ABSTRACT

The objectives of this study were to evaluate (1) the performance of an automated health-monitoring system (AHMS) to identify cows with mastitis based on an alert system (health index score, HIS) that combines rumination time and physical activity; (2) the number of days between the first HIS alert and clinical diagnosis (CD) of mastitis by farm personnel; and (3) the daily rumination time, physical activity, and HIS patterns around CD. Holstein cows (n = 1,121; 451 nulliparous and 670 multiparous) were fitted with a neck-mounted electronic rumination and activity monitoring tag (HR Tags, SCR Dairy, Netanya, Israel) from at least −21 to 80 d in milk (DIM). Raw data collected in 2-h periods were summarized per 24 h as daily rumination and activity. An HIS (0 to 100 arbitrary units) was calculated daily for individual cows with an algorithm that used rumination and activity. A positive HIS outcome was defined as an HIS of <86 units during at least 1 d from −5 to 2 d after CD. Blood concentrations of nonesterified fatty acids, β-hydroxybutyrate, total calcium, and haptoglobin were also determined in a subgroup of cows (n = 459) at −11 ± 3, −4 ± 3, 0, 3 ± 1, 7 ± 1, 14 ± 1, and 28 ± 1 DIM. The sensitivity of the HIS was 58% [95% confidence interval (CI): 49, 67] for all cases of clinical mastitis (n = 123), and 55% (95% CI: 46, 64; n = 114) and 89% (95% CI: 68, 100; n = 9) for cases of mastitis alone or concurrent with other health disorders, respectively. Among clinical cases, sensitivity was 80.7% (95% CI: 67, 97) for cases caused by *Escherichia coli* (n = 31) and ranged from 45 to 48% for cases caused by gram-positive bacteria (n = 39; *Streptococcus agalactiae*, *Streptococcus dysgalactiae*, *Streptococcus uberis*, *Streptococcus spp.*, *Staphylococcus spp.*, and *Trueperella pyogenes*), *Staphylococcus aureus* (n = 11), or cases with no bacterial growth (n = 25). Days between the first HIS <86 and CD were −0.6 (95% CI: −1.1, −0.2) for all cases of mastitis. Cows diagnosed with mastitis had alterations of their rumination, activity, HIS patterns, and reduced milk production around CD depending on the type of mastitis case. Cows with mastitis also had some alterations of their calcium and haptoglobin concentrations around calving. The AHMS used in this study was effective for identifying cows with clinical cases of mastitis caused by *E. coli* and cows with another disease occurring during an event of mastitis, but it was less effective in identifying cows with mastitis not caused by *E. coli*.

Key words: rumination, activity, mastitis, dairy cow

INTRODUCTION

Early postpartum health disorders negatively affect cow well-being and are associated with significant economic losses for dairy farms because of alterations to cow health, welfare, and performance (Bareille et al., 2003; Ingvartsen, 2006; Hailemariam et al., 2014). Clinical mastitis is one of the most prevalent disorders affecting cow health and performance (Kaneene and Hurd, 1990; Ingvartsen et al., 2003; Østerås et al., 2007). Mastitis leads to major milk losses (Gröhn et al., 2004; Bar et al., 2007; Schukken et al., 2009) and reduces reproductive performance (Santos et al., 2004; Ahmadzadeh et al., 2009; Hertl et al., 2010); some types of mastitis may severely compromise cow health, leading to increased culling or death (Gröhn et al., 2005; Whist et al., 2009; Hertl et al., 2011).

To detect cows with clinical mastitis, health-monitoring programs include the evaluation of milk characteristics, signs of udder inflammation, and systemic signs of illness (Nash et al., 2002; Wenz et al., 2006). Cases of mastitis caused by certain pathogens such as *Streptococcus agalactiae*, *Streptococcus dysgalactiae*, and *Streptococcus uberis* are more commonly associated with mild changes in milk and udder inflammation (Todhunter et al., 1995; Keefe, 1997; Schukken et al., 2011); cases of mastitis caused by pathogens such as
**Escherichia coli** and other gram-negative bacteria are characterized by a severe inflammatory response and systemic compromise (Harmon, 1994; Burvenich et al., 2007; Schukken et al., 2011). Thus, the severity of mastitis may range from mild changes in milk appearance to an important systemic compromise (Sargeant et al., 1998; Bradley and Green, 2001; Nikolić et al., 2003).

Multiple automated data collection systems based on sensors (e.g., daily milk weights, milk composition, electrical conductivity, somatic cell counts) have been tested and are available to detect mastitis through changes in milk production and its attributes (Kampfluis et al., 2008; Koop et al., 2015; Sørensen et al., 2016). Conversely, data about the use of automated rumination time and physical activity monitoring systems to detect cows with mastitis are scarce, because only a few studies have evaluated rumination time in cows with induced clinical mastitis or challenged with LPS (Siivonen et al., 2011; Fogsgaard et al., 2012; Fitzpatrick et al., 2013), and no studies have assessed a combination of both rumination and activity data to identify cows with mastitis. Beyond the potential of using rumination and activity monitoring alone or to complement other methods of mastitis detection, these 2 parameters may provide additional insights into overall cow health that are not provided by other sensor systems that monitor only milk or by clinical examination of cows and milk.

We hypothesized that an automated health-monitoring system (AHMS) that continuously monitors rumination and activity would be able to identify cows with mastitis. Also, we expected that changes in rumination and activity before evident clinical signs of disease would result in earlier identification of mastitis. The objectives of this study were to evaluate (1) the performance of an automated rumination and physical activity monitoring system to identify cows with mastitis; (2) the interval between the AHMS alert based on a health index score (HIS) and the day of clinical diagnosis (CD) by farm personnel; and (3) the rumination, activity, AHMS-generated alert, and milk production patterns for cows with mastitis. We also used markers of energy status [nonesterified fatty acids (NEFA) and BHB], mineral status (total calcium), and systemic inflammation (haptoglobin) were used to complement the diagnosis of mastitis and the performance of the AHMS alert.

**MATERIALS AND METHODS**

**Animals and Study Design**

All procedures were approved by the Institutional Animal Care and Use Committee of Cornell University. The study followed an observational prospective cohort design. Details about the animals and study design are provided in a companion manuscript (Stangaferro et al., 2016). Briefly, Holstein cows (n = 1,121; 451 nulliparous and 670 multiparous) were fitted with a neck-mounted electronic rumination and activity monitoring tag (HR Tags; SCR Dairy, Netanya, Israel) to monitor rumination and activity from at least 21 d before expected calving until at least 80 d after calving. Of 1,121 cows enrolled in the study, 41 (3.7%) were removed from the data set due to tag malfunction or misplacement during data collection. Thus, 1,080 cows were included in the final data set for analysis. Based on rumination and activity data, an HIS (0 to 100 arbitrary units) for each cow was generated by the system software (DataFlow, Netanya, Israel) using a series of internal algorithms (proprietary to SCR Dairy). An HIS of 100 arbitrary units represents a cow with an ideal pattern of rumination and activity; an HIS value <86 arbitrary units may be indicative of a health disorder. An HIS report was generated daily to include cows with <86 arbitrary units (as determined by SCR) and stored for evaluation by the research group. During the study, farm personnel did not have access to the HIS report or any information generated by the AHMS.

**Fresh Cow Monitoring Program and Case Definitions**

The fresh cow health-monitoring program and definition of each particular health disorder are provided in detail in a companion manuscript (Stangaferro et al., 2016). In particular, clinical mastitis was defined as swelling or pain in the udder, or milk with abnormal appearance (milk was striped onto the floor and observed for flakes or clots). Signs of udder inflammation may or may not have been accompanied by depressed attitude, anorexia, and fever. Mastitis monitoring was conducted during milking and during health monitoring of fresh cows. Milk culture was performed on all cows at the beginning of lactation (first milking) and on the day of mastitis diagnosis. Milk samples for pathogen detection were collected aseptically and shipped daily to the Quality Milk Production Services Laboratory at Cornell University (Ithaca, NY). Results were provided to the farm within 24 h of sample retrieval. Culture outcomes were grouped as follows: (1) *E. coli*; (2) *Klebsiella* spp.; (3) gram-positive bacteria (*Strep. agalactiae*, *Strep. dysgalactiae*, *Strep. uberis*, *Streptococcus* spp., *Staphylococcus* spp., *Truiperella pyogenes*); (4) *Staphylococcus aureus*; and (5) no important growth after 48 h. Two consecutive episodes of mastitis were considered separate episodes if they occurred at least 7 d apart or in a different quarter.
**Blood Collection and Laboratory Analyses**

Blood samples were collected from a subgroup of cows (n = 459) on d −11 ± 3 and −4 ± 3 prepartum, and 0, 3 ± 1, 7 ± 1, 14 ± 1, and 28 ± 1 after calving. Plasma samples were analyzed for NEFA (d −11, −4, 0, 3, 7, 14, 28), BHB (d 0, 3, 7, 14, 28), total Ca (d 0, 3, 7, 14), and haptoglobin (d −4, 0, 3, 7, 14, 28). Details about blood collection, plasma separation, and NEFA, BHB, total Ca and haptoglobin analysis are provided in a companion manuscript (Stangaferro et al., 2016).

**Statistical Analysis**

**System Performance.** The main outcome of interest for this study was the ability of the HIS to correctly identify cows with mastitis. Clinical diagnosis by farm personnel was used as the reference test. Because HIS does not confirm the occurrence of disease or indicate the type of disease, a positive outcome was defined as an HIS value of <86 arbitrary units during at least 1 d during the 5 d preceding, the day of, or 2 d after CD. The sensitivity and 95% CI for the HIS to identify cows with mastitis was calculated using PROC FREQ in SAS (version 9.4; SAS Institute Inc., Cary, NC) and was defined as the ability of the HIS to correctly identify cows with a positive CD for mastitis. To evaluate the potential confounding effect of other health disorders (i.e., all disorders of interest, pneumonia, and lameness) on the sensitivity of HIS, we conducted 3 separate analyses. The first analysis included all cows with cases of clinical mastitis; a second analysis included cows diagnosed only with mastitis during the range of interest around CD; and a third analysis included cows diagnosed with mastitis and at least one other health disorder during the range of interest around CD. Clinical cases of mastitis were also stratified by milk culture results. For this analysis, cows were separated into the following groups: (1) E. coli; (2) Klebsiella spp.; (3) gram-positive bacteria (Strept. agalactiae, Strept. dysgalactiae, Strept. uberis, Streptococcus spp., Staphylococcus spp., T. pyogenes); (4) Staph. aureus; and (5) no important growth after 48 h. We determined differences in the sensitivity of HIS between the subgroup of cows with mastitis only and cows with mastitis and at least another disorder, and for the different pathogen subgroups, by logistic regression using the events over trials option of PROC LOGISTIC in SAS.

**Interval Between the First Positive HIS Outcome and Clinical Diagnosis.** To determine if HIS was capable of identifying cows with mastitis earlier than CD by farm personnel, the interval (in days) between the first positive HIS outcome during the period of interest around CD (5 d before to 2 d after) and the day of CD was evaluated. For this analysis, which included only cows flagged by the AHMS, we compared the mean number of days from the first HIS-positive outcome and the day of CD with a paired t-test conducted using the PROC TTEST in SAS.

**Rumination, Activity, HIS, and Milk Production Relative to Clinical Diagnosis.** Daily rumination time (min/d), daily activity (arbitrary units/d), and HIS (arbitrary units) were evaluated from 5 d before to 5 d after CD (d 0) for the first event of mastitis. Milk production (kg/d) was evaluated only from 5 d before until the day of CD, because data was not available after CD for cows treated with an antibiotic that required milk withdrawal. Before statistical analysis, we assessed normality of the data for rumination, activity, HIS, and milk production using the Shapiro-Wilk statistic and graphical methods (histogram and Q-Q plot) and using PROC UNIVARIATE in SAS. Based on this analysis, no data transformations were necessary.

Different analyses were conducted using data from groups of cows created based on specific criteria. A first analysis included a nondisease group (ND) and the group with clinical mastitis (for first event of mastitis). Thereafter, cows were grouped according to the following criteria: CD-positive and HIS-positive (HI+); HIS <86 arbitrary units at least 1 d within 5 d before, the day of, and 2 d after CD); CD-positive and HIS-negative (HI−; HIS ≥86 arbitrary units from 5 d before, the day of, and 2 d after CD); CD-positive and HIS-negative (HI−; HIS ≥86 arbitrary units from 5 d before, the day of, and 2 d after CD); and CD-negative (ND; cows not diagnosed with a health disorder during the study period). For cows in the ND group, we considered the average DIM at CD for cows with mastitis “day 0.” Data were analyzed by ANOVA with repeated measurements using PROC MIXED in SAS. Models for each outcome of interest included group (e.g., ND and clinical mastitis; and ND, HI+, and HI−), time, and group-by-time interaction as explanatory variables. Parity and group-by-parity interaction were also offered to the initial models. The occurrence of another health disorder (i.e., all disorders of interest, pneumonia, and lameness) during the −5 to 5 d period after CD was offered as a categorical variable (0 = no occurrence and 1 = occurrence) to the initial models to evaluate the potential effect of multiple disorders on the parameter of interest. The final model for each variable of interest was selected by backward elimination of explanatory variables with $P > 0.10$ and determination of the lowest value for the Akaike’s information criterion. Group, time, and group-by-time interaction were forced in all models. Cow within group was included as a random effect in all models. Cow was the subject of the repeated measurements, and all models were run using an autoregressive (AR-1) covariance structure. When the main effect or interaction between explanatory variables was significant, post hoc analyses were completed using the lsmeans statement in SAS (SAS Institute Inc., Cary, NC).
variables was significant, we used the LSD post hoc mean separation test to determine differences between groups of means.

**Plasma Concentrations of NEFA, BHB, Ca, and Haptoglobin.** Cows were grouped in the same way as for evaluation of rumination, activity, and HIS. Data were analyzed by ANOVA with repeated measurements using PROC MIXED in SAS as described and using the same models as for the other parameters.

Data for proportions are presented as arithmetical means and 95% CI; quantitative data are presented as LSM ± SEM or 95% CI, unless otherwise stated. All explanatory variables and their interactions were considered significant if \( P \leq 0.10 \), and \( 0.05 < P \leq 0.05 \) was considered a tendency.

**RESULTS**

**Mastitis Incidence and System Performance**

The incidence of mastitis and DIM at CD are presented in Table 1. Of all the cows affected with clinical mastitis during the study period (\( n = 123 \)), 90.2% developed 1 event and 9.8% developed 2 events. The greatest incidence observed was for cases with a culture result for gram-positive bacteria (not including *Staph. aureus*), followed by *E. coli* and no growth after 48 h. Eleven cows with clinical mastitis (8.9% of the total clinical cases) were not included in the analysis for specific pathogens because they had the following culture results: yeast, no culture results, other, or contamination.

The sensitivity of HIS to detect cows with mastitis and the interval between an HIS-positive outcome and a CD of mastitis by farm personnel are also presented in Table 1. The overall sensitivity for clinical mastitis was 58%, with a tendency to be greater (\( P = 0.08 \)) for cows that developed another health disorder from \(-5\) to \(2\) d after CD than for cows that had mastitis only (89 vs. 55%, respectively). For cases of clinical mastitis stratified by pathogen, the sensitivity of HIS was greater (\( P = 0.04 \)) for mastitis caused by *E. coli* (81%) than for cases caused by *Klebsiella* spp. (33%), gram-positive bacteria (49%), *Staph. aureus* (46%) or no growth after 48 h (48%). Overall, all cases of clinical mastitis were flagged earlier (\( P < 0.02 \)) based on HIS than CD by farm personnel.

**Rumination, Activity, HIS, and Milk Production Relative to Clinical Diagnosis**

Daily rumination time, activity, and HIS patterns from \(-5\) to \(5\) d after CD for cows with clinical mastitis (\( n = 110 \); first event of mastitis only), and cows in the ND group (\( n = 435 \)) are shown in Figure 1. For all parameters, the effect of parity and the group-by-parity interaction were not significant. For rumination, we observed an interaction between group and day (\( P < 0.01 \)). Rumination was lower for cows in the clinical mastitis group than for cows in the ND group from \(-1\) to \(3\) d relative to CD. Cows with clinical mastitis reached their nadir (\(~397\) min/d) on \(-1\). We detected an interaction between group and day (\( P < 0.01 \)) for activity. Cows with clinical mastitis had lower activity than cows in the ND group during the entire period analyzed, reaching their nadir (\(~485\) arbitrary units/d) on \(0\). We observed an interaction between group and day (\( P < 0.01 \)) for HIS in cows with mastitis. Cows in the clinical mastitis group had lower HIS than cows in the ND group from \(-1\) to \(5\) d relative to CD, reaching the lowest value (\ (~84\) HIS units) on \(0\). In addition, HIS was affected by the occurrence of metritis. Cows with metritis had lower HIS (\( P < 0.01 \)) than cows without metritis during the period of interest around CD (88.3 ± 1.7 vs. 95.2 ± 0.1 units, respectively).

Figure 2 includes rumination, activity, and HIS patterns from \(-5\) to \(5\) d relative to CD for cows with clinical mastitis caused by *E. coli* or gram-positive bacteria and included in the HI+ and HI− groups. For all parameters, the effect of parity and the group-by-parity interaction were not significant. For cows with mastitis caused by *E. coli* (Figure 2A, B, C), we observed an interaction between group and day (\( P < 0.01 \)) for rumination. Rumination times were lower for cows in the HI+ group (\( n = 21 \)) than the ND group (\( n = 435 \)) from \(-1\) to \(3\) d relative to CD, reaching their nadir (\ (~291\) min/d) and the greatest difference with the ND group (\ (~183\) min /d) on \(0\). Cows in the HI− group (\( n = 5 \)) had higher rumination time than cows in the HI+ group from \(-1\) to \(1\) d relative to CD. We detected an interaction between group and day (\( P = 0.01 \)) for activity. Cows in the HI+ group had lower activity than cows in the HI− and ND groups from \(-1\) to \(5\) d relative to CD. We detected an interaction between group and day (\( P < 0.01 \)) for HIS, such that cows in the HI+ group had lower HIS than cows in the HI− and ND groups from \(-1\) to \(5\) d relative to CD. The HIS for cows in the HI+ group reached their lowest value (\ (~77\) units) on the day of CD. In addition, HIS was affected by the occurrence of lameness, because cows with lameness had reduced HIS (\( P < 0.01 \)) than cows without lameness during the period of interest around CD (94.6 ± 0.3 vs. 99.6 ± 1.8 units, respectively).

For cows diagnosed with mastitis caused by gram-positive bacteria (Figure 2D, E, F), we observed an
Table 1. Incidence of mastitis, DIM at clinical diagnosis, sensitivity of health index score (HIS) to detect cows with mastitis, and interval between the first HIS-positive outcome and clinical diagnosis (CD) of mastitis by farm personnel

<table>
<thead>
<tr>
<th>Item</th>
<th>Cows (no.)</th>
<th>Incidence (%)</th>
<th>DIM at event (mean ± SD)</th>
<th>Sensitivity</th>
<th>HIS-positive to CD²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cows</td>
<td>(no.)</td>
<td>(%)</td>
<td>(mean ± SD)</td>
<td>% (no./no.)</td>
<td>Days, 95% CI</td>
</tr>
<tr>
<td>Clinical mastitis³</td>
<td>123</td>
<td>11.4</td>
<td>38 ± 24</td>
<td>58 (71/123)</td>
<td>49, 67</td>
</tr>
<tr>
<td>Clinical mastitis only⁴</td>
<td>114</td>
<td>10.6</td>
<td>40 ± 24</td>
<td>55 (63/114)</td>
<td>46, 64</td>
</tr>
<tr>
<td>Clinical mastitis with other disorders⁵</td>
<td>9</td>
<td>0.8</td>
<td>20 ± 23</td>
<td>89 (8/9)</td>
<td>68, 100</td>
</tr>
<tr>
<td>Clinical mastitis by pathogen⁶</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Escherichia coli</td>
<td>31</td>
<td>25.2</td>
<td>40 ± 24</td>
<td>81 (25/31)</td>
<td>67, 95</td>
</tr>
<tr>
<td>Klebsiella spp.</td>
<td>6</td>
<td>4.9</td>
<td>37 ± 24</td>
<td>33 (2/6)</td>
<td>1, 71</td>
</tr>
<tr>
<td>Gram-positives⁷</td>
<td>39</td>
<td>31.7</td>
<td>37 ± 26</td>
<td>49 (19/39)</td>
<td>32, 65</td>
</tr>
<tr>
<td>Staphylococcus aureus</td>
<td>11</td>
<td>8.9</td>
<td>38 ± 20</td>
<td>46 (5/11)</td>
<td>17, 77</td>
</tr>
<tr>
<td>No growth⁸</td>
<td>25</td>
<td>20.3</td>
<td>37 ± 23</td>
<td>48 (12/25)</td>
<td>28, 69</td>
</tr>
</tbody>
</table>

Different superscripts indicate differences (P ≤ 0.05) between means based on mean separation with the LSD test.

¹Number of events diagnosed.
²HIS-positive to CD = interval in days between the first positive HIS outcome (positive outcomes only) and CD. For cases of mastitis caused by Klebsiella spp., HIS-positive to CD was not calculated because of lack of sufficient observations.
³All clinical mastitis events recorded.
⁴Cows diagnosed with only clinical mastitis from −5 to 2 d relative to CD.
⁵Cows diagnosed with clinical mastitis and at least another health disorder from −5 to 2 d relative to CD.
⁶Clinical mastitis events classified by the results of milk culture [11 cows not included: no culture results (n = 6); yeast (n = 2); other (n = 2); contamination (n = 1)].
⁷Gram-positives = Streptococcus agalactiae, Streptococcus dysgalactiae, Streptococcus uberis, Streptococcus spp., Staphylococcus spp., Trueperella pyogenes.
⁸No important growth after 48 h.
interaction between group and day \((P < 0.01)\) for rumination. Cows in the HI+ group \((n = 19)\) had lower rumination than cows in the ND \((n = 435)\) group from \(-1\) to \(4\) d relative to CD, reaching a nadir \((\sim 330\) min/d) on d 0. Cows in the HI− group \((n = 23)\) had higher rumination than cows in the HI+ group from 0 to \(4\) d after CD. We detected an interaction between group and day \((P = 0.05)\) for activity. Daily activity was lower in the HI+ group than in the ND group from \(-2\) to \(5\) d relative to CD; cows in the HI− group exhibited no difference compared with the HI+ or ND groups. We detected an interaction between group and day \((P < 0.01)\) for HIS. Cows in the HI+ group had lower HIS than cows in the HI− and ND groups from \(-2\) to \(5\) d relative to CD, with the lowest values \((\sim 86\) HIS units) from d 0 to 2. In addition, HIS was also affected by the occurrence of metritis, because cows with metritis had reduced HIS \((P < 0.01)\) than cows without metritis during the period of interest around CD \((88.6 \pm 2.2\) vs. \(94.8 \pm 0.2\) units, respectively).

Daily milk production \((\text{kg/d})\) from \(5\) d before until the day of CD for cows that had clinical mastitis is presented in Figure 3. Because parity had an effect \((P < 0.01)\) on milk production, results are presented separately for primiparous and multiparous cows. For primiparous cows (Figure 3A), we observed an interaction between group and day \((P < 0.01)\). Cows with mastitis regardless of HIS group (HI+ and HI−) produced less milk than cows in the ND group from \(-3\) to 0 (HI− vs. ND) and from \(-2\) to 0 d (HI+ vs. ND) relative to CD. We observed the greatest difference between cows in the HI+ and ND groups on d \(-1\) \((12.8 \text{ kg/d})\), and the greatest difference between cows in the HI− and the ND group on d 0 \((9.5 \text{ kg/d})\). We observed no differences between primiparous cows in the HI+ and HI− groups. For multiparous cows (Figure 3B), we observed an interaction between group and day \((P < 0.01)\). Multiparous cows in the HI+ and HI− groups produced less milk than cows in the ND group from \(-2\) d until the day of CD. We observed the greatest difference between the HI+ and HI− groups compared to the ND group on d 0 \((16.9 \text{ kg/d} \text{ and } 10.1 \text{ kg/d}, \text{respectively})\). Cows in the HI+ group produced less milk than cows in the HI− group on the day of CD.

**Plasma Concentrations of NEFA, BHB, Ca, and Haptoglobin**

Plasma concentrations of NEFA, BHB, Ca, and haptoglobin for cows that developed clinical mastitis before 30 DIM are presented in Figure 4. We detected only an effect of day \((P < 0.01)\) for NEFA concentrations, because NEFA were greater on d 0, 3, and 7 than the rest of the days. We detected no effect of group \((P = 0.58)\)
or day ($P = 0.64$), and no interaction between group and day ($P = 0.83$) for BHB concentrations. For total Ca concentrations, we observed a group-by-day interaction ($P = 0.03$). Cows in the HI+ group had lower Ca concentrations (~1.0 mg/dL different) than cows in the HI− and ND group at 14 DIM. For haptoglobin concent-

**Figure 2.** Rumination (A), activity (B; arbitrary units, AU), and health index score (HIS; AU; C) patterns from −5 to 5 d relative to clinical diagnosis (CD) for cows that developed clinical mastitis caused by *Escherichia coli* (HI+: $n = 21$; HI−: $n = 5$) compared with cows in the nondisease group ($n = 435$). Cows were assigned to the HI+ or HI− group if they had an HIS of $<86$ or $\geq 86$ arbitrary units, respectively, during the 5 d preceding, the day of, and 2 d after CD. For cows in the nondisease group, the average DIM at CD for cows with mastitis was considered “day 0.” Rumination (D), activity (E), and HIS (F) patterns from −5 to 5 d relative to clinical diagnosis for cows that developed clinical mastitis caused by gram-positive bacteria (*Streptococcus agalactiae*, *Streptococcus dysgalactiae*, *Streptococcus uberis*, *Streptococcus* spp., *Staphylococcus* spp., *Trueperella pyogenes*) compared with cows in the nondisease group ($n = 435$). Cows with clinical mastitis caused by gram-positive bacteria were assigned to the HI+ ($n = 19$) and HI− ($n = 18$) groups following the same criteria as for cows with clinical mastitis caused by *E. coli*. Values are presented as LSM ± SEM. Within a day, pairwise comparisons that were statistically different ($P \leq 0.05$) based on LSD are represented as follows: *control vs. HI+; ‡HI+ vs. HI−.
trations, we observed an interaction between group and day \((P = 0.02)\). Cows with clinical mastitis (both HI+ and HI− groups) had greater haptoglobin concentrations than cows in the ND group on d 14 and 28.

**DISCUSSION**

In the current study, we evaluated the performance of an AHMS that combined rumination and activity to...
identify cows with mastitis. Our results support the notion that an HIS alert generated based on rumination and activity would be able to identify cows with more severe cases of mastitis. We observed moderate sensitivity when all cases of mastitis were included in the analysis but substantial differences in sensitivity when cows were stratified based on the presence or absence of another disorder from 5 d before to 2 d after clinical diagnosis of mastitis. Moreover, when cows with mastitis caused by E. coli were evaluated individually, the sensitivity of the AHMS was more than 20 percentage points greater than when all cases were combined. This finding was expected, because intramammary infections caused by E. coli are characterized by a severe inflammatory response, including sudden shock, sepsis, and often death (Burvenich et al., 2003; 2007; White et al., 2010; Schukken et al., 2011). Indeed, cows affected by E. coli and identified by the AHMS (HI+ group) had sudden and dramatic reductions in rumination, activity, and HIS, reaching nadirs similar to those of cows affected by metabolic and digestive disorders (Stangaferro et al., 2016). As a result of the low HIS observed during the nadir (<86 points), most cows were flagged by the AHMS. Conversely, the low sensitivity of the AHMS in detecting cows with mastitis caused by Klebsiella spp. was unexpected. Clinical cases caused by this pathogen are usually characterized by clinical signs similar to those observed in mastitis caused by E. coli (Harmon, 1994; Radostits et al., 2006). Although we did not have information about the severity of the mastitis event, we speculate that the moderate sensitivity of the AHMS in these cases was due to an over-representation of mild cases among the small number of cows with mastitis caused by this pathogen.

The observed lower sensitivity for cases of clinical mastitis caused by gram-positive pathogens and Staph. aureus was also expected, because these pathogens do not cause the same level of toxemia as E. coli (Harmon, 1994; Todhunter et al., 1995; Keefe, 1997; Schukken et al., 2011). Based on the patterns of rumination, activity, and HIS for the HI+ group, however, it seemed obvious that this subgroup contained cows with more severe cases of mastitis and systemic compromise. Cows in the HI+ group had significant reductions in rumination and activity within 1 to 2 d before CD and, for some cows, the HIS was below 86 points from the day of and up to 2 d after CD. It is uncertain whether the cases of clinical mastitis with no pathogen growth in cultured milk resembled more closely those caused by E. coli or the rest of the pathogens isolated during the study. However, the sensitivity of the AHMS for this subgroup of cows and the patterns of rumination, activity, and HIS (data not shown) closely matched those of cows with mastitis caused by gram-positive pathogens and Staph. aureus.

Another interesting aspect of cows affected by clinical mastitis was the lack of difference in milk production between cows in the HI+ and HI− groups around CD (except for d 0 in the multiparous group). Both groups produced less milk on the days leading up to and day of CD than cows with no health disorders. This was in contrast to our observations for metabolic and digestive disorders (Stangaferro et al., 2016). In such cases, milk production before CD was lowest for cows in the HI+ group; cows in the HI− group had less of a reduction or no reduction before CD compared to cows in the ND group. These contrasting results for milk production differences between the HI+ and HI− groups for cows affected by metabolic and digestive disorders or mastitis were probably a reflection of the reasons underlying reduced milk production. It is likely that in cows with clinical mastitis, the decline in milk production around CD was, for the most part, due to the direct effect of inflammation on the mammary gland (Zhao and Lacasse, 2008; Akers and Nickerson, 2011). In contrast, it is likely that cows with metabolic and digestive disorders produced less milk because of a reduction in DMI and the overall effect of disease on cow health (Bareille et al., 2003; Van Winden et al., 2003).

Similar to cows with metabolic and digestive disorders (Stangaferro et al., 2016), cows with clinical mastitis and an HIS <86 points were flagged earlier than by farm personnel (approximately half a day). In this case, the difference in favor of the AHMS was smaller than for the other disorders, and we observed no statistical differences when mastitis events were separated by pathogen. The short time frame from an HIS <86 points and CD was likely a reflection of the sudden drop in rumination and activity within no more than 1 d of CD in cows flagged based on HIS. In contrast, for disorders such as displaced abomasum or ketosis, rumination and activity were below levels observed in cows with no health disorders as early as 5 d before CD. Taken together, these results suggest that there may not be a major advantage for the AHMS in terms of the timing of mastitis diagnosis for herds with similar mastitis detection programs. The characteristic changes in milk composition, udder appearance, and consistency observed in cows with clinical cases of mastitis make other direct and simple methods of detection (e.g., milk stripping, udder visual inspection, palpation) more effective than an AHMS that is based on rumination and activity only. Nevertheless, rumination and activity or alert systems generated based on these parameters (e.g., HIS) could be used as tools for diagnosing severe cases of clinical mastitis caused by pathogens such as E.
coli, which have profound systemic effects for the cow. Another potential application consists of using rumination and activity as markers of systemic compromise and as an aid in treatment decision-making, because changes in milk composition or udder status do not provide information about cows’ overall health status.

From an on-farm implementation perspective, the HIS generated by the AHMS tested in our study could be used to detect severe cases of mastitis that affect the cow systemically. The use of HIS could be complemented by other parameters not reported in this study but generated based on rumination behavior and activity (e.g., weekly rumination and activity, deviations in the last 2 h) that could be retrieved separately for individual cows or groups of cows. However, the most likely scenario is that not all cows that develop a case of mastitis will be detected. This is because the underlying health disorder does not compromise cow health sufficiently to dramatically reduce rumination and activity in all cows. Thus, it seems reasonable to suggest that for farms with proactive health-monitoring and treatment programs, the AHMS could be used in combination with other traditional methods of mastitis detection. The AHMS may be a valuable tool for providing further insights about the overall health status of the cow.

CONCLUSIONS

Our findings demonstrated that automated rumination and activity monitoring was effective for identifying cows with clinical cases of mastitis caused by E. coli and cases of mastitis concurrent with another health disorder. Conversely, the ability of the AHMS to identify cows with clinical mastitis caused by pathogens other than E. coli was moderate. Overall, cows with clinical mastitis were identified earlier than through CD by farm personnel. The patterns of rumination, activity, and HIS from −5 to 5 d after CD for cows with clinical mastitis were characterized by marked differences compared to cows in the ND group. Among cows with clinical mastitis, those not identified by HIS had rumination and activity patterns very similar to cows in the ND group and different from cows diagnosed with the disorder but flagged by HIS. We conclude that automated health-monitoring systems that use rumination and physical activity should be used in combination with or to complement traditional methods of mastitis detection. Future research is needed to evaluate the effect of management programs that combine rumination and activity monitoring with traditional methods to diagnose mastitis on cow well-being and performance.

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REFERENCES


RUMINATION AND ACTIVITY FOR HEALTH MONITORING


