

***Salmonella dublin* in Danish Dairy Herds: Frequency of Change to Positive Serological Status in Bulk Tank Milk ELISA in Relation to Serostatus of Neighbouring Farms**

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Wedderkopp A, Strøger U, Lind P: *Salmonella dublin* in Danish dairy herds: Frequency of change to positive serological status in bulk tank milk ELISA in relation to serostatus of neighbouring farms. Acta vet. scand. 2001, 42, 295-302. – Bulk tank milk from 1,429 herds were collected in 3 rounds from 19 different geographic areas. The milk samples were tested by use of indirect LPS-ELISA procedure to detect *Salmonella dublin* antibodies. From the obtained OD-values herd seroprevalence in the given area was determined and GR-scores calculated for each herd by addition of the number of positive sampling rounds by the 5 geographically closest neighbour herds. In the 19 different areas the calculated prevalence ranged from 0.01 to 0.41. Totally 3,697 GR-scores were given. The mean GR-scores in the areas ranged from 0.0 to 6.5. Higher GR-scores were found in herds changing to seropositive status compared with herds seronegative throughout the study period. The results indicate that the risk for a dairy herd to receive *S. dublin* infection increases with the disease status among the nearest neighbours and with the prevalence of seropositive herds in the geographic area.

***Salmonella dublin*; cattle; dairy herds; milk samples; LPS-ELISA; epidemiology; seroprevalence; risk.**

Introduction

Salmonella dublin is a host-adapted serotype causing infection in cattle of all ages (Richardson 1975). It has been isolated from cattle in all parts of the world (Walker 1995). In Denmark this serotype has been isolated in 60%-75% of all diagnosed *Salmonella* outbreaks in cattle since 1980 (Annual Report, Danish Veterinary Laboratory 1989-1995). *S. dublin* may persist in the cattle herds, as animals of all ages may be infected from *S. dublin* contaminated environments and some animals may become carriers excreting the organism in faeces for years, eventually for life (Wray et al. 1989). An ELISA test for IgG antibodies directed against

S. dublin LPS has been shown effective in detecting both mammary and faecal carriers (Smith et al. 1992). The prevalence of *Salmonella* infection in dairy herds has been determined by means of serologic testing and percentage of culture-positive calves and environment in California (Smith et al. 1994). It is possible to screen milk samples for *S. dublin* infection in dairy herds (Hoorfar et al. 1994, Hoorfar et al. 1995, Hoorfar et al. 1996). The LPS-ELISA has the capability to detect *S. dublin* infection in dairy herds and to point out newly infected herds by repeated testing (Wedderkopp et al. 2001). In order to control and

eventually eliminate an infection figures for risk and prevalence are beneficial. Spatial analysis of seroconversion, and spatial and temporal analysis of antibiotic resistance in cattle have been performed (Ward et al. 1996, Singer et al. 1998). The purpose of the present study was to estimate the prevalence of *S. dublin* infection for different geographic areas and to evaluate the risk of receiving the infection if the neighbouring herds have *S. dublin* infection.

Materials and methods

Test samples

A total of 4,287 bulk tank milk samples were collected from March 1994 to March 1996 in 3 test rounds from a total of 1,429 dairy herds (Wedderkopp et al. 2001). Nineteen geographic areas were represented. The radius of the areas ranged from 6 to 21 kilometres, and the number of dairy herds ranged from 16 to 258. Two geographic areas (Bornholm and Samsø) were islands. Inside the geographic areas the herds were equally distributed, and almost all dairy herds within the designated areas participated. Consequently approximately 10 percent of all Danish dairy herds participated in the study.

The milk samples were taken in connection with the Danish control program for milk quality. The samples were transported without preservatives at 5°C to the laboratory, where they were centrifuged at 200 × g for 5 min and the fat layer removed. The samples were stored in polypropylene tubes at -18°C until use.

Serology

The ELISA described by Hoorfar et al. (1995), was slightly modified for this study. Briefly: microwell plates (Cat. # 475094, Nunc, Denmark) were coated with *S. dublin* LPS antigen, and an undiluted positive, a weakly positive and a negative control samples were added in duplicate to each plate, each test milk was added undiluted (100 µL) in duplicate, and the plates were incu-

bated overnight at 5°C. The bound antibodies were detected by using a purified immunoglobulin fraction of goat antiserum to bovine IgG (γ), labelled with horseradish peroxidase (Cat. # 14-12-02, Kirkegaard & Perry Lab. Maryland). H₂O₂ was used as a substrate and OPD (1,2-Ortho-Phenyl-Diamine) as an indicator. A test set-up was considered valid if the negative milk control had an OD below 0.2 and the weak positive milk control had OD between 0.6 and 1.0. A test-sample was considered positive with OD values ≥0.30, and negative with OD-values <0.30 (Wedderkopp et al. 2001).

Statistical analyses

For the calculations, statistical and geographic analyses SAS Software (version 6.11, OS/2) was used, while @Risk (Palisade corporation, Newfield) was used for the risk analysis and for calculation of the true prevalences.

Each herd was given a geographical risk score (GR-) from 0 to 15, according to the reactions of the 5 nearest neighbouring dairy herds within its designated area: If all of the 5 neighbours tested negative in all 3 test rounds the herd was given the score 0 - if only 1 of the 5 neighbours tested positive in one test-round the herd was given the score 1- if 2 herds of the 5 neighbours tested positive each in one test-round or one tested positive in 2 test-rounds the herd was given the score 2 and so on, so that the maximum score 15 was given when all the neighbours tested positive in all 3 test-rounds. As the closest neighbours not necessary were located within the designated geographical area, a certain edge effect will confound the calculations. The size of the effect will be negatively correlated with the number of herds included.

The distribution of GR-scores in the 4 groups: repeatedly positive herds, herds changing from positive to negative, herds changing from negative to positive, and herds fluctuating from neg-

Table 1. Distribution of scores according to serological status.

	Number of Herds (%)*	Number of GR-scores (%)*	Mean GR-scores*	P-value
All Herds	1429 (100)	3697 (100)	2.6	
Herds test-negative in all rounds	1058 (74.0)	2251 (60.9)	2.1	
Herds test-positive in all rounds	143 (10.0)	689 (18.6)	4.8	0.0000
Herds changing from test-positive to test-negative	55 (3.8)	197 (5.3)	3.6	0.0003
Herds changing form test-negative to test-positive	117 (8.2)	394 (10.7)	3.4	0.0001
Herds fluctuating from test-negative to test-positive to test-negative and reverse	56 (3.9)	166 (4.5)	3.0	0.0330

*The number of herds, GR-scores, and mean GR-scores are stated according to whether the herds tested negative in all test-rounds, tested positive in all test-rounds or changed test-status during the study period. The GR-scores were divided into 4 groups: repeatedly positive herds, herds changing from positive to negative, herds changing from negative to positive, and herds fluctuating from negative to positive and reverse. The groups were compared to the repeatedly negative herds using chi-square.

ative to positive and reverse were compared to the repeatedly negative herds using chi-square from the Proc catmod procedure in SAS.

The true prevalence of *S. dublin* infection was calculated for each area using the Beta-distribution, Beta (α_1, α_2), where $\alpha_1 = (r + 1)$ and $\alpha_2 = (n - r + 1)$ and r = the true number of infected herds in all test rounds, and n = the total number of herds (Vose 1996). Where the true number of infected herds was obtained by adding the false negative and subtracting the false positive while correcting for the specificity and sensitivity of the test previously estimated to 0.89 and 0.88 respectively (Wedderkopp *et al.* 2001).

For estimation of the number of false negative samples the outcome of a negative binomial distribution with parameters (s, p), where s = number of test-positive herds + 1 and p = sensitivity was used (Vose 1996).

Spearman rank correlation was used to determine the correlation between the calculated

true prevalence, the proportion of herds changing from test-negative to test-positive, the mean GR-scores given to the group of herds, which tested constantly negative, the mean GR-scores given to the group of herds changing from test-negative to test-positive.

Results

A total of 3,697 GR-scores were given reflecting the test-results from the 5 nearest neighbours. A total of 1,058 herds (74.1%) were constantly negative in all 3 test-rounds, 90% of these herds were given less than 6 GR-scores. A total of 117 herds (8.1%) turned positive during the study, 90% of these herds were given 9 GR-scores and less.

Within the geographic areas the mean GR-scores ranged from 0.0 to 6.5 with mean GR-score = 2.6. The mean score was significantly higher for herds changing from test-negative to test-positive (GR = 3.4) during the study period than for herds that remained test-negative (GR

= 2.1) throughout the study period (chi-square: 16.26, $p < 0.0001$). The highest mean scores were, however, obtained for herds, that tested positive in all test rounds (GR = 4.8) (Table 1). For each area and for each round the true prevalence was calculated with the mean true prevalence ranging from 0.01 to 0.41. The percentage of herds changing from test-negative to test-positive ranged from 0.0 to 13.5. The mean number of GR-scores for the group of herds, which were constantly negative ranged from 0.0 to 6.8, and for the group of herds, which turned positive the number of GR-score ranged from 0.0 to 7.3.

For each of the 19 geographic areas the calculated true prevalence was compared to the proportion of herds changing from test-negative to test-positive, and to the mean GR-scores given to herds testing negative in all test-rounds and herds changing from test-negative to test-positive respectively. Spearman rank correlation between the calculated true prevalence and the

proportion of herds changing from test-negative to test-positive was 0.63 ($p = 0.004$). Spearman rank correlation between the calculated true prevalence and the mean GR-scores given to herds testing negative all round and to herds changing form test-negative to test-positive was 0.94 ($p = 0.000$) and 0.66 ($p = 0.002$), respectively (Fig. 1).

As a mean true prevalence of 0.10 divided the geographic area into two sections fairly uniform according to the numbers of geographic areas, the material was stratified according to this mean true prevalence: One group of areas ($n = 11$) with a mean true prevalence lower than 0.10 (range: 0.01 to 0.03) and one group of areas ($n = 8$) with a mean true prevalence higher than 0.10 (range: 0.11 to 0.41). The distribution of test-negative herds compared to the herds which changed form test-negative to test-positive in the two prevalence areas is shown in Fig. 2. Statistical analysis showed no significant difference in the mean scores for the herds in the

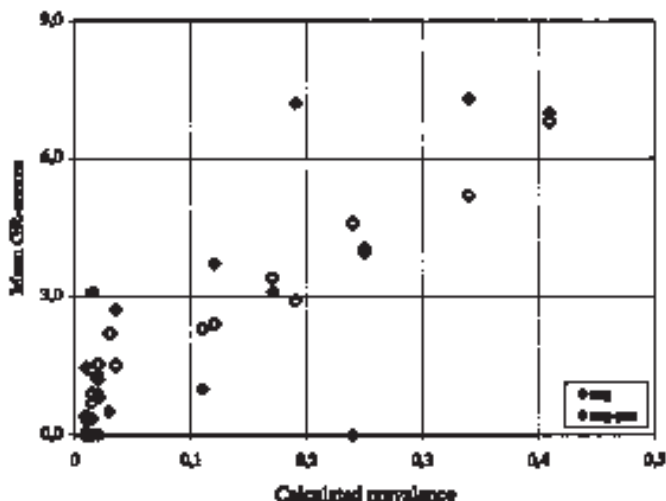


Figure 1. Comparison of prevalence and mean geographical risk-scores. For each area the calculated prevalence are compared to the mean GR-risk scores given to herds testing negative in all test-rounds, and the herds changing from test-negative to test-positive respectively.

low prevalence areas (639 herds, chi-square = 0.82, $p = 0.364$). In the high prevalence areas (790 herds) the mean score was significantly higher for the herds changing from negative to positive during the study period (mean score = 4.4), compared to the herds that remained negative (mean score = 3.5), chi-square = 4.31 ($p = 0.038$).

In the low prevalence areas ($n = 11$) the correla-

tion between the calculated true prevalence and the mean GR-scores given to the herds, which tested constantly negative was 0.67 ($p = 0.024$). In the high prevalence areas ($n = 8$) the correlation between the calculated true prevalence and the mean GR-scores given to the herds, which tested constantly negative was 0.95 ($p = 0.0003$). All the calculations were corrected for ties (Fig. 1).

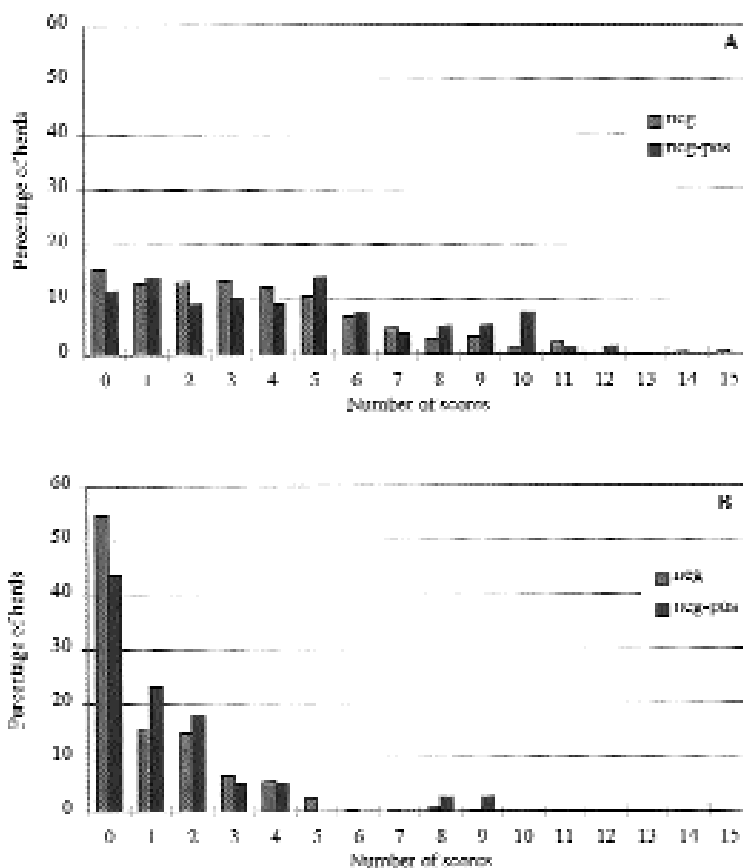


Figure 2. Comparison of test-negative herds and herds changing from test-negative to test-positive in areas with high prevalence (>0.10) and areas with low prevalence (<0.10). A: The distribution of GR-scores in the high prevalence areas where 84 herds of totally 790 herds changed from negative to positive. B: The distribution of GR-scores in the low prevalence areas where 33 herds of totally 639 herds changed from negative to positive.

For the estimation of edge effect 2 areas (Bornholm and Samsø) were left out of the calculation, as these areas were islands. For the rest of the areas no correlation between the number of herds in the designated areas and the number of GR-scores was found.

Discussion

For cattle diseases spatial analysis have been used to estimate the effect of clustering in space. Ward et al. (1996) detected clustering of bluetongue virus serotype 1 in Queensland. Singer et al. (1998) tried to determine whether ampicillin- and tetracycline-resistant strains of *Pasteurella multocida* and *P. haemolytica* were spatially and temporally clustered, and found spatial clustering of resistant isolates in California. Both studies used Cuzick and Edwards' test to detect spatial clustering by comparing coordinates of cases and controls. In the present study the coordinates were used to point out the neighbouring dairy herds within a geographic area. Queensland covers around 1,727,500 km², California covers 405,940 km², while Denmark covers 43,032 km². Thus the present study claims to yield a closer look into the areas under examination. Differences between 19 geographic areas were evaluated, and we investigated the use of a new herd parameter, the GR-score, that estimates the frequency of *Salmonella* infection in the neighbouring herds. In each area the results for the 5 nearest dairy herds were compared and each herd were given GR-scores after the results in the neighbouring dairies. Though the herds at the edge of an area are only receiving GR-scores from herds at one to 3 surrounding sides, this edge effect was left out of consideration as no correlation between number of herds, and number of mean GR-scores was found in the geographic areas.

Totally 668 herds (46.7%) were given none or one GR-score. If the GR-scores had been equally distributed between the herds studied,

the repeatedly negative herds would have received 2.736 GR-scores; instead these herds were given 485 GR-scores less (17.7%). The test-positive herds should be given only 370 GR-scores at equal distribution, instead these herds were given 319 GR-scores more (86.2%), indicating an additional risk among the test-positive herds of having a test-positive neighbour (Table 1).

That herd seroconversion is primarily determined by seroprevalence in the given geographic area is supported both by the calculated Spearman rank correlations, which was significant at 5% confidence-level, and by the calculations based on stratification into high prevalence areas and low prevalence areas (Fig. 2), where the mean number of GR-scores is 1.1 in repeatedly test-negative herds and 1.4 for the herds changing from test-negative to test-positive in the low prevalence areas, while the corresponding figures are 3.5 and 4.4 in the high prevalence areas. This difference was only statistically significant in the latter population of herds. Thus in the low prevalence areas (<0.10) acquisition of *S. dublin* infection do not appear to be directly dependent of close contact with infected neighbour herds, but may be contracted otherwise e.g. by animal trade. Indications were found for an increased risk associated with infected neighbour herds. Unfortunately we are presently not able to evaluate this effect compared to factors operating on each area in general, such as herd size, climate and management factors, all though we expect most cases of infection to occur during the grazing season, as a result of the close contact between animals at pasture. In addition the size of the areas and the number of participating herds within these were not the same in all the geographic areas examined, however the distance between neighbour herds may influence the risk for receiving the infection.

In conclusion, a higher risk for changing

serostatus from negative to positive for *S. dublin* infection was found if a dairy herd was located in a high prevalence area, and if the neighbouring herds were infected.

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Sammendrag

Salmonella dublin i malkekvægbesætninger: Ændringer til positiv serologiske status i tankmælk ELISA i forhold til serostatus i nabobesætningen.

Der blev indsamlet tankmælks prøver i 3 runder fra 1.429 besætninger placeret i 19 forskellige geografiske. Prøverne blev undersøgt ved hjælp af en indirekte LPS-ELISA med henblik på at påvise *S. dublin* antistoffer. Besætnings seroprevalence blev beregnet ud fra test-resultaterne, og der blev tildelt geografisk risiko (GR-) point på grundlag test-resultaterne fra de fem nærmeste mælkeleverende besætninger. I alt blev der tildelt 3.697 GR-point. De gennemsnitlige GR-point varierede fra 0,0 til 6,5, prævalensen varierede fra 0,01 til 0,41 i de geografiske områder, undersøgelsen omfattede. Der blev tildelt flest GR-point til besætninger, der skiftede til seropositiv status sammenlignet med besætninger, der forblev seronegative i alle tre testrunder. Resultaterne indikerer, at risikoen for at en malkekvægbesætning bliver smittet med *S. dublin*, afhænger af testresultaterne i nabobesætningerne og af seroprevalencen i det geografiske område, besætninger er placeret i.

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