Microarray analysis of toxicogenomic effects of Ortho-phenylphenol in Staphylococcus aureus

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Abstract

**Background:** Staphylococcus aureus (S. aureus), is responsible for many infectious diseases, ranging from benign skin infections to life-threatening endocarditis and toxic shock syndrome. Ortho-phenylphenol (OPP) is an antimicrobial agent and an active ingredient of EPA-registered disinfectants with wide human exposure in various agricultural, hospital and veterinary disinfectant products. Despite many uses, an understanding of a cellular response to OPP and it’s mechanism of action, targeted genes, and the connectivity between targeted genes and the rest of cell metabolism remains obscure.

**Results:** Herein, we performed a genome-wide transcriptome analysis of the cellular responses of S. aureus when exposed to 0.82 mM of OPP for 20 and 60 min. Our data indicated that OPP downregulated the biosynthesis of many amino acids, which are required for protein synthesis. In particular, the genes encoding the enzymes of the diaminopimelate (DAP) pathway which results in lysine biosynthesis were significantly downregulated. Intriguingly, we revealed that the transcription of genes encoding ribosomal proteins was upregulated by OPP and at the same time, the genes encoding iron acquisition and transport were downregulated. The genes encoding virulence factors were upregulated and genes encoding phospholipids were downregulated upon 20 min exposure to OPP.

**Conclusion:** By using microarray analysis that enables us to simultaneously and globally examine the complete transcriptome during cellular responses, we have revealed novel information regarding the mode of action of OPP on Staphylococcus: OPP inhibits anabolism of many amino acids and highly downregulates the genes that encode the enzymes involved in the DAP pathway. Lysine and DAP are essential for building up the peptidoglycan cell wall. It was concluded that the mode of action of OPP is similar to the mechanism of action of some antibiotics. The discovery of this phenomenon provides useful information that will benefit further antimicrobial research on S. aureus.
Background

The U.S. Environmental Protection Agency (EPA) has endeavored to determine the efficacy and the mode of action of antimicrobials. At EPA, 5,000 antimicrobial products are registered, and hospital-level disinfectants are being tested against pathogens such as *S. aureus*, which is responsible for many infectious diseases, ranging from benign skin infections to life-threatening endocarditis and toxic shock syndrome [1]. One of the reasons EPA has exerted such efforts is that hospital-acquired infections are a serious threat to public health. Therefore, it is important to use appropriate antimicrobial agents with clear understanding of the subsequent effects to prevent infection outbreaks in health care environments [2].

The phenolic compound, ortho-phenylphenol (OPP), is an antimicrobial agent and an active ingredient of EPA-registered disinfectant with wide human exposure in various agricultural, hospital, and veterinary disinfectant products. OPP is employed in a variety of applications, including hard surface disinfection, wood preservation, treatment of citrus fruit, vegetables before packaging to prevent microbial decay and textile production due to its bactericidal and fungicidal activity [3-5].

There have been several reports related to the exposure of OPP on humans. It has been reported that OPP increased the incidence of urinary bladder tumors in F344/DuCrj rats when administered in the diet [6]. The results of this study stimulated the initiation of additional testing of OPP for both tumor induction and possible reactivity with DNA. OPP has been found to have estrogenic or antiandrogenic activity, and binds to the androgen or estrogen receptors [7]. In spite of these effects OPP is still used in applications that simultaneously contact both humans and bacteria. It is therefore important to understand the differential effects on each so that its efficacy can be understood and even optimized.

Moreover, a lack of understanding of a cellular response to OPP hinders further development of more innovative methods for combating pathogens. Certainly, better elucidation of the molecular events responsible for establishing and maintaining pathogenicity will help to map affected cell functions and serve to delineate the mechanisms involved in the disinfectant activity.

Microarrays have been effectively employed to simultaneously and globally examine the complete transcriptional response at the genomic level in *Pseudomonas aeruginosa* and *S. aureus* upon exposure to antimicrobials [8-15].

In this study, to our knowledge, for the first time, we show that the global transcription response of *S. aureus* to OPP includes downregulation of genes involved in lysine metabolism, as well as genes involved in amino acid metabolism, by utilizing Affymetrix *S. aureus* GeneChip arrays. Our findings indicate that: (i) many cellular protective processes were upregulated, (ii) the transcription of genes involved in primary metabolic pathways was downregulated, and (iii) the transcription of genes encoding lysine and histidine biosynthesis was downregulated. Next we performed real-time PCR analysis on selected genes to validate the array results. Based on this result, it was concluded that this study may help to elucidate the mechanism of action by which OPP stops cell wall construction and thereby inhibiting *S. aureus* growth, and may facilitate the design of more effective antimicrobials.

Results and discussion

Growth inhibition by OPP

To determine the sublethal inhibitory effect of OPP on *S. aureus*, we first exposed the exponentially growing cells to different concentrations of OPP dissolved in DMSO (0 up to 1.18 mM). In figure 1, we demonstrate that 0.82 mM concentration of OPP caused a growth inhibition for about 20 min. Note that minimum inhibitory concentration (MIC50) of OPP on *S. aureus* was reportedly 500 mg/l (3 mM) [16]. In this study, to better understand how *S. aureus* initially responds to OPP, we chose the rate of cell growth inhibition with 0.82 mM OPP after 20 and 60 min exposure times compared to control (without OPP).

Transcriptional profiles in response to OPP

To investigate early transcriptional changes in response to OPP exposure, we isolated total RNA after 20 min and 60 minutes exposure to 0.82 mM OPP and conducted five independent microarray experiments in the absence (control) and the presence (experimental) of 0.82 mM OPP (see figure 1). To further identify genes with statistically marked changes in expression levels, we applied the following criteria to each of the 20 min, 60 min, and control-experimental microarray data sets: (i) a p-value for a t-test should be equal to or less than 0.05, (ii) an absolute fold change in transcript level should be equal to or greater than 2, and (iii) a gene should have a presence or marginal call [17] from 50% or more replicates on both the experimental and control replicate sets. Of the 7,775 genes represented on the *S. aureus* GeneChip, 2,348 genes showed statistical significance based on a 1-way ANOVA. We found that mRNA levels 669 genes of *S. aureus* were significantly altered in response to OPP by two fold or more upregulation or downregulation. The raw data of 7,775 genes control (0 min) and experimental genes after (20 and 60 min exposure to 0.82 mM of OPP) has been deposited in NCBI’s Gene Expression Omnibus [18] and is accessible through GEO Series accession number GSE10605 (Additional file 1).
Analysis of gene expression changes in 20 min and 60 min
To examine how genes with transcript level changes are distributed with regard to their functions, we further classified these 669 genes that were either upregulated or downregulated by a fold change of two or more according to the Gene Classification based on COG functional categories in the genome of National Center for Biotechnology information (NCBI) [19] (see also Additional file 2).

In Figure 2, the differences between the numbers of up and downregulated genes in each functional class after 20 and 60 minutes exposure to 0.82 mM of OPP are illustrated. Note that Figure 2 represents a total of 669 genes including the group of “function unknown (36), hypothetical protein (132) and general function predicted only (70)”.

Some interesting findings are as follows: (i) the genes of amino acid transport and metabolism were highly downregulated at both 20 and 60 min; (ii) the genes of inorganic ion transport and metabolism were downregulated at 20 min and decrease in the number of genes downregulated at 60 min was also observed; (iii) the genes in the class of “translation, ribosomal structure and biogenesis” were significantly upregulated at 20 min; (iv) the number of genes involved in nucleotide transport and metabolism were increased after 20 min compared to after 60 min. In general, figure 2 illustrates that the functional classes contained more downregulated and fewer upregulated genes at 20 min. This result suggests that the functional class profiles were notably different between 20 and 60 min, and this difference might explain why *S. aureus* underwent the initial growth inhibition followed by partial growth recovery upon exposure to OPP.

Functional classifications analysis
To further identify genes with similar transcription patterns during the time course, we removed 238 genes (including the group of functional unknown (36), hypothetical protein (132) and general function predicted only (70)). We categorized 431 genes with known functions into 6 groups on the basis of their transcription directions.
Briefly, group I contained 23 genes upregulated upon both exposure times, while group II had 80 genes with increased expression levels at 20 min and no significant changes upon 60 min exposure. Further, group III possessed 26 genes that were upregulated at 60 min exposure. Group IV contained 128 genes downregulated upon both exposure times, whereas 95 genes of group V exhibited downregulation after 20 min. Finally, group VI had 79 genes that were downregulated upon 60 min exposure. Figure 4 displays the number of genes (431) within groups I through VI in each functional class. As indicated above, additional file 2 contains all 669 genes including the genes classified under the functional group designated as "unknown, hypothetical, and general function prediction only".

Since most of the genes discussed in this report are in additional file 2, for further analysis of the data and for the readers convenience, we decided to make table 1 with 138 S. aureus genes that were most strongly upregulated or downregulated in response to OPP after 20 and 60 minutes exposure. These genes were also classified under seven groups based on their transcription directions.

**Group I: genes upregulated upon 20 and 60 min exposures**

Group I of table 1 contains 18 genes associated with virulence in S. aureus. Interestingly, five of these genes encode the secretory antigen precursor, SsaA. The ssaA gene potentially regulated by the YycG/YycF system encodes proteins involved in cell wall metabolism, membrane-bound transport systems, and pathogenicity, including two major antigenic proteins, SsaA and IsaA. YycF has also been shown to bind specifically to the promoter regions.
of two genes, encoding the IsaA antigen and the LytM peptidoglycan hydrolase. This is in agreement with the proposed role of this system in controlling virulence and cell wall metabolism [20].

In this study, OPP also upregulated the clfB (clumping factor B) gene expression upon both 20 and 60 min exposure. In our previous results, triclosan upregulated the expression of SA2423 encoding ClfB, which binds fibrinogen [12]. The results of this study show that the produc-

Figure 3
Groups of differentially regulated 431 genes with known functional class, which are categorized by their transcription directions upon 20 and 60 min exposures. Group I contained 23 genes upregulated upon both exposure times, while group II had 80 genes upregulated at 20 min and no significant changes upon 60 min exposure. Further, group III possessed 26 genes that were upregulated in response to 60 min exposure. Group IV contained 128 genes downregulated upon both exposure times, whereas 95 genes of group V exhibited downregulation after 20 min exposure. Finally, group VI had 79 genes that were downregulated upon 60 min exposure.
tion of virulence factors in *S. aureus* may be a secondary effect of OPP and this may provide new insight into the protective response of *S. aureus* to OPP.

**Group II: genes upregulated upon 20 min exposure**

Group II of table 1 indicates that the class of "translation, ribosomal structure and biogenesis" which is responsible for the synthesis of ribosomal proteins was upregulated after 20 minutes. In group II of table 1, for instance four genes encode 30S ribosomal proteins and 15 genes code for 50S ribosomal proteins. In addition, SA0459 (*rplY*) which encoded general stress protein was upregulated at 20 min. Ribosomal proteins are required for protein translation. Such early response of these ribosomal protein genes may reflect a stress response during exposure. The upregulation of ribosomal protein genes may reflect the translation process or help proper ribosome functioning under stress conditions as exposure to OPP. The suggestion that the expression of ribosomal proteins is activated upon exposure to OPP is surprising as this presumably reflects increased growth rate or virulence.

In group II (table 1), we also observed genes related to primary metabolism that mainly belonged to the functional classes of "purines, pyrimidines, nucleosides, and nucleotides". The gene cluster: SA1041–SA1048 (*pyr*<sub>RPB-CAABFE</sub>) which is homologous to the pyrimidine biosynthetic (*pyr*) operon of *Bacillus subtilis* [21] was upregulated at 20 min.

**Group III: genes upregulated upon 60 min exposure**

In group III of table 1, there were some genes from amino acid transport and metabolism, an ATP-binding cassette (ABC) transporters and transcription. The oligopeptide transport system (Opp) of *S. aureus* is an ABC transporter that transports amino acids, cations- and iron-carrying compounds and peptides with a broad specificity [22]. The peptides are mainly used as nutrients by the multiple amino acid auxotrophic *S. aureus*. The Opp system consists of four different proteins: OppB and OppC, OppD and OppF. Interestingly, 4 of the 26 genes, including SA0845–SA0848 that code for proteins associated with amino acid transport were upregulated (table 1). Therefore, the suggestion that the expression of these proteins is
### Table 1: List of 138 S. aureus genes that were most strongly affected by OPP and are discussed in this report categorized by their related function. The microarray results are the mean of five replicates of each gene.

| Affymetrix Probe ID | ORF no. | p-value | Fold change | p-value | Fold change | Description | Gene symbol | Functional class |
|---------------------|---------|---------|-------------|---------|-------------|-------------|-------------|-----------------|
| **Group I: Upregulation (20 min) – Upregulation (60 min) 18 genes** |
| sa_c6812s5946_a_at  | SA0265  | 8.55E-06 | 2.1         | 8.55E-06 | 3.1         | peptidoglycan hydrolase (surface antigen) | lytM         | Cell envelope biogenesis, outer membrane |
| sa_c7382s10191_a_at | SA0423* | 5.16E-07 | 8.1         | 5.16E-07 | 19.4        | hypothetical protein, similar to autolysin (N-acetylmuramoyl-L-alanine amidase) | | General function prediction only |
| sa_c7698s6703_a_at  | SA0519  | 0.00293 | 2.8         | 0.00293 | 2.5         | Ser-Asp rich fibrinogen-binding, bone sialoprotein-binding protein | sdrC         | Cell envelope biogenesis, outer membrane |
| sa_c8045s7032_at    | SA0620  | 1.27E-06 | 7.6         | 1.27E-06 | 5.6         | hypothetical protein, similar to secretory antigen precursor SsaA | | General function prediction only |
| sa_c592s9345_a_at   | SA0905  | 4.83E-08 | 5.7         | 4.83E-08 | 7.0         | N-acetylgalactosaminidase (major autolysin) | atl          | Cell envelope biogenesis, outer membrane |
| sa_c1007s793_a_at   | SA1003  | 7.81E-06 | 3.7         | 7.81E-06 | 4.2         | fibrinogen-binding protein | | hypothetical protein |
| sa_c4394s3743_a_at  | SA1898  | 7.99E-05 | 6.1         | 7.99E-05 | 5.8         | hypothetical protein, similar to SceD precursor | | hypothetical protein |
| sa_c4612s9984cs_s_at | SA1972 | 0.00267 | 2.2         | 0.00267 | 2.7         | multidrug resistance protein (efflux transporter) | | hypothetical protein |
| sa_c5066s4362_a_at  | SA2093* | 0.00025 | 6.8         | 0.00025 | 9.0         | hypothetical protein, similar to secretory antigen precursor SsaA | ssaA         | General function prediction only |
| sa_c5082s4380_a_at  | SA2097* | 3.71E-05 | 6.9         | 3.71E-05 | 11.5        | hypothetical protein, similar to secretory antigen precursor SsaA | | General function prediction only |
| sa_c342s182_a_at    | SA2142  | 9.36E-07 | 3.6         | 9.36E-07 | 3.6         | multidrug resistance protein B (drug efflux transporter) | | hypothetical protein |
| sa_c5274s4572_a_at  | SA2143  | 1.24E-07 | 3.9         | 1.24E-07 | 4.1         | multidrug resistance efflux pump | | hypothetical protein |
| sa_c5652s4904_a_at  | SA2206  | 0.00082 | 2.5         | 0.00082 | 2.8         | Immunoglobulin G binding protein A precursor | sbi          | Cell envelope biogenesis, outer membrane |
| sa_c6151s5333_a_at  | SA2332  | 0.00019 | 7.8         | 0.00019 | 8.6         | hypothetical protein, similar to secretory antigen precursor SsaA | | General function prediction only |
| sa_c6250s5428_a_at  | SA2353* | 1.91E-05 | 10.2        | 1.91E-05 | 13.8        | secretory antigen precursor SsaA | | General function prediction only |
| sa_c9402s8223_a_at  | SA2355* | 3.23E-06 | 8.5         | 3.23E-06 | 9.4         | transcriptional regulator, MARR family | | hypothetical protein |
| sa_c6259s5439_a_at  | SA2356  | 3.43E-06 | 6.5         | 3.43E-06 | 6.9         | immunodominant antigen A | isoA         | Cell envelope biogenesis, outer membrane |
| sa_c6506s5675_a_at  | SA2423  | 0.00148 | 2.9         | 0.00148 | 3.5         | fibrinogen-binding protein A, clumping factor | clfB         | Posttranslational modification, protein turnover, chaperones |
Table 1: List of 138 *S. aureus* genes that were most strongly affected by OPP and are discussed in this report categorized by their related function. The microarray results are the mean of five replicates of each gene. (Continued)

**Group II: Upregulation (20 min) – No change (60 min) 28 genes**

| Gene ID            | strain | Log2 Ratio | fold change |
|--------------------|--------|------------|-------------|
| sa_c7511s6531_a_at | SA0459 | 0.000138   | 2.3         |
| sa_c7621s6634_a_at | SA0497 | 7.95E-05   | 2.2         |
| sa_c7625s6638_at  | SA0498 | 5.07E-05   | 3.5         |
| sa_c1147s928_a_at | SA1041 | 0.00142    | 2.8         |
| sa_c1151s932_a_at | SA1042 | 0.000143   | 6.7         |
| sa_c9991s8687_a_at| SA1043 | 7.66E-05   | 6.5         |
| sa_c1155s937_a_at | SA1044 | 0.000101   | 6.0         |
| sa_c1159s942_a_at | SA1045 | 0.000161   | 5.1         |
| sa_c1165s946_a_at | SA1046 | 0.00018    | 3.3         |
| sa_c1167s950_a_at | SA1047 | 0.000788   | 3.3         |
| sa_c9989s8682_a_at| SA1048 | 0.00107    | 3.0         |
| sa_c1302s1077_a_at| SA1084 | 0.000221   | 2.7         |
| sa_c4792s4098_at  | SA2022 | 0.000417   | 2.1         |
| sa_c4824s4130_a_at| SA2029 | 2.76E-05   | 2.1         |
| sa_c4836s4142_at  | SA2032 | 0.00024    | 2.1         |
| sa_c9951s8647_at  | SA2033 | 0.000342   | 2.6         |
| sa_c4848s4156_at  | SA2035 | 0.00213    | 2.1         |
| sa_c4852s4158_at  | SA2036 | 0.00267    | 2.1         |
| sa_c10191s8871_a_at| SA2038 | 0.0026     | 2.8         |
| sa_c4860s4166_at  | SA2039 | 0.00204    | 2.5         |
| sa_c4864s4170_at  | SA2040 | 0.000749   | 2.7         |

- **ribosomal protein L25** (general stress protein Ctc)
- **50S ribosomal protein L10**; ribosomal protein L10 (BL5)
- **50S ribosomal protein L7,L12**; ribosomal protein L7,L12
- uracil phosphoribosyltransferase; Pyrimidine operon regulatory protein pyrR
- uracil permease (uracil transporter)
- aspartate carbamoyltransferase catalytic chain (Aspartate transcarbamylase) (ATCase)
- dihydroorotase, dihydroorotase-like
- carbamoyl-phosphate synthase, arginine-specific, small chain
- carbamoyl-phosphate synthase, arginine-specific, large chain
- orotidine 5-phosphate decarboxylase
- orotate phosphoribosyltransferase
- 50S ribosomal protein L19; ribosomal protein L19
- 50S ribosomal protein L17; ribosomal protein L17
- 50S ribosomal protein L15; ribosomal protein L15
- 50S ribosomal protein L18; ribosomal protein L18
- 50S ribosomal protein L6; ribosomal protein L6 (BL8)
- 50S ribosomal protein L5; ribosomal protein L5 (BL6)
- 50S ribosomal protein L24; ribosomal protein L24 (BL23)
- 50S ribosomal protein S17; ribosomal protein S17 (BS16)
- 50S ribosomal protein L29; ribosomal protein L29
- 50S ribosomal protein L16; ribosomal protein L16

- **rplY** Translation, ribosomal structure and biogenesis
- **rplO** Translation, ribosomal structure and biogenesis
- **rplR** Translation, ribosomal structure and biogenesis
- **rplF** Translation, ribosomal structure and biogenesis
- **rpsN** Translation, ribosomal structure and biogenesis
- **rplX** Translation, ribosomal structure and biogenesis
- **rpsQ** Translation, ribosomal structure and biogenesis
- **rpsC** Translation, ribosomal structure and biogenesis
- **rplP** Translation, ribosomal structure and biogenesis
Table 1: List of 138 S. aureus genes that were most strongly affected by OPP and are discussed in this report categorized by their related function. The microarray results are the mean of five replicates of each gene. (Continued)

| Gene ID | Symbol | Description | Fold Change 20 min | Fold Change 60 min | Function |
|---------|--------|-------------|--------------------|--------------------|----------|
| sa_c4868s4175_a_at | SA2041 | 30S ribosomal protein S3; ribosomal protein S3 (BS3) | 2.8 | 2.8 | Translation, ribosomal structure and biogenesis |
| sa_c4876s4184_at | SA2043 | 30S ribosomal protein S19; ribosomal protein S19 (BS19) | 2.5 | 2.5 | Translation, ribosomal structure and biogenesis |
| sa_c9959s8654_a_at | SA2044 | 50S ribosomal protein L2; ribosomal protein L2 (BL2) | 2.7 | 2.7 | Translation, ribosomal structure and biogenesis |
| sa_c10192s8875_at | SA2045 | 50S ribosomal protein L23 | 2.8 | 2.8 | Translation, ribosomal structure and biogenesis |
| sa_c4880s4187_at | SA2046 | 50S ribosomal protein L4; ribosomal protein L4 | 2.3 | 2.3 | Translation, ribosomal structure and biogenesis |
| sa_c4888s4195_a_at | SA2047 | 30S ribosomal protein S10; ribosomal protein S10 (BS13) | 2.4 | 2.4 | Translation, ribosomal structure and biogenesis |

**Group III: No change (20 min) – Upregulation (60 min) 8 genes**

| Gene ID | Symbol | Description | Fold Change 20 min | Fold Change 60 min | Function |
|---------|--------|-------------|--------------------|--------------------|----------|
| sa_c10571s9056_a_at | SA0845 | putative oligopeptide ABC transporter integral membrane protein (fragment) | 2.5 | 2.5 | Amino acid transport and metabolism, Inorganic ion transport and metabolism |
| sa_c324s166_a_at | SA0846 | probable peptide ABC transporter permease | 2.4 | 2.4 | Amino acid transport and metabolism, Inorganic ion transport and metabolism |
| sa_c328s170_a_at | SA0847 | probable peptide ABC transporter ATP-binding ABC transporter protein | 2.2 | 2.2 | Amino acid transport and metabolism, Inorganic ion transport and metabolism |
| sa_c332s172_a_at | SA0848 | probable peptide ABC transporter ATP-binding ABC transporter protein | 2.2 | 2.2 | Amino acid transport and metabolism, Inorganic ion transport and metabolism |
| sa_c5349s4625_a_at | SA0950 | ABC transporter ATP-binding protein – spermidine:putrescine transport | 2.2 | 2.2 | Amino acid transport and metabolism |
| sa_c795s596_a_at | SA0952 | ABC transporter membrane-spanning permease – spermidine:putrescine transport | 2.2 | 2.2 | Amino acid transport and metabolism |
| sa_c803s604_a_at | SA0953 | spermidine:putrescine ABC transporter, spermidine:putrescine-binding periplasmic protein (potD) homolog | 2.2 | 2.2 | Amino acid transport and metabolism |
| sa_c8848s7783_a_at | SA1601 | CRCB, CrcB-like protein | 2.3 | 2.3 | Cell division and chromosome partitioning |

**Group IV: Downregulation (20 min) – Downregulation (60 min) 27 genes**

| Gene ID | Symbol | Description | Fold Change 20 min | Fold Change 60 min | Function |
|---------|--------|-------------|--------------------|--------------------|----------|
| sa_c5061s4360_a_at | SA0229 | dipeptide ABC transporter, periplasmic dipeptide-binding protein (dppA) | -5.8 | -5.8 | Amino acid transport and metabolism |
| sa_c736s544_a_at | SA0937 | cytochrome D ubiquinol oxidase subunit I | -2.9 | -2.9 | Energy production and conversion |
Table 1: List of 138 *S. aureus* genes that were most strongly affected by OPP and are discussed in this report categorized by their related function. The microarray results are the mean of five replicates of each gene. (Continued)

| GenBank Accession | BAC clone | Log2 Fold Change | P-value | Function |
|-------------------|-----------|------------------|---------|----------|
| sa_c740548_a_at   | SA0938    | -2.9             | 0.00375 | cytochrome D ubiquinol oxidase subunit II homolog |
|                   |           |                  |         | homoserine dehydrogenase (HDH) |
|                   |           |                  |         | dhoM     |
| sa_c1659s1395_a_at| SA1164    | -8.9             | 0.000431| threonine synthase (EC 2.3.1) |
|                   |           |                  |         | thrC     |
| sa_c1665s1401_a_at| SA1165    | -8.3             | 0.00011 | homoserine kinase (thrB) |
|                   |           |                  |         | thrB     |
| sa_c1669s1406_a_at| SA1166    | -11.5            | 5.06E-05| probable peptide ABC transporter protein |
|                   |           |                  |         | opp-2C   |
| sa_c1872s1598_a_at| SA1213    | -2.8             | 0.00127 | putative oligopeptide ABC transporter integral membrane protein (fragment) |
|                   |           |                  |         | opp-2B   |
| sa_c1876s1602_a_at| SA1214    | -3.7             | 8.58E-06| aspartokinase II in bifunctional enzyme: aspartokinase II; homoserine dehydrogenase II |
|                   |           |                  |         | lysC     |
| sa_c1912s1635_a_at| SA1225    | -54.6            | 8.90E-06| aspartate-semialdehyde dehydrogenase (ASA dehydrogenase) |
|                   |           |                  |         | asd      |
| sa_c1918s1640_a_at| SA1226    | -21.5            | 1.78E-05| dihydrodipicolinate synthase (DHDPS) |
|                   |           |                  |         | dapA     |
| sa_c1922s1644_a_at| SA1227    | -27.3            | 2.75E-07| dihydrodipicolinate reductase (DHPR) |
|                   |           |                  |         | dapB     |
| sa_c1924s1648_a_at| SA1228    | -31.4            | 1.17E-07| 2,3,4,5-tetrahydroxypyridine-2,6-dicarboxylate N-succinyltransferase |
|                   |           |                  |         | dapD     |
| sa_c1928s1652_a_at| SA1229    | -23.5            | 2.55E-07| serine-pyruvate aminotransferase; alanine-glyoxylate aminotransferase |
|                   |           |                  |         | serA     |
| sa_c1936s1659_a_at| SA1231    | -18.5            | 8.94E-08| parvulin-like PPlase precursor (Peptidyl-prolyl cis-trans isomerase (PPI) (Rotamase PPI) |
|                   |           |                  |         | prsA     |
| sa_c3202s2750_a_at| SA1544    | -15.8            | 2.74E-05| succinyl-diaminopimelate desuccinylase (dapE) |
|                   |           |                  |         | ureA     |
| sa_c3603s3083_a_at| SA1659    | -2.4             | 7.54E-06| urease gamma chain (urea amidohydrolase) |
|                   |           |                  |         | ureB     |
| sa_c3603s3083_a_at| SA1659    | -6.3             | 1.99E-05| urease alpha chain (urea amidohydrolase) |
|                   |           |                  |         | ureC     |
| sa_c3603s3083_a_at| SA1659    | -3.3             | 1.99E-05| Parvulin-like PPIase precursor (Peptidyl-prolyl cis-trans isomerase (PPI) (Rotamase PPI) |
|                   |           |                  |         | prsA     |
| sa_c5029s4326_a_at| SA2083    | -3.5             | 6.38E-07| succinyl-diaminopimelate desuccinylase (dapE) |
|                   |           |                  |         | ureA     |
| sa_c5029s4326_a_at| SA2083    | -3.5             | 6.38E-07| urease gamma chain (urea amidohydrolase) |
|                   |           |                  |         | ureB     |
| sa_c5029s4326_a_at| SA2083    | -3.5             | 6.38E-07| urease alpha chain (urea amidohydrolase) |
|                   |           |                  |         | ureC     |
Table 1: List of 138 S. aureus genes that were most strongly affected by OPP and are discussed in this report categorized by their related function. The microarray results are the mean of five replicates of each gene. (Continued)

| Genbank Accession | SA Accession | Fold Change | Expression | Gene Description |
|-------------------|--------------|-------------|------------|------------------|
| sa_c5035s4334_at  | SA2085       | 1.71E-05    | -2.9       | urease accessory protein UreE |
| sa_c5039s4340_a_at| SA2086       | 1.40E-05    | -2.3       | urease accessory protein UreF |
| sa_c9293s8136_a_at| SA2088       | 0.000118    | -2.2       | urease accessory protein UreD |
| sa_c5303s4583_a_at| SA2149       | 1.28E-07    | -35.8      | probable peptide ABC transporter ATP-binding ABC transporter protein |
| sa_c5307s4587_at  | SA2150       | 9.33E-07    | -36.5      | ABC-type transporter, permease component |
| sa_c5777s5020_a_at| SA2235       | 2.82E-05    | -5.1       | putative ABC transporter; osmoprotectant-binding protein, anaerobic ribonucleoside-triphosphate reductase activating protein |
| sa_c6435s5604_a_at| SA2409       | 0.000134    | -4.6       | anaerobic ribonucleoside-triphosphate reductase activating protein |
| sa_c37s34_a_at    | SA0010       | 2.44E-06    | -4.7       | branched-chain amino acid permease |
| sa_c4055s3432_a_at| SA0201       | 0.00111     | -2.3       | RGD-containing lipoprotein |
| sa_c7055s6165_a_at| SA0331       | 0.000236    | -3.0       | probable lipoprotein |
| sa_c7100s6210_a_at| SA0344       | 2.80E-06    | -15.2      | methyltetrahydropteroylglutamate – homocysteine methyltransferase (vitamin-B12-independent methionine synthase isozyme) |
| sa_c5418s4689_a_at| SA0420       | 7.55E-05    | -2.8       | probable amino acid ABC transporter, ATP-binding protein (abc) |
| sa_c7374s6406_a_at| SA0421       | 6.63E-07    | -3.3       | putative amino acid ABC transporter, permease protein, glutamine transport system |
| sa_c5431s4700_a_at| SA0769       | 2.32E-06    | -5.4       | probable amino acid ABC transporter, ATP-binding protein permease protein of ABC transporter system |
| sa_c8512s7471_a_at| SA0770       | 1.42E-05    | -6.3       | probable D-methionine-binding lipoprotein metQ precursor (Outer membrane lipoprotein1) |
| sa_c8518s7475_a_at| SA0771       | 5.12E-06    | -7.5       | hypothetical protein |

Group V: Downregulation (20 min) – No change (60 min) 35 genes

| Genbank Accession | SA Accession | Fold Change | Expression | Gene Description |
|-------------------|--------------|-------------|------------|------------------|
| sa_c37s34_a_at    | SA0010       | 2.44E-06    | -4.7       | branched-chain amino acid permease |
| sa_c4055s3432_a_at| SA0201       | 0.00111     | -2.3       | RGD-containing lipoprotein |
| sa_c7055s6165_a_at| SA0331       | 0.000236    | -3.0       | probable lipoprotein |
| sa_c7100s6210_a_at| SA0344       | 2.80E-06    | -15.2      | methyltetrahydropteroylglutamate – homocysteine methyltransferase (vitamin-B12-independent methionine synthase isozyme) |
| sa_c5418s4689_a_at| SA0420       | 7.55E-05    | -2.8       | probable amino acid ABC transporter, ATP-binding protein (abc) |
| sa_c7374s6406_a_at| SA0421       | 6.63E-07    | -3.3       | putative amino acid ABC transporter, permease protein, glutamine transport system |
| sa_c5431s4700_a_at| SA0769       | 2.32E-06    | -5.4       | probable amino acid ABC transporter, ATP-binding protein permease protein of ABC transporter system |
| sa_c8512s7471_a_at| SA0770       | 1.42E-05    | -6.3       | probable D-methionine-binding lipoprotein metQ precursor (Outer membrane lipoprotein1) |
| sa_c8518s7475_a_at| SA0771       | 5.12E-06    | -7.5       | hypothetical protein |
Table 1: List of 138 S. aureus genes that were most strongly affected by OPP and are discussed in this report categorized by their related function. The microarray results are the mean of five replicates of each gene. (Continued)

| Gene ID          | Accession | Log2 Fold Change | Description and Function                                                                 | Related Function       |
|------------------|-----------|------------------|----------------------------------------------------------------------------------------|------------------------|
| sa_c350s191_a_at | SA0849    | 8.12E-05         | oligopeptide ABC transporter, periplasmic oligopeptide-binding protein (oppA-2)          | Amino acid transport and metabolism |
| sa_c352s195_a_at | SA0850    | 0.000757         | periplasmic oligopeptide-binding protein of oligopeptide ABC transporter                  | Amino acid transport and metabolism |
| sa_c1820s1547_a_at | SA1200    | 0.00698          | para-aminobenzoate synthetase glutamine amidotransferase component II                   | Amino acid transport and metabolism |
| sa_c1828s1551_a_at | SA1201    | 0.000803         | pir|AE0653 anthranilate synthase component II, anthranilate synthase; indole-glycerol phosphate synthase; indole-3-glycerolphosphate synthetase | trpD, trpC, trpF, Amino acid transport and metabolism |
| sa_c1832s1558_a_at | SA1202    | 0.00239          |                | trpD, trpC, Amino acid transport and metabolism |
| sa_c1836s1562_at | SA1203    | 0.00279          |                | trpD, trpC, Amino acid transport and metabolism |
| sa_c1840s1566_a_at | SA1204    | 0.000663         | tryptophan synthase beta chain; tryptophan synthase (beta subunit)                     | trpB, Amino acid transport and metabolism |
| sa_c1844s1570_a_at | SA1205    | 5.52E-05         | tryptophan synthase alpha chain; tryptophan synthase                                   | trpA, Amino acid transport and metabolism |
| sa_c1866s1587_a_at | SA1211    | 0.000158         | ATP-binding ABC transporter protein                                                     | opp-2F, Amino acid transport and metabolism |
| sa_c4209s3561_a_at | SA1858    | 0.000144         | dihydroxy-acid dehydratase (DAD)                                                       | ilvD, Amino acid transport and metabolism, Coenzyme metabolism |
| sa_c4213s3565_a_at | SA1859    | 0.000431         | acacetolactate synthase isozyme III large subunit (AHAS-III)                           | ilvB, Amino acid transport and metabolism, Coenzyme metabolism |
| sa_c9931s8627_a_at | SA1861    | 0.000255         | ketol-acid reductoisomerase (Acetohydroxy-acid isomeroreductase)                       | ilvC, Amino acid transport and metabolism, Coenzyme metabolism |
| sa_c4223s3575_a_at | SA1862    | 4.06E-05         | 2-isopropylmalate synthase                                                             | leuA, Amino acid transport and metabolism |
| sa_c4225s3576_a_at | SA1863    | 5.85E-05         | 3-isopropylmalate dehydrogenase                                                        | leuB, Energy production and conversion, Amino acid transport and metabolism |
| sa_c4229s3580_a_at | SA1864    | 1.39E-05         | 3-isopropylmalate dehydratase large subunit                                            | leuC, Amino acid transport and metabolism |
| sa_c4239s3588_a_at | SA1865    | 6.09E-05         | 3-isopropylmalate dehydratase small subunit                                             | leuD, Amino acid transport and metabolism |
| sa_c4243s3594_a_at | SA1866    | 1.58E-05         | threonine dehydratase biosynthetic (Threonine deaminase) (TD)                           | ilVA, Amino acid transport and metabolism |
Table 1: List of 138 *S. aureus* genes that were most strongly affected by OPP and are discussed in this report categorized by their related function. The microarray results are the mean of five replicates of each gene. (Continued)

| Strain ID | Gene Symbol | Description | log2Ratio | Annotation |
|-----------|-------------|-------------|-----------|------------|
| **Group VI: No change (20 min) – Downregulation (60 min) 19 genes** |
| sa_c2346s1974_a_at | SA0144 | capsular polysaccharide synthesis enzyme Cap5A | 4.02E-05 | -3.1 | copA | Cell envelope biogenesis, outer membrane |
| sa_c2385s1987_a_at | SA0145 | capsular polysaccharide synthesis enzyme Cap5B | 4.55E-06 | -2.9 | copB | Cell division and chromosome partitioning |
| sa_c2399s1991_a_at | SA0146 | capsular polysaccharide synthesis enzyme Cap8C | 0.00055 | -2.6 | copC | Cell envelope biogenesis, outer membrane, Carbohydrate transport and metabolism |
| sa_c2413s1997_a_at | SA0147 | capsular polysaccharide synthesis enzyme Cap5D | 0.0001 | -2.6 | copD | Cell envelope biogenesis, outer membrane, Carbohydrate transport and metabolism |
| sa_c9546s8318_a_at | SA0148 | putative UDP-glucose 4-epimerase (Galactoswaldenase) (UDP-galactose 4-epimerase) | 9.77E-05 | -2.8 | copE | Cell envelope biogenesis, outer membrane, Carbohydrate transport and metabolism |
| sa_c2479s2056_a_at | SA0149 | capsular polysaccharide synthesis enzyme Cap5F | 2.40E-06 | -3.1 | copF | Cell envelope biogenesis, outer membrane, Carbohydrate transport and metabolism |
Table 1: List of 138 *S. aureus* genes that were most strongly affected by OPP and are discussed in this report categorized by their related function. The microarray results are the mean of five replicates of each gene. (Continued)

| Gene ID | Description | Expression | Function | Additional Information |
|---------|-------------|------------|----------|------------------------|
| sa_c2516s2092_a_at | UDP-N-acetylglicosamine 2-epimerase (UDP-GlcNAc-2-epimerase) (Xenobiotic acetyltransferase) (XAT) | 5.92E-05 | -2.4 | capG | Cell envelope biogenesis, outer membrane |
| sa_c10086s8810_a_at | chloramphenicol acetyltransferase | 0.00025 | -2.8 | capH | General function prediction only |
| sa_c10087s8814_a_at | capsular polysaccharide synthesis enzyme CapSI | 0.00017 | -2.7 | capI | Cell envelope biogenesis, outer membrane |
| sa_c10089s8822_a_at | capsular polysaccharide synthesis enzyme CapSK | 0.00103 | -2.1 | capK | Cell envelope biogenesis, outer membrane |
| sa_c898s698_a_at | 29-kDa cell surface protein | 0.00044 | -2.7 | isdA | Cell envelope biogenesis, outer membrane |
| sa_c906s704_a_at | hypothetical protein SirD | 0.00538 | -2.2 | isdC | Cell envelope biogenesis, outer membrane |
| sa_c3380s9339_a_at | 6,7-dimethyl-8-ribityllumazine synthase (DMRL synthase) | 1.41E-05 | -2.4 | ribH | Coenzyme metabolism |
| sa_c3387s2918_a_at | probable riboflavin biosynthesis bifunctional protein | 6.15E-06 | -2.8 | ribA | Coenzyme metabolism |
| sa_c3391s2919_a_at | riboflavin synthase alpha chain | 3.10E-06 | -2.7 | ribB | Coenzyme metabolism |
| sa_c3395s2925_a_at | riboflavin specific deaminase | 5.10E-07 | -3.0 | ribD | Coenzyme metabolism |
| sa_c4369s3721_at | thiamine-phosphate pyrophosphorylase | 0.000661 | -2.3 | thiE | Coenzyme metabolism |
| sa_c4373s3725_a_at | hydroxyethylthiazole kinase | 3.05E-05 | -2.5 | thiM | Coenzyme metabolism |
| sa_c4379s3726_a_at | bifunctional enzyme: hydroxyphosphomethylpyrimidine kinase | 0.000137 | -2.8 | thiD | Coenzyme metabolism |

Group VII: Downregulation (20 min) – Upregulation (60 min) 3 genes

| Gene ID | Description | Expression | Function | Additional Information |
|---------|-------------|------------|----------|------------------------|
| sa_c9442s8255_a_at | intercellular adhesion protein IcaA | 0.0016 | -2.5 | icaA | Cell envelope biogenesis, outer membrane |
| sa_c6677s5830_a_at | IcaD | 0.0136 | -2.0 | icaD | Cell envelope biogenesis, outer membrane |
| sa_c6681s9106_a_at | intercellular adhesion protein IcaB | 3.83E-06 | -2.1 | icaB | Cell envelope biogenesis, outer membrane |

* Genes were validated by real-time PCR.
activated upon exposure to OPP is surprising as this presumably reflects increased growth rate or recovery. We found that a putative operon containing four open reading frames (ORFs) (potABCD) was upregulated (table 1). The potABCD operon encodes a periplasmic binding protein dependent ABC transport systems from Gram-positive bacteria [23]. The SA0950–SA0952–SA0953 shows homology to the genes encoding this potABCD transport system (potA, potC and potD), which are involved in the transport of spermidine and putrescine. Further, we showed the upregulation of ABC transport systems-related genes, which accompanied the growth recovery.

Of further importance was that group III contained genes related to integral membrane protein, which belonged to the functional class of "cell division and chromosome partitioning". SA1601 (crcB) is a putative integral membrane protein possibly involved in chromosome condensation (table 1).

Group IV: genes downregulated upon 20 and 60 min exposures

In group IV in table 1, we noted that genes belonging to the functional classes of "amino acid transport and metabolism", "carbohydrate transport and metabolism", "energy production and conversion", "posttranslational modification protein turnover chaperones", "transcription" classes and putative lipoproteins were downregulated upon both exposure times.

Intriguingly, we observed the high downregulation of SA2149 and SA2150 (hrt A and B), the heme-regulated transport system, which consist of a novel transport system which plays a critical role in staphylococcal heme metabolism (table 1). Among the genes in the class of "amino acid transport and metabolism", SA1225 (lysC)-SA1226 (adsl)-SA1227 (dapA)-SA1228 (dapB)-SA1229 (dapD), and SA1814 (dapE) fall within a predicted operon and are all involved in dianaminopimelate (DAP) biosynthesis (table 1). The disruption of biosynthetic pathways involved in building up bacterial cell wall components is a common mode of action of antibiotics [24]. Penicillins [25], methicillin [26], cephalosporins [27] and glycopeptide drugs such as vancomycin [28] are all drugs that inhibit major steps in the construction of the peptidoglycan layer of bacterial cell walls. Lysine or its biosynthetic precursor, DAP [29], are essential to most bacteria for the synthesis of the peptidoglycan layer of the cell wall [30-33]. Since mammals neither make nor use DAP and require L-lysine as an essential amino acid that is supplied through dietary intake, inhibitors of the DAP biosynthetic pathway will probably not result in mammalian toxicity. Decisively SA1225 (lysC)-SA1226 (adsl)-SA1227 (dapA)-SA1228 (dapB) and SA1229 (dapD) show fold highest decreases as -54.6, -21.5, -27.3, -31.4, and -23.5 folds at 20 min and -7.7, -4.3, -5.2, -5.1, and -4.4 folds at 60 min in this experiment (see also table 2). Our findings suggest that the mode of action of OPP may be related to bacterial biosynthesis of amino acids. Other genes of amino acids, including methionine, threonine, histidine and lysine were also highly down regulated at -15, -7, -19 and -54 folds. Therefore, this outcome in conjunction with the extensive downregulation of the genes encoding DAP biosynthesis suggests that OPP may inhibit construction of the peptidoglycan in cell wall of S. aureus. These genes were less downregulated at 60 min than at 20 min. These results suggest that OPP inhibits the growth of S. aureus at 20 min and that growth recovery occurs at 60 minutes, indicating a possible mechanism of action of OPP in S. aureus. In a similar study carried out using *Pseudomonas aeruginosa* treated with 0.82 mM OPP, we did not observe extensive downregulation of the genes involved in amino acid biosynthesis and specifically lysine biosynthesis (data not shown). This suggests that the mechanisms of action of 0.82 mM OPP on *P. aeruginosa* and *S. aureus* may differ.

Additional amino acid biosynthesis genes including: SA1164 (*dhoM*)-SA1165 (*thtC*)-SA1166 (*thtB*) involved in threonine biosynthesis were also in this group (table 1). Further, SA2082–SA2083–SA2084–SA2085–SA2086–SA2088 (*ureABCEFD*), which make CO₂ and NH₃ from urea and encode urea amidohydrolase and urease accessory proteins and SA1544 which codes for serine-pyruvate aminotransferase were downregulated at 20 and 60 min (table 1).

Group IV of table 1 also shows that the functional class of "cell envelope biogenesis, outer membrane" was distinctive. In particular, SA1231 which shows an -18-fold decrease after 20 min encodes an alanine racemase that catalyses the conversion of L-alanine into D-alanine, a key component of bacterial peptidoglycan [34]. Additionally, the putative lipoproteins: SA0229 (*dppA*), SA1213 (*opp-2C*)-SA1214 (*opp-2B*), SA1659 (*psA*), SA2235 (*opuCC*), and SA2409 which are cell wall anchoring surface proteins were downregulated in response to OPP (table 1).

In group IV, we also observed genes related to primary metabolism that mainly belonged to the functional classes of "energy metabolism", "lipid metabolism", and "transcription". For example, cytochrome *bd* complex: SA0937–SA0938 (*cydAB*) was downregulated upon both 20 min and 60 min exposure (table 1). Cytochrome *bd* complex is one of two terminal oxidases in the bacterial respiratory chain that reduce molecular oxygen to water, utilizing intermediates shuttled through the electron transport chain [35]. Cytochrome *d* oxidase catalyses the last step of oxygen respiration and prevails under oxygen-limiting conditions [36]. Interestingly, it was speculated
that cytochrome d oxidase is required under conditions of environmental stress and may have crucial roles in cellular physiology other than acting as an oxidase [37]. However, prior studies revealed that the cydAB genes were strongly upregulated upon exposure to hydrogen peroxide strengthens the confidence of the prior assignments about the role of cytochrome d oxidase in oxidative protection processes of both Gram positive and Gram negative bacteria [10,13].

**Group V: genes downregulated upon 20 min exposures**

In group V in table 1, the most dominant class was "amino acid transport and metabolism", which contained half of the genes in that group. Further, SA1858 (ilvD)-SA1859 (ilvB) and SA1861 (ilvC)-SA1862 (leuA)-SA1863

| Gene   | mRNA level change with microarray | mRNA level change with real-time PCR | Sense primer sequence | Antisense primer sequence |
|--------|----------------------------------|-------------------------------------|-----------------------|---------------------------|
|        | 20 min  | 60 min  | 20 min  | 60 min  | Sense primer sequence | Antisense primer sequence |
| SA0423 | 8.1     | 19.4    | 9.1(± 1.4) | 18.6(± 1.2) | 5'-CGG GTG AAT CAG | 5'-TAT TAT CCA CCA |
| SA1164 | -8.9    | -3.4    | -40.9(± 5.0) | -2.1(± 0.1) | 5'-TAT GAT CCG CCA | 5'-GAG TGT AGC AGG |
| SA1225 | -54.6   | -7.7    | -265.0(± 18.9) | -58.0(± 1.4) | 5'-GAG TGT AGC AGG | 5'-TCA TCA GTG GGA |
| SA1226 | -21.5   | -4.3    | -14.9(± 3.3) | -2.1(± 0.7) | 5'-TCA TCA GTG GGA | 5'-TCA TCA GTG CCA-3' |
| SA1227 | -27.3   | -5.2    | -58.0(± 1.5) | -2.8(± 0.1) | 5'-ACT TTA GGC AGA | 5'-ACT TTA GGC AGA-3' |
| SA1228 | -31.4   | -5.1    | -35.3(± 1.6) | -1.6(± 0.4) | 5'-AGT CTA GGC AGA | 5'-AGT CTA GGC AGA-3' |
| SA1229 | -23.5   | -4.4    | -7.4(± 2.0) | -1.2(± 0.2) | 5'-TTG TGC GTA CCA-3' | 5'-TTG TGC GTA CCA-3' |
| SA2093 | 6.8     | 9.0     | 15.7(± 1.2) | 4.6(± 0.5) | 5'-TTA AAT TTA CCA-3' | 5'-TTA AAT TTA CCA-3' |
| SA2097 | 6.9     | 11.5    | 5.5(± 1.8) | 24.5(± 1.2) | 5'-TTA AAT TTA CCA-3' | 5'-TTA AAT TTA CCA-3' |
| SA2353 | 10.2    | 13.8    | 8.0(± 1.6) | 16.4(± 1.9) | 5'-TTA AAT TTA CCA-3' | 5'-TTA AAT TTA CCA-3' |
| SA2355 | 8.5     | 9.4     | 5.3(± 1.3) | 20.1(± 1.7) | 5'-TTA AAT TTA CCA-3' | 5'-TTA AAT TTA CCA-3' |
| SA2459c | -2.5    | 2.5     | -3.1(± 0.2) | 2.0(± 0.3) | 5'-TTA AAT TTA CCA-3' | 5'-TTA AAT TTA CCA-3' |
| SA2460c | -2.0    | 4.7     | -1.8(± 0.3) | 2.5(± 0.3) | 5'-TTA AAT TTA CCA-3' | 5'-TTA AAT TTA CCA-3' |
| SA2461c | -2.1    | 3.4     | -2.6(± 0.5) | 2.1(± 0.5) | 5'-TTA AAT TTA CCA-3' | 5'-TTA AAT TTA CCA-3' |
| SA2464c | -10.5   | -       | -5.5(± 1.4) | -5.5(± 1.4) | 5'-TCA AAT TTA CCA-3' | 5'-TCA AAT TTA CCA-3' |
| SA2465c | -10.6   | -       | -13.0(± 1.4) | -13.0(± 1.4) | 5'-GCA AAT TAG CCA-3' | 5'-GCA AAT TAG CCA-3' |
| SA2466c | -14.5   | -       | -5.7(± 1.3) | -5.7(± 1.3) | 5'-GCA AAT TAG CCA-3' | 5'-GCA AAT TAG CCA-3' |
| SA2468c | -13.6   | -       | -58.1(± 1.3) | -58.1(± 1.3) | 5'-GCA AAT TAG CCA-3' | 5'-GCA AAT TAG CCA-3' |
| SA0727a | 1.00    | 1.00    | 1.00    | 1.00    | 5'-GCA AAT TAG CCA-3' | 5'-GCA AAT TAG CCA-3' |
|         |         |         |         |         | 5'-TCA AAT TTA CCA-3' | 5'-TCA AAT TTA CCA-3' |

The real time PCR results are the mean of three biological replicates with three technical replicates for each gene. The microarray results are the mean of five replicates of each gene.

1SA0727 was glyceraldehyde 3-phosphate dehydrogenase (GAPDH) and used as the house-keeping gene. 1SA2464, SA2465, SA2466, and SA2468 were downregulated at 20 min with no change at 60 min. 1SA2459, SA2460, and SA2461 were downregulated after 20 min and upregulated after 60 min exposure.
were downregulated at 60 min. These genes share homologous to the histidine biosynthesis were highly downregulated on 20 min (table 1). SA1200–SA1201 homologous to the histidine biosynthesis were highly downregulated on 20 min. First, SA0769 (metA) involved in D-methionine-binding lipoprotein (outer membrane lipoprotein 1). The proteins encoded by SA0344 (metE)-SA0769 (metN)-SA0770 (metL)-SA0420 (metN)-SA0421 (metL) are involved in D-methionine transporter of S. aureus ABC transporter (table 1). Interestingly, group V contained lipoproteins such as SA0010, SA0331, SA0849, SA0850, SA0201 and SA1211 (table 1). These results, along with downregulation of all the genes of lipoproteins of S. aureus in group IV and V, suggest that OPP exposure may decrease stability of the staphylococcal membrane. Secondly, the proteins encoded by SA2475 (chiQ)-SA2476 (chiO) are involved in cobalt and nickel transport (table 1).

**Group V: genes downregulated upon 60 min exposures**

Table 1 illustrates that the functional classes of group VI in general contained more downregulated genes at 60 min. In particular, the functional classes of "cell envelope biosynthesis outer membrane", "carbohydrate transport and metabolism", "amino acid transport and metabolism", "coenzyme metabolism", "energy production and conversion" and "posttranslational modification protein turnover chaperones" had significantly more downregulated genes at 60 min (see also figure 4). This result suggests that the functional class profiles were notably different between 20 and 60 min.

One of the characteristics of group VI of table 1 was the downregulation of 12 genes belonging to the functional class of "cell wall/membrane/envelope biogenesis". In particular, genes related to envelope biogenesis were distinctive: SA0144 (capA)-SA0145 (capB)-SA0146 (capC)-SA0147 (capD)-SA0148 (capE)-SA0149 (capF)-SA0150 (capG)-SA0151 (capH)-SA0152 (capI)-SA0154 (capK) were downregulated at 60 min. These genes share homology with the capsular polysaccharide synthesis enzyme (cap) operon which in turn is essential for virulence by impeding phagocytosis [39]. This finding is congruent with the previous outcome that triclosan downregulates several virulence factor-related genes (SA0144–SA0153 (capABCDEFGHJ)) in S. aureus [12]. Moreover, IsdAC encoded by SA0977 and SA0978, the iron-regulated surface determinant (Isd) system, was downregulated at 60 min (table 1). Identification of the Isd system in S. aureus has demonstrated the importance of cell-wall sorted proteins in heme binding and transport [40]. To date, the Isd system comprises the only known heme-iron utilization pathway in S. aureus. Cell-wall sorted proteins of the S. aureus iron-regulated surface determinant system bind human hemoproteins, remove the heme molecule, and transport heme through the cell wall and plasma membrane for accumulation in the bacterial cytoplasm [41].

Particularly important was that many of the genes in the class of "coenzyme metabolism" were also members of group VI (figure 4 and table 1). Intriguingly, the genes were all involved in the riboflavin biosynthesis. SA1586 (ribH)-SA1587 (ribA)-SA1588 (ribB)-SA1589 (ribD) was downregulated at 60 min exposure (table 1). Riboflavin (vitamin B2) is an essential component of the basic metabolism, being a precursor of coenzymes flavin adenine dinucleotide (FAD) and flavin mononucleotide (FMN). The best studied system of the riboflavin biosynthesis in bacteria is the rib operon of Bacillus subtilis encoding a pyrimidine deaminase/reductase, α-subunit of riboflavin synthase, GTP cyclohydrolase/3,4-dihydroxy 2-butanone 4-phosphate (3,4-DHBP) synthase, and β-subunit of riboflavin synthase [42]. These enzymes form a pathway that creates one riboflavin molecule from one molecule of GTP and two molecules of ribulose 5-phosphate [43]. The proteins encoded by SA1894 (thiE)-SA1895 (thiM)-SA1896 (thiD) were involved in thiamine biosynthesis of coenzyme metabolism at 60 min (table 1). Methylcillin-resistant S. aureus small colony variants are frequently auxotrophic for hemin, menadione, thiamine, and CO2 involved in biosynthesis of the electron transport chain element. This phenotype grows slowly, and forms very small, nonhemolytic colonies in routine culture, so it may lead to the misidentification of this organism. As discussed above, group IV also had SA2149 (hrtA)-SA2150 (hrtB), which exhibited expression level decreases upon 20 and 60 min exposures in chorus with the repression of the genes of thiamine biosynthesis. Therefore, this result suggests that growth inhibition was accompanied with the repression of many coenzyme metabolism-related genes.

**Group VII: genes downregulated upon 20 min and upregulated upon 60 min exposures**

Note that group VII has been included only in table 1 in order to discuss the aberrant behaviour of the ica genes.
In this study, we used S. aureus NCTC 8325 obtained from the Network on Antimicrobial Resistance in S. aureus (NARSA). As previously described [10-12], we initiated and maintained S. aureus cultures at 37°C with shaking at 250 rpm using sterilized Luria-Bertani (LB) broth. For growth inhibition, 0.14 mg/L (0.82 mM) of OPP (Aldrich Chemical Co., St. Louis, MO) was dissolved in DMSO and used for the microarray study and added immediately after OD_{600} reached 0.8. OD_{600} was measured by using Lambda 25 spectrophotometer (PerkinElmer, Inc., MA). Note that the pH of S. aureus cultures was around 7.0 at 37°C after the exposure [48].

RNA isolation
Total RNA was isolated after 20 and 60 min incubation with and without (control) OPP using the RiboPure – Bacteria kit (Ambion, Inc., Austin, TX) [11]. The quantity of eluted RNA was determined using the NanoDrop spectrophotometer (NanoDrop Technologies, Inc., Wilmington, DE). RNA 6000 Nano LabChip with an Agilent 2100 Bioanalyzer (Agilent Technologies, Palo Alto, CA).

cDNA synthesis, labeling, hybridization, staining, and scanning
cDNA synthesis, cDNA fragmentation, labeling, hybridization, staining and washing steps were performed according to the manufacturer's protocol for the Affymetrix S. aureus GeneChip arrays (Affymetrix, Inc., Santa Clara, CA).

Affymetrix S. aureus genechip analysis
The arrays were scanned with the Affymetrix GeneChip Scanner 3000. To analyze the array data, GeneChip Operating Software (GCOS) v. 1.2 (Affymetrix, Inc., Santa Clara, CA) and GeneSpring GX v. 7.3 (Agilent Technologies, Inc., Santa Clara, CA) were utilized with the following parameters: alpha 1, 0.04; alpha 2, 0.06; tau, 0.015; target signal, 500. Fold changes were calculated as the ratio between the signal averages of five biological con-
Real-time PCR analysis
To determine the validity of the array data, transcript level changes obtained with the microarray analysis were compared with those from quantitative real-time PCR. Genes and primer sequences employed for the real-time PCR analysis are listed in Table 2. The housekeeping gene, glyceraldehyde-3-phosphate dehydrogenase (GAPDH), was used as an endogenous control. The real-time PCR was performed by employing iCycler iQ Real-Time PCR Detection System with iScript cDNA Synthesis Kit and IQ SYBR Green Supermix (BioRad Laboratories, Inc., Hercules, CA). For each gene, three biological replicates with three technical replicates each were employed. Reaction mixtures were initially incubated for 3 min at 95.0°C, followed by 40 cycles of 10 s at 95.0°C, 30 s at 55.0°C, and 20 s at 72.0°C. PCR efficiencies were derived from standard curve slopes in the iCycler software v. 3.1 (BioRad Laboratories, Inc., Hercules, CA). Melt-curve analysis was also performed to evaluate PCR specificity and resulted in single primer-specific melting temperatures. In this report, relative quantification based on the relative expression of a target gene versus GAPDH gene was utilized to determine transcript level changes.

Authors’ contributions
HJ performed microarray experiments, and data analysis, and drafted the manuscript. FT initiated and supervised the study, and reviewed the manuscript. CN and WEB reviewed the manuscript.

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