IsdC from *Staphylococcus lugdunensis* induces biofilm formation in low-iron growth conditions

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ABSTRACT

*Staphylococcus lugdunensis* is a coagulase-negative *Staphylococcus* that is a commensal of humans and an opportunistic pathogen. It can cause a spectrum of infections including those that are associated with the ability to form biofilm such as endocarditis or indwelling medical devices. The genome sequences of two strains revealed the presence of orthologues of the *ica* genes that are responsible for synthesis of poly-N-acetyl glucosamine (PNAG) that is commonly associated with biofilm in other staphylococci. However, here we discovered that biofilm formed by a panel of *S. lugdunensis* isolates growing in iron-restricted medium was susceptible to degradation by proteases and not by metaperiodate suggesting that the biofilm matrix comprised proteins and not PNAG. When the iron concentration was raised to 1mM biofilm formation by all strains tested was greatly reduced. A mutant of strain N920143 lacking the entire locus that encodes iron-regulated surface determinant (Isd) proteins was defective in biofilm formation under iron-limited conditions. A IsdC null mutant was defective whereas IsdK, IsdJ and IsdB mutants formed biofilm to the same level as the parental strain. Expression of IsdC was required both for the primary attachment to unconditioned polystyrene and for the accumulation phase of biofilm involving cell-cell interactions. Purified recombinant IsdC protein formed dimers in solution and *Lactococcus lactis* cells expressing only IsdC adhered to immobilized recombinant IsdC but not to IsdJ, IsdK or IsdB. This is consistent with a specific homophilic interaction between IsdC molecules on neighbouring cells contributing to accumulation of *S. lugdunensis* biofilm *in vivo*.

INTRODUCTION

Coagulase-negative staphylococci (CoNS) and *Staphylococcus aureus* are the predominant etiological agents of medical device-related infections, largely owing to their ability to form biofilm. Biofilms are defined as communities of bacteria encased in a self-synthesized extracellular
polymeric matrix (1) growing attached to biological or abiotic surfaces. Staphylococci in biofilms are resistant to antibiotics (2) and host immune responses (3), reducing the efficacy of available antimicrobials. The formation of biofilm is a complex, multifactorial process. Initially, bacteria adhere directly to the surface of implanted device or to devices coated with the host matrix components. In *S. aureus* biofilm the major autolysin Atl mediates primary attachment to plastic surfaces by promoting release of DNA from bacterial cells (4, 5), while adherence to surfaces conditioned by host plasma proteins is promoted by surface protein adhesins such as the fibrinogen-binding clumping factor A or fibronectin binding proteins (6). This process is followed by proliferation, accumulation and intercellular interactions mediated by the icaADBC-encoded polysaccharide intercellular adhesins (PIA) (7) or surface proteins such as Bap (8), SasG (9), SasC (10), protein A (11), or fibronectin-binding proteins (FnBPs) (12, 13). Likewise, biofilm formation by *S. epidermidis* is dependent on PIA or proteinaceous components such as Aap (14, 15) or SesC (16).

*Staphylococcus lugdunensis* is a coagulase-negative species with enhanced virulence compared with the other CoNS (17). *S. lugdunensis* causes a severe form of native valve endocarditis (18, 19), infections of prosthetic heart valves (20), intravascular catheters (21), prosthetic joints (22) and ventriculoperitoneal shunts (23). This pathogenic potential is largely attributed to the ability of this bacterium to form biofilm. A previous study by Frank and Patel (24) demonstrated that despite the presence of icaADBC orthologues in *S. lugdunensis*, PIA is not the major component of the extracellular matrix of biofilms formed in vitro by this species. Rather, *S. lugdunensis* biofilms appear to be composed of proteins. *S. lugdunensis* expresses a fibrinogen-binding protein (Fbl) (25), a member of the family of MSCRAMM (Microbial Surface Components Recognizing Adhesive Matrix Molecules) family that is closely related to ClfA, which does not appear to have any role in biofilm formation (26) and a yet uncharacterized von Willebrand factor binding protein (27).
Uniquely for CoNS, *S. lugdunensis* contains a cluster of genes with similarity both in terms of organization and sequence to the iron-regulated surface determinant (*isd*) locus of *S. aureus* (28). Both systems are expressed under iron-restricted conditions (28, 29). Four of the *S. lugdunensis Isd* proteins are anchored to the cell wall peptidoglycan by sortases. In *S. aureus* and *S. lugdunensis*, the Isd proteins cooperate to capture heme and transfer it across the wall to a membrane-bound transporter, which delivers it to the cytoplasm, where heme is degraded to recover iron (30). There is evidence that surface-exposed Isd proteins may have additional roles in colonization and pathogenesis of both species. For example, IsdJ from *S. lugdunensis* (29) and IsdA from *S. aureus* (31) are multifunctional proteins which recognize and bind several host proteins and can confer resistance to skin fatty acids. In this study we investigated biofilm formation *in vitro* by a collection of *S. lugdunensis* isolates grown in low iron conditions and assessed the role of *isd* locus in biofilm formation by this important pathogen.

**MATERIALS AND METHODS**

**Bacterial strains and culture conditions.** The microorganisms used in this study are reported in Table 1. *S. aureus* V329 (8) and SA113 (32) were kindly donated by Dr. J.R. Penades (Universidad Cardenal Herrera-CEU, Moncada, Valencia, Spain). The clinical isolate of *S. epidermidis* 5179R (14) was provided by Dr. H. Rohde (University Medical Centre Hamburg-Eppendorf, Hamburg, Germany). *S. epidermidis* RP62A was originally isolated from a patient with intravascular catheter-associated sepsis by Muller *et al.* (33). Staphylococci were grown in TSB (trypticase soy broth) (Difco, Detroit, MI, USA) or in RPMI 1640 (Biowest, Nuaillé, France) supplemented with 2 mM glutamine (Lonza Srl, Bergamo, Italy) at 37°C for 16-18 h with high shaking (200 rpm). *L. lactis* transformants were grown in M17 medium (Difco) containing 0.5 % glucose and 10% lactose at 30°C.
Construction of *S. lugdunensis* mutants. *S. lugdunensis* N920143 mutants deficient in individual genes *isdB*, *isdC*, *isdI* and *isdK* and the mutant deficient in the entire *isd* locus were reported earlier (29) (Table 1). Additional mutants deficient in *srtB* and the autolysin *atlI* were created using the thermosensitive vector pIMAY and allelic exchange. A detailed protocol of the procedure is described elsewhere (34). Primers used for the construction of deletion cassettes are summarized in Supplementary Table 1.

Construction and expression of *isdC* in *S. lugdunensis Δisd*. In order to create a *S. lugdunensis* strain expressing IsdC in the absence of any other Isd proteins, the *isdC* gene was cloned into the anhydrotetracycline-inducible vector pRMC2 (35). The gene was PCR amplified with the primers isdC-pRF and isdC-pRR using pNZ8048 *isdC*-LPQTG as the template. *S. lugdunensis Δisd* is deficient in sortase B which will prevent cell wall sorting of the wild type IsdC protein harboring a NPQTS motif. The IsdC-LPQTG protein will be sorted by sortase A.

The *isdC*-LPQTG gene was cloned into pRMC2 using sequence and ligase independent cloning (SLIC) (36,37). This method involves PCR amplification of the vector backbone and of the insert. Amplification of the vector backbone generates a linear product with blunt ends at the site required for the cloning. Primers incorporate short identical DNA sequences in vector and insert. Treatment of both PCR products with T4 polymerase (3’-5’ exonuclease activity) creates single stranded, 5’-overhangs in both vector and insert. The DNA fragments are assembled in vitro (without ligation) and used to transform *E. coli*.

Primers pRA and pRB were used to amplify the pRMC2 backbone (Supplementary Table 1). Identical sequences (20-25 nucleotides) were integrated in the primers for the amplification of the insert (isdC-pRF / isdC-pRR, Supplementary Table 1). 10 ng of plasmid DNA was used as template for the amplification of the plasmid backbone with Phusion polymerase (Finnzymes, Keilaranta 16 A, 02150 Espoo, Finland). The PCR products were purified and the vector product was digested with DpnI to remove methylated template DNA. 1 μg of vector and insert DNA was
treated with T4 polymerase in a final volume of 40 μl of NEB Buffer 2 (New England Biolabs, Ipswich, MA, USA) with highly purified BSA (100 μg/ml) (New England Biolabs), 5 mM dithiothreitol (DTT), 200 mM urea and 3 units T4 DNA polymerase (New England Biolabs), incubated for 20 min at 23°C and the reaction stopped by the addition of 25 mM EDTA and subsequent incubation for 20 min at 73°C.

5 μl of vector DNA and 5 μl of insert DNA were mixed and the single-stranded overhangs were allowed to anneal. The tube was placed in a PCR machine for 10 min at 65°C followed by a slow decrease in temperature from 65°C to 25°C with a 1 min hold for each degree. 2.5 μl of the reaction was used to transform E. coli to isolate pRMC2 isdC-LPQTG which was confirmed by DNA sequencing. The plasmid was transformed into S. lugdunensis Δisd (34). Induction of IsdC-LPXTG expression from pRMC2 in S. lugdunensis Δisd was carried out by adding anhydrotetracycline (0.125 μg/ml) (Sigma) to exponential growing cultures.

Construction of Lactococcus lactis expressing IsdC. To express IsdC in the surrogate host L. lactis, the isdC gene was amplified (IsdC-F / IsdC-R) and cloned into the nisin inducible expression vector pNZ8048 (38). L. lactis was transformed with the recombinant plasmid as described earlier (39).

IsdC possesses a NPQTS sorting signal at its C-terminus and is therefore anchored to the cell wall by the transpeptidase sortase B which is encoded within the isd operon. L. lactis does not encode sortase B which will prevent sorting of IsdC to the cell wall. To allow sorting, pNZ8048-isdC was isolated and DNA encoding the NPQTS signal was exchanged to LPQTG using 5’-phosphorylated primers (LPXTG-A/LPXTG-B) and inverse PCR. The primers allowed the amplification of the entire plasmid and introduced the required nucleotide substitutions in the cell wall anchoring region of isdC. The PCR product was treated with T4 ligase to allow circularization of the plasmid and transformed into L. lactis. The resulting plasmid (pNZ8048isdC-LPQTG) was confirmed by DNA sequencing. Primers are summarized in Supplementary Table 1. Induction of IsdC-LPXTG
expression from pNZ8048 in *L. lactis* was carried out by adding nisin (0.4 ng/ml) to exponential growing cultures.

**Biofilm formation.** Overnight cultures of staphylococci were diluted 1:200 in TSB containing 0.3% glucose (TSB0.3%glucose) or RPMI supplemented with 0.3% glucose (RPMI0.3%glucose) and 2 mM glutamine. Aliquots (200 μl) of the diluted bacterial suspensions were added to 96-well flat-bottom sterile polystyrene microplates (Costar. Corning, NY, NY) and incubated statically for 24 h at 37°C. Biofilm formation was detected by the method of Christensen et al. (40). Briefly, biofilms formed on the plates were gently washed twice with phosphate-buffered saline (PBS) (137 mM NaCl, 2.7 mM KCl, 4.3 mM Na2HPO4, pH 7.4) to remove planktonic and loosely adhering bacteria. Adherent cells were fixed with 96% ethanol for 10 min and stained with 0.1% crystal violet for 15 min and, after several washings, the wells were air dried. For a quantitative estimation of biofilm density, bound crystal violet was solubilised with 10% glacial acetic acid and the absorbance of the solubilised dye was read at 595 nm in a microplate reader (model 680) (Bio-Rad Laboratories, Inc., Hercules, CA). To test the role of iron on biofilm formation, *S. lugdunensis* N920143, N940025, N940113 and N940135 were cultured in RPMI 0.3%glucose / 2 mM glutamine supplemented with 1000 μM FeCl3 in 96-well flat-bottom sterile polystyrene microplates, and treated as described above. Iron-depleted growth medium was obtained by treatment of RPMI with divalent metal chelator Chelex 100 following the manufacturer’s instructions (Bio-Rad).

**Enzymatic and chemical treatment of biofilms.** Chemical and enzymatic treatments of biofilms were carried out as described previously (41, 42). Briefly, the biofilms grown in microtiter plates were rinsed with 200 μl 0.9% NaCl and then treated for 2 h at 37 °C with 100 μl of 10 mM sodium metaperiodate (Sigma, St. Louis MO) in 50 mM sodium acetate buffer, pH 4.5. Alternatively, biofilms were incubated with 100 μl proteinase K (Sigma) at 1 mg/ml in 20 mM Tris buffer
containing 100 mM NaCl, pH 7.5 or 100 μl DNase I (Sigma) at 2 mg/ml in PBS. Enzymes or sodium metaperiodate were replaced with the appropriate amounts of buffer in the controls. To rule out the possibility that DNase I could be contaminated with proteases, the enzyme was incubated with albumin for 2 h and the mixture subjected to SDS-PAGE: in these conditions no difference in the electrophoretic mobility of DNase I-treated and untreated samples of albumin was observed.

Expression and purification of recombinant proteins. Recombinant His-tagged proteins were expressed and purified by Ni\(^{2+}\) chelate chromatography as described previously (29) (Table 1). *E. coli* strain TOPP3 (Stratagene, La Jolla, CA), used for the expression of recombinant His-tagged proteins, was grown in Luria-Bertani (LB) broth (Difco) supplemented with ampicillin (100 μg/ml) (Sigma) at 37°C for 18 h with shaking (150 rpm). Overnight cultures were diluted 1:100 in LB medium and grown at 37°C, with shaking, until the OD\(_{600}\)nm reached 0.5–0.6. Expression was induced by adding isopropyl-thio-β-D-galactoside (Inalco, Milan, Italy) to a final concentration of 1 mM. Bacteria were harvested by centrifugation at 1700xg for 20 min, and lysed by passage through a French press. The cell debris was removed by centrifugation (20000xg), and the filtered supernatant (0.45 μm membrane) was applied to a 1 ml Ni\(^{2+}\)-Sepharose His Trap HP column (GE Healthcare, Buckinghamshire, UK). Fusion proteins were eluted with twenty column volumes of 0.00-500 mM imidazole (Sigma) gradient in 20 mM sodium phosphate, 0.5 M NaCl buffer, pH 7.4. Fractions corresponding to the recombinant protein were pooled and extensively dialyzed against PBS. Protein concentrations were determined with a bicinchoninic acid protein assay kit (Pierce, Rockford, IL, USA).

Primary attachment assay. The attachment assay was performed as reported by Geoghegan *et al.* (9). Briefly, bacteria were grown overnight in RPMI/2 mM glutamine, diluted in RPMI\(_{0.3%}\)glucose/2 mM glutamine, and approximately 300 CFU in 100 μl was spread on the base of empty petri dishes. Dishes were incubated upright at 37°C for 30 min, washed three times with 5 ml of sterile PBS, and
covered with TSB agar. Bacterial plate counts were run in parallel, and the percent attachment was calculated. Each experiment was repeated three times. Statistical significance was determined with Student’s t test, using Graph Pad software.

**Aggregation assays.** The aggregation assay was based on the method described by Geoghegan *et al.* (9). Bacteria were grown overnight in RPMI/2mM glutamine and diluted in RPMI 0.3% glucose /2 mM glutamine, to an optical density at 600 nm (OD600) of 1.0. Tubes were incubated statically at 37°C up to 24 h. One ml of broth was removed from the top of the tube at the indicated times, and the OD600 was measured. The remaining culture was vortexed to resuspend the cells, and the OD600 was measured again. The percent aggregation was calculated using the following formula: 100 x [(OD600 of vortexed sample - OD600 before vortexing) / (OD600 of vortexed sample)]. Statistical significance was determined with Student’s t test, using Graph Pad software.

**Gel filtration chromatography.** The size exclusion chromatography experiment was performed using a Superose 12 HR 10/30 column (GE Healthcare) connected to an AKTA design™ chromatography system (GE Healthcare). 100 μl of 100 μM IsdC was loaded onto a gel filtration column equilibrated in PBS with or without 100 μM FeCl3 and eluted with one column volume (24 ml) at a flow rate of 0.5 ml/min. Recorded data were analyzed using UNICORN 5.10 control software (GE Healthcare). The Mr of IsdC was determined from the calibration curve (plot of Kav versus log Mr) once its Kav value was calculated from the measured elution volume.

**Preparation of bacterial lysates.** The lysates from *S. lugdunensis* and *L. lactis* strains were prepared as previously described (29) with minor modifications. Briefly, bacterial cultures were harvested by centrifugation, washed in PBS and adjusted to an optical density at 600 nm of 10. A 1 ml portion of the bacterial suspension was pelleted and resuspended in 250 μl of digestion buffer (50 mM Tris-HCl, 20 mM MgCl2, 30% (w/v) raffinose pH 7.5), containing complete mini protease...
inhibitors (Roche). Cell wall proteins were solubilised by digestion with lysostaphin (500 μg/ml) (S. lugdunensis) or a combination of mutanolysin (1000 U/ml) and lysozyme (900 μg/ml) (L. lactis) at 37° C for 30 or 15 min, respectively. Protoplasts were harvested by centrifugation (5000xg, 15 min), and the supernatants were subjected to SDS-PAGE/Western blotting.

**Western immunoblotting.** Whole cell lysates of transformants of *L. lactis* were subjected to 12.5% polyacrylamide gel electrophoresis and then electroblotted onto a nitrocellulose membrane (GE HealthCare). The membrane was treated with a solution containing 5% (w/v) dried milk in PBS, washed and incubated with anti-IsdC rabbit IgG for 1 h at 22°C. Following additional washings with 0.5% (v/v) Tween 20 in PBS, the membrane was incubated for 1 h with HRP-conjugated goat anti-rabbit IgG. The membrane was treated with ECL detection reagents 1 and 2 as recommended by the manufacturer (GE HealthCare) and exposed to an X-ray film for 30-60 sec.

**Attachment of *L. lactis* transformants to Isd proteins.** Isd proteins (1 μg/well) were coated in microtiter wells in bicarbonate buffer overnight. *L. lactis* (pNZ8048isdC) and *L. lactis* (pNZ8048) (5x10^8 cells/well) were added to the wells and incubated for 1 h at 22°C. After extensive washing with PBS adhering cells were fixed with 25% formaldehyde (Sigma), stained with 2.5% crystal violet and the A_595 measured.

**Statistical methods.** Continuous data were expressed as means and SD. Two-group comparisons were performed by student’s *t* test. One-way ANOVA followed by Bonferroni’s post-hoc tests was exploited for comparison of three or more groups. Analyses were performed using Graph Pad Prism 4.0 (Graph Pad Software Inc., San Diego, CA, USA). Two-tailed *P* values < 0.05 were considered statistically significant.
RESULTS

*S. lugdunensis* clinical isolates form proteinaceous biofilm in iron-restricted conditions

It was reported previously that *S. lugdunensis* strains growing in rich broth formed biofilm that was predominantly proteinaceous in that preformed biofilm could be disrupted by proteases but not by periodate (24). In order to investigate the nature of *S. lugdunensis* biofilm in more detail we tested a panel of clinical isolates along with *S. aureus* and *S. epidermidis* controls where the composition of the biofilm matrix is known to be composed predominantly of protein (V329, 5179R) or polysaccharide (SA113, RP62A) growing in iron replete TSB or in iron-deficient RPMI. The controls formed protein or polysaccharide dependent biofilms as previously reported.

All *S. lugdunensis* strains tested formed biofilm when growing in both media. Glucose concentrations above 0.25% supported higher levels of biofilm formation compared to lower concentrations (p <0.001, data not shown). We confirmed the previous report that biofilms formed by *S. lugdunensis* growing in TSB-glucose were susceptible to detachment by proteinase K but not sodium metaperiodate (Fig. 1). We also found that biofilms formed by *S. lugdunensis* growing in RPMI were susceptible to protease and not to periodate. Furthermore, DNase caused significant detachment of *S. lugdunensis* biofilm formed under both growth conditions. DNase also detached the control protein-dependent biofilm but not that involving a polysaccharide matrix (Fig. 1).

Involvement of the IsdC protein in *S. lugdunensis* biofilm.

To determine if *S. lugdunensis* biofilm formed during growth in RPMI was influenced by the availability of iron, three strains were tested in RPMI supplemented with 1 mM FeCl₃. Interestingly, the addition of iron reduced the biofilm density suggesting the involvement of proteins whose expression was regulated by iron (Fig. 2). *S. lugdunensis* is the only species of coagulase-negative staphylococcus that harbours an iron regulated surface determinant (Isd) locus.
It is only expressed under iron-limited conditions and is responsible for the acquisition of iron from haemoglobin and heme \textit{in vivo} (28, 29). To determine if the \textit{isd} locus is involved in biofilm formation, \textit{S. lugdunensis} N920143 wild type and a mutant where the entire \textit{isd} locus is deleted (\textit{Δisd}) were compared. A schematic representation of the \textit{isd} locus and mutations are shown in Fig. 3. The level of biofilm formed by the \textit{isd} mutant was the same as that formed by the wild-type strain in the presence of FeCl$_3$. This indicates that one or more proteins expressed by the \textit{isd} locus are involved in biofilm formation in iron limited conditions (Fig. 4). The \textit{S. lugdunensis} \textit{isd} locus expresses four cell envelope-associated proteins IsdC, IsdB, IsdJ and IsdK (29). Mutants lacking each of the Isd proteins were tested for biofilm formation in RPMI. Notably, only the IsdC mutant was defective and showed the same low level of biofilm formation as the wild-type supplemented with 1 mM FeCl$_3$ or the \textit{Δisd} mutant. To exclude the possibility that the reduction in biofilm formation was due to a difference in growth in RPMI the growth curve for the wild type and each of the mutants were compared and found to be superimposable. A revertant strain where the \textit{isdC} gene had been restored to wild-type expressed biofilm normally indicating that the \textit{isdC} mutation and not a secondary mutation is responsible.

IsdC is known to be surface-exposed and anchored to cell wall peptidoglycan by sortase B (29). Consistent with this observation, a sortase B mutant (\textit{ΔsrtB}) gene formed biofilm at a level comparable to that of the \textit{isdC} mutant. In contrast, an \textit{attl} mutant expressed biofilm at a level comparable to the wild type strain. Furthermore, the \textit{Δisd} mutant (lacking the entire \textit{isd} locus), carrying plasmid pRMC2 bearing an \textit{isdC} gene engineered to express IsdC containing a sortase A recognition sequence (pRMC2\textit{isdC}-LPXTG), expressed a similar level of biofilm to that of \textit{S. lugdunensis} N920143 wild type. This shows that IsdC alone is necessary and sufficient to promote biofilm formation in \textit{S. lugdunensis}. (Fig. 4).

To examine further the effect of iron on biofilm development, \textit{S. lugdunensis} N920143 was grown in Chelex 100-pretreated RPMI supplemented with various amounts of FeCl$_3$ and tested for
biofilm formation. The levels of IsdC detected by Western immunoblotting were the same under depleted and low-iron conditions (from 0 to 50 μM) and virtually missing in 250-2000 μM FeCl₃ (Fig. 5). A high level of biofilm formation was observed with bacteria growing in the low-iron environment (1-50 μM), whereas significantly reduced biofilm was detected either in FeCl₃-depleted conditions or in the presence of high iron concentrations (≥ 250 μM). In conclusion, for the IsdC-dependent biofilm formation critical concentrations of FeCl₃ are required.

We also tested the effect of pH on IsdC expression by planktonic cells of S. lugdunensis N920143 and on biofilm formation. The same level of IsdC was detected by Western immunoblotting of lysates of staphylococcal cells grown at pH 5-8 (Fig. 6, panel A), while the biofilm level was significantly enhanced when the pH of the medium was 6.0 or 6.5 compared to 5.5, 7.0, 7.5 or 8.0 (Fig. 6, panel B). A similar trend was found when the effect of pH on biofilm formation by the isdC mutant was examined. These conditions also promote protein-dependent biofilm formation in S. aureus (12). Together these data indicate that different pHs do not affect IsdC expression but somehow influence the levels of biofilm formation.

Role of IsdC in attachment and accumulation during biofilm formation.

Biofilm formation in static growth conditions occurs in several stages beginning with attachment of individual cells to a surface followed by growth and the accumulation of cells in a multi-layered complex held together by proteins or polysaccharide. In our experiments bacteria must attach to the surface of a polystyrene dish. To test attachment, bacterial cells were incubated in dishes, washed and immobilized in molten agar before incubation overnight to allow colonies to develop. For the wild type strain N920143 73% of the added cells attached whereas for the isdC mutant only 36.7% attached. The reverted control strain isdCr adhered at a similar level as the wild type. To investigate if IsdC contributes to cell-cell aggregation and thus possibly to biofilm accumulation, suspensions of RPMI-grown bacteria were allowed to settle for up to 24h. The
density of cells at the top of the suspension was measured periodically. The density of the *isdC*
mutant was significantly lower than the wild-type or the restored mutant after 6 and 8 h but the
densities were the same after 24h. This indicates that IsdC is involved both in the initial attachment
of bacterial cells and also in the accumulation phase.

**IsdC forms homodimers.**

To test the hypothesis that the ability of IsdC to promote the accumulation phase of biofilm
formation by iron-starved bacteria could be due to a homophilic interaction between IsdC proteins
attached to adjacent cells, the ability of the protein to dimerize in solution was examined. A
preparation of purified recombinant IsdC was subjected to gel filtration chromatography on a
Superose 12 HR column equilibrated with or without FeCl$_3$ (Fig. 7). Two major peaks were eluted
from the column equilibrated with 100 μM FeCl$_3$, an early peak corresponding to IsdC dimers (Mr
= 59 kDa) and a retarded peak with a molecular mass of 29.5 kDa which corresponds to the IsdC
monomer. Conversely, a single peak of 29.5 kDa was obtained from the column equilibrated and
eluted with PBS only. Thus an equilibrium exists between the monomeric and dimeric forms of
IsdC in solution.

To determine if IsdC protein that is attached to the surface of a bacterial cell could interact
with other IsdC molecules, the *isdC* gene was cloned into the nisin-inducible *Lactococcus lactis*
expression vector pNZ8048. The sortase B cleavage motif NPQTS was changed to LPQTG in order
to provide a substrate for sortase A in *L. lactis*. After induction, expression of IsdC was detected in
the cell wall fraction derived from protoplasts of *L. lactis* by Western immunoblotting (Fig 8, panel
A) indicating that IsdC was sorted to the cell wall. When the ability of *L. lactis* expressing IsdC to
adhere to immobilized Isd proteins was tested, *L. lactis* (pNZ8048*isdC*) adhered to immobilized
IsdC at a four-fold higher level than the control strain bearing the empty vector, but not
significantly to IsdB, IsdJ or IsdK. A slight promotion of adherence to IsdB was noted but this was
not statistically significant. These data are consistent with the conclusion that IsdC homophilic interactions contribute to the accumulation phase of *S. lugdunensis* biofilm formation under iron-restricted conditions (Fig 8, panel B). As reported for biofilm formation by *S. lugdunensis* N920143 (Fig. 6, panel B), the interaction of *L. lactis* (pNZ8048isdC) with immobilized IsdC was enhanced at pH values of 6.0 and 6.5 and decreased at pH 5.0, 5.5, 7.0, 7.5 and 8.0 (Fig. 8, panel C). Consistent with low iron-induced biofilm formation (Fig. 5), we also found that attachment of *L. lactis* pNZ8048isdC to surface-coated IsdC requires the presence of FeCl$_3$ ($\geq$ 1μM). Significantly reduced levels of IsdC homophilic interactions were observed when bacterial attachment was performed in iron-depleted conditions (Fig. 8, panel D).

**Inhibition by recombinant Isd proteins and antibodies.**

The soluble recombinant Isd proteins or specific antisera raised against each of the Isd proteins were added to the RPMI$_{0.3\%\text{glucose}/2\text{mM glutamine}}$ medium at the same time as the inoculum and incubated for 24 h before the density of biofilm formed was measured. Only IsdC protein or anti-IsdC serum caused a significant (~ 35%) reduction in biofilm formation (Fig. 9). Although complete inhibition was not achieved we can be confident of the specificity of the effect which is consistent with an important role in biofilm formation.

**DISCUSSION**

The identification of surface proteins that are able to induce biofilm development in the absence of polysaccharides is one of the most unexpected results obtained recently in the field of staphylococcal biofilm studies. The existence of alternative mechanisms promoting biofilm development suggests that different staphylococcal strains might involve a specific component to
form a particular type of biofilm matrix. *S. aureus* can opt for at least five different surface proteins, Bap (8), protein A (11), SasG (9), SasC (10) and FnBPs (12, 13) to achieve protein-mediated biofilm. Moreover, functional amyloids composed of phenol soluble modulins (PSMs) stabilize *S. aureus* biofilms (43). A similar proteinaceous biofilm has been demonstrated with *S. epidermidis* based on the expression of proteins such as Aap (14) or SesC (16, 44). Furthermore, the same staphylococcal strain can switch between a proteinaceous or polysaccharide-based biofilm depending on the growth conditions (45).

Investigating biofilm production by a collection of strains of *S. lugdunensis*, we confirmed the proteinaceous nature of biofilms in iron-rich (TSB) medium (24) and demonstrated for the first time that biofilm formed in iron-restricted conditions (RPMI) was also protein-dependent. We found that proteins expressed by the *isd* locus are involved in biofilm development and we identified the IsdC protein as the main factor responsible of cell-to-cell interactions and biofilm formation in iron-restricted conditions. IsdC contributed both to the attachment of cells to a polystyrene surface and to the accumulation phase. Thus, IsdC can be added to the growing list of proteinaceous factors involved in staphylococcal biofilm formation.

Surprisingly, for maximal biofilm formation, both IsdC expression and iron-restricted conditions were required, whereas *S. lugdunensis* grown in completely iron-depleted conditions (chelex-treated RPMI) or in RPMI supplemented with excess amounts of FeCl₃ (≥ 250 μM) developed a significantly reduced biofilm. Thus iron plays a dual role in IsdC-dependent biofilm formation. Iron restriction is required to induce IsdC expression and a low level of iron is required to support the formation of biofilm while high levels (>250 μM) are inhibitory. Given that *S. lugdunensis* biofilm is dependent on IsdC and that IsdC has been demonstrated to bind heme (29), it is also possible that IsdC-associated heme is involved in biofilm formation, although it cannot be excluded that the requirement for iron could be unrelated to heme binding. The fact that IsdC is only expressed under iron-restricted conditions and the ability of *S. lugdunensis* strains also to form
proteinaceous biofilm under iron-replete conditions implies that biofilm formation in TSB is
promoted by proteinaceous factors distinct from those involved in RPMI growth medium. It would
be interesting to determine if the protein(s) involved are anchored by sortase A.

The observation that IsdC confers primary attachment to polystyrene and production of cell aggregates in the accumulation phase of biofilm, parallels the behaviour observed for Bap (8) and SasC (10) of *S. aureus*. The aggregation activity observed for IsdC is also promoted by staphylococcal proteins such as Aap (14), SasG (9), and FnBPs (13).

The structural characterization of Aap and SasG in biofilm development has been recently defined (46, 47). Aap and SasG comprise an N-terminal A region and repeated B domains towards the C-terminus. Proteolytic removal of the A domain of Aap or limited partial cleavage within the B domains of SasG is required for biofilm formation. This presumably reduces steric hindrance by the N-terminal region and allows the B repeats from opposing cells to engage. The length of the B region was shown to be critical for biofilm development because only SasG (or Aap) constructs with five or more B-repeats supported biofilm formation, while constructs with four or fewer B-repeats did not. Structural studies revealed that Aap B-repeats of the appropriate length adopt an elongated, rope-like structure coordinated by zinc ions and wrap around one another in an antiparallel fashion to form bundles of fibres that establish homophilic interactions and have the potential to interconnect neighbour cells (46, 47).

Homophilic protein-protein interactions may be also operational in the biofilm promoted by IsdC. Indeed IsdC forms dimers in solution in the presence of FeCl₃ and immobilized recombinant IsdC specifically interacts with IsdC expressed on the surface of *L. lactis*. Moreover, as reported for biofilm formation by *S. lugdunensis*, this interaction is dependent on the pH, suggesting that a pH-induced changes in the conformation of IsdC contribute to both biofilm formation and homophilic interactions.
Due to the absence of tandem repeats in IsdC, alternative structures/mechanisms should be envisaged in IsdC-mediated biofilm formation. For example, the N-terminal subdomain N2N3 of the A region of the fibronectin-binding protein FnBPA and the N-terminal domain of SasC, but not their repetitive regions, mediate biofilm formation in *S. aureus* (13, 10). Thus, there are two different scenarios, a first one involving Aap and SasG which requires a Zn-dependent interaction between repeats and a second one where a direct Zn-dependent binding of the A domain of FnBPA on one cell to the A domain on another one is needed. The possibility that the A domain of FnBPA bound to a different ligand on the adjacent cell surface cannot be excluded. However, this was not backed up by *in vitro* studies with recombinant proteins as in the case of IsdC.

Although *in vitro* evidence indicates the involvement of homophilic interactions, it cannot be excluded that IsdC *in vivo* might establish heterophilic interactions with proteinaceous or non-proteinaceous components expressed on the surface or localized in the extracellular matrix of staphylococcal cells. Moreover, due to the reduced but not abolished ability of the Δisd mutant of *S. lugdunensis* strains to develop biofilms and the partial inhibition by soluble IsdC protein or anti-IsdC serum, it is plausible that additional, yet undefined factors may contribute. The residual biofilm in the Δisd mutant and ΔisdC mutant was the same and appears to be proteinaceous (data not shown). It would be interesting to determine whether sortase A anchored proteins (other than IsdB or IsdJ) could play a role in this process. This would require construction of a double Δisd srtA mutant.

Several questions on the role of IsdC in biofilm formation remain unanswered. For example, it should be of interest to investigate the molecular details of IsdC-mediated biofilm formation and whether the putative heme-binding NEAT domain is involved. This would require the production of recombinant mutant proteins and assessment of their ability to form dimers. Mutant strains with an altered potential to bind heme could also be constructed and concurrent evaluation of their capability to form biofilm performed. In connection with this, it could be also interesting to
investigate the reasons why only IsdC, but not IsdB, IsdJ or IsdK, is involved in biofilm formation, despite similarity between their NEAT domains (29).

*S. lugdunensis* is an important cause of infections associated with indwelling medical devices. In general, device-related staphylococcal infections, including those associated with *S. lugdunensis* infections, are difficult to treat because staphylococci form biofilm and are protected from antimicrobial agents and the host’s immune system. As a consequence, device removal can be required to resolve the infection. Thus, it will be of interest to determine if IsdC-promoted biofilm contributes to virulence in animal models of foreign body infection. This is particularly important if one considers that due to the very low concentrations of free iron in the body fluids, IsdC should be expressed on the surface of *S. lugdunensis* cells and could play a role in biofilm formation under these conditions. This study provides initial clues about a protein that could be targeted to prevent developing of biofilm in conditions resembling those present in the body fluids.

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**FIGURE LEGENDS**

FIG. 1 Characterization of *S. lugdunensis* biofilm matrix composition. Biofilms were grown in sterile 96-well polystyrene plates in TSB (panels A, B and C) or RPMI (Panels D, E and F) supplemented with 0.3% glucose, for 24 h at 37°C and the adherent cells were treated with proteinase K (panels A and D), sodium metaperiodate (panels B and E) or DNAase I (panels C and F). After washing, cells were stained with 0.1% crystal violet and the absorbance measured at 595
nm. Bars are the averages of measurements taken from four wells. Error bars represent the standard deviations. Controls for protein-dependent biofilm (S. aureus V329 and S. epidermidis 5179R) and PIA-dependent biofilm (S. aureus SA113 and S. epidermidis RP62a) were included in experiments to test sensitivity to proteinase K, DNAase and sodium metaperiodate. This experiment was performed three times with similar results.

FIG. 2 Biofilm formation by S. lugdunensis clinical isolates grown in RPMI with or without 100μM FeCl₃. Bacteria were grown for 24 hrs in RPMI and diluted 1:200 in RPMI-glucose (0.3%, w/v) in the presence or absence of 1 mM FeCl₃. Diluted bacteria (200 μl) were added to sterile 96-well polystyrene plates and statically incubated at 37°C for 24 h. Wells were washed three times with phosphate-buffered saline and fixed with 96% ethanol. Adherent cells were stained with 0.1% (w/v) crystal violet and the absorbance measured at 595 nm. Bars are the averages of measurements taken from four wells. Error bars represent the standard deviations. This experiment was performed three times with similar results. P values were determined using the two-tailed, two-sample unequal variance Student’s t test in Graph Pad.

FIG. 3 Schematic diagram of the isd locus and mutations. The open boxes denote individual genes and the arrows the direction of their transcription. Encoded NEAT motifs are shown as small black boxes. Angled dashes indicate the beginning and the end of the isd locus. Mutations are indicated by horizontal bi-headed arrows.

FIG. 4 Effect of isd locus deletion on biofilm formation by S. lugdunensis in iron-limiting conditions. S. lugdunensis N920143 WT and its deletion mutants covering the entire isd locus or single isd genes were grown overnight in RPMI. Experimental conditions for biofilm formation by
the strains and biofilm detection are identical to those reported in Fig 2. The effect of 1 mM FeCl₃ on biofilm formation is also reported. This experiment was performed four times with similar results. P values were determined using the two-tailed, two-sample unequal variance Student’s t test in Graph Pad.

FIG. 5 Iron-dependent biofilm formation by *S. lugdunensis* N920143. Bacteria grown overnight in Chelex 100-treated RPMI were diluted in the same medium supplemented with increasing amounts of FeCl₃ (final concentrations from 0 to 2000 μM), added to sterile 96-well polystyrene plates and statically incubated at 37°C for 24 h. Experimental conditions for biofilm detection are identical to those reported in Fig 2. The upper part of the figure shows the detection by Western blotting of IsdC protein in the lysates obtained from cultures grown overnight planctonically in different FeCl₃ concentrations. This experiment was performed three times with similar results. P values were determined using the two-tailed, two-sample unequal variance Student’s t test in Graph Pad.

FIG. 6 Effect of pH on the IsdC expression and biofilm formation by *S. lugdunensis* N920143. Panel A, *S. lugdunensis* N920143 was grown overnight in RPMI adjusted to the indicated pH and the corresponding lysates subjected to Western immunoblotting. IsdC protein transblotted onto nitrocellulose filter was detected with rabbit anti-IsdC IgG, followed by HRP-conjugated secondary goat anti-rabbit IgG. Panel B, *S. lugdunensis* N920143 and its isdC deletion mutant were grown overnight in RPMI, diluted 1:200 in the same medium adjusted to different pHs, added to sterile 96-well polystyrene plates and statically incubated at 37°C for 24 h. Experimental conditions for biofilm detection are identical to those reported in Fig 2. This experiment was
performed three times with similar results. P values were determined using the two-tailed, two-sample unequal variance Student’s t test in Graph Pad.

FIG. 7 Analysis of IsdC interactions by size exclusion chromatography. 100 μl of IsdC at a concentration of 100 μM was loaded onto a gel filtration column of Superose 12 HR and eluted with one column volume (Vt= 24 ml) of PBS with or without 100 μM FeCl₃ at a flow rate of 0.5 ml/min. Molecular weight determination was carried out by comparing elution volumes of the two peaks marked as IsdC1 and IsdC2, with the values obtained for several known calibration standards. The Mr of IsdC1 or IsdC2 was determined from the calibration curve (plot of Kav versus log Mr) once their Kav value were calculated from the measured elution volumes [Kav= (elution volume - void volume)/(column volume - void volume)] (inset).

FIG. 8. Homophilic and heterophilic interactions of IsdC expressed by L. lactis with surface-coated Isd proteins. Panel A, validation of IsdC expression by L. lactis. Expression of IsdC by L. lactis (pNZ8084 isdC) and L. lactis (pNZ8084) was demonstrated by Western immunoblotting. IsdC protein in the solubilised cell wall lysates was detected with rabbit anti-IsdC IgG, followed by HRP-conjugated secondary goat anti-rabbit IgG. Panel B, adhesion of L. lactis expressing IsdC to Isd proteins. Microtiter wells were coated with recombinant IsdB, IsdC, IsdK and IsdJ in a bicarbonate buffer overnight (1 μg/well). L. lactis (pNZ8084isdC) and L. lactis (pNZ8084) (5x10⁸ cells/well) were added to the wells and the mixtures incubated for 1 h. After washing with PBS, adherent cells were fixed, stained with crystal violet and measured at 595 nm. Panel C, effect of pH on the adherence of L. lactis (pNZ8084isdC) and L. lactis (pNZ8084) to immobilized IsdC protein. Bacterial cells (5x10⁸ cells/well) suspended in growth medium were adjusted to the indicated pHs and added to the wells. Panel D, effect of iron concentrations on the adherence of L.
lactis. Bacterial cells (5x10^8 cells/well) suspended in growth medium were supplemented with increasing amounts of FeCl₃ and added to the wells. Incubation of the mixtures as described in panels C and D was carried out for 1 h and detection of attached bacteria was performed as reported in panel B. Results shown in panels B, C and D are the mean values of triplicate samples. Error bars show the standard deviation. P values were determined using the two-tailed, two-sample unequal variance Student’s t test in Graph Pad.

FIG. 9. Effect of soluble Isd proteins or anti-Isd IgG on the biofilm formation by S. lugdunensis 920143. Overnight cultures of S. lugdunensis 920143 grown in RPMI were diluted into fresh RPMI-glucose and 200 μl of mixtures were allowed to form biofilm for 24 h in the presence of 5 μM Isd proteins (panel A) or 2 μg/ml of pre-immune or immune IgG raised against each Isd protein (panel B). Adherent cells were stained with crystal violet and the absorbance measured at 595 nm. Bars are the averages of measurements taken from four wells. Biofilm values are expressed as percentage of biofilm level obtained from cultures grown in the absence of any potential inhibitor. Error bars represent the standard deviations. This experiment was performed three times with similar results. P values were determined using the two-tailed, two-sample unequal variance Student’s t test in Graph Pad.

REFERENCES


\[\Delta isd\] – deletion of the entire locus
* = p < 0.001
**A**

Image showing a gel with protein bands labeled with kDa values: 100, 70, 55, 35, and 25.

**B**

Bar graph showing A595nm at different pH levels (5.5, 6, 6.5, 7, 7.5, 8) for WT and ΔisdC strains. The graph includes error bars and significance markers (* for p<0.01, ** for p<0.001).
**A**

![Bar graph showing % Biofilm formation for different treatments.

- Control
- + rIsdC
- + rIsdJ
- + rIsdB
- + rIsdK

* * p<0.01

**B**

![Bar graph showing % Biofilm Formation for different treatments.

- Control
- anti-IsdC IgG
- anti-IsdJ IgG
- anti-IsdB IgG
- anti-IsdK IgG
- Preimmune IgG

* * p<0.01
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<thead>
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<th>Strain or plasmid</th>
<th>Relevant characteristics(^a)</th>
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<td>This study</td>
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\(^a\) Ap\(^\prime\) ampicillin resistance, Cm\(^\prime\) chloramphenicol resistance