Effects of time and sampling location on concentrations of β-hydroxybutyric acid in dairy cows

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ABSTRACT

Two trials were conducted to examine factors potentially influencing the measurement of blood β-hydroxybutyric acid (BHBA) in dairy cows. The objective of the first trial was to study effects of sampling time on BHBA concentration in continuously fed dairy cows. Furthermore, we determined test characteristics of a single BHBA measurement at a random time of the day to diagnose subclinical ketosis considering commonly used cut-points (1.2 and 1.4 mmol/L). Finally, we set out to evaluate if test characteristics could be enhanced by repeating measurements after different time intervals. During 4 herd visits, a total of 128 cows (8 to 28 d in milk) fed 10 times daily were screened at 0900 h and preselected by BHBA concentration. Blood samples were drawn from the tail vessels and BHBA concentrations were measured using an electronic BHBA meter (Precision Xceed, Abbott Diabetes Care Ltd., Witney, UK). Cows with BHBA concentrations ≥0.8 mmol/L at this time were enrolled in the trial (n = 92). Subsequent BHBA measurements took place every 3 h for a total of 8 measurements during 24 h. The effect of sampling time on BHBA concentrations was tested in a repeated-measures ANOVA repeating sampling time. Sampling time did not affect BHBA concentrations in continuously fed dairy cows. Defining the average daily BHBA concentration calculated from the 8 measurements as the gold standard, a single measurement at a random time of the day to diagnose subclinical ketosis considering commonly used cut-points (1.2 and 1.4 mmol/L). Finally, we set out to evaluate if test characteristics could be enhanced by repeating measurements after different time intervals improved test characteristics only slightly. In the second experiment, we compared BHBA concentrations of samples drawn from 3 different blood sampling locations (tail vessels, jugular vein, and mammary vein) of 116 lactating dairy cows. Concentrations of BHBA differed in samples from the 3 sampling locations. Mean BHBA concentration was 0.3 mmol/L lower when measured in the mammary vein compared with the jugular vein and 0.4 mmol/L lower in the mammary vein compared with the tail vessels. We conclude that to measure BHBA, blood samples of continuously fed dairy cows can be drawn at any time of the day. A single measurement provides very good test characteristics for on-farm conditions. Blood samples for BHBA measurement should be drawn from the jugular vein or tail vessels; the mammary vein should not be used for this purpose. Key words: subclinical ketosis, β-hydroxybutyric acid, diurnal rhythm, herd health

INTRODUCTION

During the transition period, dairy cows have to adapt to increased energy demands resulting from late gestation and the onset of lactation after calving (Goff and Horst, 1997) while DMI is reduced. Subsequently, a negative energy balance (NEB) occurs in most cows (Herdt, 2000). The resulting lipolysis causes elevated levels of serum NEFA and, if their complete oxidation in the liver fails due to an overload, levels of serum ketone bodies (i.e., BHBA, acetoacetate, acetone) also increase (Krebs, 1966; Herdt, 2000). Subclinical ketosis (SCK) is one of the most important metabolic diseases in high-producing dairy cattle and is defined as the presence of elevated concentrations of circulating ketone bodies without the appearance of clinical signs (Andersson, 1988). First-parity heifers and second-parity cows have a reduced risk for SCK compared with older cows (Duffield et al., 1998). Subclinical ketosis causes economic losses (Duffield, 2000; Geishauser et al., 2001; Hogeveen, 2012) and increases the risk of subsequent diseases; for example, displaced abomasum (LeBlanc et al., 2005; Duffield et al., 2009), metritis (Duffield et al., 2009), clinical ketosis (Duffield et al., 2009; Seifi et al., 2011), and lameness (Sutthar et al., 2013). It is well documented that SCK negatively affects reproductive performance (Koller et al., 2003; Walsh et al., 2007a,b; Ospina et al., 2010b) and milk yield (Ospina et al., 2010b). Roberts et al. (2012) found an increased risk of culling within the first 60 DIM for cows with elevated BHBA concentrations.
Subclinical ketosis has been described as a threshold disease (Oetzel, 2004) and the diagnosis depends on a cut-point (i.e., 1.2 to 1.4 mmol/L; Duffield et al., 1998; Enjalbert et al., 2001; Oetzel, 2004). Therefore, diagnostic methods have to determine the concentration of ketone bodies semiquantitatively or quantitatively in different body fluids. Measurement of blood BHBA concentration in laboratories is the gold standard test for diagnosis of SCK (Oetzel, 2004). In addition, a range of cow-side ketone body tests is available. Several semiquantitative urine or milk tests have been evaluated (Geishauser et al., 1998, 2000; Carrier et al., 2004). An electronic patient-side test measuring whole-blood BHBA concentration (Precision Xceed, Abbott Diabetes Care Ltd., Witney, UK) was recently validated for use in dairy cows (Iwersen et al., 2009) and sheep (Panousis et al., 2012). Its main advantages include an immediate quantitative measurement with high sensitivity and specificity without logistical efforts for laboratory analysis (e.g., cooling, centrifugation, or shipping).

Frequently used cut-points for the concentration of BHBA in serum of dairy cows are 1.2 mmol/L (Duffield et al., 1998; Enjalbert et al., 2001; Suthar et al., 2013) and 1.4 mmol/L (Geishauser et al., 2001; Oetzel, 2004). Such cut-points, however, are biased by sampling time relative to calving and cut-point criteria such as disease prevalence, milk production, or both (Duffield et al., 2009). Several authors recommend strategic programs for monitoring dairy herds for magnitude of NEB, SCK, and subsequent diseases (Duffield, 2000; Oetzel, 2004; LeBlanc et al., 2005).

To implement screening protocols with reliable and repeatable results, information on factors potentially influencing the measurement of BHBA is essential. Oetzel (2004) supposed that the blood sampling location affects the BHBA concentration. He recommended that blood samples to measure BHBA concentration should not be collected from the mammary vein (V. epigastrica cranialis superficialis) because the mammary gland is not be collected from the mammary vein (V. epigastrica cranialis superficialis) because the mammary gland is able to utilize BHBA as an energy source (Kronfeld et al., 1968), which might affect venous BHBA concentration. One previous study observed 9.3% lower BHBA concentrations in blood samples obtained from the mammary vein compared with the jugular vein (Redetzky et al., 2003). Sample size, however, was limited (n = 12) and the difference was not significant in this study. Wilhelm et al. (2013) also found higher concentrations of BHBA in samples taken from the jugular vein and saphenous veins compared with the mammary vein in lactating dairy cows. However, the tail vessels were not considered in either of these trials. Overall, there is a dearth of information on the effect of commonly used sampling locations (i.e., jugular vein, mammary vein, tail vessels) on BHBA concentrations.

Furthermore, several authors have reported effects of diurnal patterns (Andersson, 1988; Plaizier et al., 2005) or sampling time relative to feeding (Oetzel, 2004; Nikkhah et al., 2008; Quiroz-Rocha et al., 2010) on blood BHBA concentrations. The diurnal rhythm of BHBA was found to be associated with feeding frequency (Eicher et al., 1999). The latter authors recommended collection of 2 samples (morning and afternoon) to enhance the assessment of metabolic situations. An increased feeding frequency reduces both the maximum concentration and the amplitude of daily fluctuations in concentrations of BHBA (Sutton et al., 1988). Information is lacking on diurnal patterns of BHBA concentration under conditions of a continuous TMR feeding regimen and how sampling time and frequency of testing affect test characteristics (i.e., sensitivity and specificity). Therefore, the overall objective of this study was to analyze diurnal effects on BHBA concentrations for cows with a continuous TMR feeding. Specifically, we set out to (1) study effects of sampling time on BHBA concentration in continuously TMR-fed dairy cows, (2) determine test characteristics (sensitivity, specificity, positive and negative predictive values) of a single BHBA measurement at a random time of the day to diagnose SCK, (3) evaluate if test characteristics can be enhanced by repeating measurements, and (4) compare BHBA concentrations of samples drawn from 3 different blood sampling locations.

**MATERIALS AND METHODS**

Two trials were conducted between April and July 2012 on a commercial dairy farm (Sachsen-Anhalt, Germany) that housed approximately 1,200 Holstein-Friesian dairy cows with an average annual milk yield of 10,701 kg (4.04% fat and 3.35% protein). Cows were housed in a freestall barn with slatted floors and cubicles equipped with rubber mats. Cows were fed a TMR consisting of 38.5% corn silage, 35.9% concentrate mineral mix, 22.5% grass silage, and 3.1% barley straw on a DM basis (NE\textsubscript{L} = 7.15 MJ/kg of DM). Feed was delivered over a conveyer belt system 10 times per day. Cows were milked 3 times a day at approximately 0630, 1430, and 2230 h in a 52-stall rotary milking parlor (Lemmer-Fullwood GmbH, Lohmar, Germany).

In both trials, BHBA was measured with an electronic BHBA meter (Precision Xceed, Abbott Diabetes Care Ltd.) that has been used in multiple research trials (Iwersen et al., 2009; Endecott et al., 2012; McArt et al., 2012). The test system consists of the handheld device and test strips, which are equipped with an
electrochemical reaction field at the tip. Results are displayed in millimoles per liter.

The purpose of trial 1 was to evaluate the effects of sampling time on BHBA concentrations. For this trial, the farm was visited 4 times at intervals of 3 wk each. All multiparous cows within 8 to 28 DIM at the time of a visit were considered eligible and a total of 128 cows were screened. Initial screening took place at approximately 0900 h. Blood samples were collected from the tail vessels using a 2-mL syringe (Henry Schein Inc., Melville, NY) and a 23-gauge needle (Sterican, B. Braun, Melsungen, Germany). Blood was analyzed immediately after sampling by applying a small amount of blood from the syringe onto the test strip. Displayed BHBA concentrations were recorded. Cows with a BHBA concentration ≥0.8 mmol/L at 0900 h were enrolled in the trial. Subsequent BHBA measurements took place every 3 h for a total of 24 h (i.e., 1200, 1500, 1800, 2100, 2400, 0300, and 0600 h). A total of 92 cows (DIM: 18 ± 6, parity: 3.2 ± 1.2) met the inclusion criteria (i.e., multiparous cow, DIM 8 to 28, BHBA concentration ≥0.8 mmol/L at 0900 h) and were used for the analyses.

Trial 2 was conducted to compare BHBA concentrations of samples drawn from 3 different blood sampling locations. Blood samples were taken from 116 postpartum cows (DIM: 19 ± 13, parity: 2.9 ± 1.2). Samples were taken within 3 ± 1 min in each cow from the tail vessels (A./V. coccygea), the jugular vein (V. jugularis), and the mammary vein (V. epigastrica cranialis superficialis) using an evacuated tube system (Vacutte, Greiner bio one, Kremsmünster, Austria). Whole blood samples were analyzed immediately after collection using the Precision Xceed handheld meter and test strips by connecting the reaction field at the end of the test strip with the surface of the blood. Displayed BHBA concentrations were recorded.

All data were statistically analyzed with SPSS (version 19.0, SPSS Inc., Munich, Germany). The effect of sampling time on measured BHBA concentrations was tested in a repeated-measures ANOVA repeating sampling time. In a second approach, test characteristics of a single BHBA measurement at a random time of the day to diagnose SCK were determined using 2 different cut-points (1.2 and 1.4 mmol/L) that are commonly used to define SCK. The average daily BHBA concentration calculated from the 8 measurements was used as the gold standard to define a cow as healthy or as having SCK. Cows were defined as healthy when the average daily BHBA concentration was lower than the respective cut-point (1.2 or 1.4 mmol/L) and as having SCK when the cut-point was reached. Sensitivity was calculated as the proportion of measurements that correctly diagnosed a cow with SCK as having SCK (i.e., BHBA concentration measured at the particular time of the day was equal to or above the cut-point considered and the daily average BHBA concentration was equal to or above the cut-point considered). Specificity was calculated as the proportion of measurements that correctly diagnosed a cow without SCK as healthy (BHBA concentration measured at the particular time of the day was below the cut-point considered and the daily average BHBA concentration was below the cut-point considered). The positive predictive value was determined as the proportion of measurements equal to or greater than the cut-point considered at which the cow was truly suffering from SCK (daily average BHBA concentration was equal to or above the cut-point considered). The negative predictive value was calculated as the proportion of measurements below the cut-point considered at which the cow was truly healthy (daily average BHBA concentration was below the cut-point considered). A second analysis was conducted to study if the test characteristics increased in marginal cases (i.e., BHBA concentrations from 1.0 to 1.6 mmol/L) using values obtained by repeated measurements instead of a single measurement. Cows having BHBA concentrations <1.0 mmol/L or >1.6 mmol/L at the given time of the day were classified as clearly healthy or as clearly subclinically ketotic, respectively, and the initial measured concentration was used for analysis. For measurements with marginal BHBA concentrations at the first sampling (i.e., 1.0 to 1.6 mmol/L), a second measurement 3, 6, 9, 12, 15, 18, or 21 h later was used and the average with the initial measurement calculated. Subsequently, this average was used to classify a cow as healthy or suffering from SCK. Test characteristics (sensitivity, specificity, positive and negative predictive values) were calculated as described above. Again, the average daily BHBA concentration was used as the gold standard to define a cow as healthy or as having SCK.

Relationships between BHBA concentrations measured at the 3 different sampling locations were calculated with Pearson correlation. The differences between the sampling locations were determined with a one-way ANOVA. Mean differences were determined with the least significant difference post hoc test. Because a comparison of clinical measurements can be inappropriate using Pearson correlation, the agreement between the 3 sampling locations was graphically analyzed with MedCalc (version 12.0.3.0, MedCalc Software bvba, Ostend, Belgium) as described by Bland and Altman (1986).

RESULTS

Trial 1

Of 128 eligible cows screened before the trial, 92 showed BHBA concentrations ≥0.8 mmol/L at the first
measurement at 0900 h and were included in the trial. Of all cows, 52 (40.6%) were diagnosed with SCK using a cut-point of 1.2 mmol/L. Increasing the cut-point to 1.4 mmol/L reduced the number of cows diagnosed with SCK to 38 (29.7%).

Sampling time did not affect BHBA concentrations in continuously fed dairy cows \( (P = 0.23; \text{Figure 1}) \). Test characteristics (sensitivity, specificity, positive and negative predictive values) of a single measurement of BHBA concentration at a random time of the day to diagnose SCK, using 2 different cut-points, are shown in Table 1. Furthermore, test characteristics using repeated measurements after different time intervals (3, 6, 9, 12, 15, 18, and 21 h) in marginal cases (BHBA concentrations from 1.0 to 1.6 mmol/L) are shown (Table 1).

**Trial 2**

A total of 116 postpartum cows were included in the trial. Considering BHBA concentrations measured in blood samples drawn from the tail vessels, 54 cows (46.6%) were diagnosed with SCK using a cut-point of 1.2 mmol/L. Using a cut-point of 1.4 mmol/L reduced the number of cows diagnosed with SCK to 38 (32.8%).

Concentrations of BHBA differed when comparing samples from the 3 sampling locations (tail vessels: 1.6 ± 1.2 mmol/L, jugular vein: 1.5 ± 1.2 mmol/L, mammary vein: 1.2 ± 1.1 mmol/L; \( P = 0.03 \)). Mean BHBA concentration was 0.3 mmol/L lower when measured in the mammary vein compared with the jugular vein \( (P = 0.03; \text{Figure 2A}) \). Mean BHBA concentration was 0.4 mmol/L lower when measured in the mammary vein compared with the tail vessels \( (P = 0.02; \text{Figure 2B}) \). Measured BHBA concentration did not differ significantly between the jugular vein and the tail vessels \( (P = 0.82; \text{Figure 2C}) \). Concentrations of BHBA were strongly correlated for every combination of sampling locations (jugular vein and mammary vein: \( r = 0.98 \); jugular vein and tail vessels: \( r = 0.99 \), mammary vein and tail vessels: \( r = 0.99; P < 0.05 \)).

**DISCUSSION**

In both trials conducted during this study, the prevalence of SCK was higher than the range (11.2 to 36.6% at a cut-point of 1.2 mmol/L) described in previous studies (McArt et al., 2012; Suthar et al., 2013). For
the first trial, one possible reason for this difference might be the selection of multiparous cows (parity 3.2 ± 1.2) only. First-parity heifers and second-parity cows are at lower risk of developing SCK (Duffield et al., 1998); thus, a higher prevalence of SCK is plausible in a group consisting of multiparous cows only. For the second trial, only 10 primiparous and 106 multiparous cows (parity 2.9 ± 1.2) were enrolled, which likewise increased the prevalence of SCK. Multiple factors have been identified that influence the development of SCK, leading to variation in prevalence rates between farms and countries (Suthar et al., 2013). Several authors found associations between ketosis and transition cow management, feeding strategies, genetic variation, milk yield, concomitant diseases, and body condition score patterns during the transition period and early lactation (Oetzel, 2007; Roche et al., 2009; van der Drift et al., 2006; Winsten et al., 2010). Diurnal feed alley attendance patterns and feeding behavior of TMR fed lactating dairy cows are affected by return from milking, feed push-ups, and, particularly, fresh feed delivery (DeVries et al., 2003; DeVries and von Keyserlingk, 2005; Mäntysaari et al., 2006). An increase in daily feeding time, as well as a more even distribution of feeding time during the day, was found in dairy cows when increasing the TMR feeding frequency (DeVries et al., 2005). These findings might explain a more continuous BHBA metabolism and consequently the absence of diurnal patterns, as described in this study. Our BHBA data underline the importance of feeding strategies that assure constant access to freshly delivered TMR.

To our knowledge, this is the first trial studying cows preselected by BHBA concentration as being at risk for SCK. Other studies did not implement such an inclusion criterion for the enrollment of study animals; consequently, most cows investigated had BHBA concentrations below current accepted cut-points for diagnosis of SCK.

**Test Characteristics of a Single Measurement of BHBA at a Random Time of the Day**

Eicher et al. (1999) recommended standardizing the time after feeding for sample collection and suggested 4 to 5 h after the beginning of feeding as a proper sampling time for testing for metabolic disorders. Although plausible, this hypothesis was not tested. On the contrary, the time before the morning feeding has been described as being the most useful to test cows in early lactation for parameters of NEB (Wylie et al., 2008). In the latter study, however, only primiparous cows were used and they were fed only once daily. The scope of our study was to investigate diurnal effects in multiparous cows under conditions of a continuous TMR feeding.
A single measurement of BHBA concentration at a random time of the day enabled the correct classification of a cow as suffering from SCK with a sensitivity of 0.90 (≥1.2 mmol/L) or 0.89 (≥1.4 mmol/L), respectively. Specificity was 0.88 or 0.90 using the same cut-points. These test characteristics can be considered as very good for a cow-side test. Based on our results, screening dairy cows for SCK under conditions of a continuous TMR feeding regimen can be integrated in a fresh cow protocol at any time of the day.

**Test Characteristics of Repeated Measurements**

Eicher et al. (1999) presumed the collection of 2 samples (morning and afternoon) to be more informative regarding the metabolic status of a herd. Sensitivity and specificity of repeated measurements after different time intervals to diagnose SCK (Table 1), however, increased only slightly compared with a single measurement at a random time of the day. Hence, from our data, the preferred sampling interval can be chosen. As these improvements are practically negligible, we feel that test characteristics of a single measurement are sufficient for on-farm situations.

**BHBA Concentrations of Samples Drawn from 3 Different Blood Sampling Locations**

It has been reported that blood samples for measurement of BHBA concentration should not be collected from the mammary vein (Oetzel, 2004; Wilhelm et al., 2013) but instead from the tail vessels (LeBlanc, 2006). It is known that the lactating mammary gland extracts BHBA as a precursor for milk fat from the blood, leading to an arterio-venous difference in BHBA concentration (Kronfeld et al., 1968; Guinard-Flament et al., 2011). The most important factor affecting the extent of this uptake is the arterial BHBA concentration (Miller et al., 1991).

Wilhelm et al. (2013) compared BHBA concentrations in blood samples taken from 4 different sampling locations (mammary vein, jugular vein, and left and right V. saphena externa lateralis). The authors found higher concentrations of BHBA in samples taken from the jugular vein and Vv. saphenae compared with the mammary vein in lactating dairy cows. Redetzky et al. (2003) also found higher levels of BHBA in blood drawn from the jugular vein compared with the mammary vein in lactating dairy cows. Interestingly, BHBA concentrations in nonlactating cows showed an inverse relationship (Wilhelm et al., 2013). The authors speculated that different BHBA consumption rates in the tissues drained by these veins (i.e., especially brain and udder) caused this difference. However, the tail ves-
sels were not included in both of these comparisons, although this sampling location has practical benefits (e.g., easier handling) and has been used in previous research trials (Enjalbert et al., 2001; Iwersen et al., 2009; McArt et al., 2012).

Comparing the mammary vein, jugular vein, and tail vessels, we found significant differences in blood BHBA concentrations between samples drawn from the mammary vein and both of the other locations. Concentrations of BHBA in blood samples taken from the mammary vein were 0.3 and 0.4 mmol/L lower compared with samples taken from the jugular vein and tail vessels, respectively. No significant difference was observed between samples taken from the jugular vein and tail vessels. Therefore, we recommend the use of the jugular vein or tail vessels as a sampling location for BHBA measurements.

**CONCLUSIONS**

Screening dairy cows for SCK under conditions of continuous TMR feeding can be performed at any time of the day. A single measurement provides very good test characteristics for on-farm conditions. Repeated measurements after different time intervals improved test characteristics only slightly in our study. The jugular vein or tail vessels should be used as blood sampling locations for measurement of BHBA. Blood drawn from the mammary vein contains lower BHBA concentrations compared with the jugular vein or tail vessels.

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**REFERENCES**


