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A Novel Antimicrobial Coating Represses Biofilm and Virulence-Related Genes in Methicillin-Resistant *Staphylococcus aureus*

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INTRODUCTION

*Methicillin-resistant Staphylococcus aureus* (MRSA) has become an important cause of hospital-acquired infections worldwide. It is one of the most threatening pathogens due to its multi-drug resistance and strong biofilm-forming capacity. Thus, there is an urgent need for novel alternative strategies to combat bacterial infections. Recently, we demonstrated that a novel antimicrobial surface coating, AGXX⃝, consisting of micro-galvanic elements of the two noble metals, silver and ruthenium, surface-conditioned with ascorbic acid, efficiently inhibits MRSA growth. In this study, we demonstrated that the antimicrobial coating caused a significant reduction in biofilm formation (46%) of the clinical MRSA isolate, *S. aureus* 04-02981. To understand the molecular mechanism of the antimicrobial coating, we exposed *S. aureus* 04-02981 for different time-periods to the coating and investigated its molecular response via next-generation RNA-sequencing. A conventional antimicrobial silver coating served as a control. RNA-sequencing demonstrated down-regulation of many biofilm-associated genes and of genes related to virulence of *S. aureus*. The antimicrobial substance also down-regulated the two-component quorum-sensing system agr suggesting that it might interfere with quorum-sensing while diminishing biofilm formation in *S. aureus* 04-02981.

Keywords: antimicrobial surface, MRSA, virulence, biofilm, quorum-sensing, RNA sequencing

Staphylococcus aureus is an opportunist pathogen commonly found in the human respiratory tract, nasal areas and skin. It colonizes the anterior nares of approximately 20–25% of the healthy adult population, while 60% are intermittently colonized (Kluytmans et al., 1997; Ellis et al., 2014). Methicillin-resistant *Staphylococcus aureus* (MRSA) is a crucial human pathogen causing infections ranging from skin and soft tissue infections to fatal sepsis (Marathe et al., 2015). It is one of the leading pathogens that cause nosocomial infections (Paniagua-Contreras et al., 2012; Lister and Horswill, 2014); it is resistant to methicillin and many other antibiotics (Marathe et al., 2015), and it is also known to produce thick biofilm (Paniagua-Contreras et al., 2012; Qin et al., 2014). MRSA was shown to cause catheter-associated and other medical devices-related...
infections (Arcoli et al., 2001; Paniagua-Contreras et al., 2012). Eighty percent of prosthetic infections are caused by Staphylococci (Kirmusaoglu, 2016). Its firm attachment to medical devices and host tissues, and its ability to form robust biofilms makes it a cause of chronic infections (Yarwood et al., 2004). S. aureus biofilms cause numerous infections in which the accessory gene regulator (agr) quorum-sensing system (QS) plays an important role (Yarwood et al., 2004). Around 90% of the infections caused by the bacterium are skin and soft tissue infections, and the agr QS system is associated with these infections (Sully et al., 2014).

Multiple drug resistance combined with a thick biofilm makes the treatment and eradication of S. aureus infections even more difficult. This entails the urge of development of novel antimicrobials, which could also be potential biofilm inhibitors. Virulence factors of S. aureus serve as targets for the newly developed class of biological anti-staphylococcal agents. These targets include, surface bound adhesins, immunoglobulin-binding proteins, surface-associated and secreted proteases, a family of immune-stimulatory exotoxins called 'superantigens' (SAgs), and potent leukocidal toxins (Sause et al., 2015).

Metals like copper and silver have been used as antimicrobials since a long time. The use of copper in human civilization is known since the 5th and 6th millennia B.C. Silver was officially approved for use as an antimicrobial agent in the 20th century (Chopra, 2007; Grass et al., 2011; Schäberle and Hack, 2014; Guridi et al., 2015). Copper and copper alloys have also been used as antimicrobials (Warnes and Keevil, 2013). These metals are known to kill bacteria and fungi by a phenomenon called contact killing (Grass et al., 2011) and can be used to coat medical devices as they inhibit biofilm formation of pathogens (Baker et al., 2010). In the 17th century, silver was described as an essential multipurpose medicinal product and the first scientific documentation of its medical use dates from 1901 (Maillard and Hartemann, 2013). However, in 1975, several patients died from a silver resistant Salmonella Typhimurium isolate in the Massachusetts General Hospital; this was the first report of silver resistant bacteria (Gupta et al., 1999). Excessive use of silver is questioned due to its toxicity to the environment as well as to the human body (Landsdown, 2010). Silver resistance, like antibiotic resistance in bacteria, prompts us to develop new strategies to control bacterial infections. One such novel, broad-spectrum antimicrobial agent is AGXX®.

AGXX® (Largentec GmbH, Berlin, Germany) is a combination of two transition metals, silver and ruthenium which can be galvanically electroplated on various carriers like V2A steel, silver sheets, Polydimethylsiloxane (PDMS), fleece, etc. The coating is conditioned by ascorbic acid and is active against many Gram-positive and Gram-negative bacteria (Guridi et al., 2015). It is not only an efficient antibacterial but also kills yeasts, viruses, and fungi (Landau et al., 2017a,b). The coating was used successfully for the decontamination of industrial cooling and process water (Landau, 2013). As it is only slightly cytotoxic (Bouchard, 2011), it can be incorporated into various medical applications. Although, the exact mode of action of the antimicrobial activity of the coating is not fully understood, it is known that the generation of reactive oxygen species (ROS) plays an important role in making it a potent antimicrobial.

The formation of hydrogen peroxide and hydroxyl radicals has been detected by spectroscopic methods (Clauss-Lendzian et al., 2017). Putative formation of other ROS is under investigation. ROS can damage cellular components, including, DNA, lipids and proteins. Superoxide dismutase and catalase are involved in detoxification of ROS (Paraje, 2011).

In this study, we performed total RNA-sequencing of S. aureus 04-02981 (MRSA) to investigate differential gene expression after different times of exposure of the pathogen to the antimicrobials AGXX® or Ag. Our data demonstrate that AGXX® likely reduces biofilm formation and virulence in S. aureus 04-02981 by interfering with the QS, by down-regulating the expression of toxins like leukocidins (lukE) and gamma-hemolysins (hlgA), and of genes associated with surface adhesins and capsular polysaccharide.

MATERIALS AND METHODS

Preparation of Antimicrobial Metal Sheets

Silver sheets of 0.125 mm thickness were used as a base material to prepare the antimicrobial metal sheets. Both sides of the silver sheets were etched by immersing them in half-concentrated nitric acid, for 60 s. The silver sheets were cleaned with de-ionized water and galvanically plated with a 0.16 µm ruthenium coating on both sides for 40 s. Then, the sheets were cleaned with de-ionized water, conditioned with ascorbic acid, rinsed with de-ionized water and dried with a paper towel. Prior to use, AGXX®, and Ag sheets, used as reference material, were autoclaved at 121°C for 20 min.

Bacterial Strain and Culture Conditions

Staphylococcus aureus 04-02981 (Nuebel et al., 2010) was grown at 37°C in Tryptic Soy Broth [TSB] (Carl Roth GmbH & Co. KG, Karlsruhe, Germany) with constant agitation at 150 rpm or on Tryptic Soy Agar [TSA] (Carl Roth GmbH & Co. KG, Karlsruhe, Germany). Growth inhibition tests on agar surface were performed according to CLSI guidelines for disk diffusion test (Naas et al., 2006). For this assay, 0.25 cm² sheets of Ag and AGXX® were used.

For generation of growth curves, bacteria were pre-cultured overnight, diluted in TSB to an optical density at 600 nm (OD₆₀₀) of 0.05 and incubated for further 8 h either in presence of AGXX® or in the presence of silver (Ag), 24 cm² each in 30 mL medium to obtain a sheet surface to medium volume ratio (A: V) of 0.8. Cultures grown in the absence of a metal sheet served as controls. OD₆₀₀ of the cultures was measured using the Genesys 10S UV-Vis spectrophotometer (Thermo Scientific, China). Colony forming units (CFU) per mL were determined hourly from 0 to 8 h post inoculation. Growth experiments were performed in triplicate with independent biological replicates.

Biofilm Screening Assay

To study the effect of Ag, and AGXX® on biofilm formation of S. aureus 04-02981, the Crystal Violet Assay was performed.
without any metal sheet, in presence of Ag (24 cm² uncoated silver sheet) and in presence of AGXX® (24 cm² silver sheet coated with ruthenium for 40 s). The sheet surface: medium volume ratio (A: V) was 0.8 (24 cm² metal sheet: 30 mL medium). The overnight culture of *S. aureus* 04-02981 was diluted to an initial OD₆₀₀ of 0.05. The culture was incubated at 37°C and 150 rpm for 4 h (mid-exponential phase, OD₆₀₀~1.5). Then, it was transferred to the transparent 96-well plate (Carl Roth GmbH & Co. KG, Karlsruhe, Germany) containing Ag, or AGXX®. The plate was incubated at 37°C for 24 h, then the antimicrobial metal sheets were carefully removed and OD₆₀₀ of the cultures was measured. In addition, at this stage, the CFU per mL of the planktonic cultures and the biofilms in presence as well as in absence of the metal sheets were determined. Means of five values each and two biological replicates are given. The biofilm assay was performed according to Schiwon et al. (2013). *Enterococcus faecalis* 12030, a strong biofilm former was used as a positive control (Huebner et al., 1999), and Tryptic Soy Broth (TSB) as a negative control (Schiwon et al., 2013). Biofilm formation was measured in EnSpire Multimode Plate Reader 2300-0000 (PerkinElmer, Turku, Finland) at 570 nm. Normalized biofilm formation was calculated by dividing the biofilm measurement at OD₅₇₀ by the bacterial growth at OD₆₀₀. Following criteria were used for the interpretation of the results, ODc = negative control; OD ≤ ODc = non-adherent, ODc ≤ OD ≤ (2 × ODc) = weakly adherent, (2 × ODc) < OD ≤ (4 × ODc) = moderately adherent, (4 × ODc) < OD = strongly adherent, as described in Nyenje et al. (2013). Biofilm inhibitory rates of AGXX® and Ag were calculated using the following equation, as described by Qin et al. (2014).

\[
\text{Inhibitory rate (\%) = } \frac{\text{OD}_{570} (\text{Control}) - \text{OD}_{570} (\text{Sample})}{\text{OD}_{570} (\text{Control})} \times 100
\]

Student's *t*-test was used to check if biofilm inhibition was statistically significant, using SigmaPlot version 11.0 (Systat software, Inc., San Jose, CA, United States) (Wass, 2009).

**Spinning Disk Confocal Microscopy**

*Staphylococcus aureus* 04-02981 was grown in TSB overnight at 37°C, 150 rpm, then it was diluted to an OD₆₀₀ of 0.05 and further incubated at 37°C for 4 h (mid-exponential phase, OD₆₀₀ ~1.5). Then, the culture was transferred to a μ-Dish (μ-Dish 35 mm, low, from ibidi GmbH, Martinsried, Germany) containing Ag, or AGXX® (sheet surface: medium volume ratio = 0.8) and incubated at 37°C for 24 h. The culture was removed from the μ-Dish, and the biofilm on the μ-Dish was washed three times with phosphate buffered saline (PBS). The biofilm was stained for 10 min in the dark with Hoechst 33342 (5 µg/mL) and propidium iodide (1 µg/mL) (Thermo Fisher, Eugene, OR, United States). The staining solution was then replaced with 50% glycerol to prevent movement of bacteria during imaging. Imaging was performed with a Nikon TiE-based Visitron spinning disk confocal microscope using a 100× NA1.45 objective. Fluorescent dyes were excited using 405 nm (Hoechst 33342) and 561 nm (propidium iodide) laser lines and fluorescent emission captured through appropriate filters onto an iXon888 EMCCD detector (Andor, Belfast, United Kingdom). Images were subsequently analyzed using Fiji (ImageJ) version 3.2.0.2.

**Metal Stress and RNA Extraction**

Overnight cultures of *S. aureus* 04-02981 were diluted as described above and grown until mid-exponential growth phase (4 h post dilution, OD₆₀₀~1.5). The cultures were then subjected to metal stress by exposure to AGXX® or Ag sheets (sheet-surface to medium-volume ratio of 0.8) followed by further incubation for 6, 12, 24, 80, and 120 min at 37°C with constant agitation at 150 rpm. As a control, no metal sheet was added to the culture. Cells from 30 mL culture were harvested by centrifugation for 1 min at 10,000 rpm and 4°C in a Heraeus Multifuge X3R Centrifuge (Thermo Electron LED GmbH, Osterode am Harz, Germany). Cell pellets were immediately frozen in liquid nitrogen and stored at ~80°C or directly used for RNA extraction using the ZR Fungal/Bacterial RNA MiniPrep™ Kit (ZymoResearch, Freiburg, Germany) following the manufacturer’s instructions. To recover total RNA including small RNAs, 1.5 volumes of absolute ethanol were added in step 5. Finally, total RNA was eluted with 50 µl DNase- and RNase-free water and stored at ~80°C. RNA quantity and quality were assessed with a NanoDrop 2000c UV-Vis Spectrophotometer (Thermo Scientific, Osterode am Harz, Germany) as well as on bleach agarose gels. Residual contaminating DNA was eliminated with TURBO DNA-free™ Kit Ambion (Life Technologies, Darmstadt, Germany).

**RNA Sequencing**

Total RNA sequencing was done by PrimBio Research Institute, Exton, PA, United States. The protocol was performed in five steps; RNA removal was done using the Ribo-Zero RNA Removal Kit (Bacteria) (Illumina, Cat# MRZMB126), followed by library preparation, and templating, enrichment and sequencing.

**RNA-sequencing Data Analysis**

Raw sequencing reads were aligned to the reference genome of *S. aureus* 04-02981, using Bowtie2 (Langmead and Salzberg, 2012) version 2.2.3 with optimal settings for the IonProton™ Sequence. Post-processing of the SAM files into sorted BAM files was carried out with SAMtools (Li et al., 2009, version 1.2-207). The samples AGXX®, and Ag were normalized (AGXX®-Control, Ag-Control) against the control of the respective time-points. Length normalized confidence interval RPMK (=Reads per Kilobase of transcript per Million mapped reads) values were obtained with Cufflinks (Trapnell et al., 2010). Finally, statistical analysis was carried out using the T-REx RNA-Seq analysis pipeline (de Jong et al., 2015). A gene was considered significantly differentially expressed when the fold change was ≥[2.0] and the false discovery rate (FDR) adjusted p-value ≤ 0.05. The data presented in this paper have been deposited at NCBI, and are accessible through GSE103064.

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1[http://www.systatsoftware.com](http://www.systatsoftware.com)

Reverse Transcription Quantitative PCR (RT-qPCR)

To verify the results obtained from RNA-sequencing, RT-qPCR was performed on five genes detected as highly differentially expressed via RNA-seq. To this end, RNA extracted from *S. aureus* 04-02981 cultures exposed to Ag or AGXX<sup>®</sup> for 24, and 80 min, was used. First strand cDNA was synthesized with hexamer primers. cDNA was diluted with DNase- and RNase-free water and amplified in a LightCycler<sup>®</sup>480 II (Roche Diagnostics GmbH, Mannheim, Germany).

The *agrC*, *lukE*, *sdrC*, *srrA*, and *cap5* genes were selected to verify the data obtained through RNA-seq. The gene *gyrB* was used as a control. These genes were amplified using TaqMan chemistry according to the instructions provided in LightCycler<sup>®</sup> 480 Probes Master Kit (Roche Diagnostics). All RT-qPCR reactions were carried out in a total volume of 20 µL. The amplification step was performed with ‘Quantification’ analysis mode at 95°C for 10 s, with a ramp rate of 4.4°C/s, followed by annealing at the respective annealing temperature for 50 s, with a ramp rate of 2.2°C/s and finally an extension at 72°C for 1 s, with a ramp rate of 4.4°C/s. The amplification step was performed 45 times. All primers and probes used in the study are listed in Supplementary Table S1. All RT-qPCR experiments were done in triplicate and each experiment was repeated at least twice. Data were analyzed by LightCycler<sup>®</sup> 480 Software release 1.5.0 by using the ‘Relative Standard Curve’ method; the standard curves were constructed using genomic DNA from *S. aureus* 04-02981. Data represent expression ratios, calculated by normalizing to the *gyrB* gene and relative to the untreated culture of *S. aureus* 04-02981 which served as the calibrator, as described in ‘Guide to performing Relative Quantitation of Gene expression using real time-quantitative PCR’ by Applied Biosystems. Means of five Ct values each were used to calculate the relative expression ratio.

Statistical Analysis

Statistical tests were performed to analyze the significance of the obtained data. Student’s t-test was applied to the normalized target, and normalized control values (normalized concentration). The tests were performed and analyzed using SigmaPlot version 11.0 (Systat software, Inc., San Jose, CA, United States<sup>®</sup>) (Wass, 2009).

RESULTS

**AGXX<sup>®</sup> Inhibits the Growth of *S. aureus* 04-02981**

To analyze the effect of Ag, and AGXX<sup>®</sup> on the growth of *S. aureus* 04-02981, disk diffusion tests with Ag, and AGXX<sup>®</sup> were performed in accordance with NCCLS-CLSI guidelines (Naas et al., 2006). The agar plates were monitored at 24 h intervals for 5 days to check if Ag or AGXX<sup>®</sup> exhibited an inhibitory effect on the pathogen, in the form of a zone of inhibition on the agar plate. The diameter of the inhibition zones was measured in ‘cm.’ The mean diameter of the inhibition zone was calculated to be 1.2 cm for AGXX<sup>®</sup> while no zone of inhibition was observed for Ag.

To verify the inhibitory effect of AGXX<sup>®</sup> on *S. aureus* 04-02981 as demonstrated in the agar diffusion tests, experiments in TSB medium were performed measuring the CFU/mL every hour for a period of 8 h, using the A: V ratio (metal mesh: medium volume) of 0.8, as described in Section “Materials and Methods.” As observed in the disk diffusion assay, Ag did not show a significant inhibitory effect on the growth of *S. aureus* 04-02981 in liquid cultures. In contrast, AGXX<sup>®</sup> had a profound inhibitory effect on this strain. The OD<sub>600</sub> of *S. aureus* 04-02981 in presence of AGXX<sup>®</sup> was very low, (OD<sub>600</sub> AGXX<sup>®</sup> at t8 = 0.149) as compared to Ag (OD<sub>600</sub> Ag at t8 = 3.086) and the control (OD<sub>600</sub> Control at t8 = 3.173) (Supplementary Table S2). The CFU/mL of *S. aureus* 04-02981 grown in the batch culture with AGXX<sup>®</sup> increased from 2.77 × 10<sup>6</sup> in the 1st hour to 3.99 × 10<sup>10</sup> in the 4th hour, but then decreased to 1.08 × 10<sup>7</sup> in the 8th hour. The colony counts of *S. aureus* 04-02981 + AGXX<sup>®</sup> (after 8 h of growth) were much lower than that of the same strain with Ag (1.27 × 10<sup>11</sup>) or without metal amendment (1.73 × 10<sup>11</sup>) (Table 1). These data confirm the antimicrobial effect of AGXX<sup>®</sup> on *S. aureus* 04-02981.

**AGXX<sup>®</sup> Strongly Reduces Biofilm Formation of *S. aureus* 04-02981**

The effect of AGXX<sup>®</sup>, and Ag on biofilm formation of *S. aureus* 04-02981 was analyzed using the Crystal Violet assay. *E. faecalis* 12030, a strong biofilm former served as a positive control (Huebner et al., 1999), and TSB as the negative control (Figure 1A). Figure 1B shows the biofilm formation by *S. aureus* 04-02981 for AGXX<sup>®</sup> (after 8 h of growth) in liquid cultures. In contrast, AGXX<sup>®</sup> had a profound inhibitory effect on this strain. The OD<sub>600</sub> of *S. aureus* 04-02981 in presence of AGXX<sup>®</sup> was very low, (OD<sub>600</sub> AGXX<sup>®</sup> at t8 = 0.149) as compared to Ag (OD<sub>600</sub> Ag at t8 = 3.086) and the control (OD<sub>600</sub> Control at t8 = 3.173) (Supplementary Table S2). The CFU/mL of AGXX<sup>®</sup> is calculated to be 3.99 × 10<sup>10</sup> in the 4th hour, but then decreased to 1.08 × 10<sup>7</sup> in the 8th hour. The colony counts of *S. aureus* 04-02981 + AGXX<sup>®</sup> (after 8 h of growth) were much lower than that of the same strain with Ag (1.27 × 10<sup>11</sup>) or without metal amendment (1.73 × 10<sup>11</sup>) (Table 1). These data confirm the antimicrobial effect of AGXX<sup>®</sup> on *S. aureus* 04-02981.

To determine the bacterial killing activity of AGXX<sup>®</sup> under these conditions (after 24 h of growth, prior to adding crystal violet), we measured the CFU per mL of the planktonic cultures and the biofilms in the presence as well as in absence of the two different metal sheets.

The following values were obtained for the biofilms: For *S. aureus* 04-02981 without metal sheet (control), 2.34 × 10<sup>9</sup> ± 8.49 × 10<sup>7</sup> CFU per mL, for the strain in presence of Ag, 2.13 × 10<sup>8</sup> ± 2.40 × 10<sup>6</sup>, and in presence of AGXX<sup>®</sup>, 1.80 × 10<sup>4</sup> ± 1.41 × 10<sup>3</sup>. When we measured the CFU per mL in the respective planktonic cultures, for the control, 2.55 × 10<sup>8</sup> ± 2.12 × 10<sup>7</sup>, and for the strain in presence of Ag, 2.00 × 10<sup>8</sup> ± 1.41 × 10<sup>7</sup> CFU per mL were obtained. However, no colonies were observed in presence of AGXX<sup>®</sup>. Thus, we conclude that in contrast to Ag, all planktonic bacteria were killed by AGXX<sup>®</sup> and after exposure to AGXX<sup>®</sup>, only a drastically reduced number of bacteria (1.80 × 10<sup>4</sup> CFU per mL) survived in the biofilm in comparison to Ag (2.13 × 10<sup>9</sup> CFU per mL).

In summary, the biofilm formation measures normalized to the bacterial growth show that AGXX<sup>®</sup> reduced biofilm formation of *S. aureus* 04-02981.
The values for 5th hour and 8th hour are bolded because after t = 5h, the CFU values of MRSA + Ag decreased. And until t = 8h, the CFU values for all the three samples (MRSA, MRSA + Ag, and MRSA + AGXX) decreased.

formation of S. aureus 04-02981 by 46%, whereas the inhibitory effect of Ag on biofilm formation was less pronounced (41%).

The strong reduction of biofilm formation by AGXX® was confirmed by Hoechst 33342/propidium iodide staining of biofilms grown for 24 h in presence of AGXX®, Ag and without antimicrobial sheet (Figure 2). The inhibitory effect of Ag was also clearly visible, although it was less distinct.

AGXX® Strongly Induces Stress Response and Represses Pathogenesis in S. aureus 04-02981

The raw RNA sequence data obtained were aligned to the S. aureus 04-02981 genome. High sequencing depth was achieved as a mean value of ~12.4 million reads was obtained. The numbers of reads per sample ranged from ~8.4 million reads (Ag_24) to 175 million reads (Control_120) (Supplementary Table S3 and Supplementary Figure S1). From the data, it is clear that the antimicrobial coating has a strong impact on the transcriptome of S. aureus 04-02981. In total, 2864 genes were differentially expressed in S. aureus 04-02981 on exposure to AGXX® and Ag (Supplementary Table S4). The number of differentially expressed genes in presence of AGXX® or Ag at different time-points is presented in Figure 3.

From Figure 3A, it can be seen that the number of differentially expressed genes at t24, t80, and t120 was quite similar. The maximum impact of AGXX® on the transcriptome of S. aureus 04-02981 was reached already after exposure for 24 min (723 genes up-regulated and 823 genes down-regulated) and remained nearly the same after exposure for 80 min (716 genes up- and 822 genes down-regulated), and 120 min (726 genes up- and 836 genes down-regulated). The lowest number of genes was differentially expressed at t6.

The differentially expressed genes were categorized as per Gene Ontology (GO) using the GSEA_Pro option in the RNA-Seq analysis section in the T-REx RNA-Seq analysis pipeline (de Jong et al., 2015). Several GOs were obtained via
FIGURE 2 | Confocal images of biofilm formation by *S. aureus* 04-02981 (MRSA). The pathogen was grown on sterile cover slips for 24 h with the following conditions: (A) without antimicrobial sheet, (B) with a silver sheet or (C) with an AGXX® sheet. Biofilms were then stained with Hoechst 33342 (to mark out *S. aureus* 04-02981; blue) and propidium iodide (to identify dead cells; red) followed by acquisition of small Z-stack (500 nm spacing) through the biofilms using a confocal microscope. Images show an average of Z-projection (average of 4–5 Z planes containing the biofilm) of the fluorescence signal through the biofilms with the propidium iodide staining shown alone in the images on the right (gray scale images). Scale bars = 10 µm.

GSEA_Pro, namely, oxidoreductase process, lipopolysaccharide synthesis, ATP binding, membrane transport, metabolism, metal binding, pathogenesis, transcription regulation, response to heat shock, iron-siderophore transporter activity, serine protease activity, etc. (Supplementary Table S5). In the GO “lipopolysaccharide synthesis,” the *cap* genes mediating capsular polysaccharide synthesis (*cap5A, capA, and cap8C*) were all down-regulated. Genes (*clpB, ctsR, clpC*, and *groES*) involved in response to heat shock were up-regulated. Among the genes related to virulence (pathogenesis), 10 out of 11 genes were down-regulated, while only one gene was up-regulated at t120 (staphylokinase, a plasminogen activator). Among the responding transcriptional regulator genes, nine were up-regulated and 25 were down-regulated. Figure 4 shows the differential expression of these GOs in *S. aureus* 04-02981 exposed to AGXX®.

*hlgA* (SA2981_RS09385) was the most differentially expressed gene associated with virulence, it was down-regulated at t24 (378 fold), at t80 (192 fold), and at t120 (16 fold). The protein encoded by *hlgA* functions as a two-component toxin along with leukocidins in the lysis of erythrocytes (Gouaux et al., 1997). Among the transcriptional regulators, the gene of the LysR family transcriptional regulator, *lysR* was the most significantly influenced one by AGXX®, being down-regulated about 4700 fold at t80, and about 11,000 fold at t120. One of the LysR family transcriptional regulators, HutR is involved in metabolic processes of *S. aureus* (Ibarra et al., 2013). AGXX® had the highest impact on the expression of *capA*, of all the genes mediating capsular polysaccharide synthesis. *capA* was down-regulated by 329 fold at t80. Among the most differentially expressed genes in response to heat shock was *clpB*. It is a member of the stress-induced multi-chaperone system and works with DnaK, DnaJ, and GrpE in the recovery of the cell from heat-shock damage (Frees et al., 2005). Among the genes in the GO families influenced by AGXX®, only those involved in enterotoxin (SA2981_RS09440), and staphylokinase production were also influenced by Ag, by -533 fold, and -2 fold, respectively, at t80 (Supplementary Table S6). In addition to the GO families, the effect of AGXX®, and Ag on the expression of operons in the pathogen was analyzed using the GSEA_Pro option on the T-REx pipeline. The results are presented in Supplementary Tables S7, S8, respectively.

FIGURE 3 | (A,B) Show the number of differentially expressed genes in *S. aureus* 04-02981 on exposure to AGXX® (A) and Ag (B) for the indicated time-period, compared to control (*S. aureus* 04-02981 grown without a metal sheet).
AGXX\textsuperscript{R} Represses the Expression of Biofilm and Virulence-Associated Genes

We checked the effect of AGXX\textsuperscript{R}, and Ag on the expression of genes associated with biofilm formation and virulence in S. aureus 04-02981. Many genes that are known to be crucial for biofilm formation and virulence were differentially expressed on exposure to AGXX\textsuperscript{R} while Ag had an effect on just a few of them. The genes affected by AGXX\textsuperscript{R} encode virulence factors, methicillin resistance, surface adhesins, capsular polysaccharide, two-component systems, and other biofilm-associated genes, as well as toxins (Table 2).

Upon exposure to AGXX\textsuperscript{R}, the QS system genes agrA, agrB, agrC, and agrD of S. aureus 04-02981 were all down-regulated. In general, the response of S. aureus 04-02981 to AGXX\textsuperscript{R} was clearly visible after 24 min of exposure time. Genes encoding adhesins, isdC, srtB, and sdrC were also down-regulated. The mecA gene was down-regulated at t24. The up-regulation of genes inducing biofilm formation in S. aureus, such as saeR (2.3 fold at t120), icaA (36 fold at t24, 29 fold at t80 and 27 fold at t120), icaB (8 fold at t120) and icaD (55 fold at t12, and 6 fold at t120) was intriguing. The genes icaB, icaA, and icaD are involved in ica-dependent biofilm formation. In addition, other key genes associated with biofilm formation and virulence, such as, codY, srrA, luxS, and genes for toxins like leukocidins, enterotoxins, hemolysins, were all differentially expressed at least at one of the time-points (Figure 5). Description of all locus tags and Gene IDs shown to the right of the heatmap is given in Table 2.

In general, it was observed that AGXX\textsuperscript{R} had a huge impact on the transcriptome of S. aureus 04-02981, in particular at the later time-points 24, 80, and 120 min. In contrast, the effect of Ag was much less pronounced as already visible in the growth kinetics and to a lesser extent in the biofilm assays. Although, quite a number of S. aureus 04-02981 genes were differentially expressed upon exposure to Ag, only very few belong to the group of biofilm or virulence-associated genes. Among those, which were significantly differentially expressed in the presence of Ag, were fmtC, which is associated with methicillin resistance (approximately 3 fold up-regulated at t80; in the presence of AGXX\textsuperscript{R} it was 2 fold up-regulated at t24), transcriptional regulator sarR (approximately 3 fold down-regulated at t24;
not differentially expressed in the presence of AGXX®), the gene of the holin-like protein CidA (approximately 4 fold down-regulated at t24; ∼2 and ∼8 fold up-regulated at t80 and t120, respectively, with AGXX®), the arginine deaminase gene arcA (approximately 6 fold down-regulated at t120 and 4.6 fold down-regulated with AGXX®), the hemolysin II gene (approximately 2 fold down-regulated at t24 and approximately 3 fold down-regulated at t120; ∼3.7 fold up-regulated with AGXX® at t80 and t120) and the gene of the antiholin-like protein lrgA (approximately 6 fold up-regulated at t6 with Ag, in the presence of AGXX®, it was ∼3- to 3.7 fold down-regulated at t24, t80, and t120).

**Validation of RNA-Sequencing Data Using RT-qPCR**

From the RNA-seq data, we observed that AGXX® affected genes encoding two-component systems, surface adhesins, capsular polysaccharides, and toxins. In total, five, highly

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**TABLE 2 | Differential expression of biofilm, and virulence-associated genes in S. aureus 04-02981 on exposure to AGXX®.**

<table>
<thead>
<tr>
<th>Locus tag</th>
<th>Abbreviation</th>
<th>Description</th>
<th>6 min</th>
<th>12 min</th>
<th>24 min</th>
<th>80 min</th>
<th>120 min</th>
</tr>
</thead>
<tbody>
<tr>
<td>SA2981_RS10640</td>
<td>agrD</td>
<td>Accessory gene regulator D</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SA2981_RS10645</td>
<td>*agrC</td>
<td>Histidine kinase of the competence regulon ComD</td>
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<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>SA2981_RS10635</td>
<td>agrB</td>
<td>Accessory gene regulator B</td>
<td>−5.7</td>
<td>−18.3</td>
<td>−40.7</td>
<td>−40.8</td>
<td></td>
</tr>
<tr>
<td>SA2981_RS10650</td>
<td>agrA</td>
<td>Two-component system, LytR family, response regulator AgrA</td>
<td>−2.2</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SA2981_RS05970</td>
<td>PSM-β</td>
<td>Phenol-soluble modulin Beta</td>
<td>−10.5</td>
<td>−22.7</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SA2981_RS05965</td>
<td>PSM-β</td>
<td>Phenol-soluble modulin Beta</td>
<td>−10.2</td>
<td>−12.4</td>
<td></td>
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<tr>
<td>SA2981_RS10825</td>
<td>sigB</td>
<td>RNA polymerase Sigma-B factor</td>
<td>−2.3</td>
<td>−4.9</td>
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</tr>
<tr>
<td>SA2981_RS07880</td>
<td>*smA</td>
<td>DNA-binding response regulator SmA</td>
<td>−4.6</td>
<td>−9.1</td>
<td>−4.6</td>
<td>−5.3</td>
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<tr>
<td>SA2981_RS00190</td>
<td>mecA</td>
<td>mecA-Penicillin- binding Protein 2</td>
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<td>−5.5</td>
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<td></td>
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<tr>
<td>SA2981_RS12040</td>
<td>sarR</td>
<td>Transcriptional regulator SarR</td>
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<td></td>
<td>2.4</td>
<td>3.2</td>
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<tr>
<td>SA2981_RS06390</td>
<td>codY</td>
<td>GTP-sensing transcriptional pleiotropic repressor CodY</td>
<td>−2.3</td>
<td>2.4</td>
<td>3.2</td>
<td></td>
<td></td>
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<tr>
<td>SA2981_RS00550</td>
<td>sarH1</td>
<td>Staphylococcal accessory regulator A</td>
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<td>SA2981_RS03525</td>
<td>FmtA</td>
<td>FmtA protein involved in methicillin resistance</td>
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<tr>
<td>SA2981_RS05940</td>
<td>arcD</td>
<td>Arginine/ornithine antiporter ArcD</td>
<td>2.4</td>
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<td></td>
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<td></td>
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<tr>
<td>SA2981_RS07770</td>
<td>capF</td>
<td>Capsular polysaccharide synthesis enzyme Cap5A</td>
<td>−4.8</td>
<td>−3.1</td>
<td>−2</td>
<td></td>
<td></td>
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<tr>
<td>SA2981_RS05275</td>
<td>sspB</td>
<td>Staphopain B precursor</td>
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<tr>
<td>SA2981_RS13930</td>
<td>cidA</td>
<td>Holin-like protein</td>
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<tr>
<td>SA2981_RS13925</td>
<td>arcA</td>
<td>Arginine deaminase</td>
<td>−2.6</td>
<td>−11.8</td>
<td>−114.7</td>
<td>−20</td>
<td>−4.6</td>
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<tr>
<td>SA2981_RS03620</td>
<td>saeR</td>
<td>two-component system, OmpR family, response regulator SaeR</td>
<td>2.3</td>
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<tr>
<td>SA2981_RS06960</td>
<td>FmtC</td>
<td>Protein involved in methicillin resistance/l-lysine modification of phosphatidylglycerol</td>
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<tr>
<td>SA2981_RS05900</td>
<td>hemolysin II</td>
<td>Alpha-hemolysin precursor</td>
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<td>3.7</td>
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<td>SA2981_RS13335</td>
<td>lrgA</td>
<td>LrgA-like protein</td>
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<td>−3.1</td>
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<tr>
<td>SA2981_RS00745</td>
<td>*capA</td>
<td>Capsular polysaccharide synthesis enzyme Cap5A</td>
<td>−34.9</td>
<td>−76.9</td>
<td>−61.9</td>
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<td>SA2981_RS07850</td>
<td>capB</td>
<td>Tyrosine-protein kinase EpsD/capsular polysaccharide synthesis enzyme</td>
<td>2.4</td>
<td>−4.9</td>
<td>−29</td>
<td>−38.1</td>
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<tr>
<td>SA2981_RS00755</td>
<td>capC</td>
<td>protein-tyrosine phosphatase/capsular polysaccharide synthesis enzyme</td>
<td>2.8</td>
<td>−15</td>
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<td>SA2981_RS13940</td>
<td>aur</td>
<td>Zinc metalloproteinase precursor/auraeolysin</td>
<td>−12.4</td>
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<tr>
<td>SA2981_RS02875</td>
<td>*sdcC</td>
<td>Serine-aspartate repeat-containing protein C</td>
<td>−13.4</td>
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<tr>
<td>SA2981_RS02035</td>
<td>Exotoxin 6</td>
<td>Superantigen-like protein</td>
<td>−8.7</td>
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<td>−12.6</td>
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<tr>
<td>SA2981_RS13920</td>
<td>arcB</td>
<td>Ornithine carbamoyltransferase</td>
<td>−16.3</td>
<td>−26.1</td>
<td>−8.6</td>
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<tr>
<td>SA2981_RS05715</td>
<td>isoC</td>
<td>NPQTN cell wall anchored protein IsoC</td>
<td>−5</td>
<td>−9.1</td>
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<tr>
<td>SA2981_RS05735</td>
<td>srtB</td>
<td>Sortase B</td>
<td>−8.3</td>
<td>−22.4</td>
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<tr>
<td>SA2981_RS14090</td>
<td>icaD</td>
<td>Polysaccharide intercellular adhesin (PA) biosynthesis protein</td>
<td>54.7</td>
<td>5.6</td>
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<td>SA2981_RS14085</td>
<td>icaA</td>
<td>Polysaccharide intercellular adhesin (PA) biosynthesis N-glycosyltransferase</td>
<td>35.7</td>
<td>29.9</td>
<td>26.5</td>
<td></td>
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<tr>
<td>SA2981_RS14095</td>
<td>icaB</td>
<td>Polysaccharide intercellular adhesin (PA) biosynthesis deacetylase</td>
<td>−102.4</td>
<td>7.8</td>
<td></td>
<td></td>
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<tr>
<td>SA2981_RS09385</td>
<td>*lukE</td>
<td>Leukotoxin/leukocidin</td>
<td>−378.9</td>
<td>−192.2</td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

*Genes selected for validation via RT-qPCR.*
FIGURE 5 | Heatmap of differential expression of biofilm, and virulence-associated genes in S. aureus 04-02981. The genes are clustered as indicated by the dendrograms on the left side of the heatmap. Yellow represents genes agrD, agrC, agrB, agrA, and PSM-β, red represents genes sigB, mecA, sarR, codY, and sarH1. Green color is for genes fmtA, arcD, capF, sapb, cidA, pink represents arcA while purple is for saeR, fmtC, hemolysin II, and lmgA genes. Blue represents genes mediating capsular polysaccharide synthesis, namely, capA, capB, capC, sur; *sdrC, exotoxin 5, arcB, isdC, and srtB are shown in orange. Gray represents icaD, and icaA and brown color represents icaB, and *lukE genes. *Indicates genes selected for RT-qPCR.

FIGURE 6 | Differential expression of agrC, lukE, sdrC, srrA, and cap5A in S. aureus 04-02981 on 24-min exposure to Ag or AGXX®©, obtained via RT-qPCR (A). Expression ratio of the genes of interest in S. aureus 04-02981 on exposure relative to control (untreated culture of S. aureus 04-02981) normalized to gyrB. (B) Shows differential expression of agrC, lukE, sdrC, srrA, and cap5A in S. aureus 04-02981 on 24-min exposure to Ag or AGXX®©, obtained via RNA-seq as fold change. Error bars indicate standard deviation. Asterisks indicate p-values showing statistical significance. They were obtained from t-test using SigmaPlot 11.0 (**p < 0.0001, ***p < 0.001, **p < 0.01, *p < 0.05; n.s, not significant).

differentially expressed genes encoding these functions were selected to validate the RNA-seq derived transcriptional response of S. aureus 04-02981 to exposure to Ag or AGXX®©. The validation experiment was performed on RNA extracted from S. aureus 04-02981 cultures exposed for 24, and 80 min to Ag or AGXX® since the selected genes were most differentially expressed at these time-points. The five selected genes were, agrC, and srrA which are part of the two-component systems AgrCA and SrrAB, respectively (Baker et al., 2010; Wu et al., 2015), lukE which encodes a toxin (Liu et al., 2016), sdrC specifying a surface adhesin (Barbu et al., 2014), and cap5A mediating the synthesis of capsular polysaccharides.
(Qin et al., 2014). gyrB was used as the house-keeping gene (Smith et al., 2010; Cheung et al., 2011). Figures 6, 7 show the results of these experiments.

After exposure to AGXX® for 24 min, all five genes were down-regulated both in RNA-seq analysis and in RT-qPCR studies as can be seen in Table 2, and Figure 6. However, after exposure to AGXX® for 80 min, sdrC was down-regulated in RT-qPCR assays but it was not differentially expressed in RNA-seq. All the other genes were down-regulated in both approaches as seen in Table 2 and Figure 7, respectively. On exposure to AGXX® for 24 min, sdrC was the most down-regulated gene followed by cap5A, lukE, srrA, and agrC, whereas after 80 min, agrC was the most down-regulated gene followed by srrA, lukE, cap5A, and sdrC. On exposure to Ag for 24 min, srrA was the most down-regulated gene, whereas agrC was the most up-regulated gene, and after 80 min, cap5A was the most down-regulated gene while sdrC was the only up-regulated gene, as observed in the RT-qPCR experiments.

**DISCUSSION**

Multiple drug resistant, biofilm forming nosocomial pathogens such as MRSA pose a severe threat to public health demanding the development of novel antimicrobials as well as potent biofilm inhibitors. AGXX® is an effective antimicrobial that is active against many Gram-positive and Gram-negative bacteria (Guridi et al., 2015). AGXX® has been demonstrated to kill S. aureus 04-02981 as shown here by disk diffusion assay and growth kinetics experiments. In addition, AGXX® inhibited biofilm formation of S. aureus 04-02981 by ~46%. Moreover, for all time-points examined, the number of differentially expressed S. aureus 04-02981 genes was much higher upon exposure to AGXX® (in total 2391) than to Ag (317). For t120, the time-point showing the highest number of differentially expressed S. aureus 04-02981 genes, 1562 genes were differentially expressed in presence of AGXX®, while only 96 genes were affected by Ag.

Up-regulation of genes of Gene Ontology (GOs) groups “response to heat shock” and “oxidoreductases” involved in oxidative stress response, and down-regulation of genes of GOs “pathogenesis” and “lipopolysaccharide synthesis” involving genes mediating capsular polysaccharide synthesis important for biofilm formation, point to a role of AGXX® as an antimicrobial and potent biofilm inhibitor. Together with results of a recent study where we have shown that the QS system of S. aureus 04-02981, agr was completely repressed after 4 h of exposure to AGXX® (Probst et al., 2016), we propose that AGXX® acts as a potential biofilm inhibitor. In S. aureus, two main mechanisms of biofilm formation are known, namely ica-dependent biofilm formation, which involves the production of polysaccharide intercellular adhesin (PIA), and ica-independent biofilm formation (Kirmusaoglu, 2016). Here we show that, in the presence of AGXX®, icaA, icaD were up-regulated and icaB was down-regulated. icaA and icaD contribute to the production of PIA (polymer). icaA transfers PIA to the cell surface of the bacteria while icaB deacylates PIA by fixing PIA to the outer surface of the bacteria (Kirmusaoglu, 2016). In our study, intercellular adhesion biosynthesis N-deacetylase, icaB gene was down-regulated at t80 by ~100 fold. The structural development of exopolysaccharide-based biofilm requires deacetylation of PIA (Aricola et al., 2015). Since icaB was strongly down-regulated at t80, deacetylation of PIA probably does not occur which would obstruct the development of an exopolysaccharide-based biofilm. Fitzpatrick et al. (2005) showed that biofilm formation was unaffected in an icaA/DBC operon-deleted MRSA strain, while the same mutation in a methicillin sensitive strain of S. aureus (MSSA) impaired biofilm formation, suggesting strain-specificity in ica-dependent biofilm formation.

A two-component system associated with ica-dependent biofilm formation is SrrAB that acts as an autoregulator of biofilm formation. Deletion of srrAB inhibited S. aureus biofilm formation under oxic as well as microaerobic conditions (Wu et al., 2015). In our study, srrA was down-regulated 4 to 5 fold after 24, 80, and 120 min of exposure to AGXX®.

Global regulatory systems such as the agr QS system are among the best-studied factors involved in ica-(PIA) independent biofilm formation. Other proteins involved in such biofilms are SasG, SasC, Protein A, FnbB, FnbA, ATLA or ATLE, SdrG, SdrC, SdrD, biofilm associated protein (Bap) and lipoteichoic acid (Kirmusaoglu, 2016). We observed that two of these genes were down-regulated when AGXX® was present, namely sdrD and sdrC, sdrC was down-regulated 13- to 10 fold at t24 and t120, while sdrD was down-regulated 2 to 3 fold at t24 and t80. Moreover, the expression of lipoteichoic acid synthase, an enzyme responsible for the synthesis of lipoteichoic acid (Karatsa-Dodgson et al., 2010) was down-regulated approximately 4 fold after 24, 80, or 120 min of AGXX® presence. These data suggest that AGXX® might be working in an ica-independent manner to inhibit biofilm formation.

The agr locus contains five genes, agrA, agrB, agrC, agrD, and hld. On exposing S. aureus 04-02981 to AGXX®, only hld was not differentially expressed at any time-point, while all the other four genes were significantly down-regulated. The agr gene cluster regulates the expression of virulence factors such as phenol soluble modulins (PSMs), proteins that are closely associated with human skin and soft tissue infections (SSTIs) (Sully et al., 2014). “AgrD is a precursor peptide of autoinducer peptide (AIP)” (Quave and Horswill, 2014), AgrB is a membrane protease, which is involved in proteolytic processing and export of AgrD. It is also involved in AIP production (Njoroge and Sperandio, 2009; Quave and Horswill, 2014). AgrBD produce and secrete AIPs. AgrC, a sensor histidine kinase is activated when AIPs bind to AgrC. As a consequence, AgrC undergoes phosphorylation to activate Agra, which is a DNA-binding response regulator (Njoroge and Sperandio, 2009). In our study, the agrB gene was the most down-regulated, at t80, and t120 (approximately 41 fold in both cases), while agrA was differentially expressed only at t24 (2 fold down-regulated). At t12, only agrB was differentially expressed, approximately 6 fold down-regulated. None of the agr genes was differentially expressed at 16. PSMs are staphylococcal toxins playing a role in acute infection (Kirmusaoglu, 2016); they are required for maturation and detachment of biofilm (Ma...
et al., 2012). PSMs were also down-regulated in presence of AGXXR by ~10 fold at t80, and by 12 and 23 fold at t120. agr also regulates the expression of sspB which encodes a cysteine protease. sspB is positively associated with biofilm formation (Ma et al., 2012). It was down-regulated by 2.3 fold at t80. Inactivation of the alternative sigma factor SigB decreases biofilm formation in S. aureus (Ma et al., 2012). In presence of AGXXR, sigB was down-regulated 2–5 fold at the longer exposure times (t24, t80, and t120). In summary, down-regulation of all of the genes mentioned in this paragraph will likely reduce biofilm formation by S. aureus.

The two component systems, AgrCA and SaeRS influence biofilm formation in S. aureus, by the production of PSMs and by suppressing the synthesis of extracellular proteases, respectively (Baldry et al., 2016). The extracellular proteases degrade proteins that are important for biofilm formation (Baldry et al., 2016). In S. aureus, the saeRS system regulates the production of many virulence factors such as leukocidins, superantigens, proteases, surface proteins, and hemolysins (Liu et al., 2016). The gene for LukE, which enables S. aureus evasion from phagocytic cells by damaging the phagocytes was strongly down-regulated at t24 (379 fold) and t80 (192 fold). SplA is a serine protease, which is directly controlled by the saeRS system. splA was down-regulated 135 fold after 80 min of AGXXR presence. Mutations in genes for extracellular proteases (splABCDEF) in S. aureus SH1000 induced an increase in extracellular protease activity, which was associated with a reduction in biofilm formation (Chen et al., 2013). These facts taken together with saeRS not being differentially expressed at any time-point in the presence of AGXXR, except for a slight 2.3 fold up-regulation of saeR at t120, might suggest that saeR is not expressed in the mid exponential phase of growth of S. aureus 04-02981.

Capsular polysaccharides are also possible targets of the saeRS system (Liu et al., 2016). They play an important role in the virulence of the organism (Tuchscher et al., 2010). The synthesis of capsular polysaccharides is mediated by the cap5ABCDF genes (Qin et al., 2014). Among these genes, only capG was not differentially expressed, all other genes were significantly down-regulated, especially at t24, t80, and t120, suggesting a role of AGXXR in repression of virulence in S. aureus 04-02981.

Another QS system, which significantly influences biofilm formation and virulence in Staphylococci is the luxS system. luxS impacts biofilm formation in a similar way as agr does, but by regulating different factors. luxS negatively regulates biofilm formation via cell–cell interactions based on autoinducer 2 secretion (Xu et al., 2006). The gene was 2.9 fold up-regulated at t24 in the presence of AGXXR.

In addition, the genes isdC, srtB, sdrC, encoding adhesins, were all down-regulated in the pathogen exposed to AGXXR. Iron regulated surface determinant IsdC is necessary for the primary attachment of S. aureus to surfaces such as polystyrene, as well as for the accumulation phase of biofilm formation; as such, it induces biofilm formation (Missineo et al., 2014). IsdC is anchored to the cell wall by sortase B (Hammer and Skaar, 2011). Serine-aspartate repeat containing protein C precursor (SdrC) assists bacteria in adhering to surfaces and promotes biofilm formation (Barbu et al., 2014). In S. aureus 04-02981 exposed to AGXXR, isdC was down-regulated by 5 and 9 fold at t80 and t120, respectively. The sortase B gene srtB was also down-regulated in cells treated with AGXXR, at t80 (8 fold) and t120 (22 fold). sdrC, too, was down-regulated some 10 to 13 fold at t24 and t120. Thus, we suggest that AGXXR inhibits biofilm formation in S. aureus 04-02981, also by repressing the expression of adhesins.

Reverse transcription quantitative PCR assays were performed on RNA extracted from S. aureus 04-02981 cultures exposed to Ag or AGXXR for 24 min, and 80 min to validate the RNA-seq data. In RT-qPCR, on exposure to AGXXR for 24 min, agrC, sdrC, srrA, and cap5A were statistically significantly down-regulated, whereas the down-regulation of lukE was not statistically
significant. In agreement with these data, the five genes were also significantly down-regulated in RNA-seq. By contrast, none of the five genes was significantly differentially expressed after 24 min in presence of Ag, as determined by RNA-seq, whereas RT-qPCR revealed a statistically significant down-regulation of srA and a statistically significant up-regulation of agrC. The difference in expression of the other three genes lukE, sdrC, and cap5A was statistically not significant. When S. aureus 04-02981 was exposed to AGXX® for 80 min, all the five genes were down-regulated in RT-qPCR. The effect was statistically significant while in RNA-seq all genes were significantly down-regulated except sdrC. On exposure to Ag for 80 min, only sdrC was non-statistically significantly up-regulated. Thus, the trends in gene expression of S. aureus 04-02981 on exposure to AGXX® observed in RNA-seq and in RT-qPCR were similar.

In previous studies by others, differential gene expression of S. aureus in planktonic and biofilm mode has been examined. Resch et al. (2005) observed that in biofilms, genes encoding polysaccharide intercellular adhesin, and enzymes associated with cell envelope synthesis were significantly up-regulated (Resch et al., 2005). To combat biofilms, many metals have been tested for their capacity to inhibit bacterial biofilm formation. Specifically, silver nanoparticles have received much attention with respect to their antimicrobial nature. However, the minimum concentration of silver nanoparticles (AgNPs) required to eliminate biofilm formation is considered to have toxic effects on mammalian cells (Loo et al., 2016). They studied the effect of AgNPs and curcumin nanoparticles (Cur-NPs) on S. aureus and discovered that the combination of both nanoparticles was more effective than the individual AgNPs or Cur-NPs. Curcumin interferes with the QS system as was observed by the down-regulation of genes involved in QS, upon exposure to the substance (Loo et al., 2016). Ma et al. (2012) investigated the effect of two novel anti-virulence compounds on growth and biofilm formation of S. aureus. The compounds inhibited biofilm formation by repressing genes associated with biofilm formation such as icaA, sdrD, sspB, sigB, codY, which were also down-regulated in our studies at least at one of the five time-points (Ma et al., 2012).

In summary, based on our findings, we conclude that AGXX® is an effective antimicrobial substance which might also act as a biofilm inhibitor based on our molecular data. The mechanism of inhibition is likely ica-independent without the production of PIA, by interfering with the QS system and by repressing genes associated with surface adhesin and lipopolysaccharide synthesis. In addition, the antimicrobial might also reduce pathogenesis of S. aureus 04-02981 by down-regulating the synthesis of toxins and virulence factors.

**AUTHOR CONTRIBUTIONS**

AV performed all the microbiological and molecular experiments, drafted the manuscript, and designed the figures. ADJ supervised and discussed bioinformatics analyses of RNA-seq, and prepared and deposited the RNA-seq data at NCBI. DW performed the confocal microscopy and analyzed the data. JK drafted part of the discussion and gave insightful suggestions on molecular biology of Gram-positive pathogens. EG designed the project and supervised all the experiments. All authors discussed and corrected the manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: [https://www.frontiersin.org/articles/10.3389/fmicb.2018.00221/full#supplementary-material](https://www.frontiersin.org/articles/10.3389/fmicb.2018.00221/full#supplementary-material)

**FIGURE S1** | Library sizes of all the RNA samples (S. aureus 04-02981, S. aureus 04-02981 + Ag , and S. aureus 04-02981 + AGXX), at different time periods. The image indicates the read depth of each sample. The X-axis represents the experiment names as used in the factors file, and gene counts file during RNA-seq analysis via T-REx. The sample names comprise the metal sheet used, followed by the time of exposure. For example, sample ‘AGXX_06’ represents S. aureus 04-02981 exposed to AGXX for 6 minutes. The Y-axis represents the total number of mapped reads.

**TABLE S1** | Primer and probe sequences used for RT-qPCR.

**TABLE S2** | AGXX®-mediated growth inhibition of S. aureus 04-02981 in batch cultures.

**TABLE S3** | Alignment rates of the RNA-sequences of S. aureus 04-02981.

**TABLE S4** | Differentially expressed genes in S. aureus 04-02981 on exposure to Ag and AGXX®.

**TABLE S5** | Gene Ontology assignments on exposing S. aureus 04-02981 to AGXX® for 6 minutes. Rate = The rating values (1 to 5) reflect binned values based on: (TopHits/ClassSize) * −log2(adj-pvalue).

**TABLE S6** | Gene Ontology assignments on exposing S. aureus 04-02981 to Ag for 80 minutes. Rate = The rating values (1 to 5) reflect binned values based on: (TopHits/ClassSize) * −log2(adj-pvalue).

**TABLE S7** | Expression of operons in S. aureus 04-02981 on exposure to AGXX® for 6 minutes. Rate = The rating values (1 to 5) reflect binned values based on: (TopHits/ClassSize) * −log2(adj-pvalue).

**TABLE S8** | Expression of operons in S. aureus 04-02981 on exposure to Ag for 80 minutes. Rate = The rating values (1 to 5) reflect binned values based on: (TopHits/ClassSize) * −log2(adj-pvalue).


Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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