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Tricarboxylic Acid Cycle-Dependent Regulation of Staphylococcus epidermidis Polysaccharide Intercellular Adhesin Synthesis∗†

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Staphylococcus epidermidis is a major nosocomial pathogen primarily infecting immunocompromised individuals or those with implanted biomaterials (e.g., catheters). Biomaterial-associated infections often involve the formation of a biofilm on the surface of the medical device. In S. epidermidis, polysaccharide intercellular adhesin (PIA) is an important mediator of biofilm formation and pathogenesis. Synthesis of PIA is regulated by at least three DNA binding proteins (IcaR, SarA, and σB) and several environmental and nutritional conditions. Previously, we observed the environmental conditions that increased PIA synthesis decreased tricarboxylic acid (TCA) cycle activity. In this study, S. epidermidis TCA cycle mutants were constructed, and the function of central metabolism in PIA biosynthesis was examined. TCA cycle inactivation altered the metabolic status of S. epidermidis, resulting in a massive derepression of PIA biosynthetic genes and a redirection of carbon from growth into PIA biosynthesis. These data demonstrate that the bacterial metabolic status is a critical regulatory determinant of PIA synthesis. In addition, these data lead us to propose that the TCA cycle acts as a signal transduction pathway to translate external environmental cues into intracellular metabolic signals that modulate the activity of transcriptional regulators.

Staphylococcus epidermidis is the most frequently isolated nosocomial pathogen from the blood cultures of adult and pediatric intensive care unit patients in the United States of America (48). Frequently, S. epidermidis infections involve the formation of a biofilm on the surfaces of biomaterials (e.g., catheters) (77). A biofilm is a complex aggregation of bacteria usually encapsulated by an adhesive exopolysaccharide matrix. The exopolysaccharide matrix provides structural stability to the biofilm, enhanced adhesion to surfaces, and protection from host defenses and antibiotics (2, 50, 79). In S. epidermidis, the exopolysaccharide facilitating bacterial adherence in a biofilm is polysaccharide intercellular adhesin (PIA). PIA is an N-acetylglucosamine polymer (44) of at least 130 residues (42) whose synthesis requires the enzymes encoded by genes in the ica operon (icaADBC) (27). The synthesis of PIA is required for the maturation of biofilms (27), important for evasion of the host innate immune system (79, 80), and critical in biomaterial-associated infections (55, 57).

Bacteria live in environments subject to rapid changes in the availability of the nutrients necessary to provide energy and biosynthetic intermediates for the synthesis of macromolecules. Consequently, bacterial survival depends on the ability to regulate expression of genes coding for enzymes required for growth in the altered environment. In pathogenic bacteria, adaptation to an altered environment often includes activating transcription of virulence genes; hence, synthesis of many virulence determinants is regulated by environmental/nutritional signals (e.g., nitrogen, iron, and calcium) (47). Because PIA is the most important virulence determinant of S. epidermidis (2, 7, 38, 56, 57, 84), it is not surprising that PIA synthesis is regulated by environmental and nutritional signals (13, 16, 18, 22, 53).

Regulation of the icaADBC operon is complex, involving at least two DNA binding proteins (IcaR and SarA) and the alternative sigma factor σB. IcaR binds immediately S’ to the icaA transcriptional start site and represses transcription of the ica operon (9, 29). SarA is an essential positive effector of icaADBC transcription that binds to the icaA promoter region (26, 74). σB affects PIA synthesis indirectly by regulating the expression of icaR (34, 35, 52). In addition, the spa quorum-sensing system has a minimal effect on PIA synthesis and biofilm formation; however, the mechanism remains to be determined (82).

In Staphylococcus aureus, σB is activated during stress conditions and growth phase transitions (3, 81). Because S. epidermidis icaADBC transcription and PIA synthesis are induced by environmental and nutritional stresses, stress-dependent activation of σB has been a focal point of research into the environmental regulation of PIA synthesis and biofilm formation (10, 26, 35). Importantly, σB does not respond directly to environmental signals; therefore, a second area of research interest has focused on identifying the regulatory network controlling σB activation, which is involved in responding to environmental signals. The activation of σB involves an anti-sigma factor (RsbW) and an anti-anti-sigma factor (RsbV) and resembles the process in the closely related organism Bacillus subtilis (35). In the absence of environmental stimuli, σB is bound in a complex with RsbW. Stress-inducing stimuli are hypothesized to activate the RsbU phosphatase to dephosphorylate (activate) the anti-anti-sigma

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factor RsbV, which then binds RsbW in a competitive manner to increase the concentration of free α F (34, 35). Based on several published studies, it seems likely that the α F posttranslational regulatory network is responsible for responding to some environmental stresses (i.e., osmotic stress) but is only minimally involved in responding to nutritional signals (e.g., glucose), suggesting that there are other means for S. epidermidis to “sense” nutritional signals (18, 35, 52).

Previously, we observed the environmental and nutritional signals that enhance PIA synthesis also repress tricarboxylic acid (TCA) cycle activity, leading us to hypothesize that repression of TCA cycle activity would increase PIA accumulation (78). Fluorocitrate is a highly toxic compound when metabolized to 4-hydroxy-PIA produced increased proportionally (78). Fluorocitrate is a minimally inhibited TCA cycle activity, and determined the expression of TCA cycle activity would increase PIA accumulation. The genes coding for citrate synthase and isocitrate dehydrogenase are present in a two-gene operon and cotranscribed. To inactivate the citZC operon, a gene splicing by overlap extension technique was used to replace a 2-kb internal DNA fragment; Ampr Camr Ermr 51. Citrate synthase (citZ) and isocitrate dehydrogenase (citC) double mutant. The genes coding for citrate synthase and isocitrate dehydrogenase are present in a two-gene operon and cotranscribed. To inactivate the citZC operon, a gene splicing by overlap extension technique was used to replace a 2-kb internal DNA region of citZC by the ermB gene. The ermB gene was amplified from pEC4 (5) using primers citC-r ermB2 and citZ2-ermB (Table 2), containing sequences homologous to the citC and citZ genes. Genomic DNA from S. epidermidis strain
TABLE 2. Oligonucleotide primers used in this study

<table>
<thead>
<tr>
<th>Primer</th>
<th>Nucleotide sequence (5′→3′)</th>
</tr>
</thead>
<tbody>
<tr>
<td>663 forward</td>
<td>GCCGCGCGGCGGCGGATATAAA</td>
</tr>
<tr>
<td>664 reverse</td>
<td>GCCGCGCGGCAGACGCAAGCC</td>
</tr>
<tr>
<td>SERP1762(glM)-f</td>
<td>CACCTGAATTGCGTCTTTAAC</td>
</tr>
<tr>
<td>SERP1762(glM)-r</td>
<td>TTAACCAAGATTGCGTCTTTAAC</td>
</tr>
<tr>
<td>SERP1760(glM)-f</td>
<td>GTGTGAGATTGCGTCTTTAAC</td>
</tr>
<tr>
<td>SERP1760(glM)-r</td>
<td>TGCC</td>
</tr>
<tr>
<td>glmU-f</td>
<td>TGGGCCATGAAATGGAATGTGGACC</td>
</tr>
<tr>
<td>glmU-r</td>
<td>GTTGAATGCATGAAATGTGGACC</td>
</tr>
<tr>
<td>icaProbeforward</td>
<td>GACAGTCGCTACGAAAAAA</td>
</tr>
<tr>
<td>icaProbereverse</td>
<td>CCGAAATTTGTAATTTTACCAAG</td>
</tr>
<tr>
<td>sigB-f</td>
<td>CGAAGATGTCGAATGCTCATAAG</td>
</tr>
<tr>
<td>sigB-r</td>
<td>GGTCAAGTTGCTACTAATTC</td>
</tr>
<tr>
<td>Sarl-SD-acnA-f</td>
<td>CTCTAAGCTGCAATAATACGGAACGGAATATAAAGATTGCTC</td>
</tr>
<tr>
<td>Sarl-acnA-r</td>
<td>GGTACCTTGGTACGTTGTTAT</td>
</tr>
<tr>
<td>pCL15-f</td>
<td>GAGGATCGCTACTGCTTATT</td>
</tr>
<tr>
<td>pCL15-r</td>
<td>ACCG</td>
</tr>
<tr>
<td>citc-c-ermB2</td>
<td>GACGTTGCTACCTTCTTTTGCAGGGAATATAAAGATTGCTC</td>
</tr>
<tr>
<td>citc-f-ermB2</td>
<td>GCCATGAACTGCGAATGCGGGAATATAAAGATTGCTC</td>
</tr>
<tr>
<td>phoP-BamHI</td>
<td>GCAGAATGTCGTCATGTTGACT</td>
</tr>
<tr>
<td>ermB-citC-f2</td>
<td>CACCATGAAATTTGTAATGCGGGAATATAAAGATTGCTC</td>
</tr>
<tr>
<td>ansP-SacI</td>
<td>GCCAAGCTCGCATCGTCTTTAAT</td>
</tr>
<tr>
<td>ermB-citZ-r</td>
<td>GACGATTGCTACCTTCTTTGCAACGAGAATATAAAGATTGCTC</td>
</tr>
</tbody>
</table>

1457 was used as a template for PCR amplification of the DNA flanking the internal citZC region. Primers ansP-SacI and ermB-citZ-r (Table 2) were used for amplification of a 1.9-kb region of the citZ upstream region. A 1.7-kb region of the citZ downstream region was amplified using phoP-BamHI and ermB-citC-f2 primers (Table 2). The resulting 4.6-kb PCR product consisted of a 1-kb ermB gene with DNA flanking the internal citZC regions. The 4.6-kb PCR product contained SacI and BamHI sites that were used for ligation into pTS1-d digested with SacI and BamHI to generate pMR55. The temperature-sensitive plasmid pTS1-d is a derivative of pTS1 (24) with deletion of the plasmid-encoded 3′ region of the ermC gene. It was constructed by PCR amplification of pTS1 DNA using primers pTS1dErm-f and pTS1dErm-r, and the resulting PCR product was self-ligated as described previously (45). Plasmid pMR55 was used to construct a strain 1457 citZC double mutant (1457citZC:ermB) (19). Allelic replacement of the internal region of citZC genes by the ermB cassette was verified by PCR and Southern blot analysis.

Construction of the complementation plasmid. Plasmid pCL15 (a kind gift of Chia Lee), containing a P_pesc promoter, was used for the construction of an acnA complementation plasmid. The pMR52 complementation plasmid was constructed by amplifying the 2.7-kb promoterless acnA gene from S. epidermidis strain 1457 using Sall-SD-acnA-f and SacI-acnA-r primers, followed by ligation of the PCR product into plasmid pCL15 digested with SacI and SacI. Plasmid pMR52 was isolated from S. aureus strain RN4220 and electroporated into S. epidermidis strain 1457 (59). Plasmid pMR52 was introduced into strain SE1457-acnA using transducing phase 71 (49). Induction of expression from the P_pesc promoter was performed by supplementing bacterial cultures with isopropyl-β-D-thiogalactopyranoside (IPTG) (1 mM).

Measurement of acetic acid and glucose concentrations in culture medium. Aliquots of bacteria (2 ml) were centrifuged for 3 min at 14,000 rpm at 4°C, and the supernatants were removed and stored at −20°C until use. Acetate and glucose concentrations were determined with kits purchased from R-Biopharm and used according to the manufacturer’s protocol.

Determination of NAD+, NADH, and ATP concentrations. Intracellular NAD+ and NADH concentrations were determined with an enzymatic cycling assay kit (Biovision). Briefly, aliquots (5 to 20 ml) were harvested at 2, 3, 4, 5, and 6 h by centrifugation at 4°C for 10 min at 4,000 rpm. Bacterial pellets were suspended in 1 ml of extraction buffer (Biovision) and lysed using lysis matrix B tubes (Qbiogene) in a FastPrep instrument (Qbiogene). The lysate was centrifuged at 4°C for 5 min at 14,000 rpm. NAD+ and NADH concentrations were determined according to the manufacturer’s protocol and normalized to the cell density. All assays were performed in duplicate for two independent experiments.

Intracellular ATP concentrations were determined using the Enliten ATP assay (Promega). Briefly, 2 × 10^8 bacteria were collected at 2, 3, 4, 5, and 6 h and washed three times with 1 ml of phosphate-buffered saline, and suspended in 100 μl of phosphate-buffered saline. Bacteria were lysed by the addition of 900 μl of dimethyl sulfoxide (23). ATP concentrations were determined according to the manufacturer’s protocol and normalized to the cell density. All assays were performed in duplicate for two independent experiments.

Determination of aconitase activity. Cell-free lysates of S. epidermidis were prepared as follows: aliquots of bacteria (2 ml) were harvested by centrifugation and suspended in 750 μl of lysis buffer containing 90 mM Tris (pH 8.0) and 100 μM fluorocyanate. The samples were lysed using lysis matrix B tubes (Qbiogene) in a FastPrep instrument (Qbiogene). The lysate was centrifuged for 5 min at 14,000 rpm and 4°C. Aconitase activity was assayed in the resulting cell-free lysate as previously described (63). One unit of aconitase activity is defined as the amount of enzyme necessary to give a ΔA_{420} min⁻¹ of 0.0033 (1). Protein concentrations were determined by the method of Lowry (41).

PIA immunoblot assay. PIA accumulation was determined as described previously (78). Briefly, TSB medium containing 0.25% glucose was inoculated with equal numbers of bacteria from overnight cultures. The cultures were grown at 37°C, with a flask-to-medium ratio of 7:1, and aerated at 225 rpm. Every 2 h, equal numbers of bacteria were harvested by centrifugation, and the PIA was extracted in 0.5 M EDTA (pH 8.0) by boiling for 5 min. Aliquots of PIA were applied to a polyvinylidene difluoride membrane (Immobilon-P; Millipore) and blocked with 5% skim milk overnight. The polyvinylidene difluoride membrane was incubated for 2 h with PIA-specific antiserum (generously provided by Michael Otto), which was followed by a 2-h incubation with an anti-rabbit immunoglobulin G peroxidase conjugate. The presence of PIA was detected using SuperSignal West Pico chemiluminescent substrate (Pierce). The integrated density values of bands on autoradiographs were determined with the TotalLab software (Nonlinear Dynamics Ltd.).

Northern blot analysis. Northern blot analysis of transcripts was performed as described previously (78), except that total RNA was isolated using the FastRNA Pro Blue kit (Qiagen) and purified using an RNAeasy kit (Qiagen). Oligonucleotide primers (Table 2) used in making DNA probes were designed using the S. epidermidis Rf2A genome sequence (GenBank accession number CP000029). Probes for Northern blotting were generated by PCR amplification of unique internal regions of glmM, glmS, glmU, icaD, icaC, pgi, sarA, and sigB genes and labeled using the NorthzSouth random prime labeling kit (Pierce). Electrophoresis, transfer of the RNA to the Nytran SPC nylon membrane (Whatman), and hybridization were done using the NorthernMax kit (Ambion). Detection was performed using the chemiluminescent nucleic acid detection module (Pierce). When necessary, reprobing of blots was done according to the manufacturer’s protocol.

NMR data collection, processing, and analysis. Nuclear magnetic resonance (NMR) samples were prepared from three independent cultures (50 ml) of S. epidermidis strains 1457 and 1457-acnA during the exponential growth phase (2 h) and postexponential growth phase (6 h) using [13C6]glucose (Cambridge
The intracellular concentration was considered to be increased when the percent difference in the NMR peak intensities was fivefold greater than the percent error observed in the peak intensities between the replicate NMR spectra. Consequently, only compounds that are metabolic products of \([13C_6]\)glucose were detected.

**Statistical analysis.** The statistical significance of changes between wild-type and mutant strains (e.g., ATP concentrations) was assessed with the Student’s *t* test. To determine whether a correlation existed between two physiological parameters, a correlation coefficient (\(\rho\)) was calculated.

### RESULTS

#### Construction and characterization of an *S. epidermidis* strain 1457 aconitase mutant.

Partial biochemical inhibition of TCA cycle activity in *S. epidermidis* increased the accumulation of PIA (78). Because fluorocitrate is very toxic, we were unable to completely block TCA cycle activity without killing the bacteria. To overcome this obstacle, we inactivated the TCA cycle in *S. epidermidis* strain 1457 by insertion of a tetM cassette into the gene encoding aconitase (acnA; also known as citB) and then assessed the effect on PIA synthesis. In *S. epidermidis*, aconitase is present as a sole copy on the chromosome; hence, inactivation of aconitase results in a nonfunctional TCA cycle.

Inactivation of the aconitase gene in *S. epidermidis* strain 1457 by allelic replacement was genotypically verified by PCR and Southern blot analysis (data not shown) and phenotypically by aconitase enzymatic activity assays (Fig. 1A). In addition, overall TCA cycle function was assessed by measuring the accumulation and depletion of acetic acid from the culture medium of the wild-type, mutant, and complemented strains (Fig. 1D). (Acetic acid catabolism in staphylococci requires a fully functional TCA cycle [64, 65]). As expected, aconitase inactivation inhibited the postexponential oxidative catabolism of acetic acid (Fig. 1D). Of importance, when the concentration of acetic acid in the culture medium was plotted as a function of growth, the exponential growth phase rates of acetic acid accumulation in strains 1457 and 1457-\(acnA\) were equivalent (data not shown), suggesting that carbon utilization in both strains was proceeding through glycolysis and the acetate kinase/phosphotransacetylase pathway at a similar rate. The equivalent rate of acetic acid accumulation in strains 1457 and 1457-\(acnA\) was also supported by NMR metabolomic analysis (Table 3). If carbon flow through glycolysis were equivalent,
then it is reasonable to predict that the growth rate of the wild-type and aconitase mutant strains would also be equivalent. Surprisingly, the growth rate of strain 1457-<i>acnA</i> was lower than the growth rate of the isogenic wild-type strain, and this was reflected in the temporal depletion of glucose (Fig. 1C). This result was surprising because TCA cycle activity is repressed during the exponential growth phase when nutrients are abundant; thus, inactivating a repressed metabolic pathway should have minimal consequences. Restoration of the growth rate and the ability to catabolize acetate was accomplished by complementation of the strain 1457-<i>acnA</i> mutation with a plasmid (pMRS2) containing the <i>acnA</i> gene under the control of a P<sub>spac</sub> promoter (Fig. 1A to D). As a whole, these data demonstrate that <i>acnA</i> was inactivated, and this resulted in a metabolic block in the TCA cycle.

**TCA cycle inactivation redirects carbon from growth into UDP-N-acetylglucosamine biosynthesis.** When grown in the presence of a rapidly catabolizable carbon source, gram-positive bacteria repress the TCA cycle; hence, there is an accumulation of acidic secondary metabolites (e.g., acetate) in the culture medium. Derepression of the TCA cycle occurs upon depletion of the readily catabolizable carbon source(s) and coincides with the depletion of acetate from the culture medium. The processing of glucose through glycolysis and the resulting accumulation of acetate during substrate-level phosphorylation were equivalent in <i>S. epidermidis</i> strains 1457 and 1457-<i>acnA</i>; hence, the lower growth rate of strain 1457-<i>acnA</i> relative to the wild-type strain was likely due to a redirection of carbon and energy from growth into other cellular processes or metabolic pathways. If carbon and/or energy were diverted from growth to another metabolic pathway or process, then the number of bacteria produced per unit of glucose consumed should be lower in the aconitase mutant strain than in the wild-type strain. Plotting the glucose concentrations in the culture medium as a function of optical density at 600 nm for three independent experiments (Fig. 2) confirmed that fewer bacteria are produced per unit of glucose consumed by strain 1457-<i>acnA</i> than by strain 1457. These data strongly suggest that carbon and/or energy were diverted from growth into other cellular processes. To determine the metabolic pathways into which carbon might be redirected, we utilized NMR metabolomics to assess the metabolic status of strains 1457 and 1457-<i>acnA</i> in the exponential growth phase (2 h) and the postexponential growth phase (6 h) (see Fig. S1 and S2 in the supplemental material) (Tables 3 and 4). As expected, TCA cycle inactivation decreased the intracellular concentration of amino acids derived from TCA cycle intermediates (Asn, Asp, Gln, and Glu) and increased accumulation of the PIA biosynthetic precursor UDP-N-acetylglucosamine. Interestingly, TCA cycle inactivation resulted in an exponential-growth-phase accumulation of branched-chain amino acids relative to the wild-type strain (see Fig. S1 in the supplemental material).
the bacterium. As an indicator of the energy status, the intracellular metabolic block in the TCA cycle will alter the energy status of the TCA cycle. It generates ATP and reducing potential; therefore, a biosynthesis-driven metabolomic analysis was performed to determine whether TCA cycle inactivation altered the transcription of genes coding for the UDP-N-acetylglucosamine biosynthetic enzymes would be increased. To determine whether TCA cycle inactivation altered the transcription of UDP-N-acetylglucosamine biosynthetic genes, we probed total RNA for \( pgI \) (glucose-6-phosphate isomerase), \( glmS \) (glucosamine-6-phosphate synthase), \( glmM \) (phosphoglucomutase), and \( glmU \) (UDP-N-acetylglucosamine pyrophosphorylase) (Fig. 3). Consistent with metabolomic data indicating an increased intracellular concentration of fructose-6-phosphate, transcription of \( pgI \) was slightly increased in \( S. epidermidis \) strain 1457-\( acnA \) relative to strain 1457 during the exponential growth phase. (Glucose-6-phosphate isomerase catalyzes the second step in glycolysis, specifically, the reversible isomerization of glucose-6-phosphate into fructose-6-phosphate.) Fructose-6-phosphate is the biosynthetic precursor of UDP-N-acetylglucosamine, whose biosynthesis is encoded within the genes \( glmS, glmM, \) and \( glmU \). The exponential-growth-phase mRNA levels of \( glmS, glmM, \) and \( glmU \) were increased in strain 1457-\( acnA \) relative to strain 1457. Taken together, these data demonstrate that TCA cycle inactivation results in transcriptional changes that redirect carbon into N-acetylglucosamine biosynthesis.

In addition to providing biosynthetic intermediates, the TCA cycle generates ATP and reducing potential; therefore, a metabolic block in the TCA cycle will alter the energy status of the bacterium. As an indicator of the energy status, the intracellular concentrations of NAD\(^+\), NADH, and ATP were determined for \( S. epidermidis \) strains 1457 and 1457-\( acnA \). The intracellular concentration of ATP was lower in the aconitase mutant strain relative to the wild-type strain during both the exponential and postexponential growth phases (Fig. 4C). This decreased ATP concentration was likely due to decreased oxidative phosphorylation, as the intracellular concentrations of NAD\(^+\) and NADH were significantly \((P < 0.01)\) lower in the aconitase mutant strain than in the wild-type strain (Fig. 4A and B).

**TCA cycle inactivation increases PIA accumulation.** To determine whether TCA cycle inactivation resulted in increased synthesis of PIA, we assessed the relative amounts of cell-associated PIA in the wild-type strain, the complemented \( acnA \) mutant, and the wild-type strains using a PIA immunoblot assay (Fig. 5A and B). In addition, to facilitate comparison of the effect of TCA cycle inactivation on PIA accumulation with the inactivation of well-established regulators (i.e., IcaR, sigB, and SarA), we determined the relative amounts of cell-associated PIA for isogenic strains that had \( icaR \), \( sarA \), or \( sigB \) genetically inactivated. In agreement with previous studies (9, 10, 26, 35, 74), we observed that mutations in the genes of positive regulators \( sigB \) and \( sarA \) decreased PIA accumulation relative to the wild-type strain, while inactivation of \( icaR \) increased PIA accumulation (Fig. 5A and B). Surprisingly, aconitase inactivation had a much greater effect on PIA synthesis than did inactivation of the well-studied repressor \( IcaR \) (Fig. 5A and B). Complementation of the \( acnA \) mutation restores PIA accumulation to that seen in the parental strain (Fig. 5A and B).

In \( B. subtilis \), aconitase inactivation results in the accumulation of citric acid, which is partially responsible for blocking sporulation and causing a slight decrease in the growth rate (12). NMR metabolomics revealed an accumulation of citric acid in the aconitase mutant strain relative to the wild-type strain (see Fig. S1 and S2 in the supplemental material) (Tables 3 and 4), raising the possibility that the increased accumulation of PIA and the decreased growth rate of \( S. epidermidis \) strain 1457-\( acnA \) relative to strain 1457 might be due to citric acid-induced stress. To address these...
possibilities, we inactivated both citrate synthase (citZ) and isocitrate dehydrogenase (citC) in *S. epidermidis* strain 1457 and assessed growth and assayed for PIA accumulation (data not shown). Similar to the aconitase mutant strain, the citCZ double mutant dramatically increased PIA accumulation. Additionally, the citCZ double mutant had a decreased growth rate relative to the wild-type strain, but the growth rate was increased relative to strain 1457-\textit{acnA} (data not shown), suggesting that citric acid accumulation was partially responsible for the slower growth of the aconitase mutant. Taken together, these data demonstrate that TCA cycle activity represses *S. epidermidis* PIA accumulation and that this effect is independent of citric acid.

**TCA cycle activity represses icaADBC transcription.** TCA cycle inactivation alters the metabolic status of *S. epidermidis* (Fig. 1A to D and 3 and Tables 3 and 4), raising the possibility that an icaADBC transcriptional regulator could respond to the altered metabolic environment to increase icaADBC transcription. To determine whether the increased accumulation of PIA by the \textit{acnA} mutant strain was due to altered icaADBC transcription, we examined icaADBC mRNA levels in *S. epidermidis* strains 1457 and 1457-\textit{acnA}. In addition, icaADBC mRNA levels were determined for strain 1457 with mutations in sigB, sarA, or icaR. Consistent with the PIA immunoblot data (Fig. 5A and B), aconitase inactivation resulted in a very large increase in icaADBC transcription, or mRNA stability, relative to the isogenic wild-type strain (Fig. 6A and B), demonstrating that TCA cycle activity represses icaADBC transcription. In agreement with previous results, we observed that sigB and sarA inactivation repressed icaADBC transcription and that icaR inactivation derepressed icaADBC transcription (9, 11, 26). Interestingly, the extent of derepression in the 1457-\textit{acnA} mutant strain was dramatically higher than seen in an isogenic icaR mutant (Fig. 6A and B), suggesting TCA cycle activity has a greater role in regulating icaADBC transcription than does IcaR. These data demonstrate that, like reg-

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**FIG. 4.** Aconitase inactivation alters the redox and energy status of *S. epidermidis*. (A) At the indicated times during the growth cycle, intracellular concentrations of NAD and NADH were determined in *S. epidermidis* strain 1457. (B) Similarly, the concentrations of reduced and oxidized NAD were determined for strain 1457-\textit{acnA}. (C) ATP concentrations were determined for strains 1457 and 1457-\textit{acnA}. The results are presented as the means plus standard errors of the means (error bars) of duplicate determinations for at least two independent experiments.

**FIG. 5.** TCA cycle inactivation increases PIA accumulation. (A) Relative amounts of cell-associated PIA were determined using a PIA immunoblot assay for *S. epidermidis* strains 1457, 1457-\textit{acnA}, 1457-sigB, 1457-sarA, 1457-icaR, and 1457-\textit{acnA} containing pMRS2 (1457-\textit{acnA} + pMRS2). To determine the relative amounts of PIA, we defined the amount of PIA produced by strain 1457 at 2 h in one experiment as 100% and expressed all other amounts as relative to the amount at this 2-h point. The results are presented as the means plus standard errors of the means (error bars) for three independent experiments. (B) Representative immunoblot used for determining spot volumes. Strain 1457-\textit{ica}, which does not synthesize PIA, was used to determine the background density.
ulation of the lac operon, icaADBC transcription is regulated by a catabolic process, specifically, the TCA cycle. TCA cycle inactivation alters transcription of icaR, sarA, and sigB. As stated, transcription of icaADBC requires at least two positive effectors, SarA and SigB, and one negative regulator, IcaR. The dramatic increase in icaADBC mRNA levels in the aconitase mutant strain relative to the wild-type strain could be due to an increased number of positive effectors or a decreased number of negative effectors. To determine whether aconitase inactivation altered transcription of the known regulators of icaADBC transcription, we probed total RNA isolated throughout the growth cycle for changes in the transcription of sigB, sarA, and icaR (Fig. 7). Surprisingly, aconitase inactivation increased transcription or mRNA stability of both positive regulators sigB and sarA and the negative regulator icaR. These data suggest that TCA cycle inactivation can alter the mRNA levels of the known regulators of PIA synthesis; however, they fail to establish that these regulators are involved in TCA cycle-mediated regulation of PIA synthesis. To determine whether IcaR, SarA, or SigB is regulating PIA synthesis in response to TCA cycle-associated metabolic changes, icaR, sarA, and sigB regulatory mutants of strain 1457 were incubated with 400 μM of the aconitase inhibitor fluorocitric acid, and PIA accumulation was assessed relative to the parental strain 1457 (Fig. 8). The addition of fluorocitric acid to the culture medium of S. epidermidis strains 1457-sigB and 1457-icaR significantly (P < 0.01) increased the accumulation of PIA relative to the untreated control cultures. In strain 1457-sarA, the accumulation of PIA was similar irrespective of the presence of fluorocitric acid in the culture medium (P > 0.05). Overall, these data suggest that TCA cycle-mediated derepression of ica transcription and PIA synthesis occur independently of IcaR and SigB. Furthermore, these data confirm that PIA synthesis requires SarA (10, 74) and at least one additional TCA cycle-responsive negative regulator.
DISCUSSION

Metabolic effect of TCA cycle activity on PIA biosynthesis. Glycolysis is the catabolic pathway that converts glucose into pyruvate, with the catabolic fate of pyruvate being determined by the availability of oxygen. Interestingly, the rate of glucose utilization by the wild-type *S. epidermidis* strain 1457 appears greater than the rate by strain 1457-acnA (Fig. 1C); however, these data fail to account for the number of bacteria in the medium. When this is taken into account, glucose utilization was greater in strain 1457-acnA than in strain 1457 (Fig. 2), suggesting that carbon flow through glycolysis might be greater in a TCA cycle mutant. If the amount of carbon flowing through glycolysis were greater in a TCA cycle mutant than in the parental strain, then the amount of acetate in the culture medium at the time when glucose was depleted should be greater in the mutant. Glucose was depleted in the wild-type and complemented strains after 5 h of growth; however, in strain 1457-acnA, glucose was depleted after 7 h of growth (Fig. 1C). Upon depletion of glucose, the concentrations of acetate in the culture medium were similar (14 to 16 mM) for all strains (Fig. 1D), suggesting that carbon flow through glycolysis was similar in the aconitase mutant and the wild-type strain. Additionally, because the concentrations of acetate were below 24 mM (if all glucose is processed through the glycolytic pathway, then a 2:1 stoichiometry of acetate to glucose should be maintained), the data confirm that carbon are flowing into other pathways. Consistent with these data, NMR metabolomics indicated that the exponential-growth-phase concentrations of acetic acid in strains 1457 and 1457-acnA were similar (Table 3). In addition to glycolysis, another major pathway staphylococci use for glucose catabolism is the pentose phosphate pathway (8, 70). The presence of glucono-1,5-lactone in the NMR metabolomic data (Table 3) suggests that carbon flow through the pentose phosphate pathway is increased in the TCA cycle mutant strain relative to the wild-type strain. One function of the pentose phosphate pathway is to provide the five-carbon intermediate for purine biosynthesis; hence, the increased carbon flow through the pentose phosphate pathway is likely linked to the decreased intracellular concentration of ATP in strain 1457-acnA relative to strain 1457 (Fig. 3). (De novo biosynthesis of ATP requires 5-phosphoribosyl-1-pyrophosphate. An important function of the pentose phosphate pathway is to provide biosynthetic intermediates for purine biosynthesis, namely, ribose-5-phosphate that is converted into 5-phosphoribosyl-1-pyrophosphate.)

NMR metabolomic data revealed an increased intracellular concentration of fructose-6-phosphate in the TCA cycle mutant relative to the wild-type strain (Table 3). Fructose-6-phosphate can undergo a transamination reaction to generate glucosamine-6-phosphate, a reaction catalyzed by glucosamine-6-phosphate synthase (*glmS*; EC 2.6.1.16). Phosphoglucomutase (*glmM*; EC 5.4.2.10) catalyzes the reversible isomerization glucosamine-6-phosphate to glucosamine-1-phosphate. Glucosamine-1-phosphate is the substrate for the bifunctional enzyme UDP-N-acetylgalactosamine pyrophosphorylase (*glmU*; EC 2.7.7.23 and EC 2.3.1.157) that catalyzes the acetylation and UTP-dependent activation of glucosamine-1-phosphate to generate UDP-N-acetylgalactosamine. NMR metabolomics confirmed that *S. epidermidis* strain 1457-acnA has a greater intracellular concentration of UDP-N-acetylgalactosamine than strain 1457 does. In addition, transcription of *glmS*, *glmM*, and *glmU* is increased more during the exponential growth phase in the TCA cycle mutant than in the wild-type strain (Fig. 3). In total, these data demonstrate that TCA cycle inactivation increased the availability of the PIA monomer, UDP-activated N-acetylgalactosamine and that this was partially due to transcriptional derepression of N-acetylgalactosamine biosynthetic genes.

In addition to PIA synthesis, UDP-N-acetylgalactosamine is critical for biosynthesis of the cell wall components teichoic acid and peptidoglycan. Inactivation of aconitase in *S. epidermidis* decreased the growth rate (Fig. 1B) and the number of bacteria produced per unit of glucose (Fig. 2), suggesting that UDP-N-acetylgalactosamine was redirected from cell wall biosynthesis to PIA synthesis (Fig. 5A). Thus, the decreased growth rate of the aconitase mutant strain was likely due to a combination of increased competition for substrate (i.e., UDP-N-acetylgalactosamine) by PIA biosynthetic enzymes and an accumulation of citric acid.

Regulatory effect of TCA cycle activity on PIA biosynthesis. Two common mechanisms by which bacteria control carbon flow are feedback inhibition of enzyme activity and regulation of enzyme synthesis. Undoubtedly, some of the metabolic changes arising from TCA cycle inactivation are the result of feedback inhibition of enzyme activity; however, by definition, the derepression of icaADBC transcription (Fig. 6A and B) must be at the regulatory level. Although the prospect that TCA cycle enzymes directly regulate icaADBC transcription is a possibility, it is more likely that regulation is being mediated by regulatory proteins responding to changes in the metabolic status of the bacteria. Metabolic regulation was one of the first bacterial regulatory modalities identified, resulting in thousands of studies addressing all aspects of regulation. This long history of research into metabolic regulation has identified a large number of regulatory proteins capable of responding to changes in the intracellular concentrations of molecules associated with TCA cycle activity, such as NADH (e.g., Rex [4], NmrA [39], and CcpA [21]), citric acid (CcpC [30, 33]), glutamate (GadX and GadW [75]), branched-chain amino acids (Cody [61]), and ATP (KinA [69]). These observations lead us to hypothesize that a regulatory protein or proteins respond to TCA cycle-associated metabolites and regulate icaADBC transcription and PIA synthesis.

TCA cycle inactivation increased the transcription or mRNA stability of the known regulators of icaADBC transcription, specifically, sigB, sarA, and icaR (Fig. 7). These data raise the possibility that TCA cycle-induced alterations in the metabolic status of the bacterium might be “sensed” by σB, SarA, or IcaR. Incubation of *S. epidermidis* strain 1457-sigB with a low concentration of the TCA cycle inhibitor fluorocitric acid restored PIA synthesis to wild-type levels (Fig. 8). This fluorocitrate-induced derepression of PIA synthesis in the sigB mutant strain is similar to that observed in an *S. epidermidis* rsbU mutant strain (RsbU is an activator of σB) treated with a PIA synthesis-stimulating concentration of ethanol (34). In that study, the authors found that ethanol could restore cell-associated PIA accumulation in an rsbU mutant strain to a level equivalent to that of the nonstimulated wild-type strain. Because ethanol is an inhibitor of TCA cycle activity (6, 37), we
hypothesize that the PIA synthesis-stimulating properties of ethanol may be due to its TCA cycle inhibitory properties.

Inactivation of sarA in S. epidermidis dramatically decreases icaADBC transcription (74) and PIA accumulation (26, 74) and results in a biofilm-negative phenotype (10). Because SarA is thought to be essential for PIA synthesis (74), the more likely outcome of incubating S. epidermidis strain 1457-sarA with fluorocitic acid would be equivalent PIA accumulation between the wild-type and sarA mutant strains. As expected, the amount of PIA produced by strain 1457-sarA was similar irrespective of the presence of fluorocitic acid (Fig. 8). These data are consistent with SarA being essential for PIA synthesis and/or being required for responding to TCA cycle-associated signals.

TCA cycle inactivation increased transcription or mRNA stability of icaR relative to the wild-type strain (Fig. 7). IcaR is a transcriptional repressor of icaADBC that was originally hypothesized to be involved in environmental regulation of PIA synthesis (9). For these reasons, IcaR appeared to be a possible candidate for responding to TCA cycle-induced metabolic changes. If IcaR were responding to TCA cycle-associated metabolic changes, then in an icaR mutant background, the synthesis of PIA should be unaffected by the presence of fluorocitic acid. Growth of strain 1457-icaR in the presence of a low concentration of fluorocitic acid significantly increased the accumulation of PIA (Fig. 8), demonstrating that a second repressor is involved in regulating PIA synthesis. It is likely that this second repressor is responding to TCA cycle-associated metabolic signals.

NMR metabolomics (Table 3) indicated that the intracellular concentrations of branched-chain amino acids were greater in the aconitate mutant strain than in the wild-type strain. In gram-positive bacteria, the intracellular concentrations of branched-chain amino acids control the expression of numerous genes, a mechanism requiring the CodY regulatory protein (25, 61, 66, 72). The affinity of CodY for its operator site is enhanced in response to high intracellular concentrations of GTP and branched-chain amino acids (54, 61). The more likely explanations for our data are that CodY is an activator of icaADBC transcription, CodY-mediated repression is not functioning in S. epidermidis during the exponential growth phase, or a positive effector, potentially SarA, can overcome CodY-mediated repression. We recently demonstrated that CodY is a repressor of PIA synthesis in the closely related species S. aureus (45); thus, it seems unlikely that CodY is an activator of icaADBC transcription in S. epidermidis. Investigations are under way to determine the function of CodY in regulating S. epidermidis PIA synthesis.

In gram-positive bacteria, carbon catabolite repression is primarily mediated by CcpA (28). Recently, CcpA was shown to activate transcription of icaADBC and increase the accumulation of PIA in S. aureus (60). In B. subtilis, CcpA regulatory activity is controlled by interactions with phosphorylated HPt or Crh and fructose-1,6-bisphosphate or glucose-6-phosphate (67). NMR analysis indicated that TCA cycle inactivation increased the intracellular concentration of fructose-6-phosphate (Table 3). Fructose-6-phosphate is the biosynthetic precursor of UDP-N-acetylgalcosamine. In addition to its importance in PIA biosynthesis, fructose-6-phosphate can be reversibly isomerized to glucose-6-phosphate by glucose-6-phosphate isomerase, resulting in CcpA activation. Taken together, these data lead us to hypothesize that TCA cycle inactivation increases CcpA-mediated activation of icaADBC.

Conclusion. TCA cycle activity is repressed during the exponential growth phase; however, a basal level of activity is present. As evidenced by the data in this study, disruption of this basal activity can have pronounced metabolic consequences. TCA cycle activity is regulated by the availability of nutrients, oxygen, and iron (8, 62, 65, 71, 76) and by certain stress-inducing stimuli, such as heat, ethanol, and antibiotics (6, 36, 73); thus, numerous external conditions can interfere with the basal TCA cycle activity and produce metabolic changes. The linkage of TCA cycle activity and exopolysaccharide synthesis and the susceptibility of the TCA cycle to environmental inactivation lead us to speculate that one mechanism by which staphylococci perceive external environmental change is through alterations in TCA cycle activity. Thus, we hypothesize that the TCA cycle acts as a novel signal transduction pathway to translate external stimuli/conditions into intracellular signals that can stimulate or repress the activity of regulatory proteins like CodY and CcpA.

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References

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Supplemental Figure 1. NMR overlay spectra of *S. epidermidis* strain 1457 and strain 1457-
*acnA* grown for 2 hours. The spectrum for strain 1457 is displayed in red and the spectrum for
strain 1475-*acnA* is displayed in blue. Peaks of metabolites of interest are indicated within the
figure. The data are representative of spectra collected from three independent cultures.
Supplemental Figure 2. NMR overlay spectra of *S. epidermidis* strain 1457 and strain 1457-"acnA" grown for 6 hours. The spectrum for strain 1457 is displayed in red and the spectrum for strain 1475-"acnA" is displayed in blue. Peaks of metabolites of interest are indicated within the figure. The data are representative of spectra collected from three independent cultures.