placenta) (Underwood and Suttle 1999). Selenium is a natural antioxidant. It protects organellae and cell membranes containing lipids from oxidation by reactive oxygen species (ROS). ROS include for instance superoxide anion, hydroxyl radical, perhydroxyl radical, hydrogen peroxide and singlet oxygen (Baldi 2005). Functional forms of selenium are selenoproteins. The most important selenoproteins are glutathione peroxidase (GSH-Px), thioredoxine reductase (TR), iodothyronine-5-deiodinase, selenoprotein P and tens of others (Underwood and Suttle 1999).
Many researchers have demonstrated a positive correlation between selenium concentrations and GSH-Px activity in the blood.

In cattle, the correlation was demonstrated by Pavlata et al. (2001) for example, who determined a correlation coefficient of 0.93, using 44 heads of cattle, and Pavlata et al. (2000) who received a correlation coefficient of $r = 0.90$ with 326 animals. The results of their studies imply that the determination of selenium status based on GSH-Px activity measurement can be used in herds where all the animals receive equal diets.

Because selenium is incorporated in erythrocytes in the form of selenocysteine only during erythropoiesis, GSH-Px activity becomes an indicator of long-term selenium status in the body. On the contrary, Se concentrations in whole blood reflect a momentary selenium status (Harapin et al. 2000; Enjalbert et al. 1999).

Selenium passes the placental barrier and is important for the intrauterine development of calves. Obviously, this Se function is very important, because even if the cow is moderately Se deficient, the calf receives a sufficient Se supply. However, this is at the expense of increasing Se deficiency in the cow. However, when the cow suffers from severe Se deficiency, the calf is affected, too. The Se deficiency in the calf will become more severe in the course of time because Se levels in the colostrum and milk are correlated with the selenium status of the cow (Enjalbert et al. 1999; Gunter et al. 2003). This, however, was not demonstrated by Pavlata et al. (2003), who monitored the correlation between the selenium status of the cow and selenium levels in her colostrum. However, only a few animals were included in the study.

In the body of a pregnant female, mainly in the last third of pregnancy, the uterus is given priority to other organs in the distribution of some substances, according to the homeorhesis principle (VanSaum et al. 1989).

The basic selenium level in beef cattle diets, recommended by NRC (National Research Council) is 0.3 mg Se per kg dry matter (NRC 2001).

Stowe and Herdt (1992) considered 70 - 100 µg/l as normal, and 40 - 70 µg/l as marginal levels. Nevertheless, they pointed out that production performance must be taken into account when evaluating Se levels. According to Gerloff (1992), Se levels under 40 µg/l indicate Se deficiency in cattle. Pavlata et al. (2000) reported slightly different values (see below).

The objective of this study was to clarify the situation concerning supplying of selenium in extensive raising of beef cattle in a region where this system of agricultural production is very widespread.

Materials and Methods

Animals included in this study were chosen from 3 beef cattle herds in the Šumava region. In total, 36 or 54 animals in three different production stages were included. The average altitude of the farms was 450 m.

Herd 1: Piemontaise breed (20 animals in the herd), Herd 2: Simmental (45 animals), Herd 3: Hereford (86 animals).

In the spring of 2004, before the start of the grazing season the herds were subjected to performance checks, and veterinary treatments were performed. Blood samples were taken as a part of the veterinary control.

In each herd, two groups of animals were randomly selected, each including 6 animals. The first group included non-pregnant young heifers only. The second group included late pregnancy cows that were due very soon. Calves born to these cows were blood-sampled at an average age of three weeks. All the calves ingested colostrum spontaneously in the first hours of life. None of them were administered a selenium-vitamin supplement parenterally between birth and blood collection.

The animal health status was examined, too. However, we did not observe marked or significant mass occurrence of signs typical of mineral deficiencies.

Animals in all the herds had free access to the mineral lick BIOXANON®; the composition of which is given in Table 1. In all the herds, the feeding of the vitamin premix VITAMIX S8 (BIOFAKTORY) was continued. The premix composition is given in Table 2. Its intake was irregular, and the mean daily intake was less than 20 g per head (the manufacturer recommends 50 - 150 g per head per day), as calculated from the total consumption in a given period.

In the summer, animals graze in the pasture, and in the winter they are fed hay and grass silage that is produced in the region. In the winter, in herd 3, corn silage is fed occasionally. Samples of the winter diet were collected, i.e., of hay and grass silage for each herd, and the average selenium contents were determined.
All the blood collections for one herd were performed at one time. Blood was withdrawn from the coccygeal and jugular veins, using the HEMOS® system, both to obtain serum and, using heparin, to determine GSH-Px activity.

After the GSH-Px determination, the heparinised whole blood samples were placed in a thermal box, and then in a refrigerator when we returned to the laboratory. The blood samples intended for serum selenium determination were allowed to coagulate at room temperature. The samples were processed in the laboratory on the next day after each collection. The laboratory analyses were made, using the COBAS MIRA analyser. Selenium levels were measured by the atomic absorption spectrophotometer (AAS) using the hydride method. The activity of glutathione peroxidase (GSH-Px) in the erythrocytes was measured using the method described by Paglia and Valentine (1967) and the RANDOX-RANSEL RS 505 kit was used.

Selenium concentration in the serum was determined by hydride generation and atomic absorption spectrophotometry (HG-AAS) as described by Sturman (1985). Mineralization was performed in a microwave oven.

For GSH-Px activity and selenium evaluation, the following reference values were used (Pavlata et al. 2000): < 70 µg/l, or < 472 µkat/l - deficiency from 70 to 100 µg/l or from 472 to 665 µkat/l - marginal status 100 µg/l or < 665 µkat/l - adequate status.

Data were processed using Microsoft Excel. Mean value (x) and standard deviation (S.D.) were calculated, and t-test was performed to compare groups of late pregnancy cows and their calves (P < 0.05).

### Results and Discussion

The mean selenium concentration in all (n = 54) animals under study was 30.6 µg/l +/- 2.91 in blood serum and mean GSH-Px activity was 167.01 µkat/l +/- 92.39 in whole blood. The mean values of Se concentration in GSH-Px activity in different animal categories are given in Table 3.
The results show that the selenium status in the animals under study was poor. Three herds were monitored and in none of them did mean Se contents and GSH-Px activities achieve marginal values. Only in herd 3, some of the individual measured values approached the marginal status level. However, in neither herd the marginal levels were exceeded.

In feed samples collected in herd 1, mean dietary selenium content was 0.1 mg/kg dry matter (DM), in herd 2, mean Se content was 0.093 mg/kg DM, and in herd 3, mean Se content was 0.088 mg/kg DM.

In the Šumava region, beef cattle is mostly grazed, with emphasis placed on extensive rearing with minimum costs. Except for mineral licks and sporadic provision of mineral feed, animals receive no mineral supplements. Our results clearly show that the Se contained in hay and grass in the pasture cannot meet their requirements.

Nevertheless, Stowe and Herdt (1992) and Gerloff (1992) reported that it is important to take into account the milk yield, which is much lower in beef cattle. On the other hand, beef cows have higher requirements for muscle growth. Specific Se levels for beef cattle in the serum, tissues and other body fluids have not been determined. However, the mean values obtained indicate severe deficiency as compared with Gerloff (1992). Only in herd 3, did individual values in some animals exceed the Se serum content of 40 µg/l.

In accordance with data reported (Enjalbert et al. 1999; Gunter et al. 2003), highest concentrations were found in heifers, i.e. non-pregnant animals. Significantly lower values were received from the t-test analysing differences in Se concentrations ($P < 0.05$) and GSH-Px activities ($P < 0.01$) between heifers and late pregnancy cows. We did not manage to find significant differences in Se concentrations between late pregnancy cows and their calves, although absolute values in calves were higher than those in their mothers. There was a significant difference in GSH-Px activity ($P < 0.01$) between the groups.

Pavlata et al. (2003) demonstrated a correlation between Se serum concentrations in pregnant cows and their calves. This is in accordance with the results obtained by Illek et al. (2002), who reported higher Se concentrations in calves than in their mothers. However, in all the groups under study, selenium deficiency at the time of calving was only moderate or marginal. However, the above studies did not analyse whether selenium contents in the newborn calves were really significantly higher than in severely Se-deficient mothers. The results of the present study indicate that the homeorhesis principle cannot be confirmed when the mother is severely Se-deficient. High GSH-Px values in calves do not correspond to the correlation coefficients determined by Pavlata et al. (2000, 2001). In both studies, GSH-Px was analysed in whole blood. Thus, it was GSH-Px that is present in cytosol of erythrocytes. For more detailed elucidation of the

| Table 3. Mean plasma Se concentrations and GSH-Px activities in the animals under study. (x - mean, S.D. - standard deviation of the mean, P - t-test) |
|---|---|---|---|---|---|---|
| Heifers (n = 18) | Cows in late pregnancy (n = 18) | Calves (n = 18) |
| Se | GSH-Px | Se | GSH-Px | Se | GSH-Px |
| µg/l | µkat/l | µg/l | µkat/l | µg/l | µkat/l |
| x   | 34.81  | 186.96 | 26.58  | 94.55  | 30.41  | 219.54 |
| S.D. | 13.84  | 112.15 | 8.01   | 35.72  | 12     | 64.41  |
| Max. | 69.81  | 509    | 43.74  | 182.3  | 59.12  | 303.4  |
| Min. | 19.13  | 82.21  | 13.49  | 48.3   | 11.06  | 15.3   |
| $p^a$ | *      | **     |        |        | -      | **     |
| $p^b$ |        |        |        |        |        |

$P < 0.05$

$**P < 0.01$

$p^a$ t-test between the groups of heifers and the cows in late pregnancy

$p^b$ t-test between the groups of the cows in late pregnancy and their calves
contradiction between the studies, it would be useful to know the haematocrit values and other haematological indicators, both in cows and their calves. It is known that newborn calves show higher haematocrit values. Foetal erythropoiesis that differs a little from that of a newborn can play a role, too. Some researchers expressed the activity of GSH-Px in E.U./kg, i.e. enzyme units per gram of haemoglobin, defined as enzyme activity level required for oxidation of 1 μmol NADPH per minute at 21 °C (Rowntree et al. 2004), taking into account the amount of haemoglobin and thereby erythrocyte counts, which is not considered if the enzyme activity is expressed in μkat per l.

Rowntree et al. (2004) found some irregularities in GSH-Px concentration. They ascribed them to the effects of breed or environment. Awadeh et al. (1998) also reported discrepancies in GSH-Px activity and Se concentration, however, in their opinion these were due to the irregular intake of Se from a lick. This fact could have played a certain role in heifers and late pregnancy cows in this study. The results suggest that the use of GSH-Px as an indirect measure of selenium status in the production stages of cattle investigated is somewhat disputable.

The severe selenium deficiency ascertained is not surprising because the dietary selenium level recommended by NRC (2001) is 0.3 mg/kg DM and the values we measured were lower than onethird of the recommended level. Such levels (including Se supplemented by mineral licks) cannot cover even the marginal requirements of the animal. The difference between the required and toxic levels, however, is very small.

It can be stated that when cows are raised extensively on pasture and their diet is not regularly supplemented with concentrated feeds in which a mineral premix could be added, they suffer from selenium deficiency. A mineral lick with increased selenium contents is not sufficient to help achieve an adequate selenium status.

Acknowledgement

The study was supported by the Ministry of Agriculture of the Czech Republic (grant No. QF 4005)

References