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Brea D. Duval, Emory University
Anselmo Mathew, Emory University
Sarah Satola, Emory University
William M Shafer, Emory University

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Altered Growth, Pigmentation, and Antimicrobial Susceptibility Properties of *Staphylococcus aureus* Due to Loss of the Major Cold Shock Gene *cspB*\(^\dagger\)

Brea D. Duval, Anselmo Mathew, Sarah W. Satola, and William M. Shafer\(^1,3,*\)

*Departments of Microbiology and Immunology*\(^1\) and Medicine, Emory University School of Medicine, Atlanta, Georgia 30322, and *Laboratory of Bacterial Pathogenesis and Antimicrobial Resistance*\(^2\) and Emerging Infections Program, VA Medical Center Research Service, VA Medical Center (Atlanta), Decatur, Georgia 30033

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An insertional mutation made in the major cold shock gene *cspB* in *Staphylococcus aureus* strain COL, a methicillin-resistant clinical isolate, yielded a mutant that displayed a reduced capacity to respond to cold shock and many phenotypic characteristics of *S. aureus* small-colony variants: a growth defect at 37°C, a reduction in pigmentation, and altered levels of susceptibility to many antimicrobials. In particular, a *cspB* null mutant displayed increased resistance to aminoglycosides, trimethoprim-sulfamethoxazole, and paraquat and increased susceptibility to daptomycin, teicoplanin, and methicillin. With the exception of the increased susceptibility to methicillin, which was due to a complete loss of the type I staphylococcal cassette chromosome *mec* element, these properties were restored to wild-type levels by complementation when *cspB* was expressed in *trans*. Taken together, our results link a stress response protein (CspB) of *S. aureus* to important phenotypic properties that include resistance to certain antimicrobials.

*Staphylococcus aureus* is a major global public health problem causing serious, often life-threatening infections in the community and hospital settings that are becoming more difficult to manage with current antibiotic therapy regimens (7). The emergence of methicillin-resistant *S. aureus* (MRSA) in the hospital and community settings, coupled with the increasing number of persistent MRSA infections (25) and multidrug-resistant strains, is a growing problem not just for immunocompromised patients but also for otherwise healthy individuals. The virulence of *S. aureus* strains is multifactorial and involves the production of extracellular toxins, surface structures that mediate interaction with host cells and resistance to host defenses, transcriptional regulatory processes that control virulence gene expression, and metabolic schemes that allow for adaptation to stresses imposed by the local environment within and outside the human or animal host.

The capacity of *S. aureus* to respond to environmental stress conditions has been the subject of recent investigations (1, 33) and is imperative for its survival in hostile environments such as extreme temperature. *S. aureus* can effectively respond and adapt to a decrease in temperature by the expression of cold shock proteins (CSPs). This cold shock response likely plays a significant role in the ability of *S. aureus* to survive refrigeration and subsequently cause food-borne illnesses. Constitutive and inducible expression of CSPs is linked to a bacterial response to lower temperatures (16). CSPs have been extensively studied in both *Escherichia coli* and *Bacillus subtilis*, and their roles in DNA and RNA binding have been investigated (15, 16). These proteins belong to several diverse classes, some of which are constitutively produced while others are induced upon cold stress (16). While the role of each of these proteins is unclear, the major CSP in *E. coli* acts as an RNA chaperone that prevents the formation of undesired secondary structures during cold shock and actively promotes transcription (4). In previous communications (20, 21), we reported that mutations within or upstream of the cold shock gene *cspA* decreased pigment production by *S. aureus* strain COL through a SigB-dependent mechanism and increased bacterial resistance to a cationic antimicrobial peptide (CAP) of human lysosomal cathepsin G. The decrease in the production of the carotenoid pigment staphyloxanthin by the *cspA* mutant was of interest, as it acts as an antioxidant, protecting staphylococci from neutrophil killing (24).

At the transcriptional level, *cspB* is the major cold shock gene in *S. aureus* (1) and its expression was impacted in *S. aureus* strain A2222M, a clinical osteomyelitis isolate, by a mutation in *hemB*, which is required for hemin biosynthesis (40). Mutations in *hemB* have been linked to the small-colony variant (SCV) phenotype often displayed by *S. aureus* strains isolated from sites of persistent or antibiotic-resistant infections (36). These naturally occurring SCV subpopulations have been isolated from patients with a wide variety of infections such as device-related infections, skin and soft tissue infections, osteomyelitis, and persistent airway infections in cystic fibrosis patients (19, 35, 44). SCVs frequently require exogenous hemin or menadione for growth, which has been implicated in their reduced membrane potential (26). Other hallmark features of SCVs include their reduced level of pigmentation, resistance to aminoglycosides, and reduced production of virulence factors (19, 35, 41, 44, 46). We created a nonpolar insertional mutation in the coding sequence of *cspB* and introduced this mutation into *S. aureus* COL. A *cspB* null mutant was found to exhibit many properties previously observed with SCVs, and most of these properties were reversed

\(^\dagger\) Corresponding author. Mailing address: Research Service (VAMC), Room 5A181, 1670 Clairmont Road, Decatur, GA 30033. Phone: (404) 728-7688. Fax: (404) 329-2210. E-mail: wshafer@emory.edu.

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The optical density at 600 nm (OD600). At mid-logarithmic phase, the culture was of TSB at 37°C with shaking (200 rpm), and growth was measured by determining DNA was eluted in 100 l of AE buffer (Qiagen, Inc.), and the eluate was then with ethidium bromide, and imaged using the ChemiDoc XRS (Quantity One quantitation software; Bio-Rad, Hercules, CA), and scanning densitometry was PCR products from all transcripts were separated on a 1.5% agarose gel, stained with the transcript for an alternative sigma factor in S. aureus. Strain BD2 contains an mec element (SCCmec).

** MATERIALS AND METHODS**

**Bacterial strains, plasmids, and growth conditions.** The bacterial strains and plasmids used in this study are listed in Table 1. S. aureus COL is a clinical MRSA isolate containing the type I SCCmec (13, 18); it is the parent strain of BD1 and BD2. Strain BD1 contains an aphA-3 insertion in cspB and was created by electroporation using pBD1 as previously described by Katzif et al. (20). Strain BD2 is a complemented version of BD1 and contains a wild-type copy of mec.

**Cold shock and gene expression analysis.** S. aureus strains were grown in 50 ml of TSA at either 30°C for maintenance of the plasmid or at 42°C to select for the resulting plasmid, pBD1, was maintained in S. aureus strain RN4220 and moved into strain COL for allelic exchange. Following the method of Brückner for allelic exchange in S. aureus (5), strains containing plasmid pBD1 were grown on TSA at either 30°C for maintenance of the plasmid or at 42°C to select for the

**Materials and methods.** The bacterial strains and plasmids used in this study are listed in Table 1. S. aureus COL is a clinical MRSA isolate containing the type I SCCmec (13, 18); it is the parent strain of BD1 and BD2. Strain BD1 contains an aphA-3 insertion in cspB and was created by electroporation using pBD1 as previously described by Katzif et al. (20). Strain BD2 is a complemented version of BD1 and contains a wild-type copy of mec.

**Cold shock and gene expression analysis.** S. aureus strains were grown in 50 ml of TSA at 37°C with shaking (200 rpm), and growth was measured by determining the optical density at 600 nm (OD600). At mid-logarithmic phase, the culture was split into two 10-ml samples and these were incubated at 15°C (cold shock) or 37°C (control) for 1 h. Growth at both temperatures was monitored by OD measurement, and viability was determined by dilution plating onto TSA. In order to determine the expression of csp genes, RNA from control and cold-shocked cultures of strain COL was prepared as described by Katzif et al. (21). For reverse transcription (RT)-PCR analysis, SuperScript II reverse transcriptase (Invitrogen, Inc.) was used with 500 ng of RNA from each sample according to the manufacturer’s instructions for cDNA synthesis. AmpliTaq (Applied Biosystems, Foster City, CA) was used to generate transcriptional PCR products from the cDNA templates. The primers used to generate all transcriptional products are summarized in Table 2. The control transcripts in each strain were sigB, the transcript for an alternative sigma factor in S. aureus, and asp23, the transcript for the alkaline shock protein, which is a SigB-regulated transcript (14, 21).

**Isolation of chromosomal and plasmid DNAs.** Chromosomal and plasmid DNAs were both isolated as previously described by Katzif et al. (20). For isolation of chromosomal DNA, the DNasey Tissue kit (Qiagen, Valencia, CA) was used with the following modifications. Lysostaphin was used at a concentration of 50 µg/ml instead of lysozyme, a 1-h incubation at 37°C was used, 4 µl of RNase A (final concentration of 0.4 µg/ml) was added to each sample after lysis, and the sample was incubated at room temperature for 10 min. Chromosomal DNA was eluted in 100 µl of AE buffer (Qiagen, Inc.), and the eluate was then applied to a spin column and eluted after a 5-min incubation. High-copy plasmids were isolated from 5-ml overnight cultures of E. coli TOP10 cells (Invitrogen) using the Qiagen minipreparation technique. Low-copy-number plasmids propagated in E. coli were also isolated using the Qiagen minipreparation technique as described by Katzif et al. (21).

**Plasmid construction and genetic exchange procedures.** Plasmid pBD1 was used to introduce a nonpolar insertion using the aphA-3 cassette encoding kanamycin resistance into the open reading frame of cspB. Plasmid pBD2 was constructed to reintroduce cspB on a low-copy-number vector in trans. Plasmids were constructed essentially as previously described (2, 5, 17, 20). To construct pBD1, a 1.4-kb region containing cspB was PCR amplified from genomic DNA from strain COL. Primers cspB5241 and cspB3743S were used to amplify a fragment containing cspB with a 3′ SmaI restriction site. All restriction enzymes were obtained from Promega (Madison, WI). Primers cspB723S and cspB31617 were used to amplify a 3′ region containing cspB with an internal SmaI site. The two fragments were ligated together, and the entire fragment was cloned into pCR2.1 (Invitrogen). This plasmid was transformed into E. coli, isolated, and digested with KpnI and XbaI. The fragment containing cspB was gel purified and ligated onto plasmid pUC19 digested with KpnI and XbaI. The aphA-3 cassette was released from plasmid pUC18K by SmaI and then ligated into the Smal site of cspB coding region on plasmid pUC19. The entire cspB fragment containing the kanamycin resistance cassette was isolated, purified, and then ligated onto a temperature-sensitive E. coli-S. aureus shuttle vector, pBT2. The resulting plasmid, pBD1, was maintained in S. aureus strain RN4220 and moved into strain COL for allelic exchange. Following the method of Brückner for allelic exchange in S. aureus (5), strains containing plasmid pBD1 were grown on TSA at either 30°C for maintenance of the plasmid or at 42°C to select for the

**TABLE 1. Bacterial strains and plasmids used in this study**

<table>
<thead>
<tr>
<th>Strain or plasmid</th>
<th>Genotype</th>
<th>Source, reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>E. coli TOP10</td>
<td>F- mcrA Δ(mrr-hsdRMS-mcrCB) Δ800lacZ Δ15ΔΔlacY74 deoR recA1 araD139 Δ(lac-lev)7697 gaiU galK rpsL, (Strr) endA1 supF</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>S. aureus strains</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RN4220</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BM2284-4 (Φ11)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BD1</td>
<td>Nonpolar aphA-3 Km’ cassette insertion into 5’ end of cspB coding region</td>
<td>J. Iandolo, 23</td>
</tr>
<tr>
<td>BD2</td>
<td>Complement of strain BD1 using plasmid pBD1</td>
<td>J. Iandolo, 32</td>
</tr>
</tbody>
</table>

**Plasmids**

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Description</th>
<th>Source, reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>pBT2</td>
<td>Low-copy-number E. coli-S. aureus shuttle vector with Amp’ in E. coli and Km’ in S. aureus with temperature-sensitive origin of replication</td>
<td>5</td>
</tr>
<tr>
<td>pBD1</td>
<td>pBT2 construct containing 2.4-kb region with aphA-3 cassette inserted into 5’ end of cspB coding region</td>
<td>This study</td>
</tr>
<tr>
<td>pUC91</td>
<td>High-copy-number E. coli host; Amp’</td>
<td>47</td>
</tr>
<tr>
<td>pUC18K</td>
<td>pUC18 with aphA-3 nonpolar Km’ cassette; Amp’</td>
<td>27</td>
</tr>
<tr>
<td>pCR2.1</td>
<td>High-copy-number PCR cloning vector; Amp’ and Km’ in E. coli</td>
<td>Invitrogen</td>
</tr>
</tbody>
</table>

**TABLE 2. Oligonucleotide primers used in this investigation**

<table>
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<th>Primer</th>
<th>Sequence (5’-3’)</th>
<th>Reference</th>
</tr>
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<tbody>
<tr>
<td>asp233512</td>
<td>GACAGGAGCATCAGCAATTTT</td>
<td>This study</td>
</tr>
<tr>
<td>asp2351</td>
<td>GAAAACTTCTGGTCCACGAG</td>
<td>This study</td>
</tr>
<tr>
<td>mecA5</td>
<td>GTTGGTAGGTTGGGTTTGG</td>
<td>This study</td>
</tr>
<tr>
<td>mecA4S</td>
<td>ATATTGCGATTCATTCCCA</td>
<td>This study</td>
</tr>
<tr>
<td>mecA5</td>
<td>GTTGGTAGGTTGGGTTTGG</td>
<td>This study</td>
</tr>
<tr>
<td>mecA5</td>
<td>CGTATGGAATTGATCTCGAATAC</td>
<td>This study</td>
</tr>
<tr>
<td>mecA5</td>
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<td>This study</td>
</tr>
<tr>
<td>mecA5</td>
<td>CTTATCATGGTGGTCTTCTT</td>
<td>This study</td>
</tr>
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<td>mecA5</td>
<td>GCCGCGGATCCAGCAAATTTT</td>
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<td>mecA5</td>
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<td>mecA5</td>
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</tr>
<tr>
<td>mecA5</td>
<td>GCACGAGCAGCATAACAAAGGA</td>
<td>This study</td>
</tr>
<tr>
<td>mecA5</td>
<td>CGTATGGAATTGATCTCGAATAC</td>
<td>This study</td>
</tr>
</tbody>
</table>
double crossover with the appropriate antibiotic. Liquid cultures of \textit{S. aureus} were prepared using TSB and were grown at the appropriate temperature for either maintenance or loss of the plasmid. A similar procedure was used to create plasmid pBD2, which was used for complementation of strain BD1, but the \textit{aphA-3} cassette was not inserted into the SmaI site. Staphylococcal phage \Phi11 was used for transduction of pBD2 into strain BD1 as previously outlined by Schafer and Iandolo (42). Phage \Phi11 was induced using mitomycin C (1 \mu g/ml) and used to infect RN4220(pBD2) to obtain a transducing lysate. Transductants of strain pBD1 were selected on TSA plates containing 25 \mu g of chloramphenicol per ml. The presence of pBD2 in representative transductants was confirmed by digesting isolated plasmid DNA with KpnI and Xbal, followed by agarose gel electrophoresis.

\textbf{PFGE and Southern hybridization.} Pulsed-field gel electrophoresis (PFGE) was performed as outlined by PulseNet, the National Molecular Subtyping Network for Foodborne Disease Surveillance of the CDC (Atlanta, GA) (43). Agarose plugs containing cells were prepared from 175 \mu l of overnight cultures of each strain grown in brain heart infusion broth (Becton Dickinson Microbiology Systems, Cockeysville, MD) and digested with SmaI (New England Biolabs, Ipswich, MA). Plugs were placed into the well of a 1\% (wt/vol) SeaKem Gold (Cambrex Biosciences Rockland Inc., Rockland, ME) agarose gel and run with the following parameters: 200 V (6 V/cm), 14\(^\circ\)C, 5-s initial switch, 40-s final switch for 21 h. The gel was stained using ethidium bromide, destained, and then visualized using UV light. For Southern hybridization, a 391-bp fragment of \textit{mecA} was PCR amplified from \textit{S. aureus} COL genomic DNA using primers mecASF and mecASR and used as a probe in Southern blot analysis. This \textit{mecA} probe was prepared using the DIG DNA labeling kit (Roche Diagnostics Corporation, Indianapolis, IN) and following the manufacturer's instructions. The digoxigenin (DIG)-labeled \textit{mecA} fragment was confirmed by agarose gel electrophoresis. Southern blot analysis was performed as described by Satola et al. (39).

\textbf{Antimicrobial susceptibility and pigment determinations.} To determine the susceptibility of \textit{S. aureus} strains to aminoglycosides (amikacin and tobramycin), methicillin, and parapquat, a modified disc diffusion protocol from Chen and Morse (9) was used. Briefly, strains were grown in TSB overnight and then diluted 10-fold in TSB. Samples of 200 \mu l were then plated onto TSA and incubated at 37\(^\circ\)C for 1 h. Whatman filter paper discs (1.0 cm in diameter; Biomedia, Goettingen, Germany) were soaked in a solution of the antimicrobial agent. The discs were then placed on the surface of the TSA plate, and incubation was continued at 37\(^\circ\)C for 24 h. Zones of growth inhibition were determined by measuring the diameter of the growth inhibition. All disc diffusion assays were performed in triplicate. To determine the MIC values of daptomycin, gentamicin, teicoplanin, and trimethoprim-sulfamethoxazole (TMS), Etest strips for each of these antibiotics were prepared using TSB and were grown at the appropriate temperature for 1 h. Etest strips for daptomycin, gentamicin, or TMS were then applied in accordance with the manufacturer's instructions, with plates from each temperature being allowed to continue to grow at 37\(^\circ\)C or 15\(^\circ\)C for 1 h. The extraction was repeated once, the supernatants from each extraction were pooled into one tube, methanol was added to yield a final volume of 1 ml, and the absorbance at 465 nm was measured.

\section*{RESULTS AND DISCUSSION}

\textit{cspB} is the major cold-inducible gene in \textit{S. aureus} strain COL. Previous work by Anderson et al. (1) indicated that \textit{cspB} is the major cold-inducible gene in strain UAMS-1. Although CSPs have been implicated in maintaining the fidelity of bacterial gene expression during exposure to low temperature (4, 18), certain \textit{csp} genes are expressed at 37\(^\circ\)C and may have functions during normal growth. Earlier work by Katzif et al. (20) showed that production of CspA at 37\(^\circ\)C can influence levels of staphylococcal susceptibility to CAPs and pigment production. Interestingly, a \textit{hemB} mutant of \textit{S. aureus} strain A22223I that displayed an SCV phenotype had altered levels of \textit{cspB} expression compared to those of its \textit{hemB} \textsuperscript{+} parent (40). Accordingly, we sought to define the functions of \textit{cspB} in this pathogen. As our previous studies (20, 21) on CAP resistance and pigment production in \textit{S. aureus} were performed with strain COL, we first examined the levels of \textit{csp} transcripts in this strain when a logarithmically growing culture was shifted from 37\(^\circ\)C to 15\(^\circ\)C. Using RNA extracted from control and cold-shocked cultures of strain COL and RT-PCRs to detect transcripts from the three main \textit{csp} genes (\textit{cspA}, \textit{cspB}, and \textit{cspC}) and two control genes (\textit{sigB} and \textit{asp23}) not differentially expressed during cold shock, we determined that \textit{cspB} is the major cold shock gene expressed by \textit{S. aureus} COL (Fig. 1). We found that the \textit{csp4} transcript was more abundant than either the \textit{cspB} or the \textit{cspC} transcript in the culture maintained at 37\(^\circ\)C, but the \textit{cspB} transcript predominated in the 15\(^\circ\)C culture; the ratios obtained when comparing the peak values at 15\(^\circ\)C to 37\(^\circ\)C were 1.34 for \textit{cspA}, 2.79 for \textit{cspB}, and 1.58 for \textit{cspC}.

\textbf{Loss of \textit{cspB} in \textit{S. aureus} COL leads to a severe growth defect and reduced pigmentation.} In order to study the function of \textit{cspB} in \textit{S. aureus} COL, we created a null mutant (strain BD1) that contained a nonpolar \textit{aphA-3} insertion in the \textit{cspB} coding sequence. To verify that phenotypic differences (see below) were linked to this mutation, we also created a complemented strain (BD2) that had \textit{cspB} expressed in \textit{trans} from pBD2. Strain BD1 demonstrated many phenotypic differences compared to parent strain COL and complemented strain BD2. The most obvious differences were the smaller pinpoint colonies formed by BD1 that were less pigmented than the larger colonies from either parental strain COL or complemented

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig1.png}
\caption{Densitometry analysis of selected \textit{S. aureus} transcripts before and after cold stress. RNA was prepared as previously described, and a Bio-Rad Chemidoc XI was used to measure the density of electrophoresed cDNA. Intensity is reported in proprietary units as a function of location on the gel. The peak intensity is reported for each transcript at 37\(^\circ\)C (A) and after cold shock for 1 h at 15\(^\circ\)C (B).}
\end{figure}
strain BD2 (data not presented). The reduced level of pigmentation seen with colonies or cell pellets (data not presented) of BD1 compared to those of COL and BD2 was confirmed by measuring the level of methanol-extractable carotenoids (Fig. 2). When the growth of these strains in TSB at 37°C was monitored, we observed that the parent and complemented strains grew similarly but BD1 had a severe growth defect (Fig. 3A). Strain BD1 was also less proficient in responding to cold shock than was parent strain COL (Fig. 3B), a difference which was reversed by complementation (data not presented). The growth defect and reduced pigment properties of BD1 were stable, as spontaneous revertants could not be isolated (data not presented). We also found that additional cspB::aphA-3 mutants of COL expressed growth and pigment production phenotypes resembling those of BD1 (data not presented).

Although complementation with the wild-type cspB gene expressed from pBD1 was able to restore normal growth and levels of pigment production, we were concerned that second-site mutations in hemB and/or menD, which have been associated with the SCV property of S. aureus (45), might exist in BD1 and contribute to some of its SCV-like phenotypes. However, we found that the coding sequences of hemB and menD were identical in all three strains (data not presented). Since BD1 exhibited growth and pigment production properties resembling those of previously reported SCVs (26, 35, 46) yet had wild-type hemB and menD sequences, we concluded that multiple mechanisms contribute to the appearance of SCVs in S. aureus, including expression of cspB.

Loss of cspB impacts levels of antimicrobial resistance.

Since SCVs have been reported to display increased resistance to certain antimicrobials (19, 26, 35), notably, aminoglycosides, and BD1 exhibited SCV-like characteristics, we examined antimicrobial resistance levels of BD1 and compared them to those of parental strain COL and complemented strain BD2. BD1 was more resistant than COL or BD2 to a panel of aminoglycosides (gentamicin, amikacin, and tobramycin), TMS, and pararquat (Table 3 and Fig. 4). In contrast, BD1 was more sensitive than COL and BD2 to daptomycin, which is an anionic antimicrobial lipo-peptide whose activity depends on the presence of calcium ions (6, 10, 12), and teicoplanin (Table 3). Interestingly, however, compared to COL, both BD1 and BD2 were very sensitive to methicillin, which we subsequently found to be due to loss of the type I SCCmec (see below).

Since loss of cspB influenced the antimicrobial susceptibility profile of staphylococci, we tested if exposure of S. aureus COL to cold shock would change its level of susceptibility to antibiotics. For this purpose, a mid-logarithmic-phase culture was shifted from 37°C to 15°C and the MICs of daptomycin, gentamicin, and TMS were determined by Etest incubated at 15°C and 37°C for both the control (37°C) and cold-shocked cultures (15°C) (Table 4). For the control culture grown and maintained at 37°C, there was a nearly 4-fold increase in susceptibility to both daptomycin and TMS when the Etest assay was incubated at 15°C compared to that obtained when the Etest assay was incubated at 37°C (P = 0.0021 and 0.002, respectively). For the cold-shocked culture, there was an increase in

TABLE 3. MICs of antimicrobials used in this study

<table>
<thead>
<tr>
<th>Strain</th>
<th>Daptomycin</th>
<th>Gentamicin</th>
<th>Teicoplanin</th>
<th>TMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>COL</td>
<td>1.5</td>
<td>0.38</td>
<td>3.0</td>
<td>2.0</td>
</tr>
<tr>
<td>BD1</td>
<td>0.064</td>
<td>32</td>
<td>0.75</td>
<td>&gt;32</td>
</tr>
<tr>
<td>BD2</td>
<td>2.0</td>
<td>0.38</td>
<td>2.0</td>
<td>1.0</td>
</tr>
</tbody>
</table>

*Values were determined by using Etest strips according to the manufacturer’s instructions.
mupirocin led us to investigate the presence of the \( mecA \) gene and the full type I SCCmec in these strains. Using PCR primers to amplify the \( mecA \) gene, a product corresponding to the \( mecA \) gene (~1.6 kb) was obtained from genomic DNA of parent strain COL but not BD1 or BD2, indicating that \( mecA \) was absent in the mutant and complement. To examine the extent of the deletion in this region, PCR was used to amplify several genes (e.g., \( orfX \), \( mecA \), \( ccrA \), and \( pls \)) that spanned the entire SCCmec in COL (Fig. 5B). While these genes (\( orfX \), \( mecA \), \( ccrA \), and \( pls \)) were readily amplified from COL chromosomal DNA, PCR products were not obtained when BD1 and BD2 DNA preparations were used as templates (Fig. 5A). To further examine the extent of this deletion, PFGE and Southern blot hybridization analyses were employed. The PFGE analysis demonstrated a unique SmaI pattern of strain COL versus strains BD1 and BD2 in that the latter strains lacked an approximately 200-kb fragment that was present in strain COL (arrow in Fig. 5C). As analysis of the COL genome sequence (cmr.jcvi.org) showed that the type I SCCmec is harbored on a 204-kb SmaI fragment, we hypothesized that this roughly 200-kb fragment in COL contained the \( mecA \) cassette. Indeed, Southern blot hybridization analysis on the gel shown in Fig. 5C that used a 391-bp \( mecA \) gene fragment probe showed that this was the case and that no bands in the SmaI digests of BD1 and BD2 hybridized to the probe. Considering these findings together with our PCR analysis (Fig. 5A), we conclude that the entire type I SCCmec was deleted in BD1 and BD2, indicating that these strains have at least a 30-kb deletion compared to parental strain COL. As BD1 and BD2 have a unique band corresponding to approximately 170 kb (asterisk in Fig. 5C); we tentatively conclude that it was generated due to deletion of the complete type I SCCmec.

We have confirmed that \( cspB \) is the major cold shock gene in \( S. aureus \) COL during growth at low temperature (Fig. 1). Not unexpectedly, a \( cspB \) null mutant (BD1) of this MRSA strain had a reduced capacity to grow at 15°C. Unexpectedly, however, it displayed several properties previously reported by others (26, 35, 46) for SCVs of clinical and laboratory strains of \( S. aureus \). SCVs are phenotypically distinct subpopulations of \( S. aureus \) that have been implicated in persistent and drug-resistant infections (36). While it has been previously shown...
that mutations in certain genes (e.g., hemB and menD) involved in components of the electron transport chain (8, 22, 30, 40, 45, 46) can lead to the phenotypes seen in SCVs, the exact genetic mechanisms that allow S. aureus to accomplish this phenotypic shift remain unclear. Although BD1 shows many properties previously described in SCVs (e.g., a slow growth rate, decreased pigmentation, and resistance to aminoglycosides), there are important differences, notably, the absence of hemB or menD mutations in BD1. Given the many similarities between our cspB insertional mutant and SCVs, we propose that cspB could play a part in this phenotype shift. In this respect, Seggewiss et al. (40) noted that expression of cspB (SA2494) in S. aureus A22223I was upregulated in a model SCV hemB insertional mutant of this strain compared to its isogenic parent. Although our results seem at variance with those of Seggewiss et al. (40), it is important to note that our groups worked with different strains (COL versus A22223I). Moreover, our work was performed in a hemB+ background, while that of Seggewiss et al. was done with a hemB mutant, which might respond differently to cold shock at the transcriptional level. The altered antimicrobial susceptibility profile of BD1 compared to parental strain COL was of interest due to its increased resistance to aminoglycosides and paraquat and susceptibility to daptomycin and methicillin. While the aminoglycoside/paraquat resistance and daptomycin susceptibility properties of BD1 were reversed by complementation with the wild-type cspB gene expressed in trans, complementation failed to reestablish methicillin resistance. For reasons that remain unclear, BD1 (and other cspB null mutants of strain COL [data not presented]) had an apparent deletion of the type I SCCmec (Fig. 5). We do not yet know if CspB contributes to the maintenance of this cassette or if the deletion of the cassette was a secondary event that helps to reduce a fitness cost associated with loss of CspB production. It has been hypothesized (31) that SCCmec deletion reflects an attempt by staphylococci to gain a competitive advantage over those that still harbor an intact SCCmec.

Resistance to aminoglycosides in staphylococcal SCVs has
been previously attributed to a diminished or inadequate membrane potential (34, 36). It is unknown if this is the reason for the resistance phenotype seen in BD1 (Table 3 and Fig. 4). However, the extreme resistance to paraquat (Table 3) seen in BD1 is suggestive of this mechanism. Paraquat is a redox cycling agent that exerts its toxicity by producing superoxide anions in the presence of oxygen, which can then form other reactive oxygen species such as hydrogen peroxide and hydroxyl radicals (3). In bacteria, it has been shown that protection against paraquat toxicity can be imparted by increasing the cellular levels of superoxide dismutase (28) and that the reduced form of superoxide dismutase binds with high affinity to the ATGGG- and CCAAT sequences in single stranded oligonucleotides. FEBS Lett. 338:157–160.

These observations suggest that either (i) the positively charged reduced form of paraquat is unable to cross the bacterial membrane (37). These observations suggest that either (i) the positively charged reduced form of paraquat is unable to cross the bacterial membrane and exert its normal activity in the cytoplasm or (ii) the protection seen in BD1 is due to increased levels of superoxide dismutase. If paraquat is unable to cross the bacterial membrane due to improper membrane potential, this likely explains the increased resistance to aminoglycosides seen in BD1.

Although the role of the CSPs in S. aureus remains unclear, previous work on these proteins in E. coli suggests that they play roles as RNA chaperones (4, 18), as transcriptional antiterminators (4), and as an alternative initiation factor during translation. We are currently determining the mechanism(s) by which CspB functions in S. aureus under normal growth conditions and during cold shock so as to understand how it regulates the production of virulence factors and levels of bacterial susceptibility to antimicrobials.

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