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Short communication: Association between udder health status and blood serum proteins in dairy cows

T. Bobbo,* P. L. Ruegg,† E. Fiore,‡ M. Gianesella,‡ M. Morgante,‡ D. Pasotto,‡ L. Gallo,* G. Bittante,* and A. Cecchinato*¹

*Department of Agronomy, Food, Natural Resources, Animals and Environment (DAFNAE), University of Padova, Viale dell'Università 16, 35020 Legnaro, Padova, Italy

†Department of Dairy Science, University of Wisconsin-Madison, 1675 Observatory Drive, Madison 53706

‡Department of Animal Medicine, Production and Health, University of Padova, Viale dell'Università 16, 35020 Legnaro, Padova, Italy

ABSTRACT

The aim of this study was to investigate the association between udder health (UH) status and blood serum proteins (i.e., total protein, albumin, globulin, and albumin-to-globulin ratio) in dairy cows. Blood and milk samples were collected from 1,508 cows of 6 different breeds (Holstein Friesian, Brown Swiss, Jersey, Simmental, Rendena, and Alpine Grey) that were housed in 41 multibreed herds. Bacteriological analysis was performed on milk samples with somatic cell count (SCC) >100,000 cells/mL and bacteria identification was confirmed by multiplex-PCR assays. Milk samples were grouped into 7 clusters of UH status: healthy (cows with milk SCC <100,000 cells/mL and not cultured); culture-negative samples with low, medium, or high SCC; and culture-positive samples with contagious, environmental, and opportunistic intramammary infections. Data of blood serum proteins were analyzed using a linear mixed model that included the fixed effects of stage of lactation, parity, breed, herd productivity (high or low production) and UH status, and the random effect of herd-date within herd productivity. Culture-negative samples with high milk SCC, which were most likely undergoing a strong inflammatory response and whose pathogens could not be isolated because they were engulfed by macrophages or because they had already cleared, and milk samples infected by contagious and environmental bacteria were associated with greater globulin concentrations (and lower albumin-to-globulin ratio) in blood. Variation in blood serum proteins seems to be associated with inflammatory status rather than infection, as serum globulin significantly increased in UH status groups with the highest milk SCC and no differences were observed

among intramammary infections pathogens. Blood serum proteins can be a mammary gland inflammation indicator, but cannot be used to differentiate among different UH status groups.

Key words: subclinical mastitis, intramammary infection, blood serum proteins, dairy cattle

Short Communication

Udder health (UH) represents a critical issue for milk production, and monitoring the health status of the animals is of great importance for dairy farm economics. Bovine mastitis, an inflammatory status of the mammary gland in response to an infection, is the most prevalent production disease and strongly affects dairy herd income by decreasing milk yield and quality, cow welfare, fertility, and longevity (Seegers et al., 2003). Thus, a successful immune response of the animal to infection can improve milk production and reduce the treatment costs.

Blood serum proteins have been used as possible markers for assessing the immune status in dairy cows (Piccinini et al., 2004). In a previous study (Bobbo et al., 2017a), we characterized the variation in blood serum proteins [i.e., total protein, albumin, globulin and albumin-to-globulin ratio (**A:G**)] in dairy cattle housed in multibreed herds and concluded that environmental factors (e.g., herd productivity), breed, and individual cow factors (stage of lactation and parity) must be considered to appropriately interpret blood serum proteins as cow health indicators. After adjusting for those factors, a linear relationship between blood serum proteins and milk SCS was identified. Cows with high SCC in milk had greater total protein and globulin concentrations in blood serum, an expected result given that α - and γ -globulins are involved in the immune response of the mammary gland.

Besides the increased milk SCC related to the inflammatory status of the mammary gland, specific

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¹Corresponding author: alessio.cecchinato@unipd.it

IMI pathogens can elicit differential immune responses (Bannerman et al., 2004) and differences in SCC variation (de Haas et al., 2002). To our knowledge, variation in blood serum proteins among various UH status groups has not been investigated yet. Therefore, the objective of the present study was to assess the association between UH status and blood serum proteins in dairy cows. Rather than simply focus on microbiologically positive infections, we evaluated both the IMI and the resulting inflammatory response.

This study is part of the Cowplus Project, described in detail in Bobbo et al. (2017a,b). Briefly, blood and milk samples were collected from 1,508 cows belonging to 3 specialized dairy breeds (Holstein Friesian, Brown Swiss, and Jersey) and 3 dual-purpose breeds of Alpine origin (Simmental, Rendena, and Alpine Grey), that were housed in 41 multibreed herds of Trentino region (northeast Italy). In a calendar year, 1 herd per day was visited once and only clinically healthy cows at the moment of the visit were sampled during an evening milking. Details on procedures used to collect and analyze blood samples for serum proteins determination can be found in Bobbo et al. (2017a). From each selected cow, a composite milk sample (40 mL) for bacteriological analyses and a second sample (50 mL) for SCC determination were collected. Details on milk samples collection, storage, and analyses were reported in Bobbo et al. (2017b). Cows with milk SCC <100,000 cells/mL were considered potentially healthy and were not subjected to microbiological examination. Bacteriological analysis was performed on milk samples with SCC >100,000 cells/mL according to the guidelines of National Mastitis Council (NMC, 1999). Briefly, each sample was plated twice by streaking 0.01 mL of milk onto Columbia Blood Agar containing 5% defibrinated sheep blood (Oxoid Ltd., Basingstoke, UK), incubated aerobically at $37 \pm 1^\circ\text{C}$, and examined after 24 and 48 h. Grown colonies were characterized mainly according to their morphology, hemolytic features, Gram staining, catalase, oxidase, coagulase reactions, and biochemical properties (NMC, 1999). Gram-positive microorganisms were differentiated as staphylococci and streptococci by the catalase reaction. The coagulase tube test in rabbit plasma (bioMérieux Italia S.p.A., Grassano, Italy), was used to differentiate coagulase positive from coagulase-negative *Staphylococcus* spp.

The Christie, Atkins, Munch-Petersen (CAMP) test was used for the presumptive identification of *Streptococcus agalactiae*. *Enterococcus* spp. was confirmed plating suspected colonies onto Bile Esculin Agar (Oxoid Ltd.). *Lactococcus lactis* and *Aerococcus viridans* were identified by using API system (bioMérieux Italia S.p.A.). Gram-negative bacteria were identified by

their morphological features on MacConkey agar, eosin methylene blue agar, Kligler iron agar, urea agar, and SIM agar (Oxoid Ltd.). Bacterial genus confirmation and species identification of *Staphylococcus*, *Streptococcus*, and *Escherichia coli* strains was confirmed by multiplex-PCR assays, as previously described by Shome et al. (2011) with minor changes related to Taq DNA polymerase (KAPA2G FastMultiplex PCR Kit, Kapa Biosystems, Wilmington, MA) and cycling protocol.

Contaminated plates presented 3 or more different colony types, with no prevalence of a single colony type (NMC, 1999). Samples were classified as culture-negative when no bacteria were isolated or no significant growth (<1,000 cfu/mL) was observed within 48 h of incubation, with the exception of potential contagious IMI cases, for which identification was performed even when 1 colony (≥ 100 cfu/mL) was isolated.

As reported in Stocco et al. (2017), farms were classified as high or low production. Briefly, the net energy content of milk (kcal/kg) yielded the day of sampling by each cow was estimated, converted to kilojoules per kilogram and multiplied by the individual daily milk production (kg/d) to obtain the individual daily milk energy production of each cow (kJ/d). Data were then analyzed using the GLM procedure of SAS (SAS Institute Inc., Cary, NC), including herd, breed, parity, and DIM of cows, to estimate the least squares means (LSM) of the average daily milk energy production of each herd. The 41 farms were ranked according to the estimated LSM of their average daily milk energy yield, and were classified as low- ($n = 21$) or high-producing ($n = 20$) based on the median value of their LSM.

Seven clusters of UH status were identified (Bobbo et al., 2017b): healthy (cows with milk SCC <100,000 cells/mL and not tested for the presence of bacteria); culture-negative samples with low (**No Growth_L**), medium (**No Growth_M**), and high (**No Growth_H**) SCC (divided on the basis of the SCS 25th and 75th percentiles); and culture-positive samples with contagious, environmental, and opportunistic IMI.

Data of blood serum proteins were analyzed using the MIXED procedure of SAS with the linear mixed model applied also in Bobbo et al. (2017b):

$$y_{ijklmno} = \mu + DIM_i + Parity_j + Breed_k + UH\ status_l + HP_m + HTD_n(HP)_m + e_{ijklmno}$$

where $y_{ijklmno}$ is the investigated trait (blood serum proteins); μ is the overall mean; DIM_i is the fixed effect of the i th class of DIM ($i = 6$ classes of 60-d intervals); $Parity_j$ is the fixed effect of the j th parity ($j = 1$ to ≥ 4); $Breed_k$ is the fixed effect of the k th breed ($k =$ Holstein

Table 1. Results of the bacteriological analysis of 639 composite milk samples with SCC >100,000 cells/mL¹

Udder health status group	No.	%
Contagious	172	26.9
<i>Staphylococcus aureus</i>	151	23.6
<i>Streptococcus agalactiae</i>	11	1.7
<i>Staphylococcus aureus</i> + <i>Streptococcus dysgalactiae</i>	5	0.8
<i>Staphylococcus aureus</i> + <i>Streptococcus agalactiae</i>	2	0.3
<i>Staphylococcus aureus</i> + <i>Enterococcus</i> spp.	1	0.2
<i>Staphylococcus aureus</i> + <i>Streptococcus uberis</i>	1	0.2
<i>Staphylococcus aureus</i> + other streptococci	1	0.2
Environmental	102	16.0
Other streptococci	26	4.1
<i>Enterococcus</i> spp.	21	3.3
<i>Streptococcus dysgalactiae</i>	16	2.5
<i>Streptococcus uberis</i>	14	2.2
<i>Proteus</i> spp.	9	1.4
<i>Aerococcus viridans</i>	5	0.8
<i>Escherichia coli</i>	5	0.8
<i>Klebsiella</i> spp.	2	0.3
<i>Bacillus</i> spp.	1	0.2
<i>Enterobacter</i> spp.	1	0.2
<i>Lactococcus lactis</i>	1	0.2
<i>Aerococcus viridans</i> + CNS	1	0.2
Opportunistic	59	9.2
CNS	59	9.2
No growth	245	38.3
Contaminated	61	9.5

¹Adapted from Bobbo et al. (2017b).

Friesian, Brown Swiss, Jersey, Simmental, Rendena, and Alpine Grey); UH_{status_l} is the fixed effect of the l th group of infection status (l = healthy, No Growth_L, No Growth_M, No Growth_H, contagious, environmental, opportunistic); HP_m is the fixed effect of the m th herd productivity [m = high or low production]; $HTD_n(HP)_m$ is the random effect of the n th herd-date (n = 41 levels) within the m th herd productivity; and $e_{ijklmno}$ is the random residual. A normal distribution was assumed for herd-date and residuals, with a mean of zero and a variance of σ_h^2 and σ_e^2 , respectively. Proportion of variance explained by herd-date was determined by dividing the corresponding variance component by the total variance. Pairwise comparisons between UH status groups were performed using the Tukey correction ($P < 0.05$).

Results of the bacteriological analysis were reported in Table 1. Healthy animals represented about 58% of the samples and had a mean milk SCS of 1.48. Due to the low frequency of recovery of some pathogens, we classified them as contagious, environmental, and opportunistic, a classification commonly used based on reservoir and mode of transmission. Contagious IMI (mean milk SCS = 4.81), identified if at least 1 colony (≥ 100 cfu/mL) of *Staphylococcus aureus* or *Strep. agalactiae* was isolated, were the most frequent, corresponding to about 27% of the cultured samples.

Environmental and opportunistic pathogens were considered to cause an IMI if at least 10 grown colonies/type (1,000 cfu/mL) were isolated in the plate. Environmental IMI (16%, mean milk SCS = 4.70) were caused by *Enterococcus* spp., *Streptococcus dysgalactiae*, *Streptococcus uberis*, *Proteus* spp., *A. viridans*, *E. coli*, *Klebsiella* spp., *Bacillus* spp., *Enterobacter* spp., *L. lactis*, and other streptococci. Opportunistic IMI (9%, mean milk SCS = 4.60) were related to the presence of CNS in milk. Sixty-one samples were contaminated and thus excluded from the analysis, whereas 245 were culture-negative (no growth), even if the mean milk SCS was relatively high (4.38). Three hypotheses were considered to explain the high SCC in samples with no bacterial growth, namely (1) healing process and spontaneous elimination of the infection at the moment of the sampling, (2) high level of inflammation and internalization of the pathogens by phagocytes, and (3) presence of false-negative results due to a dilution effect of composite samples. Culture-negative samples were thus divided into 3 subgroups on the basis of SCS 25th and 75th percentiles as culture-negative samples with low SCS (No Growth_L; 61 samples with mean SCS = 3.21), medium SCS (No Growth_M; 122 samples with mean SCS = 4.07), and high SCS (No Growth_H; 62 samples with mean SCS = 6.13).

Coefficient of variation of blood serum proteins ranged from 7 to 16% (Table 2). Serum total protein averaged 74.17 g/L, resulting from the sum of albumin and globulin (30.77 and 43.40 g/L, respectively). Albumin-to-globulin ratio had a mean value of 0.72.

In accordance to our previous results (Bobbo et al., 2017a), the proportion of variance explained by herd-date was approximately 20% and all explanatory variables were important sources of variation of the blood serum proteins, with the exception of herd productivity, which was only associated with albumin concentration in blood (Table 3). In particular, the association previously observed between blood serum proteins and mammary gland inflammation, determined by the measure of SCC in milk, is now supported by the evidence of an association also with UH status (Table 3).

Table 2. Descriptive statistics of serum proteins (n = 1,447)¹

Serum protein	Mean	CV, %	P1	P99
Total protein, g/L	74.17	7.5	61.29	88.66
Albumin, g/L	30.77	6.9	24.84	35.22
Globulin, g/L	43.40	13.5	32.83	61.03
Albumin:globulin	0.72	15.9	0.43	0.99

¹P1 = 1st percentile; P99 = 99th percentile.

Table 3. Results of ANOVA (*F*-value and *P*-value) for serum proteins

Effect	Total protein, g/L		Albumin, g/L		Globulin, g/L		Albumin:globulin	
	<i>F</i>	<i>P</i>	<i>F</i>	<i>P</i>	<i>F</i>	<i>P</i>	<i>F</i>	<i>P</i>
Herd productivity	2.4	0.133	14.8	<0.001	0.0	0.945	1.8	0.187
HTD, ¹ %	22.0		22.5		20.3		21.4	
Breed	7.2	<0.001	7.1	<0.001	8.9	<0.001	12.4	<0.001
DIM	3.2	0.007	7.6	<0.001	3.3	0.006	4.7	<0.001
Parity	28.7	<0.001	2.8	0.040	23.2	<0.001	15.6	<0.001
UH status ²	6.9	<0.001	3.2	0.004	8.4	<0.001	8.2	<0.001
RMSE ³	4.75		1.80		5.11		0.10	

¹HTD = herd-date effect expressed as proportion of variance explained by herd-test date calculated by dividing the corresponding variance component by the total variance.

²UH status = udder health status groups, defined according to Bobbo et al. (2017b): healthy (cows with milk SCC <100,000 cells/mL and not tested for the presence of bacteria), culture-negative samples divided into 3 classes on the basis of SCS 25th and 75th percentiles: with low (No Growth_L; 100–137 cells × 10³/mL), medium (No Growth_M; 137–425 cells × 10³/mL), and high (No Growth_H; >425 cells × 10³/mL) SCC, culture-positive samples with contagious, environmental, and opportunistic IMI.

³RMSE = root mean squared error.

Greater total protein concentrations were measured in blood of cows infected by contagious pathogens and in animals whose milk was characterized by medium to high SCC, but no bacteria were isolated (No Growth_M and No Growth_H) in comparison to healthy cows (Figure 1a). As serum albumin concentration did not differ among UH status groups (Figure 1b), with the exception of a slightly decrease during environmental IMI in comparison to healthy animals, the increase in total protein can be explained by an increase in the globulin fraction. Culture-negative samples with high milk SCC (NoGrowth_H), which were most likely undergoing a strong inflammatory response and pathogens could possibly not be isolated because they were engulfed by macrophages (Bobbo et al., 2017b), and milk samples infected by contagious and environmental bacteria were associated with greater globulin concentration in blood (Figure 1c). As a consequence, lower A:G was reported for those 3 UH clusters (Figure 1d). Variation in serum proteins was not observed with opportunistic IMI. Because CNS are commonly found on udder skin, some of the culture-positive results may be due to teat skin contamination during the collection of composite samples, rather than a real mammary gland infection (Thorberg et al., 2009). Thus, as the increase in globulin concentration was observed in UH status groups with the highest milk SCS (6.13, 4.80, and 4.70, for No Growth_H, contagious, and environmental IMI, respectively) and no differences were observed among IMI pathogens, variation in serum proteins seems to be associated with inflammation rather than infection. These findings support the results previously reported in Bobbo et al. (2017b), in which the greatest impairment in milk composition and technological properties related to cheese production was observed in the

2 UH status groups with the highest milk SCC (i.e., contagious IMI and culture-negative samples with high SCC).

Somatic cells and blood serum proteins seem to be 2 key components of the immune response. Once inside the teat cistern, the contact between pathogens and leukocytes and epithelial cells induces the activation of the innate immune system. After pathogen recognition, realized through the expression of specific receptors such as the cytokines toll-like receptors and CD14, bacterial growth is inhibited through the recruitment of circulating neutrophils from the bloodstream to the mammary tissue (Wellnitz and Bruckmaier, 2012). Leukocytes release several cytokines, including tumor necrosis factor- α and IL-8, which induce the synthesis of positive acute phase proteins, such as the α -globulins serum amyloid A and haptoglobin (Viguier et al., 2009). Alpha-globulins are produced in the liver and migrate to the site of infection to inhibit growth of microbes; thus, higher levels can be detected in the blood during inflammatory status, partially explaining the increase in globulin concentration observed in the serum of cows with high SCC in milk (Figure 1c). Serum amyloid A and haptoglobin have already been recognized as possible mastitis indicators, in association with the traditional SCC (Petersen et al., 2004).

The γ -globulin fraction is actively involved in the defense system. In fact, different immunoglobulin isotypes prevent colonization by bacteria (IgA) and support phagocytosis by neutrophils (IgG and IgM; Korhonen et al., 2000). In particular, animals with a successful immune response to IMI pathogens should have rapid and effective neutrophils recruitment ability and high levels in blood serum of the opsonizing antibodies IgG2, with subsequent leakage into milk (Burton and Ers-

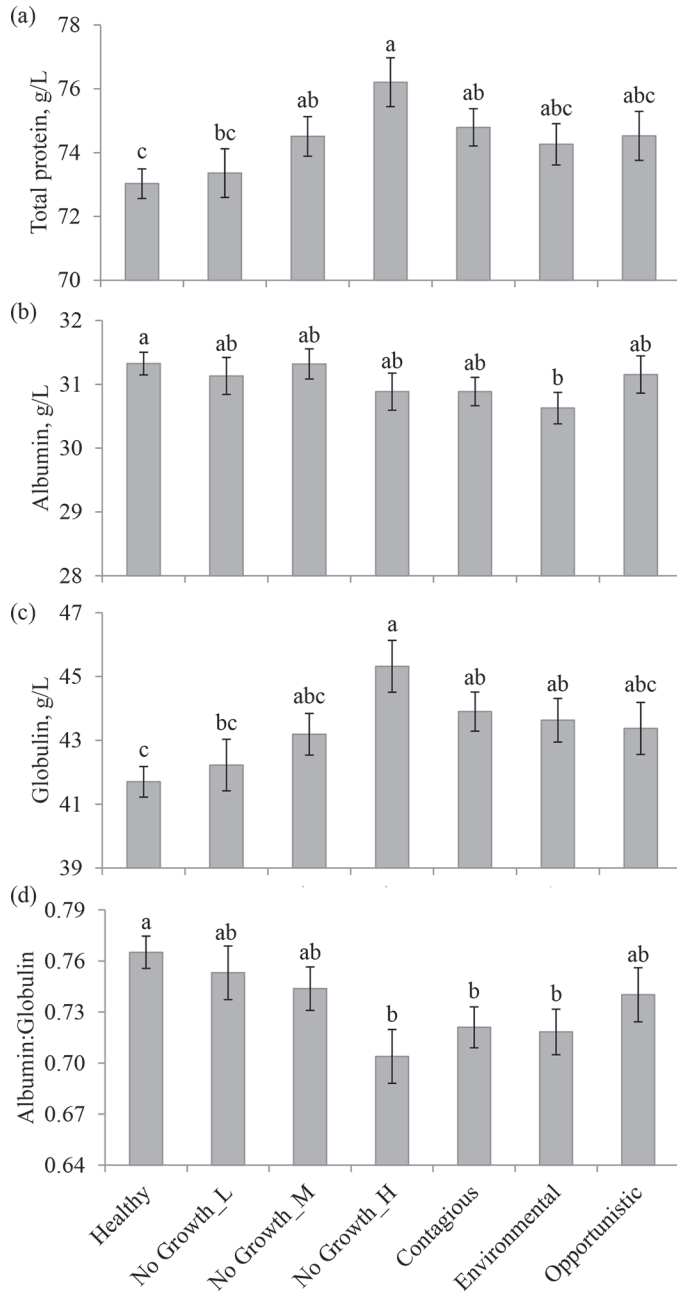


Figure 1. Least squares means and SE of serum proteins by different udder health status groups. Groups were defined according to Bobbo et al. (2017b): healthy (cows with milk SCC <100,000 cells/mL and not tested for the presence of bacteria); culture-negative samples were divided into 3 classes on the basis of SCS 25th and 75th percentiles with low (No Growth_L; 100–137 cells $\times 10^3$ /mL), medium (No Growth_M; 137–425 cells $\times 10^3$ /mL) and high (No Growth_H; >425 cells $\times 10^3$ /mL) SCC, culture-positive samples with contagious, environmental, and opportunistic IMI. The LSM with different letters (a–c) statistically differ (Tukey adjusted $P < 0.05$).

kyne, 2003). Thus, when appropriate antibodies bind bacteria, pathogen recognition and destruction by neutrophils is facilitated. A positive correlation between

IgG2 level in serum and clinical mastitis has been observed in Holstein Friesian cows (Mallard et al., 1983). Conversely, a genetic deficiency in IgG2 synthesis was associated in Danish Red cattle with a higher frequency of disease caused by pyogenic bacteria, including mastitis (Nansen, 1972). To synthesize pathogen-reactive IgG2 antibodies, B lymphocytes need IFN- γ and other cytokines, secreted by T cells only after recognition of specific antigens during infection (Burton and Erskine, 2003). Thus, the increased synthesis of γ -globulins during mammary gland inflammation may further explain the greater globulin concentration in serum of cows with high milk SCC.

Results of this study suggest that blood serum proteins (i.e., total protein, albumin, globulin, and A:G) can be a mammary gland inflammation indicator, but cannot be used to differentiate among different UH status groups. However, as different etiological agents can induce differential immune responses and stimulate the synthesis of different proinflammatory cytokines (Oviedo-Boyso et al., 2007), further studies will be required to deeply investigate variation in specific globulin fractions among different clusters of UH status.

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