The relationship between antibody status to bovine corona virus and bovine respiratory syncytial virus and disease incidence, reproduction and herd characteristics in dairy herds

Anna Ohlson*, Ulf Emanuelson, Madeleine Tråvén and Stefan Alenius

Abstract

Background: Bovine respiratory syncytial virus (BRSV) and bovine corona virus (BCV) affects cattle worldwide. Our objective was to evaluate the effects of these infections on general health and reproduction parameters measurable on herd level and to explore the association between antibody status and some herd characteristics.

Methods: We collected a pooled milk sample from five primiparous cows from 79 Swedish dairy herds in September 2006. The samples were analysed for immunoglobulin G antibodies to BCV and BRSV with indirect enzyme-linked immunosorbent assays. Herd level data from 1 September 2005 to 30 August 2006 were accessed retrospectively. The location of the herds was mapped using a geographical information system.

Results: Ten herds were antibody negative to both viruses and were compared with 69 herds positive to BCV or BRSV or both. Positive herds had a higher (P = 0.001) bulk tank milk somatic cell count (BMSCC) compared with negative herds. The medians for all other analyzed health and reproductive parameters were consistently in favour of the herds negative to both viruses although the differences were not statistically significant. A higher proportion (P = 0.01) of herds used professional technicians for artificial insemination, rather than farm personnel, amongst the 33 herds negative to BCV compared with the 46 positive herds.

Conclusions: Our result shows that herds that were antibody positive to BCV and/or BRSV had a higher BMSCC compared with herds negative to BCV and BRSV. There was also tendency that negative herds had a better general herd health compared with positive. A higher proportion amongst the BCV negative herds used external technicians for AI instead of farm personnel, indicating that it is possible to avoid infection although having regular visits. Negative herds were located in close proximity to positive herds, indicating that local spread and airborne transmission between herds might not be of great importance and that herds can stay free from these infection transmission although virus is circulating in the area.

Background

Bovine corona virus (BCV) and bovine respiratory syncytial virus (BRSV) are two worldwide distributed viruses [1,2]. BCV causes diarrhoea in calves, winter dysentery in adults and various degrees of respiratory symptoms [3-5]. BRSV is regarded as one of the most important causes of respiratory tract disease, especially in young calves. An infection can cause respiratory distress, fever, anorexia and subcutaneous emphysema and can lead to secondary bacterial pneumonia and death [6,7]. Outbreaks of BCV and BRSV occur mainly in autumn and winter [8,9]. These infections are common in dairy herds; in a nationwide survey in England and Wales the prevalence of antibodies to these viruses in bulk tank milk (BTM) was 100% [10]. Swedish studies have shown a prevalence of 70-100% for BCV and 41-89% for BRSV, with the higher prevalence in southern parts [7,11]. In a more recent study in a high animal-density area in south-west
Sweden, the prevalence in BTM was 100% for both BCV and BRSV [12].

Previous studies have shown that BRSV and BCV infections are effectively spread within the herd [5,6,13]. It has also been shown that acquired antibodies remain detectable for years, even without reinfection [5,7], whereas maternal antibodies are only detectable for a few months. Spot samples from a few young animals can thus be used to reflect recent infections of BRSV and BCV in a herd, whereas bulk tank milk samples mirror the long-term history. Spot sampling has previously been described for bovine virus diarrhea virus (BVDV) [14,15].

Despite the importance of these viruses and the fact that they are widely spread, little is known about transmission routes and management risk factors. Introduction of new animals and indirect spread via people and equipment are believed to be important and airborne transmission has been shown to occur for BRSV, at least under experimental conditions [16]. Studies have been carried out to determine the relationship between herd health, reproduction efficiency and milk production and seropositivity to other viruses, for example bovine viral diarrhoea virus and bovine leukemia virus [17-20]. Similar studies for BRSV and BCV have, as far as we know, not been conducted and it is therefore difficult to quantify their effect on the farm efficiency and economy. The purpose of this study was to explore if there were any associations between antibody status to BCV and BRSV and disease incidence, reproduction and some herd characteristics in dairy herds. A secondary aim was to investigate if there were any differences in proportion antibody positive herds between two neighbouring areas.

Methods
Study population and sampling
We used dairy herds from two neighbouring areas in central-eastern Sweden as study population. The areas approximately correspond to two veterinary districts. The herds were members of the local livestock association (Svenska Husdjur) and enrolled in the National Animal Disease Recording System (NADRS) [21] and the Swedish Official Milk Recording Scheme (SOMRS) [22]. All herds were free from BVDV according to the rules in the Swedish eradication program [23].

A convenience sample of 44 herds was taken from area 1 and of 35 herds from area 2, corresponding to 85% and 71% of the existing dairy herds in area 1 and 2, respectively. Sampling was performed by personnel from Svenska Husdjur. Herds were included if the farmer agreed to participate in the study, in the order they were visited by the personnel. The sampling period was from September 1st to October 31st 2006. The geographical locations of the herds were mapped using a geographic information system (ArcGIS, 2005).

A pooled milk sample from five home-bred primiparous cows was collected from each herd [24]. We used 10-ml test tubes containing 1.5 mg of the preservative agent Bronopol (2-bromo-2-nitropropane-1,3-diol). The milk samples were stored at -20°C until analysis.

Antibody detection and cut off
The milk samples were analysed for presence of immunoglobulin G antibodies to BCV [5] and BRSV [25] by commercially available indirect enzyme-linked immunosorbent assays (ELISA; Svanova Biotech). The same batch was used for all analyses. The samples were not diluted or centrifuged. The optical density (OD) at 450 nm was corrected by subtraction of the negative control antigen OD. To adjust for day-to-day variations we calculated the percent positivity (PP) as (corrected OD/positive control corrected OD) × 100. A PP-value of <20 was regarded as negative for the pooled samples, closely corresponding to the corrected OD of 0.20 which is the cut off for negative individual milk, both for BCV and BRSV, recommended by the manufacturer.

Outcome variables
Herd level data on number of cows, AI-strategy, milk production, reproductive performance and health status from September 1st 2005 to August 31st 2006 were obtained from NADRS and SOMRS. Disease incidences included treated cases of mastitis, any treatment for fertility problems, diseases in young stock and udder disease score (UDS) ≥6. UDS describes the udder health based on individual SCC corrected for milk yield, breed, number of parity and time in lactation [26]. The scale is from 0 to 9 and expresses the probability that a cow has mastitis. UDS class ≥6 implies a 60% probability of infection and corresponds to having an SCC of approximately ≥300 000 cells/ml. Mastitis, treatment for fertility problems, culling and diseases in young stock were calculated as the yearly incidence rates (IR), i.e. number of cases divided by number of cow days or young stock days. Early and late calf mortality was calculated as the mean of 12 monthly IR; for which each monthly IR was calculated as number of calves dead at birth or within 24 hours divided by number of calvings, and number of calves dead between 1 day and 3 months of age divided by number of calves surviving 24 hours, respectively. The incidence of UDS ≥6 was calculated as a daily mean, i.e. number of cases with UDS ≥6 divided by number of cow days. Milk yield was expressed as the mean milk yield per cow and year in kg and BMSCC as the geometric mean of 12 monthly measurements. The data on reproductive performance included average calving interval, time from calving to first insemination and number of inseminations per service period.

Explanatory variables
The main predictor was antibody status, defined as NEG for herds that were antibody negative to both BRSV and
BCV, and POS for herds positive to either BRSV or BCV or both, based on the result from the pooled milk samples. Herd characteristics were also compared separately for BCV and BRSV. The models also included the effects of herd size, milk production, breed and AI-strategy according to table 1. Herd size and milk production was dichotomized by the median, 43 cows and 8967 kg milk per cow and year respectively. In 2006 the mean herd size in Sweden was 48 cows and the mean milk yield was 8175 kg per cow and year [27]. Breed was classified into three categories: > 80% Swedish Red and White breed (SRB), > 80% Swedish Holstein breed (SH) and mixed/other breeds. SRB and SH are the two main dairy-cow breeds in Sweden. Finally, AI-strategy was defined as insemination performed by farm personnel (AIF) or by professional technicians (AIT).

**Statistical analysis**

We used t-test to compare herd size and Fisher’s exact test to compare breed and AI strategy between NEG and POS herds. Fisher’s exact test was also used to compare the proportion of antibody positive herds between the areas.

The outcome variables mastitis, treatment for fertility problems, number of calves dead between one day and three months and diseases in young stock had more than 10% missing values and were not analysed because the remaining sample of NEG herds was considered too small. The continuous outcome variables were analysed with a linear regression model whereas logistic regression models were used for the binomial outcome variables. The eight models are shown in Table 1. We chose to keep the models intact because we considered the predictor variables included in each model as biologically important for the outcome. In order to achieve normal distribution of the residuals we transformed number of AI and calving interval to the natural logarithmic scale (ln). BMSCC was transformed by taking the square root before analysis.

The predictor variables were tested for collinearity by pair-wise Spearman rank correlation test; a test result of <0.6 was considered as negative i.e. no strong correlation. There were no collinearities found between the explanatory variables. No interactions were tested in the statistical model because of the small sample size. To evaluate the fit of the models with significant outcome we plotted the studentised residuals against the predicted observation. The statistical analyses were done using Stata Software (StataCorp 2006; Stata Statistical Software: Release 9.0; College Station, TX, USA: StataCorp LP).

**Results**

**Study population and immunity**

Number of herds that were antibody positive in the pooled milk samples was for BCV 46 (58.2%) and for BRSV 62 (78.5%). Ten herds were classified as NEG and 69 herds as POS. The proportion of antibody positive herds did not differ significantly between the two areas under study for both BCV and BRSV (table 2). Geographical locations of the herds stratified by antibody status are shown in figure 1.

**Analyses of herd data**

The NEG group had a lower mean BMSCC (P < 0.001) compared with the POS group. No other significant (P > 0.05) differences were found between NEG and POS herds in the remaining outcome variables. The median and interquartile range of each variable is presented in table 3.

Herd size, milk production, breed and AI strategy did not differ significantly between the POS and NEG herds. Of all sampled herds 72% used AIT; 80% in NEG group

<table>
<thead>
<tr>
<th>Table 1: Statistical models used to study associations between outcome variables and herd antibody status</th>
</tr>
</thead>
</table>
| **Outcome variable** | **Data type** | **Model type** | **Explanatory variables**
<table>
<thead>
<tr>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>Milk yield</td>
<td>Continuous</td>
<td>Linear</td>
<td>a, b, n</td>
</tr>
<tr>
<td>BTMSCC</td>
<td>Continuous</td>
<td>Linear</td>
<td>a, b, n, p</td>
</tr>
<tr>
<td>Calving interval</td>
<td>Continuous</td>
<td>Linear</td>
<td>a, b, n, p, i</td>
</tr>
<tr>
<td>Calving to first AI</td>
<td>Continuous</td>
<td>Linear</td>
<td>a, b, n, p, i</td>
</tr>
<tr>
<td>Number of AI</td>
<td>Continuous</td>
<td>Linear</td>
<td>a, b, n, p, i</td>
</tr>
<tr>
<td>Dead calves 0-24 hours</td>
<td>Binomial</td>
<td>Logistic</td>
<td>a, b, n</td>
</tr>
<tr>
<td>Culling</td>
<td>Binomial</td>
<td>Logistic</td>
<td>a, b, n, p</td>
</tr>
<tr>
<td>UDS ≥6</td>
<td>Binomial</td>
<td>Logistic</td>
<td>a, n</td>
</tr>
</tbody>
</table>

1 a = antibody status, b = breed, n = herd size, p = milk production level, i = artificial insemination strategy
2 bulk tank milk somatic cell count.
3 artificial insemination
4 udder disease score, see main text for explanation.
and 71% in the POS group. There was a significantly (P < 0.01) higher proportion using AIT (29/33) amongst the herds negative to BCV, compared with the proportion amongst BCV positive herds (28/46). For BRSV, 12 out of 17 negative herds used AIT; corresponding numbers for the positive herds was 45 out of 62. For the distribution of breed, 56% of all herds had mainly SRB, 4% SHB and 40% had mixed/other breeds.

**Discussion**

The only herd health parameter with statistically significant differences between POS and NEG herds was BMSCC for which POS herds had a higher BMSCC than

**Table 2: Antibody status to bovine respiratory syncytial virus (BRSV) and bovine coronavirus (BCV)**

<table>
<thead>
<tr>
<th>BRSV</th>
<th>BCV</th>
<th>Area 1, n = 44</th>
<th>Area 2, n = 35</th>
<th>All, n = 79</th>
</tr>
</thead>
<tbody>
<tr>
<td>-</td>
<td>-</td>
<td>4 (9.1)</td>
<td>6 (17.1)</td>
<td>10 (12.6)</td>
</tr>
<tr>
<td>+</td>
<td>-</td>
<td>13 (29.5)</td>
<td>10 (28.6)</td>
<td>23 (29.1)</td>
</tr>
<tr>
<td>-</td>
<td>+</td>
<td>2 (4.5)</td>
<td>5 (14.3)</td>
<td>7 (8.9)</td>
</tr>
<tr>
<td>+</td>
<td>+</td>
<td>25 (56.8)</td>
<td>14 (40.0)</td>
<td>39 (49.4)</td>
</tr>
</tbody>
</table>

Antibody status of 79 Swedish dairy herds in two neighbouring areas based on pooled milk samples from five home-bred primiparous cows, measured by an ELISA and collected in September-October 2006

**Figure 1 Geographical distribution** Geographical distribution of 79 Swedish dairy herds, stratified by antibody-status to bovine coronavirus (BCV) and bovine respiratory syncytial virus (BRSV) as measured by an ELISA test in pooled milk samples of 5 primiparous cows sampled in September-October 2006. Star = Negative for BRSV and BCV Circle = Positive for BRSV, negative for BCV Square = Negative for BRSV, positive for BCV Triangle = Positive for BRSV and BCV
NEG herds. Despite the difficulties to evaluate causality in cross-sectional studies it is notable that, although not statistically significant, the medians for all analysed variables were consistently in favour of the NEG herds. One explanation for the lower BMSCC in NEG herds may be that an outbreak of BCV or BRSV increases the susceptibility to clinical or subclinical udder infections. Another explanation can be that these herds are better managed with better hygienic routines also at cow level. The information on mastitis incidence had too many missing values to be analyzed. The incidence of high UDS pointed in the same direction as the BMSCC, although the difference was not statistically significant. A high BMSCC is of economic importance for the farmers because the dairy (Arla foods in this area) pays 1% extra for milk with a cell count of ≤300 000 cells/ml and 2% extra when SCC is ≤200 000/ml whereas a deduction of 4% is made for milk with a cell count of ≥401 000 cells/ml and 10% for ≥501 000 cells/ml. Studies have also shown that a high SCC is related to a loss in milk yield [28-30] and that BMSCC is correlated with prevalence of subclinical mastitis [31]. The tendency to poorer reproductive performance and higher culling rates amongst the POS herds could be a consequence of the disease outbreaks by for example delaying AI. These findings may also be related to better general herd management.

A surprising result was that a higher proportion of herds used professional AI technicians rather than AI performed by farm personnel amongst the herds negative to BCV compared with the herds positive to BCV. This finding confirms the results the study by Bidokhti et al [32] where AI-technicians were more common in antibody negative herds compared to positive regarding both BCV and BRSV. Having visitors should not be a protective factor, especially persons that visit more than one herd each day. For example; a typically 50-cow AIT herd in study area 1 and 2 has two AI-technician visits per week, and each AI-technician in these areas visits approximately six herds per day (Rose-Marie Winroth, personal communication). An explanation could be that technicians educate the farmers continuously in how to protect their herd against contagious diseases, thus keeping updated routines in handing protective clothing for visitors as well as providing cleaning and disinfection possibilities. Another explanation could be that these farmers manage the herd more efficiently and therefore use professionals for AI instead of doing it themselves, and well managed herds are also more likely to avoid infection. In all cases, our result indicates clearly that it is possible to avoid infection even with regular visitors in the herd.

Several negative herds were located in close proximity to positive herds, e.g. one of the NEG herds was located between two herds positive to both viruses, at a distance of 1.3 km and 1.9 km. This indicates that local spread and airborne transmission between herds are not of great importance for these infections and that a herd can stay negative although virus is circulating in the area.

Table 3: Summary of the analysed variables

<table>
<thead>
<tr>
<th>Variable</th>
<th>NEG herds</th>
<th>POS herds</th>
</tr>
</thead>
<tbody>
<tr>
<td>Herd size, cow-years</td>
<td>57</td>
<td>43</td>
</tr>
<tr>
<td>Median</td>
<td>40-75</td>
<td>32-62</td>
</tr>
<tr>
<td>Milk yield, kg/cow-year</td>
<td>90/13</td>
<td>89/64</td>
</tr>
<tr>
<td>Median</td>
<td>86/40-99/82</td>
<td>88/315-87/59</td>
</tr>
<tr>
<td>BTMSCC1, 1000 cells/ml</td>
<td>163</td>
<td>218</td>
</tr>
<tr>
<td>Median</td>
<td>140-187</td>
<td>164-283</td>
</tr>
<tr>
<td>Calving interval, days</td>
<td>390</td>
<td>402</td>
</tr>
<tr>
<td>Median</td>
<td>38/1-413</td>
<td>38/7-415</td>
</tr>
<tr>
<td>Calving to first AI2, days</td>
<td>84</td>
<td>91</td>
</tr>
<tr>
<td>Median</td>
<td>76-98</td>
<td>80-104</td>
</tr>
<tr>
<td>Number of AI2</td>
<td>1.7</td>
<td>1.8</td>
</tr>
<tr>
<td>Median</td>
<td>1.5-2.3</td>
<td>1.6-2</td>
</tr>
<tr>
<td>Dead calves3 0-24 h</td>
<td>0.040</td>
<td>0.047</td>
</tr>
<tr>
<td>Median</td>
<td>0.033-0.052</td>
<td>0.028-0.093</td>
</tr>
<tr>
<td>Culling3</td>
<td>0.26</td>
<td>0.36</td>
</tr>
<tr>
<td>Median</td>
<td>0.23-0.39</td>
<td>0.26-0.42</td>
</tr>
<tr>
<td>UDS3,4 ≥ 6</td>
<td>1.3</td>
<td>1.7</td>
</tr>
<tr>
<td>Median</td>
<td>0.97-1.4</td>
<td>1.2-2.3</td>
</tr>
</tbody>
</table>

Comparison between 10 Swedish dairy herds antibody negative (NEG) to bovine coronavirus (BCV) and bovine respiratory syncytial virus (BRSV) and 69 herds positive (POS) to either BRSV or BCV or both, measured by an ELISA test in pooled milk samples from five home-bred primiparous cows, collected in September-October 2006.

1 Bulk tank milk somatic cell count, geometric mean of 12 monthly measurements. P < 0.001 estimated in a linear regression model.
2 AI = artificial insemination
3 One-year incidence rate
4 UDS = udder disease score, see main text for explanation.
The cows included in the pooled milk sample were primiparous, which means that positive herds have experienced an infection during the last two years, approximately. Some herds in the POS group could have had the infection in the year before the investigated year, which may lead to smaller differences between NEG and POS herds. Herds in the NEG group, however, have been free from infection during the year under investigation.

There is a risk of selection bias because the herds were selected by personnel from the local livestock association and only farmers willing to participate were included in the study. The antibody status of the herds was, however, not known at the time of sampling. The disease and reproduction data in the NADRS and SOMRS are reported by farmers, AI technicians and veterinarians. The reports from veterinarian and technicians should not differ between the study groups, but there could be differences in the willingness to report amongst the farmers causing information bias. It is possible that well managed herds with high biosecurity are also better in reporting, which may lead to higher incidences of diseases in NEG herds and thus an underestimate of the differences between the groups. BMSCC, UDS and milk yield are objective measurements from the dairies and from the milk-recording schemes and are not influenced by information bias.

It would be of interest to further investigate the association between BCV/BRSV infections and herd health parameters on a larger scale. It would be beneficial to follow herds over time and compare uninfected herds with herds that have shown primary infection during the study period.

Conclusion
In this study we found that herds that were antibody positive to BCV and/or BRSV had a higher BMSCC compared with herds negative to BCV and BRSV. There was a tendency that negative herds had a better general herd health compared with positive. A higher proportion amongst the BCV negative herds used external technicians for AI instead of farm personnel, indicating that it is possible to avoid infection although having regular visits. Negative herds were located geographically close to positive, indicating that local spread and airborne transmission between herds is not the major transmission routes for BCV and BRSV infections.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
AO carried out the statistical analysis and drafted the manuscript. UE assisted the statistical analysis. UE, MT and SA conceived the study. All authors participated in its design and coordination, were involved in revising the manuscript and read and approved the final manuscript.

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References


27. (Svensk Mjölk), Retrieved 07/11/15, from [http://www.svenskmjolk.se]


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