Transcriptional downregulation of agr expression in Staphylococcus aureus during growth in human serum can be overcome by constitutively active mutant forms of the sensor kinase AgrC

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Abstract

The temporal and cell density-dependent regulation of expression of virtually all the Staphylococcus aureus virulon is under the control of the agr (accessory gene regulatory) operon. The expression of the agr operon is subject to transcriptional regulation by the AgrA/C two-component response regulator/sensor kinase pair. During bacteraemia, a frequent syndrome caused by methicillin-resistant S. aureus (MRSA), the transcriptional downregulation of agr expression has been attributed to the sequestration of the quorum-signalling molecule auto-inducing peptide (AIP) by the human serum component apolipoprotein B as part of an innate immune response to infection. However, it is not known whether transcriptional downregulation of agr expression during growth in human serum is additionally subjected to regulation by transcription regulatory proteins that either directly or indirectly affect transcription from the agr operon promoters. Here, using chromosomal fluorescence reporters of agr expression in S. aureus, we show that the transcriptional downregulation of agr expression in human serum can be overcome using constitutive active mutant forms of AgrC. Therefore, it seems that the sequestration of the AIP is likely to be the only mechanism by which the host innate immune response limits agr expression at the transcriptional level to maintain the host–pathogen balance towards a noninvasive outcome.

Introduction

The cell density-dependent modulation of gene expression often manifests itself at the transcriptional level and involves a two-component signal transduction system. The cell density or quorum-signalling molecule, the autoinducer, is sensed by the receptor kinase, which modulates the activity of the response regulator, leading to alterations in the transcription patterns of target genes (Stock et al., 2000; West & Stock, 2001) Gram-positive bacteria use secreted peptides as autoinducers (Novick & Geisinger, 2008). The Gram-positive bacterial pathogen Staphylococcus aureus causes a wide variety of life-threatening invasive infections in humans. The pathogenic success of S. aureus can be attributed to the diverse array of virulence factors, involving a large number of cell-surface bound proteins (e.g. adhesins, fibrinogen/fibronectin binding proteins) that are expressed during colonisation of the host, and secreted proteins (e.g. haemolysins, proteases, lipases) that are required for acute infections (Dunman et al., 2001; Cheung et al., 2011).

An auto-inducing quorum-sensing system encoded within the agr operon in part coordinates the phenotypic change in S. aureus during infection from adhesive and colonising to tissue damaging and invasive (Novick & Geisinger, 2008). Therefore, the coordinated regulation of agr operon expression is an important criterion for the pathogenic success of S. aureus, at least during the acute stage of infection (Cheung et al., 2011). However, it should be noted that the role of agr after infection is established is less clear and mutations that inactivate agr are sometimes found in clinical isolates (Traber et al., 2008; Shopsin et al., 2010).

The agr operon, conserved in all Staphylococcus isolates examined thus far (Novick & Geisinger, 2008; Wuster & Babu, 2008), is expressed from divergent promoters, P2 and P3; where P2 encodes a quorum-sensing system and
P3 encodes a pleiotropic effector of the virulon (Fig. 1a; Koenig et al., 2004). Consistent with the central role of the agr operon in regulating the expression of the S. aureus virulon, a vast array of transcription regulatory proteins either directly or indirectly control transcription from P2 and P3 (Novick & Geisinger, 2008; Reyes et al., 2011). In addition, AgrA is also subjected to post-translational modification under specific growth conditions, and agrA mRNA is also post-transcriptionally regulated in some S. aureus strains (Sun et al., 2012; Kaito et al., 2013).

Recent evidence indicates that elements of host innate immunity regulate the agr operon-mediated phenotypic changes in S. aureus during infection, from adhesive and colonising to tissue damaging and invasive, thereby contributing to the maintenance of the host–pathogen balance in favour of a noninvasive outcome (Rothfork et al., 2004; Peterson et al., 2008; Malachowa et al., 2011; Hall et al., 2013). The sequestration of AIP by apolipoprotein B (which is present in blood that extravasates to site of acute infection) represents a primary host innate immune mechanism to downregulate agr operon expression at the transcriptional level to limit invasive infections caused by S. aureus (Peterson et al., 2008; Hall et al., 2013). The sequestration of AIP by apolipoprotein B renders the AIP unavailable to interact with its membrane-bound receptor AgrC and thereby compromises the efficiency by which AgrC activates its cognate response regulator AgrA. This subsequently leads to downregulation of transcription from P2 and P3 by AgrA (Peterson et al., 2008; Hall et al., 2013). AIP can also be inactivated by oxidants, for example NADPH (Rothfork et al., 2004). However, it is not known whether downregulation of agr expression is limited to the sequestration of AIP by apolipoprotein B or whether it additionally involves regulation of P2 and P3 activity by the other transcription regulatory proteins that affect transcription from P2 and P3 or through post-translational regulation rendering AgrA unavailable or inactive for the activation of transcription from P2 and P3. Here, we present results from experiments in which we have addressed this issue in the context of the community-associated methicillin-resistant S. aureus (CA-MRSA) strain USA300 LAC* (hereafter referred to as USA300). The USA300 lineage is the most frequent cause of CA-MRSA bacteraemia in the United States and causes the most invasive forms of infection (Klevens et al., 2007).

**Materials and methods**

**Bacterial strains, plasmids and DNA manipulation**

The bacterial strains and plasmids used in this study are listed in Table 1. *Escherichia coli* and *S. aureus* were grown in Luria broth (LB) and tryptic soy broth (TSB), respectively. The sequences of primers used for DNA manipulation and cloning are listed in Supporting Information, Table S1. Human serum (derived from male AB plasma, sterile-filtered) was purchased from Sigma. Further details of reagents, bacterial growth conditions and DNA manipulation techniques can be found in the Supporting Information, Data S1 (see also Fig. 1b and 1c for information regarding construction of transcription reporters).

**Blood agar haemolysis assay**

Strains were grown for 16 h in TSB culture, then 5 μL aliquots were subcultured onto tryptic soy agar containing 5% sheep’s blood and left to grow for 16 h at 37 °C.

**Real-time quantitative reverse transcription PCR (qRT-PCR)**

Details of RNA extraction and cDNA synthesis can be found in the Data S1. qRT-PCR was performed using primers and Taqman probes corresponding to *hld* (delta toxin), *agrA* and *gyrB* (gyrase B) with QPCR core kit, no ROX (Eurogentec) according to the manufacturer’s instructions. Reactions were performed in an ABI PRISM 7700.

**Bacterial growth and GFP expression assays**

These were conducted as described previously for *E. coli* (Shadrin et al., 2012). Simultaneous growth (OD₆₀₀) and GFP fluorescence measurements (with excitation and emission filters of 485 and 520 nm respectively) were performed in 96-well black microtitre plates with clear bottoms (Corning) in a POLARstar Omega multiwell plate reader (BMG Labtech). Three biological replicates (each defined as a single colony) were performed for each growth curve. Further details can be found in the Data S1.

**Western blotting to determine intracellular levels of AgrA**

Details of the experimental set-up and sample preparation can be found in the Data S1 (see also Fig. 3). Western blotting was performed using polyclonal antibodies against AgrA (raised in rabbits against untagged recombinant AgrA by Eurogentec; used at 1:500 dilution) and anti-rabbit–horseradish peroxidase-conjugated antibodies (Dakocytomation; used at 1:10 000 dilution) as primary and secondary antibodies, respectively, following standard laboratory protocols.
**Fig. 1.** (a) Schematic representation of the *agr* operon organisation and regulation in *Staphylococcus aureus*. (b) Schematic representation showing single-site chromosomal integration of GFP transcriptional fusion reporters for P2 and P3. (c) Sequence of P2 and P3 promoter regions used to generate the *gfp* transcriptional fusions for P2 and P3 (the consensus /C0 10 and /C0 35 sites are outlined, the transcription start sites are shown with arrows and the RBS is underlined. (d) Sheep blood agar haemolysis assay with *S. aureus* USA300WT (1), USA300 *agrA*::Tn [agrA transposon mutant from NARSA library (Fey et al., 2013)] (2), USA300 *agr IR P3-GFP* (3) and USA300 *agr IR P2-GFP* (4). (e) Graphs showing relative hld, agrA and gyrB mRNA levels in 16 h cultures of *S. aureus* USA300WT and USA300 *agr IR P3-GFP* as determined by Taqman qRT-PCR. Values are shown for each gene with respect to USA300WT levels. (f) Graph showing growth curves (OD600) of *S. aureus* USA300WT, USA300 *agr IR P2-GFP* and USA300 *agr IR P3-GFP* strains grown in TSB. (g) Graphs showing GFP expression [as GFP fluorescence units (GFP-FU)] and GFP-FU as a function of growth (OD600) over time for *S. aureus* USA300 *agr IR P2-GFP* and USA300 *agr IR P3-GFP* strains grown in TSB. (h) As in (g) but with *S. aureus* SH1000 *agr IR P2-GFP*, SH1000 *agr IR P3-GFP*, SH1001 *agr IR P2-GFP* and SH1001 *agr IR P3-GFP* Strains grown in TSB. Data for (d–h) were obtained from three biological replicates.
Table 1. Bacterial strains and plasmids used

| Strain/plasmid | Genotype/characteristics* | Source/reference |
|----------------|---------------------------|------------------|
| **Strains**    |                           |                  |
| *E. coli*      |                           | Stratagene       |
| XL1-blue       | Efficient cloning strain  |                  |
| S. aureus      |                           |                  |
| RN4220         | NCTC8225-4, restriction mutant | Peng et al. (1988) |
| SH1000         | Functional rsbU derivative of 8325-4 rsbU | Horsburgh et al. (2002) |
| SH1001         | SH1000 agr::tet           | Horsburgh et al. (2002) |
| NE1532         | USA300 LAC agrA transposon mutant, here called USA300agrA::Tn | NARS library Fey et al. (2013) |
| USA300 LAC*    | Erm-sensitive CA-MRSA LAC* strain; hereafter called USA300<sub>WT</sub> | Boles et al. (2010) |
| SH1000<sup>agr</sup> IR P3-GFP | pCL55<sub>agr</sub> IR P3-GFP integrated into SH1000 | This study |
| SH1000<sup>agr</sup> IR P2-GFP | pCL55<sub>agr</sub> IR P2-GFP integrated into SH1000 | This study |
| SH1001<sup>agr</sup> IR P3-GFP | pCL55<sub>agr</sub> IR P3-GFP integrated into SH1001 | This study |
| SH1001<sup>agr</sup> IR P2-GFP | pCL55<sub>agr</sub> IR P2-GFP integrated into SH1001 | This study |
| USA300<sup>agr</sup> IR P3-GFP | pCL55<sub>agr</sub> IR P3-GFP integrated into USA300<sub>WT</sub> | This study |
| USA300<sup>agr</sup> IR P2-GFP | pCL55<sub>agr</sub> IR P2-GFP integrated into USA300<sub>WT</sub> | This study |
| **Plasmids**   |                           |                  |
| pCL55         | Single-site integration vector for *S. aureus*. Amp<sup>β</sup> in E. coli, Cm<sup>β</sup> in *S. aureus* | Lee et al. (1991) |
| pCN34         | Shuttle vector for Gram-positive bacteria. Amp<sup>β</sup> in E. coli, Kan<sup>β</sup> in *S. aureus* | Charpentier et al. (2004) |
| pCL55<sup>agr</sup> IR P3-GFP | pCL55 carrying P3GFP | This study |
| pCL55<sup>agr</sup> IR P2-GFP | pCL55 carrying P2GFP | This study |
| pCN34<sup>agr</sup> WT | pCN34 carrying P2<sup>agrC</sup> WT | This study |
| pCN34<sup>agr</sup> M234L | pCN34 carrying P2<sup>agrC</sup> M234L | This study |
| pCN34<sup>agr</sup> R238H | pCN34 carrying P2<sup>agrC</sup> R238H | This study |
| pCN34<sup>agr</sup> Q305H | pCN34 carrying P2<sup>agrC</sup> Q305H | This study |

*<sup>Amp</sup>, ampicillin; <sup>Cm</sup>, chloramphenicol; <sup>tet</sup>, tetracycline; <sup>Kan</sup>, kanamycin; <sup>Erm</sup>, erythromycin.

**Results**

Development of a near real-time fluorescence-based system to monitor transcription from *agr* operon promoters P2 and P3

To study how transcription from P2 and P3 is affected during growth in serum, two *S. aureus* USA300 reporter strains were created by 'ectopically' placing transcriptional fusions of P2 or P3 to GFP at the *geh* locus on the *S. aureus* chromosome using the integration plasmid pCL55 whilst leaving the native *agr* operon intact (Fig. 1b; Lee et al., 1991). Each transcriptional fusion construct consisted of either the P2 or the P3 promoter sequence and the *agr* IR sequence up to, but not including, the consensus —35 promoter element of the divergent promoter upstream of the gene encoding GFP with an optimal RBS sequence (Fig. 1c).

As we wished to conduct downstream assays with the USA300 reporter strains in the absence of antibiotic chloramphenicol (pCL55 carries the *cat* gene that confers resistance to chloramphenicol), we initially determined the stability of the integrated reporter constructs after 24 h growth in TSB (without chloramphenicol), bacteria containing P2 and P3 reporter constructs were cultured on TSA in the absence of antibiotic and 28 colonies from each plate (n = 3) were examined for chloramphenicol resistance on TSA containing chloramphenicol. In each colony, chloramphenicol resistance was maintained and all colonies emitted green fluorescence when exposed to blue light (Supporting Information, Fig. S1). We therefore conclude that the reporter constructs are extremely stable under the experimental conditions used. We next tested whether the presence of an additional copy of the *agr* IR sequence in the *S. aureus* USA300<sup>agr</sup> IR P2-GFP reporter strains affected the *agr* operon expression (e.g. by titrating away regulatory interactions at the native *agr* IR) and the growth of *S. aureus* USA300 strain in any way: as *agr* dysfunction is associated with β-haemolytic activity, we used a blood agar plate haemolysis assay to establish that the *S. aureus* USA300<sup>agr</sup> IR P3-GFP reporter strains are not compromised for β-haemolysis (Fig. 1d). Further, we also isolated total RNA from *S. aureus* USA300<sup>agr</sup> IR P2-GFP and USA300<sup>agr</sup> IR P3-GFP reporter strains and determined the mRNA levels of *agrA* (reports native P2 transcription) and *hld* (reports native P3 transcription) relative to that of *gyrB* (reports
transcription of the constitutively expressed gene encoding gyrase) by qRT-PCR. As shown in Fig. 1e (and data not shown), and consistent with the results from Fig. 1d, no obvious differences in the levels of agrA and hld mRNA were detected between S. aureus USA300WT and reporter USA300agr IR P3-GFP strains. In addition, the growth rate of S. aureus USA300WT, USA300agr IR P2-GFP and USA300agr IR P3-GFP strains did not significantly differ in TSB (Fig. 1f). Having established that the presence of an additional copy of the agr IR sequence with either the P2 or P3 did not detectably affect native agr expression, agr-mediated downstream effects and the growth characteristics of S. aureus USA300WT, we measured the GFP expression as a function of growth in S. aureus USA300agr IR P2-GFP and USA300agr IR P3-GFP reporter strains. The GFP signal, originating from the ectopically placed P2 and P3, increases concomitantly with the increase in cell density in both reporter strains grown in TSB (Fig. 1g). The rate of GFP expression from P2 and P3 is very similar, thus indicating that transcription from the ectopically placed P2 and P3 occurs at a similar rate under our experimental conditions. However, in contrast, a previous study using plasmid-based GFP reporters of P2 and P3 activity in S. aureus strain SH1000 (Reyes et al., 2011) showed that GFP expressed from P2 accumulates at a much faster rate than GFP expressed from P2, thus implying that P3 is a stronger promoter than P2. To rule out any strain-specific factors and/or unregulated transcription from the P2 and P3 transcriptional fusion constructs at the geh locus accounting for similar activity of P2 and P3 under our experimental conditions, we placed transcriptional fusions of P2 or P3 to GFP at the geh locus on the chromosome of S. aureus strain SH1000 and the agr operon-deficient mutant strain SH1001 (Horsburgh et al., 2002; Table 1). As with the USA300 reporter strains, the GFP signal originating from P2 and P3 increased concomitantly with an increase in cell density and at a similar in the SH1000 reporter strain (Fig. 1h). However, as expected, no transcription form P2 and P3 is detected in the SH1001 reporter strain. Therefore, even though we are unable to differentiate the specific activities of P2 and P3 during growth in the context of our chromosomal reporter strains and under our experimental conditions (but see later, Fig. 2c), we are confident that the GFP signal originating from the ectopically placed P2 and P3 faithfully mirrors the regulatory events that occur at the native agr operon.

**Downregulation of transcription from agr operon promoters in response to human serum**

To examine the effect of human serum on P2 and P3 transcription in the S. aureus USA300agr IR P2-GFP and USA300agr IR P3-GFP reporter strains, bacteria were grown in TSB supplemented with a range of concentrations of human serum [5-50% (v/v) final concentration] and measured GFP expression as a function of growth. As expected, the reduction in GFP expression positively correlated with the increase in human serum concentration in the growth medium (Fig. 2a). As 25% (v/v) human serum did not detectably affect growth yet still caused a significant downregulation of transcription from P2 and P3 activity (Fig. 2a), we used this concentration of human serum in further experiments.

To test whether the transcriptional downregulation P2 and P3 transcription can be ‘induced’ by human serum, bacteria were grown to late-exponential phase in TSB to allow GFP expression (indicating transcription from P2 and P3) to reach a approximately one-fifth of the maximal level, then added either 25% (v/v) human serum or 25% (v/v) fresh TSB media to the cultures. As shown in Fig. 2b, whereas the growth rate of the S. aureus USA300agr IR P2-GFP and USA300agr IR P3-GFP reporter strains remains unaffected, GFP expression, indicating transcription from P2 and P3, became detectably reduced. Having established that the addition of human serum to exponentially growing S. aureus resulted in downregulation of agr operon expression, the time difference between the responses of downregulation of P2 and P3 transcription to human serum was investigated. As P3 transcription is dependent upon AgrA and agrA is transcribed from the P2 promoter, we expected that the downregulation of P3 activity would follow downregulation of P2 activity. In agreement with this, the results in Fig. 2c show that the transcriptional activity of P2 drops after c. 20 min, whereas P3 drops c. 50 min after addition of human serum to the cultures. Overall, the results in Figs 1 and 2 are consistent with the view that sequestration of AIP by apolipoprotein B in human serum leads to the downregulation of transcription from agr operon promoters P2 and P3 and unambiguously demonstrate that S. aureus USA300agr IR P2-GFP and USA300agr IR P3-GFP strains report transcription regulatory events at the native P2 and P3 promoters of the agr operon in a sensitive and faithful manner (Peterson et al., 2008; Hall et al., 2013).

**Intracellular levels of AgrA remain unchanged upon addition of human serum to exponentially growing S. aureus cells**

As AgrA can be subjected to post-transcriptional and post-translational regulation (Sun et al., 2012; Kaito et al., 2013), we considered the possibility that AgrA is subjected to regulation that renders it unavailable or inactive for the activation of transcription from P2 and P3 during growth in human serum. We therefore
Fig. 2. (a) Graphs showing GFP expression [as GFP fluorescence units (GFP-FU)] as a function of growth (OD<sub>600</sub>) over time for <em>Staphylococcus aureus</em> USA300<sub>agr</sub> IR P2-GFP and USA300<sub>agr</sub> IR P3-GFP strains grown in TSB media containing 0–50% (v/v) human serum. (b) Graph showing GFP expression (as GFP fluorescence units (GFP-FU)) as a function of growth (OD<sub>600</sub>) over time for USA300<sub>agr</sub> IR P2-GFP and USA300<sub>agr</sub> IR P3-GFP strains grown in TSB media upon addition of 25% (v/v) human serum to late-exponentially growing cells (indicated with a green arrow). (c) As in (b), however, readings for OD<sub>600</sub> and GFP expression values were taken every 30 min before the addition of human serum and every 10 min thereafter. Data for graphs (a–c) were obtained from three biological replicates.
compared intracellular AgrA levels in *S. aureus* USA300WT cells grown in the presence and absence of human serum. The bacterial cells were grown to late-exponential phase in TSB, and downregulation of agr operon expression was induced by adding human serum. At the same time, 10 μg mL⁻¹ of tetracycline was added to the cells to prevent *de novo* translation of AgrA (see schematic Fig. 3). As the results in Fig. 2c indicated that downregulation of transcription from P3 occurred after c. 50 min after the addition of human serum, cell samples were taken for analysis by Western blotting using polyclonal anti-AgrA antibodies immediately before (T0) and 2 (T2) and 16 (T16) hours after the addition of human serum. We reasoned that if the addition of human serum to exponentially growing *S. aureus* USA300WT made AgrA unavailable (e.g. by reduction of its half-life), then the intracellular levels of AgrA should be detectably lower in cells to which human serum and tetracycline are added than in cells to which only tetracycline is added. Initially, we confirmed that our anti-AgrA antibody specifically recognises AgrA using whole-cell extracts prepared from *S. aureus* SH1000 strain and the AgrA-deficient mutant *S. aureus* SH1001 (Horsburgh et al., 2002; Fig. S2). As expected, in the absence of tetracycline and human serum, AgrA is detected in the T0 sample obtained from late-exponentially growing cells (Fig. 3, lane 1); a moderate increase in AgrA levels is seen in the T2 and T16 samples as AgrA is already expressed as maximal level in late-exponentially growing cells at T0 (Fig. 3, compare lane 1 with lanes 2 and 3). Also as expected, the addition of tetracycline leads to no significant increase in AgrA levels in the T2 and T16 sample compared with the T0 sample (Fig. 3, compare lane 1 with lanes 4 and 5). Similarly, the addition of human serum to exponentially growing *S. aureus* USA300WT cells did not detectably reduce intracellular levels of AgrA in the T2 and T16 sample compared with the T0 sample (compare lane 4 and 5 with lanes 6 and 7). The reduction in AgrA levels observed between T0 and tetracycline-containing samples at T2 and T16 likely reflects the half-life of AgrA in the absence of *de novo* protein synthesis. In summary, we conclude that the half-life of AgrA is not detectably affected by human serum.

Fig. 3. Western blot indicating intracellular AgrA levels in *Staphylococcus aureus* strain USA300WT during growth in TSB medium in the absence and presence of human serum and/or tetracycline. A schematic representation of the experimental steps is shown on the top of the blot (see text for details). Samples of whole-cell extracts were prepared for the analysis immediately before and 2 and 16 h after the addition of human serum (T0, T2 and T16 respectively). Lane 8 contains purified recombinant HMK-tagged AgrA as a positive control marker (Reynolds & Wigneshweraraj, 2011). Data shown are representative from three independent experiments (biological repeats). AgrA signal intensity values were calculated by densitometry using a Typhoon FLA 7000 and indicated relative to the signal intensity at T0.

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**Lane**

| MW (kDa) | 31 | 28 |

**Tet (10 mg/mL)**

- - - + + HMK-AgrA

**Serum (25% (v/v))**

- - + +
Constitutively active mutants of AgrC overcome downregulation of agr expression in human serum

Having shown that intracellular levels of AgrA remain unchanged during growth in human serum, we investigated whether AgrA was subjected to any post-translational modifications that render it inactive for the activation of transcription from P2 and P3 promoters. We reasoned that if sequestration of AIP is the only mechanism by which the agr operon is downregulated during growth in human serum, then a constitutively active form of AgrC (that does not require AIP binding for autophosphorylation) should allow transcription from P2 and P3 to occur and thereby indicating that (i) AgrA is not subjected to any post-translational modification during growth in human serum that renders it inactive and (ii) transcription form P2 and P3 is not subject to an additional level of transcriptional regulation during growth in human serum through the action of other transcription regulatory proteins that affect transcription from P2 and P3. To experimentally test this, we introduced wild-type and previously identified constitutively active mutant forms of agrC into our S. aureus USA300agr IR P3-GFP reporter strain on a low copy number plasmid and under control of the P2 promoter (Geisinger et al., 2009). As the production of AgrC required AgrA to be produced (to activate P2), we first allowed AgrA levels to increase to a moderate level (approximately one-fifth of the maximal level) and then ‘induced’ the downregulation of agr expression by adding human serum to late-exponentially growing bacterial cells. As shown in Fig. 4, GFP expression (indicating P3 transcription) was downregulated upon addition of human serum in the S. aureus USA300agr IR P3-GFP reporter strain containing the empty plasmid or plasmid harbouring the wild-type agrC, whilst

Fig. 4. Graphs showing GFP expression [as GFP fluorescence units (GFP-FU)] as a function of growth (OD600) over time for Staphylococcus aureus USA300agr IR P3-GFP strain containing plasmids encoding for mutant and wild-type forms of AgrC (as indicated) grown in TSB with 25% (v/v) human serum added after cells reached late-exponential growth phase (indicated with a green arrow). Data shown were obtained from three biological replicates.

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the cells containing plasmids encoding constitutively active forms of AgrC had no detectable downregulation of GFP expression upon addition of human serum. This result indicates that transcriptional downregulation of agr expression during growth in human serum is restricted to sequestration of the AIP by apolipoprotein B and thus is not subjected to regulation through transcription regulatory factors that either directly or indirectly affect transcription form P2 and P3.

**Discussion**

Quorum-dependent regulation of virulence gene expression is a common strategy used by bacterial pathogens to ensure that the appropriate set of virulence genes are expressed at the correct time during infection. Thus, sequestration of the quorum-signalling molecule is an effective mechanism used by the host innate defence system to limit the quorum-dependent expression of bacterial virulence genes (Gordon et al., 2013). In the case of S. aureus, the agr operon mediates the quorum-dependent expression of the S. aureus virulon, and the level of agr operon expression is downregulated during growth in human serum, due to the sequestration of the agr quorum-signalling molecule by apolipoprotein B as part of the innate host defence mechanism. In this study, we have shown that transcriptional downregulation of agr expression during growth of S. aureus in human serum is restricted to sequestration of the AIP by apolipoprotein B and thus is not subjected to regulation through transcription regulatory factors that either directly or indirectly affect transcription form P2 and P3 (e.g. by transcription repressors that occlude the RNA polymerase from binding to P2 and P3 promoters or reduce the efficiency by which transcription initiates from P2 and P3). Importantly, we have shown that the transcriptional downregulation of agr expression in human serum can be overcome using constitutive active mutant forms of the quorum sensor AgrC, and therefore, it seems that AgrA is unlikely to be subjected to post-translational modifications that render it unavailable or inactive. Targeting bacterial quorum-sensing systems that are nonessential for growth but essential for virulence has been suggested as a novel strategy to curtail and manage bacterial infections, including those caused by S. aureus, E. coli and Pseudomonas aeruginosa (Gray et al., 2013; Melamed Yerushalmi et al., 2013). A study by Shopsin et al. revealed that the agrAC locus is a ‘hotspot’ for acquiring mutations in clinical isolates of S. aureus (Shopsin et al., 2010). Therefore, it is conceivable that mutations that confer constitutively active phenotypes could occur with increased frequency in the presence of an appropriate selection pressure (such as quorum-sensing inhibitors). Therefore, our results are relevant to the strategies that consider using quorum-sensing inhibitors since mutations that might occur in agrC or agrA through natural selection, and confer the constitutive active phenotype, could limit the effectiveness of such strategies in the long term. In addition, as agr dysfunction has been associated with persistent bacteremia in MRSA (Chong et al., 2013), it is conceivable that downregulation of agr activity, as part of host, the innate immune response could in fact contribute to the formation of staphylococcal persisters leading to chronic infections.

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

Fig. S1. Photograph of a bacterial growth plate (one of three biological replicates) taken on blue light transilluminator to determine the stability of transcriptional reporters used in this study.

Fig S2. Western blot showing specificity of anti-AgrA polyclonal antibody to AgrA.

Table S1. Primers used for cloning, colony PCR, qRT-PCR and site-directed mutagenesis (SDM).

Data S1. Materials and methods.