Distribution of coagulase-negative Staphylococcus species from milk and environment of dairy cows differs between herds

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ABSTRACT

In many parts of the world, coagulase-negative staphylococci (CNS) are the predominant pathogens causing intramammary infections (IMI) in dairy cows. The cows’ environment is thought to be a possible source for CNS mastitis and this was investigated in the present paper. A longitudinal field study was carried out in 6 well-managed dairy herds to determine the distribution and epidemiology of various CNS species isolated from milk, causing IMI and living freely in the cows’ environment, respectively. In each herd, quarter milk samples from a cohort of 10 lactating cows and environmental samples from stall air, slatted floor, sawdust from cubicles, and sawdust stock were collected monthly (n = 13). Isolates from quarter milk samples (n = 134) and the environment (n = 637) were identified to species level using amplified fragment length polymorphism (AFLP) genotyping. Staphylococcus chromogenes, S. haemolyticus, S. epidermidis, and S. simulans accounted for 81.3% of all CNS milk isolates. Quarters were considered infected with CNS (positive IMI status) only when 2 out of 3 consecutive milk samples yielded the same CNS AFLP type. The species causing IMI were S. chromogenes (n = 35 samples with positive IMI status), S. haemolyticus (n = 29), S. simulans (n = 14), and S. epidermidis (n = 6). The observed persistent IMI cases (n = 17) had a mean duration of 149.4 d (range 63.0 to 329.8 d). The CNS species predominating in the environment were S. equorum, S. sciuri, S. haemolyticus, and S. fleurettii. Herd-to-herd differences in distribution of CNS species were observed in both milk and the environment, suggesting that herd-level factors are involved in the establishment of particular species in a dairy herd. Primary reservoirs of the species causing IMI varied. Staphylococcus chromogenes and S. epidermidis were rarely found in the environment, indicating that other reservoirs were more important in their epidemiology. For S. haemolyticus and S. simulans, the environment was found as a reservoir, suggesting that IMI with these species were possibly environmental in origin.

Key words: amplified fragment length polymorphism, coagulase-negative staphylococci, environment, cow milk

INTRODUCTION

In the last 2 decades, mastitis prevention programs (National Mastitis Council, 2009) have resulted in better control of transmission of contagious mastitis pathogens in lactating dairy cows. In many well-managed dairy herds, the obtained decrease in prevalence of IMI caused by contagious major pathogens is, however, associated with a relatively higher proportion of subclinical or mild clinical infections caused by CNS (Tenhagen et al., 2006; Bradley et al., 2007; Piepers et al., 2007). Moreover, CNS are the major cause of IMI in heifers, the future milk producers of every dairy herd (Fox, 2009; Piepers et al., 2010). The change in distribution of mastitis pathogens suggests that current mastitis control measures are less effective in reducing CNS IMI.

In general, CNS induce only a mild inflammatory reaction in infected quarters, as indicated by the modest increase in the milk SCC (Schukken et al., 2009). However, they can be a cause of (mild) clinical mastitis (Waage et al., 1999; Taponen et al., 2006), and quarters infected by CNS have been reported to be at greater risk for infection by major mastitis pathogens (Hogan et al., 1988; Lam et al., 1997). On the other hand, in some studies, CNS IMI protected quarters against new infections by major mastitis pathogens (Rainard and Poutrel, 1988; Matthews et al., 1991). It has also been demonstrated that teat apex colonization with CNS protected heifers against high SCC and new IMI by major pathogens in early lactation (De Vliegher et al., 2003; Piepers et al., 2011). The effect on quarter SCC is species-specific (Supré et al., 2011) and some CNS...
species are able to persist in the udder for long periods
(Chaffer et al., 1999; Gillespie et al., 2009), prolonging
their influence on the mammary gland. Whether this
influence is harmful or advantageous remains unres-
solved and subject to debate.

A potential drawback of many studies in the past is
the classification of CNS as a homogeneous group (Ho-
gan et al., 1988; Nickerson and Boddie, 1994), which
could partly explain the aforementioned contradictory
findings on the pathogenic or protective role of CNS.
In fact, CNS originating from cows include a variety
of species with differences in antimicrobial suscepti-
bility (Lüthje and Schwarz, 2006; Sawant et al., 2009)
and virulence factors (Park et al., 2011). Therefore,
evaluating the epidemiology of individual CNS species
is of great importance to understand their respective
significance. Based on phenotypic characteristics, spe-
cies identification of CNS isolates from cows is difficult
and often inaccurate (Capurro et al., 2009; Sampimon
et al., 2009). In the last few years several genotypic
methods have been developed that enable more accu-
rate identification of CNS species (Supré et al., 2009;
Piessens et al., 2010; Braem et al., 2011).

The high prevalence of prepartum CNS infections in
heifers (Fox, 2009) that have not yet been exposed to
the milking process suggests that sources other than
the milking machine exist. As CNS are abundantly
free-living in dairy herds (Rendos et al., 1975; Matos et
al., 1991), we can hypothesize that the cows’ environ-
ment is a possible source for CNS causing IMI. The
aims of the present study were to determine the species
in the freestall, slatted floors, used sawdust bedding
and unused sawdust from the stock. For
isolation of staphylococci from the air in the freestall,
blank air strips (Biotest, Dreieich, Germany) were filled
aseptically with mannitol salt agar (MSA, BD Biosci-
eences, San Jose, CA), and kept at 4°C until further
use. At the time of sampling, an MSA-filled strip was
removed from the sterile package, placed in the RCS
standard air sampler (Biotest), and exposed for 1 min,
which is equal to sampling of 40 L of air. The slatted
floor was sampled by walking over a whole corridor
while wearing polypropylene Sekuroka overshoes (Fi-
ers, Kuurne, Belgium), after which both overshoes were
transferred to a sterile stomacher bag (180 × 300 mm,
Medical Lab Service, Menen, Belgium). Used sawdust
samples were collected from the back one-third of 10%
of the cubicles. Samples of the sawdust stock were col-
lected at 5 random places in front of the stock using a
sample bore (5.6 cm diameter, 50 cm long). Per herd
and per sample type, collected sawdust was commingled
and a subsample was transferred to a sterile stomacher.

At the beginning of the study, a cohort of 10 clini-
ically healthy cows was randomly selected in each herd
within parity blocks (4 heifers, 3 cows of second parity,
and 3 cows of third or higher parity). Before the end
of the study, 20 out of the 60 cohort cows were culled
(ranging from 0 to 5 per herd) for diverse reasons, in-
cluding problems with lameness, udder health, fertility,
or milk production. All cohort cows culled before the
12th month of the study (n = 14) were replaced by
randomly selected herdmates of the same parity. The
cohort cows were, on average, 11.2 mo under study.

Sample Collection

Quarter milk samples were collected aseptically at
monthly intervals (n = 13) from the cohort cows ac-
cording to standard procedures (National Mastitis
Council, 1999). Samples were frozen and transported
to the laboratory of the Milk Control Centre Flanders
(MCC, Lier, Belgium) for bacteriological examination.
In addition to the monthly milk samples taken from the
cohort cows, extra samples were taken when the farmer
noticed clinical signs in a quarter (any visual abnor-
mality of milk or the udder). Clinical milk samples were
taken both from cows of the cohort and from cows out-
side the cohort. The average overall incidence rate of
clinical mastitis (IRCM; all pathogens) was calculated
as described by Barkema et al. (1998).

Environmental samples were taken once a month (n
= 13) in each farm for isolation of CNS. Four different
sample types were chosen in proximity to the cows: air
in the freestall, slatted floors, used sawdust bedding
from cubicles, and unused sawdust from the stock. For
isolation of staphylococci from the air in the freestall,
blank air strips (Biotest, Dreieich, Germany) were filled
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sample bore (5.6 cm diameter, 50 cm long). Per herd
and per sample type, collected sawdust was commingled
and a subsample was transferred to a sterile stomacher.
In one herd (herd F), no sawdust samples were collected during the first 5 mo of the study, as the farmer did not use any bedding material in the cubicles at that time. All environmental samples were stored at 4°C and transported on the same day to the laboratory for processing.

**Sample Processing and Isolation of CNS**

**Quarter Milk Samples.** Bacteriological culture of quarter milk samples and bacterial identification was done in the Milk Control Centre Flanders as recommended by the National Mastitis Council (1999). Briefly, 0.01 mL of each quarter milk sample was spread on a quadrant of a blood-esculin agar plate and incubated aerobically at 37°C ± 1°C for 36 h ± 12 h. A quarter was considered culture-positive when growth of ≥1 colony was detected. A sample was considered contaminated when 3 or more dissimilar colony types were observed. Phenotypic differentiation of bacterial species was done as described by Piepers et al. (2007). *Staphylococcus aureus* was differentiated from other *Staphylococcus* spp. based on morphology, pigmentation, hemolysis, and DNase activity. All non-*S. aureus* staphylococci were a priori considered as CNS. For milk samples yielding at least 3 CNS colonies (≥300 cfu/mL), 2 colonies were picked and transferred to tryptone soy agar (TSA, Oxoid Ltd., Basingstoke, UK) for further identification. When more than one type of CNS colony was present, more colonies were picked. The TSA plates were incubated at 37°C for 18 h.

**Environmental Samples.** On the same day of sample collection, exposed air strips were incubated at 37°C for 24 to 48 h. Of each sawdust sample, 25 g was weighed and put in a new sterile stomacher bag. Overshoes were transferred to separate sterile stomacher bags. To each sawdust sample and overshoe, 225 mL of brain heart infusion broth (Oxoid Ltd.) + 7.5% NaCl was added. Sawdust samples were homogenized for 1 min in a stomacher, and overshoes were massaged by hand for 1 min for equal distribution of the medium. Samples were then incubated for 20 h ± 4 h at 37°C for selective enrichment of staphylococci. The next day, dilution series of the enrichment media were made in Ringers solution (Oxoid Ltd.), 100 μL of dilutions 10^{-2} to 10^{-5} were spread by use of sterile glass beads (2.5–3.5 mm, VWR International, West Chester, PA) on MSA plates, which were incubated for 20 h ± 4 h at 37°C. After incubation, *Staphylococcus* -like colonies were picked up from MSA plates and strips and were streaked onto TSA. Choice of colonies was random but

<table>
<thead>
<tr>
<th>Herd</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>F</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average herd size (n)</td>
<td>76</td>
<td>87</td>
<td>92</td>
<td>102</td>
<td>110</td>
<td>115</td>
</tr>
<tr>
<td>Average production per cow per year (kg)</td>
<td>8,559</td>
<td>9,955</td>
<td>8,507</td>
<td>10,372</td>
<td>11,665</td>
<td>7,147</td>
</tr>
<tr>
<td>Average bulk milk SCC (cells/mL of milk)</td>
<td>195,462</td>
<td>202,154</td>
<td>85,615</td>
<td>175,000</td>
<td>120,846</td>
<td>185,000</td>
</tr>
<tr>
<td>Incidence rate of clinical mastitis cases</td>
<td>0.180</td>
<td>0.236</td>
<td>0.347</td>
<td>0.485</td>
<td>0.290</td>
<td>0.149</td>
</tr>
<tr>
<td>Milking parlor</td>
<td>2 × 4 open tandem</td>
<td>2 × 6 herringbone</td>
<td>2 × 5 herringbone</td>
<td>2 × 4 herringbone</td>
<td>2 × 8 herringbone</td>
<td>2 × 8 herringbone</td>
</tr>
<tr>
<td>Bedding material</td>
<td>Sawdust</td>
<td>Sawdust</td>
<td>Sawdust</td>
<td>Sawdust</td>
<td>Sawdust</td>
<td>Sawdust or none</td>
</tr>
<tr>
<td>Use of pre-dip (yes/no)</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Postmilking dip/spray</td>
<td>Spray</td>
<td>Dip</td>
<td>Dip</td>
<td>Dip</td>
<td>Spray</td>
<td>Spray</td>
</tr>
<tr>
<td>Post-dip main component</td>
<td>Iodine</td>
<td>Iodine</td>
<td>Chlorhexidine digluconate</td>
<td>Iodine</td>
<td>Lactic/caprylic/capric acid</td>
<td>Iodine or iodine</td>
</tr>
</tbody>
</table>

1Average herd size during the 13 mo of study.
2Calculated as the number of quarter cases per 365 cow-days at risk.
3The farmers of herds E and F switched to an iodine post-dip product in mo 7 of study.
based on dissimilar colony morphology and pigmentation. More colonies were picked from agar plates and strips with more diverse bacterial growth. The TSA plates were further incubated for 18 h at 37°C and checked for purity the next day.

Identification of CNS Isolates

Confirmation of CNS Identity. A crude DNA preparation was made for all isolates from milk and for a random selection of 4 isolates per environmental sample with dissimilar colony morphology and pigmentation. A few fresh colonies were suspended in 45 μL of sterile HPLC water plus 5 μL of lyostaphin (1 mg/mL; Sigma-Aldrich, St. Louis, MO) and incubated at 37°C for 10 min. Next, 150 μL of Tris-HCl (0.1 M, pH 8.0) plus 5 μL of proteinase K (2.5 mg/mL; Promega Corporation, Madison, WI) were added, and the suspensions were incubated at 60°C for 10 min followed by 90°C for 5 min. Cell lysates were vortexed for 1 min, centrifuged at 14,000 × g for 1 min, and kept at −20°C until further use. To confirm the isolates as *Staphylococcus* species and to overrule *S. aureus* identity, a duplex PCR was performed. Two primer pairs described previously by Mason et al. (2001) were used, the first pair targeting the *Staphylococcus* genus-specific 16S rRNA gene (fragment of 791 bp) and the second primer pair targeting the *S. aureus*-specific *clfA* gene (fragment of 638 bp). The PCR was performed in a 25-μL reaction mixture containing 1 μL of cell lysate, 50 pmol of each primer (Eurogentec, San Diego, CA), 1 × PCR buffer II (Applied Biosystems, Foster City, CA), 1.5 mM MgCl₂, 0.1 mmol of each dNTP, and 1 U of AmpliTaq Polymerase (Applied Biosystems). Thermal cycling conditions were 1 min at 95°C, 30 cycles of 15 s at 95°C, 15 s annealing at 60°C, and 30 s elongation at 72°C, and a final elongation step at 72°C for 8 min. Fragments were analyzed by electrophoresis on 1.5% (wt/vol) agarose gels. In each executed PCR run, a positive *S. aureus* control (*S. aureus* ssp. DSM 20315), a positive CNS control (*S. auricularis* ATCC 33753), and a negative control (water) were co-analyzed.

Selection of Isolates for AFLP Genotyping. All isolates considered as CNS by PCR were subsequently analyzed by a rapid fingerprinting technique to avoid redundancy in the CNS collection. Random amplification of polymorphic DNA (RAPD)-PCR was done on 1 μL of cell lysate using the primer D11344 and PCR conditions as described by Fitzgerald et al. (1997). The RAPD-PCR fragments were separated on 2% (wt/vol) Seakem LE agarose gels (Lonza, Basel, Switzerland) at 100 V for 75 min, and RAPD fingerprints of isolates originating from the same milk sample or the same environmental sample were compared visually. When identical RAPD fingerprints were observed, only one isolate per sample was retained in the final CNS collection. Selected CNS isolates were subcultured on TSA for another 18 h at 37°C and stored as frozen stocks in brain heart infusion broth (Oxoid) with 15% (wt/vol) glycerol at −80°C for further species identification.

Genotyping and Identification of CNS Species. The DNA was prepared and amplified fragment length polymorphism (AFLP) genotyping was done on the final CNS collection as described previously (Piessens et al., 2010). The BioNumerics software version 6.01 (Applied Maths, Sint-Martens-Latem, Belgium) was used for normalization of fingerprints, and the library and identification module was used for calculation of genetic similarities and identification of the field isolates. Species identification was done based on similarity of AFLP fingerprints to entries in the staphylococcal AFLP library, which contained fingerprints of 54 CNS type and reference strains, representing 49 different CNS species and subspecies, and 247 well-identified bovine CNS isolates belonging to 18 CNS species common in cattle (Piessens et al., 2010). Similarities were calculated based on the Pearson product-moment correlation coefficient, and 50% similarity to a library entry was used as a cut-off for species identification. When an isolate showed less than 50% similarity to all library entries in the numerical analysis, its fingerprint was visually compared with library strains in a dendrogram constructed with the unweighted pair group method with arithmetic mean (UPGMA) clustering algorithm. Due to different intensities of fingerprints, similarity can be <50%, although fingerprints visually match. When clearly matching AFLP fingerprints were found in the library, the unknown field isolate was assigned to the corresponding CNS species.

Sequencing of the rpoB Gene. Isolates showing AFLP fingerprints that could not be assigned to any CNS species by numerical analysis or visual comparison were further analyzed by sequencing of the *rpoB* gene. Per unknown cluster in the AFLP dendrogram, identification of at least one isolate was done based on *rpoB* gene sequencing as described elsewhere (Supré et al., 2009).

Definition of CNS IMI

Quarters were diagnosed to have CNS IMI when at least 2 out of 3 consecutive quarter milk samples were found to be culture-positive for the same CNS species with ≥300 cfu/mL and when consecutive isolates had a similar AFLP type. In all other cases, a quarter was considered to be noninfected or of unknown IMI status when no sample was taken in one of the adjacent months (e.g., first sampling, dry period, culling). An
DISTRIBUTION OF COAGULASE-NEGATIVE STAPHYLOCOCCI

IMI that was present at only 1 sampling was considered a transient case. All other IMI were considered persistent in nature. Duration of persistent IMI cases was calculated by assuming that infections started at the midpoint between the first detection of IMI and the previous sampling, and ended at the midpoint between the last detection of IMI and the next sampling, respectively. When a cow entered or left the study infected, the extra period was calculated as the average number of days between 2 samplings for that herd (n = 13), divided by 2.

RESULTS

CNS in Milk

Based on bacteriological examination, 154 out of 2,580 cohort quarter milk samples (6.0%) were culture-positive for CNS (≥100 cfu/mL). Only CNS isolates from milk samples yielding ≥300 cfu/mL were further considered for identification. After duplex PCR (for confirmation of CNS identity) and RAPD analyses (for exclusion of duplicate CNS isolates per sample), 134 CNS isolates originating from 59 different quarters of 37 cows were retained. Six quarter milk samples showed mixed growth of 2 CNS species. All CNS isolates from milk were identifiable to species level by numerical comparison to the AFLP library (n = 126), by visual comparison of clusters in an AFLP dendrogram (n = 2), or by AFLP and rpoB gene sequencing combined (n = 6). In total, 13 species were differentiated, ranging from 5 to 7 different species per herd (Table 2). The predominating CNS species in milk with ≥300 cfu/mL were S. chromogenes (30.6% of all isolates), S. haemolyticus (27.6%), S. epidermidis (11.9%), and S. simulans (11.2%). Staphylococcus chromogenes (isolated from 12 different cows) and S. haemolyticus (13 cows) were the only species isolated in all herds. Staphylococcus epidermidis was isolated in 4 herds (9 cows). Remarkably, S. epidermidis was isolated at least once from 6 different cows in herd E. Staphylococcus simulans was isolated in 2 herds, but originated mostly from 1 persistently infected cow in herd E. The more rarely isolated CNS species were restricted to one or a few herds.

CNS IMI

In the course of the study, 84 individual CNS IMI in 18 quarters of 14 cohort cows (range 0 to 4 cows per herd) were detected (Table 3). They were caused by S. chromogenes (n = 35 IMI), S. haemolyticus (n = 29), S. simulans (n = 14), and S. epidermidis (n = 6) (Table 3). In total, 17 cases of persistent IMI in 16 quarters were detected for which the same causative CNS AFLP-type was repeatedly isolated, and 4 IMI cases in 3 quarters were transient in nature. The mean duration of persistent cases was 149.4 d (range 63.0 to 329.8 d). The longest persistent CNS infection (329.8 d) was caused by an S. chromogenes AFLP type that was isolated 11 consecutive times from the same quarter [Figure 1, herd E, cow 6, left hind quarter (LH)]. Another cow was persistently infected with S. chromogenes in a quarter in first lactation (134.2 d) and became infected with another S. chromogenes AFLP type (102.5 d) in its second lactation in the same quarter (Figure 1, herd A, cow 1, LH). Five other quarters of 5

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Table 2. Species distribution of CNS isolates from quarter milk samples taken each month (n = 13) from 10 randomly selected cohort cows per herd in 6 herds

<table>
<thead>
<tr>
<th>CNS species</th>
<th>Total no. of isolates (%)</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>F</th>
</tr>
</thead>
<tbody>
<tr>
<td>Staphylococcus chromogenes</td>
<td>41 (30.6)</td>
<td>9 (2)</td>
<td>6 (4)</td>
<td>1</td>
<td>7 (1)</td>
<td>12 (2)</td>
<td>6 (2)</td>
</tr>
<tr>
<td>S. haemolyticus</td>
<td>37 (27.6)</td>
<td>2 (2)</td>
<td>6 (1)</td>
<td>3 (2)</td>
<td>20 (3)</td>
<td>2 (2)</td>
<td>4 (3)</td>
</tr>
<tr>
<td>S. epidermidis</td>
<td>16 (11.9)</td>
<td>1</td>
<td>5 (1)</td>
<td>1</td>
<td>9 (6)</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>S. simulans</td>
<td>15 (11.2)</td>
<td>5 (3.7)</td>
<td>1</td>
<td>1</td>
<td>14 (1)</td>
<td>3 (2)</td>
<td>4 (2)</td>
</tr>
<tr>
<td>S. hominis</td>
<td>5 (3.7)</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S. cohnii</td>
<td>4 (3.0)</td>
<td>3 (2)</td>
<td>1</td>
<td>2 (2)</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>S. saprophyticus</td>
<td>4 (3.0)</td>
<td>3 (2)</td>
<td>1</td>
<td>2 (2)</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S. auricularis</td>
<td>3 (2.2)</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S. warneri</td>
<td>3 (2.2)</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S. devriesei</td>
<td>2 (1.5)</td>
<td>1</td>
<td></td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S. equorum</td>
<td>2 (1.5)</td>
<td>1</td>
<td></td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Macrococcus caseolyticus3</td>
<td>1 (0.7)</td>
<td></td>
<td>1</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S. sciuri</td>
<td>1 (0.7)</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>134 (100.0)</td>
<td>18</td>
<td>19</td>
<td>8</td>
<td>31</td>
<td>39</td>
<td>19</td>
</tr>
</tbody>
</table>

1Total number of isolates per species originating from quarter milk samples with ≥300 cfu/mL, and respective proportion (%) of all isolates (n = 134).
2Number of cows per herd from which the isolates originated is given between brackets when >1 isolate was found.
3Staphylococcus caseolyticus has been reclassified as M. caseolyticus and is sporadically isolated from cattle.
cows with persistent \emph{S. chromogenes} IMI were detected with a duration ranging from 63.2 to 290.5 d. \emph{Staphylococcus haemolyticus} was the only CNS species found to cause both transient and persistent infections. In herd D, a heifer was infected with \emph{S. haemolyticus} in 3 quarters [Figure 1, herd D, cow 1, left front (LF), right front (RF), and right hind quarter (RH)], experiencing 2 persistent (74.4 d in LF, 100.4 d in RH) and 3 transient IMI (1 RH, 2 RF). The \emph{S. haemolyticus} isolates from the LF quarter showed an AFLP type diverging from the \emph{S. haemolyticus} library strains and the isolates \emph{S. haemolyticus} from the LF quarter showed an AFLP type diverging isolates sient IMI (1 RH, 2 RF). The \emph{S. haemolyticus} infection was observed in a multiparous cow of herd F and 1 persistent infection in a heifer of herd B that lasted for 222.1 d (data not shown). The 2 quarters with persistent \emph{S. simulans} IMI belonged to the same heifer and were both infected for 214.8 d with the same AFLP type (Figure 1, herd E, cow 3, LH). Persistent \emph{S. epidermidis} IMI were found twice, 1 in herd B lasting for 117.6 d (Figure 1, herd B, cow 9, RF) and 1 in herd E for 63.0 d (Figure 1, herd E, cow 5, LF).

**Clinical Cases**

During the study period, 26 quarters with clinical signs were detected from cohort cows and 83 quarters from noncohort herdmates. The average IRCM (all cases) in the herds was 0.281 quarter cases per 365 cow-days at risk (ranging from 0.149 to 0.485 between herds, Table 1). In total, 7.5% of clinical quarter milk samples (7 out of 94 sampled quarters) were positive for CNS growth. Three samples yielded CNS exclusively, and the other 4 showed mixed growth with esculin-positive cocci (n = 3) or \emph{Corynebacterium bovis} (n = 1). Species distribution of CNS isolates from clinical samples was 2 \emph{S. chromogenes}, 1 \emph{S. epidermidis}, 1 \emph{S. equorum}, and 3 unknown (no isolate was preserved). Both \emph{S. chromogenes} isolates came from clinical cases yielding CNS exclusively.

**CNS in the Cows’ Environment**

Coagulase-negative staphylococci were isolated from 75 of 78 (96.2%) floor samples, from 77 of 78 (98.7%) air samples, from 65 of 73 (89.0%) used bedding samples, and from 50 of 73 (68.5%) bedding stock samples. After PCR and RAPD analyses, 637 CNS isolates were preserved for further identification. In total, 612 isolates (96.1%) were identified to species level and 25 isolates (3.9%) remained unidentified. Identification results and species distribution over herds are given in Table 4. Most environmental isolates were readily identifiable by numerical comparison to the AFLP library; namely, 531 out of 637 isolates (83.4%). Another 81 isolates could be identified by visual comparison of AFLP fingerprints in a cluster (n = 40) or by a combination of \emph{rpoB} gene sequencing and AFLP clustering (n = 41). The CNS species predominant in the environment over all herds were \emph{S. equorum} (19.0% of the isolates), \emph{S. sciuri} (17.9%), and \emph{S. haemolyticus} (16.6%). The species \emph{S. cohnii} (5.7%), \emph{S. simulans} (5.3%), \emph{S. xylosus} (3.1%), \emph{S. devriesii} (2.8%), and \emph{S. arlettae} (2.5%) were also isolated in the environment of each herd, but less frequently. Some notable herd-to-herd differences were observed in relation to species distribution. Several CNS species were chiefly isolated in 1 or 2 herds. \emph{Staphylococcus fleurettii} was isolated predominantly in the environment of herd E, \emph{S. cohnii} in herds A and F, \emph{S. simulans} in herds C and E, and \emph{S. saprophyticus} in herd A. In herd E, \emph{S. sciuri} (3.7% of environmental CNS isolates) was less common compared with the other herds (11.0 to 28.3%). \emph{Staphylococcus haemolyti-

### Table 3. Number of detected IMI caused by CNS species in monthly sampled cows in 6 herds (10 cows/herd)

<table>
<thead>
<tr>
<th>Causative CNS species</th>
<th>Positive IMI status</th>
<th>No. of IMI cases</th>
<th>Mean duration (range) of persistent IMI cases (d)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Transient</td>
<td>Persistent</td>
<td>Cows</td>
</tr>
<tr>
<td>\emph{Staphylococcus chromogenes}</td>
<td>4</td>
<td>35</td>
<td>8</td>
</tr>
<tr>
<td>\emph{S. haemolyticus}</td>
<td>25</td>
<td>25</td>
<td>9</td>
</tr>
<tr>
<td>\emph{S. simulans}</td>
<td>14</td>
<td>14</td>
<td>2</td>
</tr>
<tr>
<td>\emph{S. epidermidis}</td>
<td>6</td>
<td>6</td>
<td>2</td>
</tr>
<tr>
<td>Total</td>
<td>40</td>
<td>80</td>
<td>21</td>
</tr>
</tbody>
</table>

1Number of quarter milk samples with positive CNS IMI status, assigned when at least 2 out of 3 consecutive quarter milk samples were culture-positive for the same CNS species (≥300 cfu/mL), a single positive IMI status was classified as a transient IMI, and recurrent positive IMI status was considered as persistent IMI.

2Cases of transient (n = 4) or persistent (n = 17) IMI established by the same CNS amplified fragment length polymorphism (AFLP)-type in a single quarter.
cus was less common in herds C (6.9%) and E (8.3%) compared with the other herds (15.7 to 26.4%).

**Distribution of CNS Species over the Different Niches**

When distribution of CNS species over the different niches was compared, certain CNS species tended to favor particular niches (Table 5). *Staphylococcus chromogenes* and *S. epidermidis* were predominantly isolated from milk (78.8 and 57.1% of all isolates, respectively) and less often from environmental samples. *Staphylococcus haemolyticus* and *S. simulans* were regularly isolated both from milk and the environment (74.1 and 69.4% isolates from environment, respectively). Other CNS species were sporadically isolated from milk, but not causing IMI according to our definition. Common species with a mainly environmental origin (90.0 to 100.0% of all isolates) were *S. equorum*, *S. sciuri*, *S. fleurettii*, *S. cohnii*, *S. devriesei*, *S. xylosus*, *S. arlettae*, and *S. succinus*. The CNS species in the environment were distributed differently over the sampled locations. *Staphylococcus haemolyticus* and *S. equorum* were isolated mostly from stall air (40.6 and 49.6% of all isolates, respectively), whereas *S. sciuri* and *S. simulans* were isolated on a regular basis from slatted floors and sawdust of the cubicles (72.2 and 67.3%, respectively). *Staphylococcus xylosus* and *S. succinus* were isolated mostly from sawdust stock (55.0 and 80.0%, respectively).

**DISCUSSION**

In this longitudinal field study, distribution of CNS species isolated from quarter milk samples and the cows’ environment was determined in 6 herds. The
CNS isolates were genotyped and identified to species level by means of AFLP, a well-validated and reproducible whole-genome based method (Piessens et al., 2010). This is in contrast to the majority of previous field studies on CNS in which phenotypic methods were generally used for species differentiation (Matos et al., 1991; Thorberg et al., 2009). In several studies, however, it has been demonstrated that phenotypic methods are insufficient for identification of CNS isolates from the bovine (Bes et al., 2000; Capurro et al., 2009; Sampimon et al., 2009). Currently, genotyping is the preferred method to gain accurate information on CNS at the species level (Zadoks and Watts, 2009). In the past, CNS from cows’ environment have only been studied by using phenotypic methods (Matos et al., 1991) and as far as we know, their genotypes have never been compared with those of CNS causing IMI. Amplified fragment length polymorphism genotyping enables not only species identification, but also genotypic comparison of CNS isolates and tracking of the spread of specific CNS genotypes, which provides valuable information on epidemiology of these udder pathogens. Because different control practices are needed when either contagious or environmental pathogens cause a significant proportion of all IMI in a herd, determining the epidemiology of individual CNS species is useful for the dairy industry to prevent infections with particular more pathogenic CNS species.

According to our postulated IMI definition, only the CNS species *S. chromogenes*, *S. haemolyticus*, *S. simulans*, and *S. epidermidis* caused IMI in the studied cows. This is in line with other studies, where these species have also been isolated from IMI (Aarestrup and Jensen, 1997; Chaffer et al., 1999; Taponen et al., 2007), although other CNS species; for example, *S. xylosus* (Thorberg et al., 2009) and *S. hyicus* (Gillespie et al., 2009), have also been reported. Most CNS IMI detected in our study were persistent in nature, although this was largely due to the rather stringent definition for CNS IMI based on 3 consecutive monthly samples. No standard criteria exist for diagnosis of CNS IMI in the bovine udder, although attempts to define a general IMI definition have recently been published (Dohoo et al., 2011). This recent consensus is, however, based on weekly samplings, whereas we used monthly data. Because of the relatively large time interval between samplings and the drawbacks of bacterial culturing, the real number of IMI in our study is most likely underestimated, especially for IMI of short duration. Nevertheless, the effect of these transient

<table>
<thead>
<tr>
<th>CNS species</th>
<th>Total no. of isolates (%)</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>F</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Staphylococcus equorum</em></td>
<td>121 (19.0)</td>
<td>18</td>
<td>21</td>
<td>29</td>
<td>12</td>
<td>20</td>
<td>21</td>
</tr>
<tr>
<td><em>S. sciuri</em></td>
<td>114 (17.9)</td>
<td>28</td>
<td>10</td>
<td>28</td>
<td>32</td>
<td>4</td>
<td>12</td>
</tr>
<tr>
<td><em>S. haemolyticus</em></td>
<td>106 (16.6)</td>
<td>21</td>
<td>24</td>
<td>7</td>
<td>28</td>
<td>9</td>
<td>17</td>
</tr>
<tr>
<td><em>S. fleurettii</em></td>
<td>47 (7.4)</td>
<td>5</td>
<td>10</td>
<td>5</td>
<td>10</td>
<td>1</td>
<td>17</td>
</tr>
<tr>
<td><em>S. cohnii</em></td>
<td>36 (5.7)</td>
<td>12</td>
<td>2</td>
<td>1</td>
<td>6</td>
<td>3</td>
<td>12</td>
</tr>
<tr>
<td><em>S. simulans</em></td>
<td>34 (5.3)</td>
<td>3</td>
<td>4</td>
<td>9</td>
<td>4</td>
<td>10</td>
<td>4</td>
</tr>
<tr>
<td><em>S. xylosus</em></td>
<td>20 (3.1)</td>
<td>3</td>
<td>5</td>
<td>1</td>
<td>4</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td><em>S. devriesii</em></td>
<td>18 (2.8)</td>
<td>5</td>
<td>1</td>
<td>4</td>
<td>2</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td><em>S. arletiae</em></td>
<td>16 (2.5)</td>
<td>1</td>
<td>6</td>
<td>3</td>
<td>4</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><em>S. saprophyticus</em></td>
<td>15 (2.4)</td>
<td>12</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td><em>S. epidermidis</em></td>
<td>12 (1.9)</td>
<td>4</td>
<td>4</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td><em>S. chromogenes</em></td>
<td>11 (1.7)</td>
<td>1</td>
<td>3</td>
<td>1</td>
<td>3</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td><em>S. hominis</em></td>
<td>11 (1.7)</td>
<td>4</td>
<td>2</td>
<td>3</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><em>S. succinus</em></td>
<td>10 (1.6)</td>
<td>3</td>
<td>2</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td><em>S. lentus</em></td>
<td>8 (1.3)</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td><em>S. auricularis</em></td>
<td>7 (1.1)</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td><em>S. capitis</em></td>
<td>7 (1.1)</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td><em>S. vitulinus/S. pulvereri</em></td>
<td>6 (0.9)</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>4</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td><em>S. gallinarum</em></td>
<td>5 (0.8)</td>
<td>1</td>
<td>1</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td><em>S. warneri</em></td>
<td>4 (0.6)</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><em>Macrococcus caseolyticus</em></td>
<td>2 (0.3)</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><em>S. hyicus</em></td>
<td>1 (0.2)</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><em>S. nepalensis</em></td>
<td>1 (0.2)</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>25 (3.9)</td>
<td>1</td>
<td>2</td>
<td>4</td>
<td>8</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>Total</td>
<td>637 (100.0)</td>
<td>114</td>
<td>91</td>
<td>102</td>
<td>113</td>
<td>109</td>
<td>108</td>
</tr>
</tbody>
</table>

1Total number of isolates per species and respective proportion (%) of all isolates (n = 637).
2*S. vitulinus* and *S. pulvereri* are synonyms for the same species.
3*Staphylococcus caseolyticus* has been reclassified as *M. caseolyticus* and is sporadically isolated from cattle.
DISTRIBUTION OF COAGULASE-NEGATIVE STAPHYLOCOCCI

CNS IMI on general udder health is probably minor, whereas the long duration of undetected subclinical infections increases the effect on SCC and the possibility of spread of the causative bacteria. Although all CNS species isolated from milk in our study were evaluated using the same definition, striking differences were found among CNS species and only 4 were considered of particular interest. Persistence of S. chromogenes, S. haemolyticus, S. epidermidis, and S. simulans has been diagnosed in previous studies by repeated isolation of the same species from the same quarter (Aarestrup and Jensen, 1997; Chaffer et al., 1999; Thorberg et al., 2009). Our observations further confirm that particular CNS strains within these species, as evidenced by the repeated isolation of the same AFLP type, are able to survive and persist in the udder.

Comparing the prevalence of different CNS species between studies should be done with caution, as differences often exist in the identification methods used, definitions for CNS IMI, or sampling schedule. However, it can be assumed that distribution of CNS species infecting the udder varies between herds (this study; Gillespie et al., 2009; Thorberg et al., 2009). Part of this variation could be explained by factors such as type of germicide used for teat dipping (Hogan et al., 1987) or age of the cows (Matthews et al., 1992; Taponen et al., 2006). Between herds, marked differences were also seen in CNS species distribution in the environment, although farm and management characteristics of the herds included in our study were similar. This also indicates that currently unknown herd-level factors or environmental conditions determine the establishment of staphylococcal species in a dairy herd. In 2 studies in which extramammary CNS isolates have also been differentiated, it has been demonstrated that type of housing (White et al., 1989) and type of bedding (Matos et al., 1991) influence the distribution of CNS species found on body sites of heifers and in bedding samples, respectively. Although no attempt was made to quantify individual CNS species in the environment, we conclude that each dairy farm probably harbors its own CNS microbiota. To explain this remarkable variation in occurrence of particular CNS species, further study on the influence of different herd and environmental factors on CNS prevalence and distribution is required.

Table 5. Distribution of the individual CNS species isolated from monthly sampled (n = 13) niches on 6 dairy farms (high occurrence of individual species in a particular niche is indicated in bold)

<table>
<thead>
<tr>
<th>CNS species</th>
<th>No. (%) of isolates per niche&lt;sup&gt;1&lt;/sup&gt;</th>
<th>Total no. (%)&lt;sup&gt;2&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Milk&lt;sup&gt;3&lt;/sup&gt;</td>
<td>Air</td>
</tr>
<tr>
<td>Staphylococcus haemolyticus</td>
<td>37 (25.9)</td>
<td>58 (40.6)</td>
</tr>
<tr>
<td>S. epidermidis</td>
<td>2 (1.6)</td>
<td>61 (49.6)</td>
</tr>
<tr>
<td>S. simulans</td>
<td>41 (78.8)</td>
<td>3 (5.8)</td>
</tr>
<tr>
<td>S. chromogenes</td>
<td>15 (30.6)</td>
<td>1 (2.0)</td>
</tr>
<tr>
<td>S. fleurettii</td>
<td>8 (17.0)</td>
<td>17 (39.5)</td>
</tr>
<tr>
<td>S. cohnii</td>
<td>4 (10.0)</td>
<td>11 (27.5)</td>
</tr>
<tr>
<td>S. epidermidis</td>
<td>16 (57.1)</td>
<td>1 (3.6)</td>
</tr>
<tr>
<td>S. devriesei</td>
<td>2 (10.0)</td>
<td>7 (35.0)</td>
</tr>
<tr>
<td>S. zylosus</td>
<td>5 (25.0)</td>
<td>5 (25.0)</td>
</tr>
<tr>
<td>S. saprophyticus</td>
<td>4 (21.1)</td>
<td>3 (15.8)</td>
</tr>
<tr>
<td>S. arlettae</td>
<td>6 (50.0)</td>
<td>1 (6.3)</td>
</tr>
<tr>
<td>S. hominis</td>
<td>5 (31.3)</td>
<td>3 (18.8)</td>
</tr>
<tr>
<td>S. auricularis</td>
<td>3 (30.0)</td>
<td>7 (70.0)</td>
</tr>
<tr>
<td>S. succinu</td>
<td>2 (20.0)</td>
<td></td>
</tr>
<tr>
<td>S. lentus</td>
<td>1 (12.5)</td>
<td>3 (37.5)</td>
</tr>
<tr>
<td>S. capitis</td>
<td>6 (85.7)</td>
<td></td>
</tr>
<tr>
<td>S. warneri</td>
<td>3 (42.9)</td>
<td>1 (14.3)</td>
</tr>
<tr>
<td>S. vitulinus/S. pulvereri</td>
<td>4 (66.7)</td>
<td>1 (16.7)</td>
</tr>
<tr>
<td>S. gallinarum</td>
<td>3 (60.0)</td>
<td></td>
</tr>
<tr>
<td>Macroccocus caseolyticus</td>
<td>1 (33.3)</td>
<td>1 (33.3)</td>
</tr>
<tr>
<td>S. hyicus</td>
<td>1 (100.0)</td>
<td></td>
</tr>
<tr>
<td>S. nepalensis</td>
<td>1 (100.0)</td>
<td></td>
</tr>
<tr>
<td>Unknown</td>
<td>10 (40.0)</td>
<td>12 (48.0)</td>
</tr>
<tr>
<td>Total</td>
<td>134 (17.4)</td>
<td>209 (27.1)</td>
</tr>
</tbody>
</table>

<sup>1</sup>Distribution of isolates of individual CNS species over the sampled niches given in numbers and proportion (%) of total number of isolates within the respective species.

<sup>2</sup>Total number of isolates per species and respective proportion (%) of all isolates (n = 771).

<sup>3</sup>Isolates from quarter milk samples of monthly sampled cows in the 6 herds (10 cows/herd) yielding ≥300 cfu of CNS/mL.
The findings of our study suggest that CNS species are abundantly present in the studied type of dairy herds, but that their primary reservoirs differ according to species. For *S. chromogenes* and *S. epidermidis*, the udder was found as a main reservoir. These 2 species were rarely isolated from the environment and IMI probably originated from other sources; for example, the cow’s skin, milkers’ hands, or other cows. In our study, *S. chromogenes* caused the most and longest persistent subclinical IMI and was found to be the most likely cause of 2 clinical cases. As *S. chromogenes* is a commonly isolated mastitis pathogen (Aarestrup and Jensen, 1997; Gillespie et al., 2009; Sawant et al., 2009) able to induce an elevation in SCC as high as that seen for infections with the major pathogen *S. aureus* (Supré et al., 2011), its role in udder health might not be as minor as previously thought. *Staphylococcus chromogenes* has been isolated from udder skin, teat apices, and other body sites in previous studies (White et al., 1989; De Vliegher et al., 2003; Taponen et al., 2008). According to Taponen et al. (2008), this species is a typical skin opportunist. Unlike *S. chromogenes*, *S. epidermidis* is uncommon in normal bovine skin microbiota, but it is one of the most prevalent staphylococcal species on human skin and may be transmitted from milkers to cows (Thorberg et al., 2006).

Two CNS species (*S. haemolyticus* and *S. simulans*) were found that caused IMI and also survived well in the environment. In particular, *S. haemolyticus* caused a large proportion of all IMI and appeared to have a considerable environmental reservoir. In several studies, this species was not or was only rarely isolated from IMI (Matthews et al., 1992; Gillespie et al., 2009), whereas in others it has been found frequently (Chaffer et al., 1999; Thorberg et al., 2009). The apparent differences in *S. haemolyticus* prevalence could be due to either true variation in its occurrence or the difficulty in identifying this species. Among our *S. haemolyticus* isolates, 2 very distinct AFLP types were differentiated, 1 of which could only be identified by *rpoB* gene sequencing. Possibly, the latter type could lead to identification problems when using other methods. Within herds, similar AFLP types of *S. haemolyticus* were observed among isolates originating from environment and IMI, strongly indicating potential environmental sources for IMI with this species. *Staphylococcus haemolyticus* was also commonly isolated from the teat apices of the cows under study (G. Braem, B. Verbist, V. Piessens, S. De Vliegher, and L. De Vuyst, unpublished results), indicating its adaptability to various niches in the dairy herd. *Staphylococcus simulans*, on the other hand, is uncommon on cow skin and the source of IMI with this species is still unclear; *S. simulans* tends to cause more severe mastitis and predominates in studies conducted on clinical mastitis (Myllys, 1995; Waage et al., 1999).

In this study, environmental *S. simulans* isolates mostly originated from slatted floors and used bedding, which might imply contamination of the environment by infected cows. In a study by Aarestrup et al. (1999), a variety of ribotypes of *S. simulans* have been recovered from IMI in the same herds and within different quarters of the same cows. The observed genetic diversity of infectious *S. simulans* ribotypes in that study might imply multiple environmental sources, although this has not yet been demonstrated.

The origin of all other CNS species isolated in our study was primarily environmental. In addition, 25 CNS were isolated from the environment that could not be identified by our AFLP method. These isolates could represent previously unknown CNS species; however, they were not studied further because of their overall low prevalence. In 2 other studies using molecular techniques for CNS differentiation, unidentifiable genetic profiles were also generated for several presumptive CNS isolates with transfer DNA-PCR (Supré et al., 2009) and AFLP (Taponen et al., 2006). Further study of these atypical isolates led to the characterization and description of 2 new CNS species, *S. devriesei* (Supré et al., 2010) and *S. agnetis* (Taponen et al., 2011). Surprisingly, some species that were mainly environmental in our study have been found to be significant causes of IMI in others; for example, *S. xylosus* (Thorberg et al., 2009), *S. sciuri* (Davidson et al., 1992), and *S. cohnii* (Supré et al., 2011). It can be speculated that when infection pressure is high or when immunity of cows is compromised, these CNS species may act as environmental opportunistic pathogens. In addition, it could be that some strains within these species are better adapted to cause infection. Despite prevention measures such as teat disinfection and dry cow therapy, prevalence of CNS infections in some herds is remarkably high for unknown reasons (Piepers et al., 2007; Schukken et al., 2009). In the past, only a few studies have been set up to identify possible reservoirs of CNS by comparing strains originating from mastitis with strains originating from other sources (Thorberg et al., 2006; Taponen et al., 2008). Knowledge on primary reservoirs of particular CNS species in herds (e.g., *S. xylosus* in sawdust stock) could be useful in prevention of CNS infections contracted from the environment. Our study was an attempt to identify potential environmental sources of CNS IMI. However, before we can demonstrate possible transmission routes of individual CNS species or identify reservoirs of strains causing...
IMI, clonal diversity within CNS species observed in the AFLP fingerprints should be further confirmed by (an)other molecular subtyping technique(s).

CONCLUSIONS

Distribution of CNS species in the milk and environment of cows differed among herds. Epidemiology of CNS infecting the udder varied among species. *Staphylococcus chromogenes* and *S. epidermidis* seemed to act more as host-adapted pathogens specialized in surviving in the udder, and the environment was not found as a likely source of IMI. *Staphylococcus haemolyticus* and *S. simulans*, on the other hand, had a considerable reservoir in the environment and could act as environmental opportunists.

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