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8	INTERPRETIVE SUMMARY
9	TYPEABILITY OF MALDI-TOF FOR NON-AUREUS STAPHYLOCOCCI
10	IDENTIFICATION
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12	MAHMMOD
13	MALDI-TOF has never been investigated for NAS from non-milk samples. We evaluated the
14	typeability of MALDI-TOF for identification of NAS associated with bovine IMI and teat apex
15	colonization. Proportion of NAS isolates correctly identified from milk (91%) was higher than
16	proportion of isolates correctly identified from teat skin (68%). In total, 93% of isolates were
17	successfully identified as NAS, while the remaining (7%) were shown to be other bacterial species
18	using MALDI-TOF. MALDI-TOF is efficient in NAS identification from milk but it may not always
19	be acceptable for routine identification of NAS from non-milk samples. Nucleic-acid based tools is
20	vital for accurate species identification of some NAS species.

22	TYPEABILITY OF MALDI-TOF FOR NON-AUREUS STAPHYLOCOCCI
23	IDENTIFICATION
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25	Typeability of MALDI-TOF Assay for Identification of Non-Aureus Staphylococci associated
26	with Bovine Intramammary Infections and Teat Apex Colonization
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47 ABSTRACT

Matrix-assisted laser desorption/ionization time of flight (MALDI-TOF), a culture-dependent assay, 48 has recently been implemented for routine identification of non-aureus Staphylococcus (NAS) 49 50 species from milk, but the assay has never been investigated for NAS from non-milk or environmental samples. The objective of this study was to evaluate the typeability of MALDI-TOF 51 52 assay for the identification and differentiation of bovine-associated NAS species on aseptically 53 collected quarter-milk and teat skin in dairy herds. In eight herds, 14 to 20 cows with elevated somatic cell count were randomly selected for teat skin swabs and foremilk samples from right hind 54 55 and left front quarters. Teat skin swabs and milk samples were collected aseptically for preliminary identification using bacterial culture on chromogenic and calf blood agars. Colonies from milk and 56 57 teat skin samples with suspicion of having NAS were identified to species-level by MALDI-TOF 58 assay. Out of 511 isolates from 284 quarters (142 cows), 78% (n = 399) were identified by MALDI-59 TOF. The percentage of correctly identified NAS from milk (91% 105/115) using MALDI-TOF was higher than the percentage from teat skin (68%, 268/396). Out of the identified isolates, 93% (n =60 61 373) were successfully identified as NAS, while the remaining 26 (7%) were shown to be other 62 bacterial species. Out of 26 non-NAS isolates, one originated from milk (Corynebacterium stationis), while 25 originated from teat skin representing Aerococcus viridans (n = 7), Bacillus pumilis (n = 7) 63 13), Enterococcus saccharolyticus (n = 1), Clostridum septicum (n = 1), Corynebacterium stationis 64 (n = 2), and Corynebacterium casei (n = 1). MALDI-TOF identified 85% (98/115) and 62% 65 66 (245/396) of the isolates in the first test. Isolates that were not identified to species-level at first test were subjected to a second test, and here 47% (8/17) and 32% (48/151) from milk and teat skin, 67 respectively, were identified. After two rounds of MALDI-TOF, 22% (n = 112) of the isolates were 68 69 not identified representing 103 from teat skin and nine from milk. Eighteen isolates without identification by MALDI-TOF were successfully identified to species-level using sequencing, where 70 71 16 were correctly identified as NAS, while the other two were *Corynebacterium stationis*.

In conclusion, MALDI-TOF is a reliable assay for identification and typeability of NAS species from aseptically collected quarter milk samples. The assay may be used for identification of NAS species from teat skin swabs. However, confirmation using nucleic-acid based tools is vital for accurate species identification of some species and strains.

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77 KEYWORDS: non-aureus staphylococci; bovine mastitis; teat skin colonization; phenotypic
78 identification

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INTRODUCTION

Non-aureus Staphylococci (NAS) are a heterogeneous group of bacterial species (Schukken et al., 80 2009), which regarded as a common cause of IMI in dairy herds (Zadoks and Watts, 2009). 81 82 Moreover, NAS are abundantly colonizing the teat skin, teat apex and teat canal and hence, many studies have shown that teat colonization with NAS could have a significant role in initiation or 83 development of IMI with NAS in dairy cows (Leroy et al., 2015; De Visscher et al., 2016). Recent 84 85 studies documented that some species are more important than others in relation to udder health (Supré et al., 2011; De Visscher et al., 2016). Furthermore, De Visscher et al. (2014) provided 86 87 evidence that the group of NAS species is comprised of environmental, opportunistic and hostadapted species, which differ in ecology. Additionally, the authors concluded that some of the 88 extramammary niches, such as the teat apex, might act as infection sources for IMI-causing NAS. 89 90 Except for Staphylococcus xylosus, an association was observed between teat canal colonization and 91 IMI by all NAS species, in which the majority of IMI were preceded by teat canal colonization (Quirk et al., 2012). 92

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NAS are known to be a common teat apex colonizer (Braem et al., 2013; Falentin et al., 2016), and 94 95 are among the mastitis-causing bacteria most likely to enter the mammary gland through the teat orifice (Fox and Norell, 1994) resulting in the establishment of IMI (Pyörälä and Taponen, 2009; 96 Piepers et al., 2010). Braem et al. (2012) identified staphylococci among the bacterial genera with 97 98 the highest percentage (31%) of colonized teat apices and they were detected with equal prevalence from teat apices of non-infected, subclinically infected, and clinically infected quarters. Therefore, it 99 is crucial to identify and differentiate the NAS species colonizing teat skin or inhabiting milk causing 100 101 IMI to understand their epidemiology and to evaluate the clinical relevance and feasibility of speciesspecific infection control measures (Zadoks and Watts, 2009). Furthermore, it is important for the 102 routine microbiological laboratories, because rapid and correct identification of mastitis causing 103

pathogens will influence the choice of antibiotic before the final determination of antibiotic
resistance of the isolate (Nagy et al., 2014). For many years, the species identification of NAS relied
on phenotypic characteristics, which is difficult, time consuming, laborious, and often inaccurate
(Watts et al., 1991; Vanderhaeghen et al., 2014). Although biochemical assays such as API systems
are widely used for identification of NAS, the accuracy and speed is not optimal (Taponen et al.,
2006; Capurro et al., 2009; Sampimon et al., 2009; Park et al., 2011).

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Matrix-assisted laser desorption/ionization time of flight mass spectrometry (MALDI-TOF) is a 111 112 rapid method, which is able to identify a great variety of the isolated bacteria based on the composition of conserved ribosomal proteins (Kliem and Sauer, 2012). This technique is based on 113 114 the acquisition of protein (ribosomal proteins) "fingerprints" directly from intact microorganisms, 115 since such profiles vary considerably among microorganisms (Singhal et al., 2015). The assay 116 provides a new diagnostic platform, which overcomes the limitations of traditional diagnostics for NAS being time consuming and laborious, or the need of sugar fermentation or test kits (Watts et al., 117 118 1991; Capurro et al., 2009; Vanderhaeghen et al., 2014; Taponen et al., 2016). The technique is increasingly used in human medicine and, recently, it has been expanded as a routine diagnostic tool 119 in veterinary medicine (Randall et al., 2015; Pizauro et al., 2017). In the recent decade, research 120 studies showed that MALDI-TOF is a powerful and reliable diagnostic tool for identification and 121 122 discrimination of mastitis causing pathogens including NAS from bovine mastitis samples (Tomazi 123 et al., 2014; Gonçalves et al., 2014; Cameron et al., 2017a, b), and spiked milk samples (Barreiro et al., 2017). MALDI-TOF assay was validated against other routine diagnostics for NAS such as Vitek 124 2 compact system (Elbehiry et al., 2016) and PCR (Pizauro et al., 2017) where it showed a better 125 126 performance in identification and discrimination of NAS species from bovine mastitis. To the best of our knowledge there is no available literature describing the performance of MALDI-TOF assay for 127 128 identification of NAS from non-milk cow samples or environmental samples in dairy herds. The objective of this study was to evaluate the typeability of MALDI-TOF assay for the identification
and differentiation of bovine-associated NAS species on quarter level from aseptically collected milk
(IMI) and teat skin (teat apex colonization) habitats in dairy herds.

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MATERIAL AND METHODS

134 Study population

Eight dairy herds with Danish Holstein cows were selected for participating in a project on 135 Streptococcus agalactiae and Staphylococcus aureus IMI. To be eligible for inclusion in the present 136 study, herds had to have automatic milking systems (AMS) with ≥ 3 milking robots and bulk tank 137 milk PCR cycle threshold value ≤ 32 for *Streptococcus agalactiae*. About 30 to 40 lactating dairy 138 139 cows were selected randomly from each herd on the basis of the criteria of having no clinical 140 mastitis, SCC \geq 200,000 cells/mL at the preceding milk recording, and not subjected to antibiotic therapy during the four weeks prior to sample collection. From each cow with an odd laboratory 141 142 running number, teat skin swab and aseptic milk samples were taken from right hind and left front 143 quarters (Mahmmod et al., 2018).

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145 Sampling Procedures

Each herd was visited once to collect teat swab samples and aseptically collected quarter foremilk samples for bacterial culture. The farmers were asked to separate the selected cows for sampling. Cows were fixed in head lockers or tied. Teat swab samples were collected according to the modified wet-dry method (Paduch et al., 2013). Briefly, the teat skin was sampled after cleaning with dry tissue paper. The first rayon swab (DaklaPack, Glostrup, Denmark) was moistened with ¹/₄ Ringer's solution (Merck, Darmstadt, Germany) and rotated 360° around the teat about one cm from the teat canal orifice. The same procedure was carried out with a dry swab (second). Immediately after 153 sampling, the tips of both swabs were transferred into one tube with 2 mL of sterile ¹/₄ Ringer's
154 solution.

Quarter milk samples were collected directly after harvesting the teat swab samples according to 155 156 National Mastitis Council (1999) guidelines. Briefly, the teat end was thoroughly disinfected with cotton swabs drenched with ethanol (70%). Individual quarter foremilk samples were then 157 aseptically collected in sterile screw-cap plastic tubes. New latex gloves were worn at each sampling 158 procedure and were changed after each cow. Tubes containing the teat swabs and aseptically 159 collected milk samples were stored at 5°C in ice boxes until the samples were delivered to the 160 161 microbiological laboratory within 24h. All study activities including farm visits, collection of 162 samples and laboratory examination were carried out during the period from February to May 2017.

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164 Laboratory Procedures

165 Bacteriological examination

Bacterial culture for milk samples was conducted in accordance with National Mastitis Council 166 recommendations (1999). After vortexing, 0.01 mL of the milk sample from each quarter was 167 streaked using disposable, calibrated bacterial loops on a calf blood agar plate and simultaneously 168 169 another 0.01 mL of the same sample was streaked on chromogenic agar plates selective for 170 staphylococci species (SASELECT, Bio-Rad, Marnes-la-Coquette, France). Bacterial culture of teat 171 swab samples was performed according to the procedures of Paduch et al. (2013). Briefly, the teat 172 swab sample was vortexed before removing the swab tips from the tubes. The agar plates were inoculated with 0.1 mL of a swab solution prepared with ¹/₄ Ringer's solution. The inoculum was 173 spread with a sterile Drigalski spatula onto the agar surface of a calf blood agar plate and 174 175 SASELECT media simultaneously for each quarter.

All the inoculated plates were marked by the laboratory running number and sequence of the quarter
and were incubated aerobically at 37°C for 48h and examined for growth of NAS colonies after 24

178 and 48h. Staphylococcus species were presumptively identified on the basis of the phenotypic characteristics of their colonies including shape (round, glossy) and color on the selective media. Up 179 to three different NAS species per sample, according to the colony color on the SASELECT media, 180 181 were considered for further identification at species-level. Cut-off \geq five CFU on the plate was regarded as an acceptable cut-off point for definition the positivity of NAS according to Thorberg et 182 al. (2009) from milk and teat skin samples based on bacterial culture. 183

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MALDI-TOF Assay and Species Identification

186 Isolates of NAS species that were identified based on bacterial culture were subcultured individually on new calf blood agar plates and incubated for 24 h at 37°C to be submitted freshly to MALDI-TOF 187 Bruker Biotyper software system (Microflex LT, Bruker Daltonics GmbH, Bremen, Germany) using 188 189 an Autoflex Speed for identification of NAS at species-level. The bacteria were prepared for mass spectrometry analysis according to a standard extraction protocol using formic acid (Bizzini et al., 190 2010), as recommended by the manufacturer. A sterile wooden applicator was used to pick the 191 material from a single bacterial colony followed by smearing a thin film of colony material onto a 192 MTP 384 ground steel target plate (Bruker Daltonics). Subsequently, each spot was overlaid with 1.0 193 194 μL of a saturated α-cyano-4-hydroxycinnamic acid in 50% acetonitrile, and 2.5% trifluoroacetic acid 195 solution and air-dried at room temperature. Mass spectra were obtained on an Autoflex Speed 196 calibrated with the recommended Bruker Escherichia coli Bacterial Test Standard for Mass 197 Spectrometry.

Spectra of the analyzed bacteria were obtained in positive linear mode, within the mass range of 2 to 198 199 20 kDa. Spectra were acquired using the flexControl 3.4 program, and subsequently, analyzed using 200 MALDI Bruker Real-Time Classification 3.1 software equipped with the BDAL database (Bruker 201 Daltonics) comprised with 6903 reference spectra combined with verified local spectra from National Veterinary Institute, Denmark. The software compares the 10 closest spectra in the database 202

203 to the entry and since expansion of the Bruker BDAL database is an ongoing process; local spectra were generated from isolates obtained by the Diagnostic laboratory at DTU and identified by 204 biochemical methods and/or 16S rDNA sequencing to meet the requirements for both animal and 205 206 human reference spectra in the database. Improvement of the local database was conducted and isolates submitted as reference strains for the extended database were implemented. These stains 207 were identified either by 16S rDNA sequencing or by biochemical methods as described by 208 Andresen et al. (2005). The 47 custom Main SPectra (MSPs) were created according to Nonnemann 209 et al. (2013) (Supplementary material 1). 210

The analysis for each isolate was run in triplicate. If the isolate was not identified to species-level in the first run, it was subcultured and resubmitted for MALDI-TOF to exclude the reason of "no identification" due to handling of the samples and to confirm that the non-identified isolates were either due to new species or to limitation of the database. After two rounds of MALDI-TOF, the unidentified isolates were considered as "no possible identification". A cut-off score ≥ 1.7 was regarded as a reliable threshold for the bacterial identification of NAS at species-level according to Cameron et al. (2017a, b).

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219 Identification by 16S rDNA sequencing

A set of NAS isolates without possible identification by MALDI-TOF were subjected to 16S rDNA 220 sequencing for identification. Those NAS isolates were selected on the basis of having a cut-off 221 222 threshold < 1.7 in the triplicate runs of the two rounds of MALDI-TOF. One bacterial colony was suspended in 1 mL PBS and centrifuged for 5 min at 10,000 g. The supernatant was removed and the 223 pellet resuspended in 100 µL water. The sample was boiled for 10 min and immediately placed on 224 225 ice. The lysate was centrifuged for 2 min at 20,000g. The concentration of the DNA was quantified on a NanoDrop ND-1000 Spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA) and 226 227 then was adjusted to $50 \text{ ng/}\mu\text{L}$.

228 The 16S rRNA gene was amplified by PCR in two separate products with following primers; 8f -804r (5'- AGA GTT TGA TC (AC) TGG CTC AG -3' and 5'- GTA TTA CCG CGG CTG CTG G-229 3') and (5'- CCA GCA GCC GCG GTA ATA C - 3' and 5'- GTT ACC TTG TTA CGA CTT CAC -230 231 3') The amplification program ran as follows: denaturation at 94°C for 6 min, 33 cycles 94°C for 45 sec, 56 °C for 45 sec, and 72°C for 90 sec, final extension 72°C for 10 min in a reaction volume of 50 232 µL. Amplified PCR products, 796 bp and 990 bp expected length, respectively, were verified by E-233 Gel (Invitrogen). Fourty µL PCR product was purified with MinElute PCR purification kit (Qiagen) 234 and eluted in 20 µL EB buffer. The amplified 16s rRNA gene was sequenced on an ABI 3130 235 236 Genetic analyzer using BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystem) according to manufacturer protocol. The resulting sequences were compared to the described 237 sequences by blasting with the GenBank database (http://www.ncbi.nlm.nih.gov) as previously 238 239 described (Strube et al. 2015).

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241 Statistical analysis

A logistic mixed regression model with herd and cow treated as random intercept was performed to investigate if teat apex colonization with a specific NAS species increased the odds of IMI with the given species in the corresponding quarter. Different models were therefore performed for each of the NAS IMI species recovered from the quarter milk samples. Statistical analysis was carried out in R version 3.3.3 (The R Foundation for Statistical Computing). Results were considered significant if $P \le 0.05$.

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RESULTS

Out of 150 eligible cows, 16 quarters were excluded for being dry or sampled twice. Out of 284
quarters of 142 cows, 228 quarters harbored at least one NAS species ≥ five CFU/0.01 mL identified
by culture. Table 1 shows the number and distribution of NAS isolates from milk and teat skin
samples submitted to MALDI-TOF assay. Out of total 511 culture isolates, 399 (78%) were

253 identified to species-level by MALDI-TOF, while the remaining 112 (22%) were not identified after 254 two submissions to MALDI-TOF and were labeled as "no possible identification". Out of the 399 isolates identified by MALDI-TOF, 373 (93%) were successfully identified as NAS, while the 255 256 remaining 26 isolates (7%) were diagnosed to be other bacterial species such as Aerococcus viridans, 257 Bacillus pumilus, Enterococcus saccharolyticus, Clostridum septicum and Corynebacterium casei, 258 Table 1. Out of the 26 isolates identified as other bacteria, one originated from milk and 25 from teat skin. Out of the 373 isolates, 28% were isolated from milk (n = 105) and 72% from teat skin (n =259 268). In the first round of MALDI-TOF, 85% (98/115) of the submitted isolates were identified from 260 261 milk, while the corresponding percentage was much lower for isolates from teat skin (62%, 245/396). In the second round, 47% (8/17) were identified from milk while it was 32% (48/151) for 262 teat skin isolates. The number of isolates without possible identification after two rounds of MALDI-263 264 TOF from teat skin (n = 103) was about 10 times higher than that the number of isolates from milk 265 (n = 9).

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267 In total, MALDI-TOF identified 16 different NAS species, 15 species from teat skin and 10 species from milk, Table 2. In milk samples, S. epidermidis (49.5%, 52/105) was the most frequently 268 269 isolated NAS species, while S. arlettae and S. warneri (1/105) were the least frequently isolated species. In teat skin swabs, S. equorum (43.3%, 116/268) was the most common species, while S. 270 271 vitulinus and S. warneri (1/268) were the least frequently isolated species. S. equorum was the most 272 frequently isolated NAS species from both left front (35.5%, 63/183) and right hind (30%, 57/190) quarters. Teat apex colonization with S. chromogenes, S. equorum S. cohnii, S. epidermidis, S. 273 haemolyticus and S. xylosus was not found to significantly increase the odds of IMI with these NAS 274 275 species, Table 2. Eighteen isolates without identification by MALDI-TOF were successfully identified using sequencing analysis, where 89% (n = 16) were correctly identified as NAS, while the 276 277 other 11% (n = 2) were *Corynebacterium stationis*, Table 3.

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DISCUSSION

To the best of our knowledge, this is the first study to evaluate the application of MALDI-TOF for 280 281 identification and differentiation of NAS species on quarter level from milk and teat skin in dairy cows milked with AMS. Our findings showed that MALDI-TOF was able to identify 92% (106/115) 282 of the submitted culture isolates originating from milk samples indicating that it is a reliable assay 283 284 for the rapid and accurate identification of NAS species from milk samples. This finding is similar to previous reports (Banach et al., 2016; Cameron et al., 2017a, b; Goetz et al., 2017; Savage et al., 285 2017). For instance, Elbehiry et al. (2016) showed that MALDI-TOF correctly identified 100% 286 (44/44) of NAS species isolated from milk, while Tomazi et al. (2014) demonstrated that the 287 sensitivity of MALDI-TOF for the identification of NAS isolated from milk was 95.4% in 288 289 comparison to PCR-RFLP. Similarly, MALDI-TOF showed a better performance (93.2%) for 290 identification of 234 NAS representing 20 different species than Phoenix (75.6%) and Vitek-2 (75.2%) (Dupont et al., 2010). Loonen et al., (2012) reported a good performance of MALDI-TOF 291 292 for identification of NAS with a correct-identification rate of 99.3%. A lower identification percentage (78%) was reported by Ayeni et al. (2017) based on 171 isolates (13 species) using 293 294 MALDI-TOF. However, the authors found that a drawback in identifying NAS with MALDI-TOF was the inability to identify S. gallinarum in their study because it was absent from the MALDI-TOF 295 296 database at the time of study.

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After two rounds of triplicate MALDI-TOF the overall number of isolates remaining without possible identification was 22% (112/511) representing 103 from teat skin and nine from milk. This is similar to Cameron et al. (2017b), who demonstrated that 40 isolates were not identified to species-level after the first round of duplicate MALDI-TOF, but after being subjected to a second round MALDI-TOF resulted in the identification of all except seven isolates. In line with that finding 303 Pizauro et al. (2017) reported that not all NAS found in buffalo milk could be identified by MALDI-TOF. Additionally, Banach et al. (2016) reported that six isolates of NAS were not identified to 304 species-level by MALDI-TOF but were classified by means of routine bacteriological testing, and 305 306 comprised of S. sciuri (three strains), S. xylosus (two strains), and S. equorum (one strain). This finding is in agreement with previous reports about the limitations of MALDI-TOF and could have 307 been caused by limited detection capacity of the bacteria due to a limited database of different 308 309 bacteria (Moussaoui et al., 2010; Barreiro et al., 2017; Cameron et al., 2017a). Furthermore, some studies have shown that the conditions of bacterial growth, preparation of samples, number of 310 311 reference strains, and version of the software Biotyper may be a reason for the variability of NAS 312 identification (Benagli et al., 2011; Tomazi et al., 2014).

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314 Our findings indicate that MALDI-TOF provides a valuable tool for identification and typeability of 315 NAS species from milk samples, while for isolates from non-milk samples, the assay showed a limited performance. Furthermore, in both the first and second submissions, the performance of 316 317 MALDI-TOF for identification of isolates originating from milk (85% and 47%) was higher than the assay performance in identification of isolates from teat skin (62% and 32%, respectively). One 318 possible explanation is that these unidentified bacteria from teat skin come from a natural teat skin 319 microbiota (commensal bacteria), which have not been previously included in the BDAL database. 320 321 For that reason, they may be out of interest of the microbiological diagnosticians and, therefore were 322 not considered in the database of MALDI-TOF. Moreover, our findings showed that the vast majority of the unidentifiable NAS isolates were originated from teat skin. A reasonable explanation 323 could be (a) those isolates are new NAS species. Supré et al. (2010) have classified 10 non-motile, 324 325 Gram-stain-positive, coagulase-negative staphylococci isolated from bovine milk and teat apices as S. devriesei sp. nov using 16S rRNA gene and four housekeeping genes (rpoB, hsp60, tuf and dnaJ) 326 in combination with tRNA-intergenic spacer length analysis. Another reason (b) these isolates could 327

328 be known NAS species but are not included in our BDAL database. It may be worth to mention that the database of MALDI-TOF we used in this study, is not based only on the commercial version of 329 BDAL database. Our database was updated regularly because the assay is being used on research and 330 331 routine diagnostic service purposes for identification and differentiation of bacterial pathogens in different samples types from both human and animals. We think that adding additional microbial 332 spectra, MSPs for some staphylococci species will improve the identification capacity of MALDI-333 334 TOF assay both from milk and teat skin. In line with that statement, Cameron et al. (2017b) showed that using a custom reference spectra expanded database, which included an additional 13 in-house 335 336 created reference spectra, isolates were identified by MALDI-TOF mass spectrometry with 99.2% (854/861) typeability and 99.4% (849/854) accuracy. 337

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339 Moreover, the BDAL database is mainly bacterial isolates originated from humans and then it 340 gradually extended to cover veterinary isolates (Tomazi et al., 2014). In support to that, Randall et al. (2015) stated that the majority of reference spectra included in commercial MALDI-TOF databases 341 342 are derived from human isolates and, consequently, small differences between human and animal isolates of the same bacterial species may influence the results obtained when testing some isolates 343 344 from animals. Remarkably, 89% of 18 isolates without possible identification by MALDI-TOF were correctly identified as NAS using PCR indicating that confirmation using nucleic-acid tools is 345 346 essential for those suspected NAS isolates of environmental origin. By examination of the genetic 347 diversity among NAS species using molecular typing methods, Piessens et al. (2012) identified five genotypes among S. chromogenes in six dairy farms. Cameron et al. (2017a) added that these 348 genotypes would have different spectra and would require their own entry within the database. We 349 350 updated the MALDI-TOF database on the basis of the findings in this study. Adding the spectra for eight NAS species based on both prior knowledge of the species and results of the assessment of the 351 352 database, the authors were able to eliminate the unidentified risk from 8% unidentified to 0%

unidentified. Therefore, in future studies, we will expect better agreement in the performance ofMALDI-TOF between teat skin and milk samples.

Consequently, MALDI-TOF is a valuable routine diagnostic tool for identification and 355 356 differentiation of mastitis pathogens but it has some limitations with regard to different detection capacity of the pathogens depending on the bacterial species, and limited database of pathogens 357 (Moussaoui et al., 2010; Barreiro et al., 2017; Cameron et al., 2017a). In line with that statement, our 358 findings showed that the isolates without possible identification by MALDI-TOF were successfully 359 identified by PCR, and sequencing analysis confirming the limited database of MALDI-TOF. 360 361 However, the database can continuously be expanded to accommodate new species and spectra. Furthermore, the MALDI-TOF assay depends on initial microbiological culture, which is time-362 consuming and laborious (i.e., plate preparation, sterilization of materials, time of incubation, 363 364 biochemistry tests) (Barreiro et al., 2010; Cameron et al., 2017a). Additionally, the reading is not 365 constant and differs by changing the cut-off (identification score) for species-level identification (Barreiro et al., 2017; Cameron et al., 2017a) and extra rounds of identification may be necessary for 366 367 achieving accurate identification as shown in this study. Moreover, some previous studies have shown that the conditions of bacterial growth, preparation of samples, number of reference strains or 368 spectra, database and version of the Biotyper may be a reason for the variability of NAS 369 identification (Benagli et al., 2011; Tomazi et al., 2014). Out of the initially unidentifiable isolates on 370 371 first round of MALDI-TOF, our findings showed that the assay identified 8/17 (47%; milk) and 372 48/151 (32%; teat skin) on the second round of identification. A plausable argument could be related to the variation in the conditions of bacterial growth, extraction procedure and handling and 373 preparation of samples between the first and second rounds of MALDI-TOF. MALDI-TOF may be 374 375 sensitive to the sample preparations and handling due to individual variations. This may also explain some of the deviations between different rounds, supporting previous reports (Benagli et al., 2011; 376 377 Tomazi et al., 2014).

Remarkably, we noticed that NAS isolates originating from teat skin may require extra rounds for identification by MALDI-TOF in comparison to NAS isolates originating from milk. That phenomenon is supported by the marked high number of isolates from teat skin remaining "unidentifiable" after the first and second rounds of identification. A possible explanation could be that bacterial isolates from the environment may have developed an extra layer or capsule of protein material as a mean of protection against unfavorable environmental conditions.

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386 MALDI-TOF as culture-dependent assay relies on direct analysis of proteins from bacterial cell extracts of the reference strains included in the database (Liang et al., 1996). So, when a given NAS 387 strain is tested, the species of the reference strain with the closest match is retained for identification 388 389 of the tested strain (Ayeni et al., 2017). Therefore, an up-to-date database is essential for bacterial 390 identification and more spectra of appropriate reference strains of NAS should be added to the database for accurate identification (van Veen et al., 2010; Cameron et al., 2017a). The same 391 392 conclusion was supported by Murugaiyan et al. (2014) and Randall et al. (2015), who reported that the ongoing supplementation of the Bruker database library should further improve the utility of 393 MALDI-TOF in routine veterinary diagnostic laboratories. Murugaiyan et al. (2014) demonstrated 394 that 17 isolates initially diagnosed as S. intermedius with the current content of the BDAL database 395 396 were identified as S. pseudintermedius by applying the in-house reference spectra extended version 397 indicating that updating the reference spectra library allowed species identification of NAS.

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CONCLUSION

MALDI-TOF provides a valuable and efficient platform for identification and typeability of NAS
species from aseptically collected quarter milk samples. MALDI-TOF may be used for identification

402 of NAS species from teat skin swabs. However, confirmation using nucleic-acid based tools is
403 essential for accurate identification of some NAS species and strains.

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REFERENCES

- Andresen, L.O., P. Ahrens, L. Daugaard, V. Bille-Hansen. 2005. Exudative epidermitis in pigs
 caused by toxigenic *Staphylococcus chromogenes*. Vet. Microbiol. 105: 291–300.
- Ayeni, F.A., C. Andersen, and N. Nørskov-Lauritsen. 2017. Comparison of growth on mannitol salt
 agar, matrix-assisted laser desorption/ionization time-of-flight mass spectrometry, VITEK® 2
 with partial sequencing of 16S rRNA gene for identification of coagulase-negative
 staphylococci. Microb. Pathog. 105: 255-259.
- Banach, T., M. Bochniarz, P. Łyp, Ł. Adaszek, W. Wawron, B. Furmaga, M. Skrzypczak, J. Ziętek,
 and S. Winiarczyk. 2016. Application of matrix-assisted laser desorption ionization time-offlight mass spectrometry for identification of coagulase-negative staphylococci isolated from
 milk of cows with subclinical mastitis. Pol. J. Vet. Sci. 19:627-632.

- Barreiro, J.R., C.R. Ferreira, G.B. Sanvido, M., Kostrzewa, T. Maier, B. Wegemann, V. Böttcher,
 M.N. Eberlin, and M.V. dos Santos. 2010. Short communication: Identification of subclinical
 cow mastitis pathogens in milk by matrix-assisted laser desorption/ionization time-of-flight
 mass spectrometry. J. Dairy Sci. 93:5661-7.
- Barreiro, J.R., J.L. Gonçalves, P.A. Braga, A.G. Dibbern, M.N. Eberlin, and M.V. dos Santos. 2017.
 Non-culture-based identification of mastitis-causing bacteria by MALDI-TOF mass
 spectrometry. J. Dairy Sci. 100:2928-2934.
- Benagli, C., V. Rossi, M. Dolina, M. Tonolla, and O. Petrini. 2011. Matrix-assisted laser desorption
 ionization-time of flight mass spectrometry for the identification of clinically relevant bacteria.
 PLoS ONE 6:e16424.
- Bizzini, A., C. Durussel, J. Bille, G. Greub, and G. Prod'hom. 2010. Performance of matrix-assisted
 laser desorption/ ionization-time of flight mass spectrometry for identification of bacterial
 strains routinely isolated in a clinical microbiology laboratory. J. Clin. Microbiol. 48:1549–
 1554.
- Braem, G., S. De Vliegher, B. Verbist, M. Heyndrickx, F. Leroy, and L. De Vuyst. 2012. Cultureindependent exploration of the teat apex microbiota of dairy cows reveals a wide bacterial
 species diversity. Vet. Microbiol. 157:383-90.
- Braem, G., S. De Vliegher, B. Verbist, V. Piessens, E. Van Coillie, L. De Vuyst, and F. Leroy. 2013.
 Unraveling the microbiota of teat apices of clinically healthy lactating dairy cows, with special
 emphasis on coagulase-negative staphylococci. J. Dairy Sci. 96:1499–1510.
- Cameron, M., H.W. Barkema, J. De Buck, S. De Vliegher, M. Chaffer, J. Lewis, and G.P. Keefe.
 2017a. Identification of bovine-associated coagulase-negative staphylococci by matrix-assisted
 laser desorption/ionization time-of-flight mass spectrometry using a direct transfer protocol. J.
 Dairy Sci. 100:2137-2147.

Cameron, M.J. Perry, J.R. Middleton, M. Chaffer, J. Lewis, and G.P. Keefe. 2017b. Evaluation of
 MALDI-TOF mass spectrometry and a custom reference spectra expanded database for the
 identification of bovine-associated coagulase-negative staphylococci. J. Dairy Sci. 101:1–6.

- Capurro, A., K. Artursson, K.P. Waller, B. Bengtsson, H. Ericsson-Unnerstad, and A. Aspan. 2009.
 Comparison of a commercialized phenotyping system, antimicrobial susceptibility testing, and
 tuf gene sequence-based genotyping for species-level identification of coagulase-negative
 staphylococci isolated from cases of bovine mastitis. Vet. Microbiol. 134:327–333.
- De Visscher, A., K. Supré, F. Haesebrouck, R.N. Zadoks, V. Piessens, E. Van Coillie, S. Piepers, and
 S. De Vliegher. 2014. Further evidence for the existence of environmental and host-associated
 species of coagulase-negative staphylococci in dairy cattle. Vet. Microbiol. 172:466-74.
- De Visscher, A., S. Piepers, F. Haesebrouck, and S. De Vliegher. 2016. Intramammary infection with
 coagulase-negative staphylococci at parturition: Species-specific prevalence, risk factors, and
 effect on udder health. J. Dairy Sci. 99:6457-69.
- 463 Dupont, C.1., V. Sivadon-Tardy, E. Bille, B. Dauphin, J.L. Beretti, A.S. Alvarez, N. Degand, A.
 464 Ferroni, M. Rottman, J.L. Herrmann, X. Nassif, E. Ronco, and E. Carbonnelle. 2010.
 465 Identification of clinical coagulase negative staphylococci isolated in microbiology laboratories
- by MALDI-TOF mass spectrometry and two automates, Clin. Microbiol. Infect. 16:998e1004.
- Elbehiry, A., M. Al-Dubaib, E. Marzouk, S. Osman, and H. Edrees. 2016. Performance of MALDI
 biotyper compared with Vitek[™] 2 compact system for fast identification and discrimination of *Staphylococcus* species isolated from bovine mastitis. Microbiology Open, 5:1061–1070.
- 470 Falentin, H., L. Rault, A. Nicolas, D.S. Bouchard, J. Lassalas, P. Lamberton, J.M. Aubry, P.G.
- 471 Marnet, Y. Le Loir, and S. Even. 2016. Bovine teat microbiome analysis revealed reduced alpha
- 472 diversity and significant changes in taxonomic profiles in quarters with a history of mastitis.
- 473 Front. Microbiol. 7:480.

- Fox, L.K., and R.J. Norell. 1994. *Staphylococcus aureus* colonization of teat skin as affected by
 postmilking teat treatment when exposed to cold and windy conditions. J. Dairy Sci. 77:2281–
 2288.
- Goetz, C., Y.D.N. Tremblay, D. Lamarche, A. Blondeau, A.M. Gaudreau, J. Labrie, F. Malouin, and
 M. Jacques. 2017. Coagulase-negative staphylococci species affect biofilm formation of other
 coagulase-negative and coagulase-positive staphylococci. J. Dairy Sci. 100:6454-6464.
- Gonçalves, J.L., T. Tomazi, J.R. Barreiro, P.A. Braga, C.R. Ferreira, J.P. Araújo Junior, M.N.
 Eberlin, and MV. dos Santos. 2014. Identification of *Corynebacterium* spp. isolated from bovine
 intramammary infections by matrix-assisted laser desorption ionization-time of flight mass
 spectrometry. Vet. Microbiol. 173:147-51.
- Kliem, M., and S. Sauer. 2012. The essence on mass spectrometry based microbial diagnostics. Curr.
 Opin. Microbiol. 15:397–402
- Leroy, F., E. Van Coillie, G. Braem, V. Piessens, B. Verbist, L. De Vuyst, and S. De Vliegher. 2015.
 Short communication: Subtyping of *Staphylococcus haemolyticus* isolates from milk and
 corresponding teat apices to verify the potential teat-skin origin of intramammary infections in
 dairy cows. J. Dairy Sci. 98:7893-8.
- 490 Liang, X., K. Zheng, M.G. Qian, and D.M. Lubman. 1996. Determination of bacterial protein
- 491 profiles by matrix-assisted laser desorption/ionization mass spectrometry with high-performance
 492 liquid chromatography, Rapid Commun. Mass Spectrom. 10:1219e1226.
- Loonen, A.J.M., A.R. Jansz, J.N.B. Bergland, M. Valkenburg, P.F.G. Wolffs, and A.J.C. van den
 Brule. 2012. Comparative study using phenotypic, genotypic, and proteomics methods for
 identification of coagulase-negative staphylococci, J. Clin. Microbiol. 50:1437e1439.
- Mahmmod, Y., I. Klaas, L. Svennesen, K. Pedersen, and H. Ingmer 2018. Coagulase negative
 staphylococci distribution in dairy herds with automatic milking system and their crosstalk with

498	Staphylococcus aureus from intramammary infections and teat apex. National Mastitis Council
499	57 th Annual Meeting, 31 st January to 2 nd February 2018, Tucson, Arizona, US.
500	Moussaoui, W., B. Jaulhac, A.M. Hoffmann, B. Ludes, M. Kostrzewa, P. Riegel, and G. Prévost
501	2010. Matrix-assisted laser desorption ionization time-of-flight mass spectrometry identifies 90
502	% of bacteria directly from blood culture vials. Clin. Microbiol. Infect. 16:1631–1638.
503	Murugaiyan, J., B. Walther, I. Stamm, Y. Abou-Elnaga, S. Brueggemann-Schwarze, S. Vincze, L.H.
504	Wieler, A. Lubke-Becker, T. Semmler, and U. Roesler. 2014. Species differentiation within the
505	Staphylococcus intermedius group using a refined MALDI-TOF MS database. Clin. Microbiol.
506	Infect. 20:1007–1015.
507	Nagy, E., M. Abrók, N. Bartha, L. Bereczki, E. Juhász, G. Kardos, K. Kristóf, C. Miszti, and E.
508	Urbán. 2014. Special application of matrix-assisted laser desorption ionization time-of-flight
509	mass spectrometry in clinical microbiological diagnostics. Orv. Hetil, 155:1495-503.
510	National Mastitis Council, 1999. Laboratory Handbook on Bovine Mastitis. National Mastitis
511	Council, Madison, WI.
512	Nonnemann, B., M. Tvede, T. Bjarnsholt. 2013. Identification of pathogenic microorganisms directly
513	from positive blood vials by matrix-assisted laser desorption/ionization time of flight mass
514	spectrometry. APMIS 123:871-877.
515	Paduch, J.H., E. Mohr and V. Kromker. 2013. The association between bedding material and the
516	bacterial counts of Staphylococcus aureus, Streptococcus uberis and coliform bacteria on teat
517	skin and in teat canals in lactating dairy cattle. J. Dairy Res. 80:159-164.
518	Park, J.Y., L.K. Fox, K.S. Seo, M.A. McGuire, Y.H. Park, F.R. Rurangirwa, W.M. Sischo, and G.A.
519	Bohach. 2011. Comparison of phenotypic and genotypic methods for the species identification
520	of coagulase-negative staphylococcal isolates from bovine intramammary infections. Vet.
521	Microbiol. 147:142-8.

522	Piepers, S., G. Opsomer, H.W. Barkema, A. de Kruif, and S. De Vliegher. 2010. Heifers infected
523	with coagulase-negative staphylococci in early lactation have less cases of clinical mastitis and a
524	higher milk production in their first lactation than non-infected heifers. J. Dairy Sci. 93:2014-
525	2024.

Piessens, V., S. De Vliegher, B. Verbist, G. Braem, A. Van Nuffel, L. De Vuyst, M. Heyndrickx, and
 E. Van Coillie. 2012. Intra-species diversity and epidemiology varies among coagulase-negative
 Staphylococcus species causing bovine intramammary infections. Vet. Microbiol. 155:62-71.

Pizauro, L.J.L., C.C. de Almeida, G.A. Soltes, D. Slavic, O.D. Rossi-Junior, F.A. de Ávila, L.F.
Zafalon, and J.I. MacInnes. 2017. Species level identification of coagulase negative *Staphylococcus* spp. from buffalo using matrix-assisted laser desorption ionization-time of flight
mass spectrometry and cydB real-time quantitative PCR. Vet. Microbiol. 204:8-14.

- 533 Pyörälä, S., and S. Taponen. 2009. Coagulase-negative staphylococci emerging mastitis pathogens.
 534 Vet. Microbiol. 134:3-8.
- Quirk, T., L.K. Fox, D.D. Hancock, J. Capper, J. Wenz, and J. Park. 2012. Intramammary infections
 and teat canal colonization with coagulase-negative staphylococci after postmilking teat
 disinfection: species-specific responses. J. Dairy Sci. 95:1906-12.
- 538 Randall, L.P., F. Lemma, M. Koylass, J. Rogers, R.D. Ayling, D. Worth, M. Klita, A. Steventon, K.
- Line, P. Wragg, J. Muchowski, M. Kostrzewa. and A.M. Whatmore. 2015. Evaluation of
 MALDI-ToF as a method for the identification of bacteria in the veterinary diagnostic
 laboratory. Res. Vet. Sci.101:42-49.
- 542 Sampimon, O.C., R.N. Zadoks, S. De Vliegher, K. Supré, F. Haesebrouck, H.W. Barkema, J. Sol,
- and T.J.G.M. Lam. 2009b. Performance of API Staph ID 32 and Staph-Zym for identification of
- 544 coagulase-negative staphylococci isolated from bovine milk samples. Vet. Microbio. 136:300–
- 545 305.

- Savage, E., S. Chothe, V. Lintner, T. Pierre, T. Matthews, S. Kariyawasam, D. Miller, D. Tewari,
 and B. Jayarao. 2017. Evaluation of Three Bacterial Identification Systems for Species
 Identification of Bacteria Isolated from Bovine Mastitis and Bulk Tank Milk Samples.
 Foodborne Pathog. Dis. 14:177-187.
- Schukken, Y.H., R.N. González, L.L. Tikofsky, H.F. Schulte, C.G. Santisteban, F.L. Welcome, G.J.
 Bennett, M.J. Zurakowski, and R.N. Zadoks. 2009. CNS mastitis: Nothing to worry about? Vet.
 Microbiol. 134:9-14.
- Singhal, N., M. Kumar, P.K. Kanaujia, and J.S. Virdi. 2015. MALDI-TOF mass spectrometry: an
 emerging technology for microbial identification and diagnosis. Front. Microbiol. 6:791.
- Strube, M.L., H.C. Ravn, H.C. Ingerslev, A.S. Meyer, and M. Boye. 2015. *In situ* prebiotics for
 weaning piglets: *in vitro* production and fermentation of potato galacto-rhamnogalacturonan.
 Appl. Environ. Microbiol. 81:1668–1678.
- Supré, K., F. Haesebrouck, R.N. Zadoks, M. Vaneechoutte, S. Piepers, and S. De Vliegher. 2011.
 Some coagulase-negative *Staphylococcus* species affect udder health more than others. J. Dairy
 Sci. 94:2329–2340.
- Supré, K., S. De Vliegher, I. Cleenwerck, K. Engelbeen, S. Van Trappen, S. Piepers, O.C.
 Sampimon, R.N. Zadoks, P. De Vos, and F. Haesebrouck. 2010. Staphylococcus devriesei sp.

nov., isolated from teat apices and milk of dairy cows. Int. J. Syst. Evol. Microbiol. 60:2739-44.

- Taponen, S., H. Simojoki, M. Haveri, H.D. Larsen, and S. Pyörälä. 2006. Clinical characteristics and
 persistence of bovine mastitis caused by different species of coagulase-negative staphylococci
- identified with API or AFLP. Vet. Microbiol. 115:199-207.

563

Taponen, S., S. Nykäsenoja, T. Pohjanvirta, A. Pitkälä, and S. Pyörälä. 2016. Species distribution
and in vitro antimicrobial susceptibility of coagulase-negative staphylococci isolated from
bovine mastitic milk. Acta Vet. Scand. 58:12.

- Thorberg, B.M., M.L. Danielsson-Tham, U. Emanuelson, K. Persson Waller. 2009. Bovine
 subclinical mastitis caused by different types of coagulase-negative staphylococci. J. Dairy Sci.
 92:4962-4970.
- 573 Tomazi, T., J.L. Gonçalves, J.R. Barreiro, P.A. de Campos Braga, L.F. Prada e Silva, M.N. Eberlin,

and M.V. dos Santos. 2014. Identification of coagulase-negative staphylococci from bovine

- 575 intramammary infection by Matrix-Assisted Laser Desorption Ionization-Time of Flight mass
 576 spectrometry, J. Clin. Microbiol. 52:1658–1663.
- van Veen, S.Q., E.C. Claas, and E.J. Kuijper. 2010. High-throughput identification of bacteria and
 yeast by matrix-assisted laser desorption ionizationetime of flight mass spectrometry in
 conventional medical microbiology laboratories, J. Clin. Microbiol. 48:900e907.
- 580 Vanderhaeghen, W., S. Piepers, F. Leroy, E. Van Coillie, F. Haesebrouck, and S. De Vliegher. 2014.
- Invited review: Effect, persistence, and virulence of coagulase-negative *Staphylococcus* species
 associated with ruminant udder health. J. Dairy Sci. 97:5275–5293.
- Watts, J.L., C.H. Ray, and P.J. Washburn. 1991. A convenient method for differentiation of
 coagulase-negative staphylococci isolated from bovine mammary glands. J. Dairy Sci. 74:426–
 428.
- Zadoks, R.N., and J.L. Watts. 2009. Species identification of coagulase-negative staphylococci:
 Genotyping is superior to phenotyping. Vet. Microbiol. 134:20–28.

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Table 1. Number and distribution of NAS isolated from milk and teat skin of dairy cows in eight

Sample	MALDI-TOF identification			on	No possible	Bacterial species (n) wrongly identified	
type					identificatio	as NAS on culture	
	First round		Second round		n		
	Submitte	Identified	Submitte	Identifie			
	d		d	d			
Milk	115	98	17	8	9	Corynebacterium stationis (1)	
Teat skin	396	245	151	48	103	Aerococcus viridans (7);	
						Corynebacterium casei (1); Bacillus	
						pumilus (13);	
						Clostridum septicum (1); Enterococcus	
						saccharolyticus (1); Corynebacterium	
						stationis (2)	
Overall	511	343	168	56	112	26	

590 dairy herds submitted to MALDI-TOF assay for species identification of NAS.

Table 2. Description of 16 species of NAS analyzed by MALDI-TOF and their association from

aseptic quarter milk and teat skin habitats collected from 142 cows (284 quarters) in eight dairy herds

594 with automatic milking systems.

NAS species ^a	Number (%)	Sample type (%)		OR ° (95% CI)	P-value*
		Milk	Teat skin	_	
S ^b . arlettae	12 (3.2)	1 (0.9)	11 (4.1)		
S. capitis	3 (0.8)		3 (1.1)		
S. cohnii	43 (11.5)	5 (4.8)	38 (14.2)	2.23 (0.11 - 15.6)	0.48
S. epidermidis	60 (16.1)	52 (49.5)	8 (3.0)	0.88 (0.05 - 5.07)	0.90
S. haemolyticus	58 (15.6)	16 (15.2)	42 (15.7)	1.13 (0.17 - 4.24)	0.55
S. hominis	17 (4.6)	3 (2.9)	14 (5.2)		
S. piscifermentans	2 (0.5)		2 (0.8)		
S. saprophyticus	5 (1.3)		5 (1.9)		
S. sciuri	9 (2.4)		9 (3.4)		
S. simulans	2 (0.5)	2 (1.9)			
S. succinus	2 (0.5)		2 (0.8)		
S. vitulinus	1 (0.3)		1 (0.4)		
S. warneri	2 (0.5)	1 (0.9)	1 (0.4)		
S. chromogenes	16 (4.3)	11 (10.5)	5 (1.9)	NA ^d	NA
S. equorum	122 (32.7)	6 (5.7)	116 (43.3)	NA	NA
S. xylosus	19 (5.1)	8 (7.6)	11 (4.1)	NA	NA
Other NAS (S. chromogenes,	157 (42.1)	25 (23.8)	132 (49.3)	0.40 (0.09 - 1.69)	0.21
S. equorum, S. xylosus) ^e					
Total	373 (100)	105 (100)	268 (100)		

^a Staphylococcus arlettae, S. warneri, and S. hominis were not considered in the statistical analysis

because of the few number of observations (< 5), while *S. capitis*, *S. piscifermentans*, *S.*

saprophyticus, *S. sciuri*, *S. simulans*, *S. succinus*, and *S. vitulinus* were not isolated from milk and/or teat skin.

^b S= *Staphylococcus*, ^c OR= Odds ratio; , ^d NA= not applicable; ^{*} significance at P < 0.05

600 ^e Other NAS (S. chromogenes, S. equorum, S. xylosus): they are grouped together for a valid

statistical analysis because the mixed model does not work for each species separately.

Table 3. Collection of bacterial isolates (n=18) from bovine teat skin suspected to be NAS on culture
and with no possible identification by MALDI-TOF after two rounds of submissions and were
correctly identified by 16S sequencing.

1 (5 ()
1 (5.6)
2 (11.1)
7 (38.9)
1 (5.6)
3 (16.7)
2 (11.1)
2 (11.1)
18 (100)