Existence of two groups of *Staphylococcus aureus* strains isolated from bovine mastitis based on biofilm formation, intracellular survival, capsular profile and agr-typing

Marjorie Bardiau\(a, b, \ast\), Jonathan Caplin\(b\), Johann Detilleux\(c\), Hans Graber\(d\), Paolo Moronie\(e, f\), Bernard Taminiau\(g\), Jacques G. Mainil\(h\)

\(a\) Bacteriology, Department of Infectious Diseases, Faculty of Veterinary Medicine and Institute for Fundamental and Applied Research in Animal Health (FARAH), University of Liège, Sart-Tilman, Bât. 43a, B–4000 Liège, Belgium

\(b\) Environment & Public Health Research Group, School of Environment & Technology, University of Brighton, Cockcroft Building, Lewes Road, Brighton BN2 4GJ, United Kingdom

\(c\) Biostatistics, Bioinformatics and Animal Selection, Department of Animal Production, Faculty of Veterinary Medicine, University of Liège, Sart-Tilman, Bât. 43a, B–4000 Liège, Belgium

\(d\) Agroscope, Institute for Food Sciences IFS, Schwarzenburgstrasse 161, 3003 Berne, Switzerland

\(e\) Università degli Studi di Milano, Department of Health, Animal Science and Food Safety, via Celoria 10, 20133 Milano, Italy

\(f\) Cornell University, Animal Health Diagnostic Center, Quality Milk Production Services, 240 Farrier Road, Ithaca, NY 14853, USA

\(g\) Quantitative Genetics Group, Department of Animal Production, Faculty of Veterinary Medicine, University of Liège, Sart-Tilman, Bât. 43a, B–4000 Liège, Belgium

\(h\) Corresponding author at: Bacteriology, Department of Infectious Diseases, Faculty of Veterinary Medicine and Institute for Fundamental and Applied Research in Animal Health (FARAH), University of Liège, Sart-Tilman, Bât. 43a, B–4000 Liège, Belgium. Fax: +32 4 366 42 61.

E-mail address: mbardiau@ulg.ac.be (M. Bardiau).

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**A B S T R A C T**

*Staphylococcus* (S.) *aureus* is recognised worldwide as an important pathogen causing contagious acute and chronic bovine mastitis. Chronic mastitis account for a significant part of all bovine cases and represent an important economic problem for dairy producers. Several properties (biofilm formation, intracellular survival, capsular expression and group agr) are thought to be associated with this chronic status. In a previous study, we found the existence of two groups of strains based on the association of these features. The aim of the present work was to confirm on a large international and non-related collection of strains the existence of these clusters and to associate them with case history records. In addition, the genomes of eight strains were sequenced to study the genomic differences between strains of each cluster. The results confirmed the existence of both groups based on capsular typing, intracellular survival and agr-typing: strains cap\(\ast\)-positive, belonging to agr group II, showing a low invasion rate and strains cap\(\ast\)-positive, belonging to agr group I, showing a high invasion rate. None of the two clusters were associated with the chronic status of the cow. When comparing the genomes of strains belonging to both clusters, the genes specific to the group "cap\(\ast\)-agr" would suggest that these strains are better adapted to live in hostile environment. The existence of these two groups is highly important as they may represent two clusters that are adapted differently to the host and/or the surrounding environment.

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**1. Introduction**

*Staphylococcus aureus* (S. *aureus*) is an important bacteria causing contagious bovine mastitis (Watts, 1988). *S. aureus* strains can cause acute, usually clinical, and chronic, usually subclinical, mastitis. Subclinical mastitis is characterised by a non-alteration of the milk but high somatic cell count, making the milk inappropriate for the consumers if the cell count is too high. This type of mastitis is often chronic and account for up to 30% of all bovine cases (Halasa et al., 2007), which represents an important economic problem for dairy producers with reduction in milk quantity and quality, prolonged costly antibiotic treatments and premature culling. Several properties are thought to be associated to some extent with these chronic infections: biofilm formation, intracellular survival, capsular expression and group accessory gene regulator (agr).

Biofilm formation is one of the most important survival mechanisms of bacteria living in the extracellular niche. It impairs
the action of both the host immune system and antimicrobial agents (Costerton et al., 1999; Melchior et al., 2006). Second, S. aureus can be an intracellular pathogen of a large variety of eukaryotic cells, including epithelial cells of the mammary glands and immune cells (Almeida et al., 1996; Kerro Dego et al., 2002).

Therefore, the bacteria are not only protected from the action of commonly used antibiotics in mastitis treatment (mainly β-lactams), and also able to persist in the host without causing any apparent inflammation (Boulanger et al., 2003; Garzoni and Kelley, 2009). The absence of capsular expression enhances the adherence to and the invasion of eukaryotic cells by S. aureus (Pohlmann-Dietze et al., 2000; Buzzola et al., 2007; Tuchscherr et al., 2005).

In addition, S. aureus bacteria that do not express capsule induce chronic mastitis in mice, suggesting that the absence of capsule synthesis may help the bacteria to persist in the mammary glands (Tuchscherr et al., 2005). Finally, agr group I is associated with a persistence or with features that help the bacteria to persist in the udder: strains belonging to agr group I are more likely to be isolated in epithelial cells, to persist in murine mammary glands (Buzzola et al., 2007) and to be associated with penicillin resistance (Melchior et al., 2011) than the strains belonging to the other groups.

In a previous study (Bardiau et al., 2014), we correlated agr-typing, capsular expression, biofilm formation, and intracellular survival in a collection of Belgian S. aureus strains from bovine mastitis (without case history records). We found the existence of two groups based on the association of these features: caps-positive strains belonging to agr group I, which in vitro test negative for CP5 ELISA and show a high invasion rate in MAC-T cells, and cap8-positive strains belonging to agr group II, which express CP8 in vitro and show a low invasion rate in MAC-T cells. We hypothesised that the first group may correspond to strains adapted to the intracellular niche leading to chronic infection and that the second group may correspond to strains better adapted to the extracellular niche leading to acute infection.

The aim of this work was therefore to confirm on a large European and non-related collection of strains the existence of these two groups based on features described to be associated with long-lasting infections and to find out if these two clusters are associated with a persistence of the disease. We therefore investigated the association between agr-typing, capsular antigen identity and expression, biofilm formation, intracellular survival and the case history data of a collection of S. aureus strains isolated from cases of chronic and acute bovine mastitis in four countries (Belgium, Italy, Canada and Switzerland). Moreover, the genomes of eight strains were sequenced and compared to assess of the genomic differences between the formed clusters.

2. Materials and methods

2.1. Bacterial isolates

A total of 168 bovine S. aureus isolates, were collected from chronic and acute bovine intramammary infection in four countries. One hundred and two isolates were collected from chronic cases in Belgium (n = 7), Canada (n = 45), Italy (n = 25) and Switzerland (n = 25). Sixty-six isolates were collected from acute cases in Canada (n = 46) and Italy (n = 20). Strains have been isolated and identified as S. aureus according to the protocol described in Ote et al. (2011). Chronic mastitis was defined as recurring isolates in the same quarter of the same animal. Four S. aureus reference strains were included in this collection: ATCC 29740 (N305), ATCC 31885 (NL6), ATCC 49521 (Lowenstein) and ATCC 49525 (Wright).

2.2. Capsular genotyping and serotyping

Capsular genotyping was performed using PCR detection of the capsule-encoding genes cap5 and cap8 genes as previously described (Ote et al., 2011). Capsular serotyping was performed in in triplicates and in two independent experiments by enzyme-linked immunosorbent assay (ELISA) using specific monoclonal and polyclonal antibodies (kindly provided by GSK Biologicals, Belgium) against CAP5 and CAP8 as previously described (Bardiau et al., 2014). For the serotyping, OD values were compared to those obtained with S. aureus CP reference strains, namely the CP5-positive strain ATCC 49521 and CP8-positive strain ATCC 49525, and isolates that tested negative for CP5- and CP8- ELISA were defined as non-typeable (NT).

2.3. Invasion assay

Bovine mammary epithelial cells (MAC-T) were used for in vitro bacterial internalisation assays as previously described (Bardiau et al., 2014; Boulanger et al., 2007; Brouillet et al., 2003). Briefly, cell monolayers (~2.5 × 10⁵ cells/well) were inoculated with 10⁷ CFU of S. aureus (MO1–40) and incubated at 37 °C in 5% CO₂ for three hours. After the removal of extracellular bacteria by phosphate buffered saline (PBS) washing and lysozyme treatment, the MAC-T cells were detached and lysed by addition of 900 μl/well of sterile distilled water containing 0.025% Triton X-100. The cell lysates were carefully suspended, serially diluted, and plated on Columbia sheep blood agar plates to quantify intracellular staphylococci. Results are expressed as a percentage of the initial inoculum and classified using the same criteria as in our previous work (Bardiau et al., 2014).

2.4. Biofilm production

Biofilm formation was evaluated by spectrophotometry in microplates using safranin staining as previously described (Bardiau et al., 2014). Briefly, overnight cultures were diluted 1:100 in tryptic soy broth (TSB) containing 0.25% glucose (TSBglc), transferred into wells of sterile 96-well polystyrene tissue culture (TC) plates and incubated at 37 °C. TSBglc without bacteria served as negative control. After 24 h, the plates were stained with safranin 0.1% (w/v) for 10 min. A mixture of 50% ethanol and 50% acetic acid was added to each well and plates were incubated at room temperature for 15 min. Finally, the OD of each well was measured at 490 nm using a microplate reader. The results were collected from at least two independent experiments in which the biofilm formation of each culture tested was evaluated in triplicate. The quantitative classification of biofilm production based on cut-off value (ODc) and average OD values was carried out leading to four categories of strains: no biofilm producer (OD ≤ ODc); weak biofilm producer (ODc < OD ≤ 2 × ODc); moderate biofilm producer (2 × ODc < OD ≤ 4 × ODc); strong biofilm producer (4 × ODc < ODc) (Stepanovic et al., 2007).

2.5. agr-typing

agr-groups were determined by multiplex PCR as previously described (Gilot et al., 2002). In brief, multiplex PCRs were performed with the following primers: Pan (5’-ATG CAC ATG TGT CAC ATG C-3’), agr1 (5’-GTC ACA AGT ATA AGC TGC GAT), agr2 (5’-TAT TAC TAA TTG AAA AGT GGC CAT AGC-3’), agr3 (5’-GTA ATG TAA TAG CTT GTA TAA TAC TAC G-3’), and agr4 (5’-CGA TAA TGC GG TCT AAT ACC CG-3’). Amplifications were performed with the following PCR program: 1 cycle at 94 °C for 1 min; 26 cycles at 94 °C for 30 s, 55 °C for 30 s, and 72 °C for 1 min; and finally 1 cycle at
2.6. Statistical analysis

For calculation of the statistical significance of the observed frequency distributions, contingency tables of the expected values were determined and Chi-square tests were performed. P values lower than 0.05 were considered significant. K-means clustering method, adapted for the clustering of categorical data, was used to group similar strains into homogeneous groups (procedure fastclus, sas 9.1).

2.7. Genomes sequencing

Eight strains (four from group 1, named “cap5-agr1”, isolated in Belgium and in Italy and four from group 2, named “cap8-agr1”, isolated in Belgium and in Italy) were sequenced to assess of the genomic differences between strains of both groups. MiSeq next-generation sequencing NGS was performed according to the manufacturer’s instructions using Nextera Mate Pair Library Preparation kit with 2 × 250 bp paired-end (Illumina). Assembly was performed de novo using Platanus genome assembler. Annotations were performed using RAST server (Aziz et al., 2008). This Whole Genome Shotgun project has been deposited at DDBJ/EMBL/GenBank under the accession LRMX00000000, LRMY00000000, LRMZ00000000, LRNA00000000, LRNB00000000, LRNC00000000, LRND00000000 and LRNE00000000. The version described in this paper is version XXXX1000000. Genomic analyses (Multi-Locus Sequences Typing-MLST and virulence factors) were performed using the Center for Genomic Epidemiology website platform (Larsen et al., 2012; Zankari et al., 2012; Joensen et al., 2014; Kaas et al., 2014).

3. Results

3.1. Capsular serotyping, invasion assay, biofilm formation and agr-typing (Table 1)

Ninety-seven strains (58%) and 71 strains (42%) harboured the cap5 and cap8 genes, respectively. We then assessed the expression of the CP5 and CP8 polysaccharides by ELISA analysis. Eighty-two (49%) S. aureus isolates expressed CP5, 69 (41%) expressed CP8, and 17 (10%) were defined as non-typeable.

We chose to divide the distribution of the strains in the collection into 2 populations as we did in our previous study (Bardiau et al., 2014). Thus, we determined that 74 strains (44%) showed a lower invasion rate, while 94 strains (56%) showed a higher invasion rate.

Eighteen strains (11%) did not produce any biofilm, 85 (51%) produced weak biofilms, 49 (29%) produced moderate biofilms, while 16 (10%) produced strong biofilms.

In the collection of strains, three of the four agr-groups were detected by PCR, with two (agr group I and II) representing 99% of the strains: 97 (58%) strains belonged to agr group I, 69 (41%) belonged to agr group II, and two strains (1%) belonged to agr group IV.

3.2. Statistical associations and clustering method

We could find the following statistical associations (Table 1): (i) we first highlighted that more cap8-positive strains expressed CP8, whereas more cap5-positive strains were defined as non-typeable; (ii) we then observed that more cap8/CP8-strains showed a low invasion rate, whereas more cap5/NT or cap5/CP5 strains showed a higher invasion rate; (iii) furthermore, the intracellular survival capacity and the capsular serotype of the strains were associated

Table 2

<table>
<thead>
<tr>
<th>Clusters</th>
<th>I</th>
<th>II</th>
<th>III</th>
</tr>
</thead>
<tbody>
<tr>
<td>Capsular profile</td>
<td>cap5/NT</td>
<td>0</td>
<td>15</td>
</tr>
<tr>
<td>cap5/CP5</td>
<td>7</td>
<td>75</td>
<td></td>
</tr>
<tr>
<td>cap8/NT</td>
<td>2</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>cap8/CP8</td>
<td>61</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>Intracellular survival</td>
<td>Low</td>
<td>52</td>
<td>22</td>
</tr>
<tr>
<td>High</td>
<td>18</td>
<td>76</td>
<td></td>
</tr>
<tr>
<td>Biofilm formation</td>
<td>No</td>
<td>15</td>
<td>3</td>
</tr>
<tr>
<td>Weak</td>
<td>38</td>
<td>47</td>
<td></td>
</tr>
<tr>
<td>Moderate</td>
<td>16</td>
<td>33</td>
<td></td>
</tr>
<tr>
<td>Strong</td>
<td>1</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>agr-groups</td>
<td>I</td>
<td>0</td>
<td>97</td>
</tr>
<tr>
<td>II</td>
<td>68</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>III</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>IV</td>
<td>2</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Anamnesis</td>
<td>Chronic</td>
<td>41</td>
<td>61</td>
</tr>
<tr>
<td>Acute</td>
<td>29</td>
<td>37</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>70</td>
<td>98</td>
<td></td>
</tr>
</tbody>
</table>

Significant correlations (P < 0.001) are highlighted in bold. Capsular profile: cap, capsule-encoding gene; CP, capsular polysaccharide; NT, non-typeable. Intracellular survival: presented as the % of the initial inoculums. agr-typing: I, II, III, IV, agr-groups.

Table 1

<table>
<thead>
<tr>
<th>Capsular profile, intracellular survival, biofilm formation and agr-groups frequency distribution.</th>
<th>Low</th>
<th>High</th>
<th>No</th>
<th>Weak</th>
<th>Moderate</th>
<th>Strong</th>
<th>agr-groups</th>
</tr>
</thead>
<tbody>
<tr>
<td>Capsular profile</td>
<td>cap5/NT</td>
<td>4</td>
<td>11</td>
<td>0</td>
<td>8</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>cap5/CP5</td>
<td>20</td>
<td>62</td>
<td>2</td>
<td>41</td>
<td>26</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>cap8/NT</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>cap8/CP8</td>
<td>49</td>
<td>20</td>
<td>16</td>
<td>35</td>
<td>16</td>
<td>2</td>
</tr>
<tr>
<td>Intradimensional survival</td>
<td>Low</td>
<td>14</td>
<td>41</td>
<td>14</td>
<td>5</td>
<td>2</td>
<td>22</td>
</tr>
<tr>
<td></td>
<td>High</td>
<td>4</td>
<td>44</td>
<td>35</td>
<td></td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>Biofilm formation</td>
<td>No</td>
<td>3</td>
<td>15</td>
<td>0</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Weak</td>
<td>47</td>
<td>38</td>
<td>0</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Moderate</td>
<td>32</td>
<td>15</td>
<td>0</td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Strong</td>
<td>15</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Significant correlations are highlighted in bold. cap, capsule-encoding gene; CP, capsular polysaccharide; NT, non-typeable.
with the agr groups: strains belonging to agr group I showed a higher internalisation rate and were mainly cap5-positive, whilst strains belonging to agr group II showed a lower internalisation rate and were mainly cap8-positive; (iv) no association could be observed between the biofilm formation and any other results (intracellular survival, capsular serotype and agr-typing).

All the strains grouped into two clusters by the K-Means statistical method (Table 2). The cluster 1 grouped strains that were cap8-positive, belonged to agr group II, showed a low invasion rate in MAC-T cells and formed weak or no biofilm in TSBgic. The cluster 2 grouped strains that were cap5-positive, belonged to agr group I, showed a high invasion rate in MAC-T cells and formed weak, moderate or strong biofilm in TSBgic.

These two groups were associated with the case history record (chronic or acute mastitis). However, no association was seen between the clusters and the chronic status of the infection (Table 2).

### 3.3. Cluster-associated variable gene content

In order to further evaluate the genomic differences between the strains belonging to both clusters, we sequenced eight genomes: four strains belonging to the group “cap5-agrII” (two isolated in Belgium and two isolated in Italy) and four strains belonging to the group “cap8-agrII” (two isolated in Belgium and two isolated in Italy). The size of the sequenced genomes varied from 2.55 Mb to 2.77 Mb and the number of coding sequences from 1944 to 2012. MLST and virulence gene profiles were determined for all sequenced strains (Table 3). Six strains belong to five different ST (ST115, ST504, ST8, ST97, ST479) and the ST of two strains could not be identified. Some virulence genes could not be found in any strain: aap coding for the accumulation-associated protein, sea,sec, seg, sei, sel, sen, seo, respectively, coding for the enterotoxins A, C, G, I, L, N and O, etb coding for the exfoliative toxins B, cna coding for the collagen-binding protein. Some virulence genes were detected in all strains: the cap operon coding for the capsule, lukF-PV coding for the Panton–Valentine leukocidin chain F, hlb coding for the beta-haemolysin precursor, hlg coding for the gamma-haemolysin precursor, atl coding for the bifunctional autolysin Atl, eta coding for the exfoliative toxin A, spla coding for the serine protease SplA, sspa coding for the serine V8 protease, vwb coding for the von Willebrand factor-binding protein, eap coding for the extracellular adherence protein, spa coding for the protein A. When comparing the presence of the virulence genes in the strains regarding their groups, lukM, the gene coding for the leukocidin LukM, was present in most of the “cap8-agrII” strains and none of the “cap5-agrII” strains and three genes, hld, clfA and sdrC, respectively, coding for the delta-haemolysin precursor, the clumping factor A and the Bone sialoprotein-binding protein, were present in most if not all “cap5-agrII” strains and in only one if not none “cap8-agrII” strains.

When comparing all genomes, nine genes appeared to be exclusively specific to the “cap5-agrII” strains and one to the “cap8-agrII” strains. When relaxing the stringency of the comparison (presence in minimum three out of four strains in one group and absence in minimum three out of four strains in the other one), we could identify 51 (and therefore a total of 60 genes when including the nine exclusively specific) and 13 (and therefore a total of 14 genes when including the one exclusively specific) genes more specific to cluster 1 “cap5-agrII” and cluster 2 “cap8-agrII” isolates, respectively (Fig. 1). The 60 genes over-represented in the cluster 1 “cap5-agrII” were assigned to the following functional categories: antibiotic and heavy metal resistance (β-lactamase, cadmium and arsenic resistance proteins), capsular proteins (cap5), DNA-related proteins (DNA-invertase, transposase, putative primase, transcriptional regulator, mutator mutT, cl-like repressor), putative membrane protein, proline/glycine betaine transporter, pathogenicity island (Staphylococcal Pathogenicity Island 2), fibrinogen-binding protein, δ-haemolysin and hypothetical proteins. The 14 genes over-represented in the cluster 1 “cap8-agrII” were assigned to the following functional categories: exotoxin homologue (genomic island nu Sa alpha2), DNA-related proteins (DNA-cytosine methyltransferase, transcriptional regulator), intracellular protease/amidase (ThI/PPl family), pathogenicity island (Staphylococcal Pathogenicity Island 1 Orf21), phage minor head and hypothetical proteins.

<table>
<thead>
<tr>
<th>Table 3</th>
<th>MLST and virulence factors genes contents of all sequenced strains.</th>
</tr>
</thead>
<tbody>
<tr>
<td>MLST</td>
<td>cap8-agrII strains</td>
</tr>
<tr>
<td>120</td>
<td>ST504</td>
</tr>
<tr>
<td>Biofilm formation</td>
<td>ica operon</td>
</tr>
<tr>
<td></td>
<td>aap</td>
</tr>
<tr>
<td></td>
<td>cap operon</td>
</tr>
<tr>
<td>Secreted toxins</td>
<td>lukE</td>
</tr>
<tr>
<td></td>
<td>lukD</td>
</tr>
<tr>
<td></td>
<td>lukM</td>
</tr>
<tr>
<td></td>
<td>lukF-PV</td>
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<tr>
<td></td>
<td>lukG</td>
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<tr>
<td></td>
<td>hla</td>
</tr>
<tr>
<td></td>
<td>hlb</td>
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<td>hlg</td>
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<td>hld</td>
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<tr>
<td></td>
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<tr>
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<td>tsp</td>
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<tr>
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<td>splE</td>
</tr>
<tr>
<td></td>
<td>sspA</td>
</tr>
</tbody>
</table>

NC: not identified; Genes coding for: ica, Intercellular adhesin protein A; aap, accumulation-associated protein; cap, capsular polysaccharide; lukE, leukocidin LukE; lukD, leukocidin LukD; lukF, leukocidin LukF; lukF-PV, Panton-Valentine leukocidin chain F; lukS, leukocidin chain S; hla, Alpha-haemolysin precursor; hlb, Beta-haemolysin precursor; hlg, Gamma-haemolysin precursor; hld, Delta-haemolysin precursor; atl, Bifunctional autolysin; eta, Exfoliative toxin A; etb, Exfoliative toxin B; sea, enterotoxin A; sec, enterotoxin C; seg, enterotoxin G; sei, enterotoxin I; sel, enterotoxin L; sen, enterotoxin N; seo, enterotoxin O; splA, Serine protease SplA; splE, Serine protease SplE; sspa, Serine V8 protease; vwb, Secreted von Willebrand factor-binding protein precursor; clfA, Clumping factor ClfA; clfB, Clumping factor ClfB; fnbA, Fibrinectin-binding protein A; fnbB, Fibrinectin-binding protein B; eap, extracellular protein; sak, staphylokinase; spa, Protein A (IgG-binding protein); cna, Collagen-binding protein; ebp5, Cell surface elastin-binding protein; sdrC, serine-aspartate repeat protein.
4. Discussion

In a previous study (Bardiau et al., 2014), it was found that S. aureus strains isolated from Belgian bovine mastitis could be divided into several clusters based on features potentially associated with long-lasting infections (biofilm formation, capsular profile, agr-typing and intracellular survival). In this present work, we aimed to confirm using a large international and non-related collection of strains the existence of these clusters and to find out whether they could be associated with a persistence of the infection of the udder. Therefore, we studied the association between these features and the case history of strains isolated from chronic and acute bovine mastitis from four countries (Belgium, Canada, Italy, Switzerland). We also investigated the genomic characteristics of eight strains belonging to the different groups.

First, we find the same kind of associations than in our first study when comparing results two by two, except for the biofilm formation that was not associated with any other features. When correlating the results with the country of origin, the repartition is evenly distributed (data not shown); therefore the country of origin does not introduce any bias in the statistical analysis of the results.

When strains were clustered (Bardiau et al., 2014), similar correspondences were observed: one cluster of strains that are cap8-positive, belong to agr group II, show a low invasion rate in MAC-T cells and form weak or no biofilm in TSBgly, and one cluster of strains that are cap5-positive, belong to agr group I, show a high invasion rate in MAC-T cells and form weak, moderate or strong biofilm in TSBgly. Our previous hypothesis (Bardiau et al., 2014) was that the first group of “cap8-agrII” strains are better adapted to an extracellular niche and therefore could be associated with acute mastitis and that the second group of “cap5-agrI” strains are better adapted to an intracellular niche and therefore could be associated with chronic mastitis. However, no correlation is found in this study between any feature of the strain and the case history data. The fact that those case history data were determined on an individual basis in each country of origin and not as part of the same study using the same criteria may of course have introduced a bias in this study and therefore explain in part this absence of correlation.

We analysed the presence of virulence genes in the eight sequenced strains. The relative presence of the genes vary from 0% to 100% and no clear genotype subtype regrouping several strains appears as previously shown by our group (Ote et al., 2011). Nevertheless, four genes are specific to one group or the other (“cap5-agrI” or “cap8-agrII”): lukM to “cap8-agrII” and hld, clfA and sdrC to “cap5-agrI”. These results are in accordance with the study published by Peton et al. (2014) in which they have compared two S. aureus bovine strains, N305 that produces mild and chronic mastitis and RF122 that produces severe mastitis. In their study, lukM is present in RF122 and absent in N305. lukM in association with lukF-PV form LukM/F, a protein involved in cytokotaxis against polynuclear neutrophils, mainly described during a strong inflammatory reaction in the mammary gland and therefore in severe clinical mastitis. In contrast, in their study, the gamma-haemolysin is more produced in N305 than in RF122. In our study, we observe that the gamma-haemolysin gene is more present in the “cap5-agrI” than in the “cap8-agrII”. The protein encoded by the sdrC gene promotes both bacterial adherence to surfaces and biofilm formation (Barbu et al., 2014). The clumping factor A promotes bacterial attachment to eukaryotic cells, induces formation of bacterial clumps and decreases the phagocytosis. Both proteins could be implicated in the persistence of the infection by helping the bacteria to survive in hostile environment.

Finally, when analysing the genes over-represented in either group of the eight strains (four from either group) that were sequenced, we can hypothesise that the strains “cap5-agrI” are better adapted to live and persist in the environment due to the presence of heavy metal resistance genes and proline/glycine betaine transporter. It has indeed been shown that the proline/glycine betaine confers osmotic protection to various bacterial species including Staphylococcus sp. (Amin et al., 1995), thus it may help the bacteria to survive in high osmolality environments. It is interesting to note that high level of glycine betaine has recently been associated with planktonic versus biofilm producer staphylococci (Junka et al., 2013). The presence of heavy metal resistance
genes could also help the bacteria to survive in hostile environments and can facilitate the persistence and dissemination of the bacteria. Antibiotic resistance genes (especially production of β-lactamase), here associated with the strains "cap5-argI", have previously been linked with argI-positive strains (Melchior et al., 2011). However several of the genes over-represented in both groups are hypothetical proteins. It would be interesting to investigate their presence in a larger collection of strains to target the ones that seem to be specific to either group and to study their functions.

5. Conclusion

In conclusion, we confirm the existence of two groups of S. aureus strains isolated from bovine mastitis based on capsular typing, intracellular survival and agr-typing. Strains cap8-positive, belonging to agr group II, show a low invasion rate and strains cap5-positive, belonging to agr group I, show a high invasion rate. Despite the fact that we could not correlate these groups with the case history data, the existence of these two groups is highly important as they may represent two clusters that are adapted differently to the host and/or the surrounding environment. When comparing the genomes of strains belonging to both clusters, only 14 genes are over-represented in the group “cap8-argI”, but 60 genes are over-represented in the group “cap5-argI”. The genes specific to the group “cap5-argI” would suggest that these strains are better adapted to live in the environment than the strains belonging to “cap8-argI”. The first perspective of this study would be to check for the presence of these genes in a larger collection of strains to assess of their specificity for each cluster. In addition, it would be beneficial to study these features (intracellular survival, biofilm formation, capsular typing and agr-typing) in S. aureus strains isolated from other hosts and other pathologies to specify if these clusters are specific to bovine mastitis S. aureus strains or are present in all S. aureus strains. Finally, a broader study collecting worldwide strains from mastitis with anamnesis done with the same criteria would be ideal to assess of the existence of groups regarding the chronic status of the cow.

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