The \( \omega \) Subunit Governs RNA Polymerase Stability and Transcriptional Specificity in *Staphylococcus aureus*

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**ABSTRACT** *Staphylococcus aureus* is a major human pathogen that causes infection in a wide variety of sites within the human body. Its ability to adapt to the human host and to produce a successful infection requires precise orchestration of gene expression. While DNA-dependent RNA polymerase (RNAP) is generally well characterized, the roles of several small accessory subunits within the complex have yet to be fully explored. This is particularly true for the omega (\( \omega \) or RpoZ) subunit, which has been extensively studied in Gram-negative bacteria but largely neglected in Gram-positive counterparts. In *Escherichia coli*, it has been shown that ppGpp binding, and thus control of the stringent response, is facilitated by \( \omega \). Interestingly, key residues that facilitate ppGpp binding by \( \omega \) are not conserved in *S. aureus*, and consequently, survival under starvation conditions is unaffected by *rpoZ* deletion. Further to this, \( \omega \)-lacking strains of *S. aureus* display structural changes in the RNAP complex, which result from increased degradation and misfolding of the \( \beta' \) subunit, alterations in \( \delta \) and \( \sigma \) factor abundance, and a general dissociation of RNAP in the absence of \( \omega \). Through RNA sequencing analysis we detected a variety of transcriptional changes in the *rpoZ*-deficient strain, presumably as a response to the negative effects of \( \omega \) depletion on the transcription machinery. These transcriptional changes translated to an impaired ability of the *rpoZ* mutant to resist stress and to fully form a biofilm. Collectively, our data underline, for the first time, the importance of \( \omega \) for RNAP stability, function, and cellular physiology in *S. aureus*.

**IMPORTANCE** In order for bacteria to adjust to changing environments, such as within the host, the transcriptional process must be tightly controlled. Transcription is carried out by DNA-dependent RNA polymerase (RNAP). In addition to its major subunits (\( \alpha\beta\beta'\)\( \delta \)) a fifth, smaller subunit, \( \omega \), is present in all forms of life. Although this small subunit is well studied in eukaryotes and Gram-negative bacteria, only limited information is available for Gram-positive and pathogenic species. In this study, we investigated the structural and functional importance of \( \omega \), revealing key roles in subunit folding/stability, complex assembly, and maintenance of transcriptional integrity. Collectively, our data underline, for the first time, the importance of \( \omega \) for RNAP function and cellular harmony in *S. aureus*.

**KEYWORDS** RNA polymerase subunit omega, RpoZ, *Staphylococcus aureus*, gene regulation

Transcription in all forms of life is a tightly controlled process, necessitated by the essentiality of correct temporal and spatial expression of genes for survival. All transcriptional activity within a cell is maintained by the DNA-dependent RNA polymerase (RNAP). This multiprotein complex is structurally and functionally similar in distant forms of life, displaying only minor variations in composition, e.g., the presence/absence of certain subunits (1, 2). In bacteria RNAP consists of four main subunits, i.e.,
two α (RpoA) subunits and one subunit each of β (RpoB) and β′ (RpoC), forming the α₂ββ′ complex. Together they facilitate transcriptional elongation, but they require a σ factor to initiate the process. Interestingly, a large number of bacteria, and particularly the Firmicutes, possess several other accessory RNAP subunits (3–6). These are considerably smaller than the major subunits, ranging from 8.5 to 21.5 kDa, and include the δ (RpoE), ε (RpoY), and ω (RpoZ) subunits. Deletion of these subunits does not result in lethality for the cell, and thus their diminutive size and nonessential nature have resulted in their being classified as the “small accessory subunits” (reviewed in reference 6). Nevertheless, for the δ factor it has been shown that, in various species, deletion is accompanied by a deregulation of the transcriptional process, leading to decreased fitness and impaired virulence in pathogenic organisms (7, 8). While there is an extensive history of research for δ, ε has only recently been described as an RNAP subunit, with only a single study thus far performed in Bacillus subtilis (5), which suggested a role in phage immunity. While the presence of these two subunits is largely confined to the Firmicutes, homologs of ω, the smallest of the three subunits, can be found not only in bacteria but also in eukaryotes (RPB6) and archaea (RpoK) (6, 9). Although this conservation might suggest a vital role and perhaps similar function across widely different species, there are in fact marked differences in how this subunit influences cells across the various kingdoms (6, 10). Most strikingly is the observation that while it is accessory in bacteria, the ω subunit is essential in eukaryotic organisms (11).

The majority of studies on ω have been carried out in Gram-negative bacteria, with a focus on the model organism Escherichia coli (10). In this organism it has been shown that ω influences the transcriptional machinery, and thus the transcriptional process, in a variety of ways. Most notably, ω is known to interact with β′ to ensure correct folding of the subunit, as well as to facilitate docking to the α₂β complex (9, 12–14). Accordingly, deletion of ω leads to misfolding as well as degradation of β′ (13, 15), which is also observed in Mycobacterium smegmatis (16). The crystal structure of ω in complex with the other RNAP subunits of both E. coli and Thermus aquaticus has been solved and confirms the binding of ω to β′ (17, 18). Interestingly, these structures also reveal species-specific differences in the interaction of these two proteins, again highlighting the heterogeneous nature of ω function in different organisms. In contrast, no such crystal structure is available for ε or RNAP at large, in Gram-positive organisms.

Quite strikingly, in terms of functional difference, is the finding that in E. coli, the stringent response, and thus adaption to nutrient-limiting conditions, is dependent on ω. Specifically, the stringent response-inducing molecule ppGpp is recognized by and binds to the interface of β′ and ω (19–21), which in turn leads to the adjustment of transcriptional profiles to promote survival under nutrient-limiting conditions. While this role is true for E. coli, the ω protein in B. subtilis has been suggested to have no such role in the stringent response, due to an alternative mechanism of ppGpp recognition. Instead of binding to ω, the adaption to limiting conditions in B. subtilis is mediated by ppGpp-induced alterations of GTP concentrations within the cell (22, 23). Subsequently, these changes lead to alterations in gene expression, driven by the sensitivity of certain promoters to GTP availability as an initiating nucleotide. In line with this model, whereas onset of the stringent response does not require the interaction of ppGpp with the RNAP complex, is the observation that the conserved E. coli residues required for ppGpp binding to RNAP are largely absent in the B. subtilis ω and β′ subunits (20).

The final major function of ω described in the literature is a putative role in facilitating σ factor binding to the RNAP complex. For E. coli and cyanobacteria, it has been reported that depletion of ω can lead to increased binding of alternative σ factors and in turn to increased expression of genes within alternative σ factor regulons (24, 25). Again, structural differences within the ω-depleted RNAP have been implicated in this alteration of σ factor affinity for the complex. As with many of the other ω phenotypes, no in-depth studies have been performed in Gram-positive bacteria regarding this role, further underscoring the need to characterize this diverse protein.

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For Gram-positive organisms, only a limited number of studies exist, detailing a few phenotypic effects resulting from the abrogation of \( \omega \) activity. Indeed, none of these studies have unraveled the molecular basis for alterations in mutant strains, meaning that the role of \( \omega \) in Gram-positive species is still relatively elusive. Those effects that have been detailed for \( rpoZ \) mutants include alterations in cell wall morphology, cell motility, protein secretion, and biofilm formation (16, 26–29). Importantly, the role of this subunit in the virulence of pathogenic species has yet to be evaluated. Therefore, in this study we explored the role of \( \omega \) in \textit{Staphylococcus aureus}, demonstrating that it is not involved in the stringent response but instead mediates structural integrity of the RNAP complex. Deletion of this factor leads to individual RNAP subunit degradation and an induction of cellular stress responses. The latter effect was characterized by global transcriptional analyses, revealing that a significant number of the observed changes were due, at least in part, to altered \( \sigma \) factor abundance in the RNAP complex. Finally, we demonstrate that deletion of \( rpoZ \) influences the ability of \textit{S. aureus} to form biofilms, a process that mediates persistent infections and the capacity to resist antibiotic treatment. We suggest that collectively, our data underline the importance of \( \omega \) for RNAP stability, function, and cellular physiology in \textit{S. aureus}.

**RESULTS**

\( \omega \) is cotranscribed with \textit{gmk} and is highly expressed throughout growth. Analysis of RNA sequencing data previously generated by our laboratory (31) reveals that the \( \omega \)-encoding gene, \textit{rpoZ}, is highly expressed during all phases of growth and is seemingly cotranscribed with its upstream gene, \textit{gmk}, encoding a guanylate kinase (see Fig. S2A in the supplemental material). To validate the latter observation, we performed Northern blot analysis on RNA extracted from wild-type \textit{S. aureus} cells grown to mid-exponential phase using \textit{rpoZ}-specific probes. In so doing, a band of \(~\sim 1100\) nucleotides (nt) was observed, indicating that \textit{rpoZ} and \textit{gmk} are indeed organized in a bicistronic operon (Fig. S2B). In order to study the role of \( \omega \) in \textit{S. aureus}, we next created an unmarked deletion mutant, removing the majority of the \textit{rpoZ}-encoding gene and leaving \textit{gmk} intact. Validation of this arrangement and that no deleterious effects were observed on \textit{gmk} expression was obtained by RNA sequencing analysis of the mutant strain and by Northern blotting (Fig. S2A and B). RNA sequencing data for the \( \omega \)-depleted strain showed the expected deletion of the \textit{rpoZ} gene but confirmed that there were no unintended effects on transcript abundance for \textit{gmk} (see Table S1 in the supplemental material).

The \textit{S. aureus} \( \omega \) and \( \beta' \) proteins lack conserved residues required for ppGpp binding. In \textit{E. coli} the interaction of ppGpp with RNAP is mediated by conserved residues within the \( \omega \) and \( \beta' \) subunits, thus controlling the stringent response. Such residues are absent in \textit{B. subtilis}, suggesting an alternate mechanism for stringent response regulation in Gram-positive organisms (20, 32). In corroboration of this, alignment analysis of \( \omega \) and \( \beta' \) subunits from \textit{E. coli}, \textit{B. subtilis}, and \textit{S. aureus} (Fig. 1A) reveals that, although there is partial conservation of the \( \beta' \) and \( \omega \) proteins for all three organisms, the residues for ppGpp binding are almost entirely absent (a single aspartic acid is conserved in \( \beta' \) region 2) at the primary sequence level and in the biochemical characteristics of each amino acid. This finding is further validated by the observation that growth of a \( \Delta rpoZ \) strain is not impaired under standard, nutrient-limiting, or stringent response-inducing conditions. Specifically, in addition to growth in complex medium (tryptic soy broth [TSB]) (Fig. 1B), we investigated growth during amino acid limitation (Fig. 1C) as well as in amino acid-limiting medium completely depleted of valine and leucine (Fig. 1D). Importantly, the \( rpoZ \) mutant showed growth rates similar to those of the wild type in both TSB and amino acid-limiting medium and only a minor growth defect in medium devoid of valine and leucine. This defect, however, suggests an impaired ability to adapt to changing growth conditions rather than a stringent response defect, as stringent response-deficient mutants display a characteristic stalling of growth during the stringent response, rather than delayed growth (33). To further validate this finding, we assessed sensitivity to mupirocin of the \( rpoZ \) mutant alongside...
its parental strain and a codY mutant strain, as it has previously been shown that S. aureus mutants impaired in the stringent response display decreased resistance to this stringent response-inducing antibiotic (34). In so doing, we found that the ΔrpoZ strain has a mupirocin MIC of 0.32 μg/ml similar to that of the wild type, while the codY mutant has a 2-fold decrease in MIC (0.16 μg/ml), comparable to that of stringent response-deficient mutants documented elsewhere (34).

Depletion of rpoZ leads to destabilization of the RNAP complex. We next assessed whether ω has a role in maintaining RNAP complex integrity in S. aureus and, more precisely, in the folding of its subunits, as described for M. smegmatis and E. coli (13, 16). We isolated cytoplasmic proteins from exponentially growing cells and determined the abundances of the intact RNAP complex in the wild-type and mutant strains. This was achieved by concentrating fractions using a 100-kDa-cutoff filter, ensuring that the resulting retentate harbored only proteins of >100 kDa in size. As β and β′ are the only RNAP subunits larger than 100 kDa, any subunit found in the concentrated fraction must originate from protein in complex with other RNAP subunits. Consequently, β and β′ are expected to be present independently and in complex with RNAP and thus should demonstrate no alteration within the mutant even if complex instability is observed. As expected, we consistently observed equimolar amounts of β for both the wild type and the rpoZ mutant in tested samples (Fig. 2A), which was confirmed by Western blotting using a β-specific antibody (Fig. 2B). Conversely, we found a marked decrease in β′ abundance in the mutant compared to the parent in the same samples. Given the large size of β′, these results point not only toward a release of the subunit from the complex but also to its likely degradation in the absence of ω, as observed in other organisms. Similar to β′, but to a lesser extent, the α subunit, the housekeeping σ factor, σA, and the small subunit δ were also decreased in abundance in the mutant strain. In order to investigate the fate of each subunit, we performed additional
concentration experiments using a 30-kDa-cutoff filter and again isolated proteins from the wild-type and \( rpoZ \) mutant strains (Fig. 2C). Interestingly, while only trace amounts of \( \beta^+ \) were detected in these fractions (as expected), we observed significant levels of \( \beta^+ \) in the \( rpoZ \) mutant strain, again suggesting \( \beta^+ \) instability upon \( rpoZ \) deletion. With regard to \( \delta \), which should always be found in the \( <30\text{-kDa} \) fraction, the amount of this protein was significantly higher in the \( \Delta rpoZ \) strain than in the wild type. This would tend to indicate disassembly and release of this subunit from the RNAP complex upon \( rpoZ \) depletion.

To exclude the possibility that alterations in subunit abundance were caused by differential expression patterns for RNAP-encoding genes in the \( rpoZ \)-depleted strain, we analyzed RNA sequencing data sets of the USA300 parent and its corresponding \( rpoZ \) mutant (Fig. 2D). The \( rpoZ \) gene was included as an internal control for our data set, as a portion of the locus is still present in the mutant strain (Fig. S2A), and therefore an expression value is generated from the RNA sequencing experiment. Of note, all of the subunits were shown to have a moderate increase in expression in the mutant strain. Importantly, \( \beta^+ \), which displayed the strongest decrease in protein abundance, showed the strongest upregulation, with a 2.6-fold increase in transcription. This perhaps suggests a possible attempt by the cell to counteract \( \beta^+ \) degradation by enhancing transcriptional activity of the \( rpoC \) gene within the \( rpoZ \) mutant strain.

**The \( \beta^+ \) and \( \omega \) factors of RNAP have distinct and contrasting influences on \( S. aureus \) gene expression.** When further exploring the RNA sequencing comparison of the wild-type and \( rpoZ \) mutant strains, we observed 232 genes with altered expression at \( \geq 2\text{-fold} \) and 63 genes at \( \geq 3\text{-fold} \) (Fig. 3; Table S1). In order to validate these alterations, we performed quantitative reverse transcription-PCR (RT-qPCR) for a subset of differentially expressed genes (Fig. S3A and B), revealing similar changes in this and the RNA sequencing data sets. Since our RNAP composition studies showed that \( rpoZ \) deletion also results in \( \delta \) depletion from RNAP, we next assessed whether removal of \( \delta \) from the transcription complex was the driving force behind the observed changes. This is particularly important, as our previous studies have shown that \( \delta \) is a key factor
for maintaining transcriptional specificity within the S. aureus cell (8). Upon comparing RNA sequencing-derived transcriptional changes of the rpoE and rpoZ mutants, we found an overlapping regulon of only 24 genes (Fig. 3). Comparing these 24 genes to the overall number of genes changed in both mutants, only \(~10.5\%\) (\(\Delta rpoZ\)) or 12.5\% (\(\Delta rpoE\)) of each data set is identical. This low level of regulon overlap indicates that the
changes seen in an \( rpoZ \) mutant are not the result of an inability of \( \delta \) to exert its influence on RNAP.

\( \omega \) influences \( \alpha \) factor recruitment to the \( S. aureus \) RNAP complex. In \( E. coli \) and the cyanobacterium \( Synechocystis \), it is known that depletion of \( \omega \) causes alterations in \( \alpha \) factor abundance within the RNAP complex. Therefore, we next investigated if the multifaceted transcriptional changes of the \( rpoZ \) mutant were influenced by alterations in \( \alpha \) factor binding to the \( \omega \)-less RNAP. Since \( \alpha^B \) is the major alternative \( \alpha \) factor in \( S. aureus \), we compared transcriptome changes upon \( rpoZ \) deletion to those observed in a \( \sigma^B \) mutant. This was facilitated using wild-type and \( \sigma^B \) mutant RNA sequencing data sets previously produced in our lab in the SH1000 background. SH1000 was chosen for this study as it is known to have a \( \sigma^B \)-overexpression phenotype. Subsequent comparison of these data sets detected an inverse relationship between the regulons of \( \omega \) and \( \sigma^B \). Specifically, for the 20 most strongly downregulated genes in the \( \sigma^B \) mutant, the majority (70%) demonstrated increased expression upon \( rpoZ \) disruption (Fig. 4; Tables S2 and S3). Conversely, for the 20 most strongly upregulated genes in the \( \sigma^B \) mutant, most (60%) were found to be decreased in expression in the \( \Delta rpoZ \) strain.

It is noted that the comparison of regulons between different strains is not without drawbacks. Our rationale was that by choosing a \( \sigma^B \)-overexpressing strain such as SH1000, we could obtain better resolution for the identification of \( \sigma^B \)-dependent genes. Nevertheless, there remains the possibility that additional genomic differences between SH1000 and USA300 drive strain-specific expression patterns and thereby influence their corresponding \( \sigma^B \) regulons. However, the strongly correlated inverse relationship observed suggests that there are indeed higher levels of \( \sigma^B \) within the \( \omega \)-lacking RNAP complex, which corroborates effects seen in other bacterial species. Accordingly, it appears that \( \omega \) not only is required to maintain RNAP integrity but also plays a role in association of the complex with available \( \alpha \) factors, thereby influencing transcriptional stringency in \( S. aureus \).

Depletion of \( rpoZ \) leads to decreased transcriptional specificity that influences multiple cellular processes within the \( S. aureus \) cell. We next used functional clustering to investigate alterations in gene expression upon \( rpoZ \) disruption (Fig. 5). While a variety of cellular processes were shown to be affected by \( rpoZ \) deletion, we noted major changes in genes related to central dogma. Specifically, genes related to transcription and translation, as well as cellular processes connected to DNA, RNA, or protein synthesis, were strongly affected by the loss of \( \omega \). These include changes within purine and pyrimidine metabolism, enzymes for DNA replication and repair, and amino acid biosynthesis. Additionally, genes in more general ontological categories, such as genes for sugar uptake systems (phosphotransferase system [PTS]), the general stress

![Figure 4](http://jb.asm.org/)

**FIG 4** \( \omega \) affects \( \alpha \) factor recruitment to the \( S. aureus \) RNAP complex. A comparison of RNA sequencing data sets from \( \sigma^B::tet \) and \( \Delta rpoZ \) strains is depicted. Shown are the most highly repressed (A) or highly induced (B) genes in the \( \sigma^B::tet \) strain and their concomitant change in expression upon \( rpoZ \) deletion. Genes with a change of less than 10% between the \( rpoZ \) mutant and its parental strain were considered unchanged.
response, regulators, transporters, and metabolism, displayed altered expression. Interestingly, the changes also negatively affect several well-characterized virulence determinants of *S. aureus*. These include aureolysin (*H11002* 3.7-fold) and the Panton-Valentine leukocidin-encoding operon (LukSF, −3.4-fold and −3.1-fold, respectively). Lastly, a significant number of phage genes were altered in their expression. This point is particularly interesting as we have previously described this phenomenon to be one of the hallmarks of *rpoE* deletion (8); thus, these effects could result from its depletion from the RNA polymerase complex. Therefore, although overlap of the *H9254* and *H9275* regulons is limited (as described above) and unlikely to be causative for the majority of transcriptional changes observed in the Δ*rpoZ* strain, it cannot be entirely excluded that a number of effects seen in the *rpoZ* mutant are driven, at least in part, by *H9275* depletion from the RNA polymerase complex. Disruption of the *H9275* subunit results in an impaired ability to circumvent multiple forms of environmental stress. To explore the physiological outcome of these findings, we sought to determine if the mutant was impaired in resisting the impact of environmental stressors. In so doing, we observed that the mutant shows an augmented sensitivity toward various antibacterial compounds with unrelated mechanisms of action. These include increased sensitivity to triclosan (2-fold; a fatty acid biosynthesis inhibitor), erythromycin and pyrogallol (both 4-fold; a translation inhibitor and a reducing agent, respectively), and diamide (8-fold; creates disulfide stress). As these compounds all affect different cellular targets, it would appear that depletion of *ω* leads to widespread fitness defects, highlighting its global importance within the *S. aureus* cell.

The *ω* subunit of RNA polymerase is required for biofilm formation in *S. aureus*. We next set out to investigate whether the observed changes in gene expression influence pathophysiologically relevant processes. Thus, we determined whether the formation of a bacterial biofilm, a key hallmark of *S. aureus* infection (35), was influenced by *rpoZ* deletion in *vitro*. Importantly, upon analysis we observed that loss of *ω* does indeed lead to an approximately 3-fold reduction in biofilm formation that could be restored via complementation in trans (Fig. 6). As biofilm formation is strongly influenced by extracellular proteins in *S. aureus* (36), we analyzed the secretomes of the wild-type and *rpoZ* mutant strains. Although the vast majority of proteins appeared to be unchanged in abundance, we noted a marked decrease in a major protein band at around 75 kDa in the mutant strain (see Fig. S4 in the supplemental material). In order to identify this protein, we excised the corresponding band from gels and performed mass spectrometric (MS) analysis. It was identified as two separate tributyrin lipases (the products of
SAUSA300_0320 and SAUSA300_2603), with molecular masses of 76 and 77 kDa, respectively. Interestingly, a recent report on the role of lipase activity in *S. aureus* demonstrated that deleting these enzymes results in drastically impaired biofilm formation (37). This may provide some explanation for these findings, although given the complex process of biofilm formation in *S. aureus*, it is likely that other factors are also involved.

**DISCUSSION**

The ω subunit of RNAP is a widely distributed protein that is found in all branches of life. While it is well studied in Gram-negative bacteria (10) and eukaryotes (9), a recent review by our group highlighted the need for ω to be examined in a wider range of species, and particularly Gram-positive bacteria (6). This is especially true since (i) the subunit has a variety of possible roles that have received only limited attention in Gram-positive species, including RNAP subunit folding (9, 12, 13, 15) and stability (16), complex assembly/stabilization (10, 13, 14), maintenance of σ factor specificity (24, 25), and ppGpp binding (19–21), and (ii) Gram-positive species, and particularly the *Firmicutes*, include several epidemiologically relevant human pathogens. To the latter point, an understanding of cellular factors that are required for transcription and to maintain transcriptional stringency is crucial in order to better comprehend the physiology of pathogens and thus potentially aid in counteracting and preventing the spread of disease. Here we describe the role of ω within *S. aureus* for the first time. Our study explores previously documented functions from other bacterial species, assessing its impact on the global transcriptome as well as its key role in fitness and, most importantly, virulence, which to our knowledge has not yet been examined.

It has previously been suggested that the stringent response in *Firmicutes* is independent of ppGpp recognition by ω (32). Here we demonstrate that under amino acid-limiting conditions, as well as during exposure to stringent response-inducing compounds, the ΔrpoZ strain does not exhibit typical stringent response deficiency phenotypes. This is consistent with findings in *B. subtilis*, where ppGpp, instead of directly binding to ω as in Gram-negative species, exerts its function indirectly. Instead, the generation of ppGpp from GTP and ATP (38) in *B. subtilis* leads to diminished GTP pools within the cell (39, 40), which in turn causes decreased transcription of GTP-sensitive promoters (i.e., those with GTP at position +1 [22, 41, 42]) and interferes with the role of GTP as a corepressor for CodY (43). These findings are mirrored in *S. aureus*, where CodY- and GTP-regulated genes are similarly influenced by the nutritional status of the cell in a ppGpp-dependent manner (33, 44). Therefore, we hypothesize that rather than resulting from a dysfunctional stringent response, the diminished ability of a ΔrpoZ strain to survive challenging conditions is a consequence of basal stress levels.

![FIG 6](http://jb.asm.org/)

The ω subunit of RNAP is required for biofilm formation in *S. aureus*. Biofilm formation by the wild-type, rpoZ mutant, and complemented strains was measured using a 24-well plate biofilm assay. Error bars show SEM, with statistical significance measured using Student’s *t* test (**, *P* < 0.01).
experienced by the cell upon loss of the subunit. This is supported by our findings that
*S. aureus* lacking the *rpoZ* gene displays a reduced-growth phenotype not only when
exposed to nutrient-limiting and stringent response-inducing conditions but also
during growth at elevated temperatures (data not shown).

To investigate how a lack of *ω* affects the transcriptional process and integrity of
RNAP, we performed complex stability analysis and determined that *rpoZ* deletion
causes decreased levels of intact RNAP, perhaps as a result of reduced *ω* within the
complex. This is consistent with findings in other organisms, which describe a
chaperone-like function for *ω*, assisting in the folding of *β* and its subsequent docking
to RNAP (9, 10, 12–15). Indeed, the likely misfolding of *β* leads not only to degradation
of the subunit but also to impaired assembly and/or dissolution of RNAP and the
proteolysis of several other subunits. Importantly, in addition to this apparent failure of
*β* to fully associate with RNAP, *ω* deletion leads to a jettisoning of *δ* from the
transcription complex and an increase in alternative *σ* factor binding. Indeed, the
finding that *δ* and *σA* are preferentially released from *ω*-less, and presumably structur-
ally unstable, RNAP suggests that *ω*-deficient RNAP can provide a unique insight into
the effects of stress on the transcription complex and, more broadly, the role of the
accessory subunits. Based on the current literature, as well as our findings here, we
propose a model whereby stress within the cell leads to RNAP instability, resulting in
an alteration of subunits present within the transcription complex that specifically
facilitates survival during unfavorable conditions (Fig. 7). In support of this, those sigma
factors that have increased binding to RNAP upon *ω* deletion (*α5* in *E. coli* [24], *αD*/*αE*
in cyanobacteria [25], and *αB* in *S. aureus*) are all major components of the general
stress response in their respective organism (45–49). Furthermore, as with *σ* factors, the
release of *δ* from RNAP has the potential for profound alterations in gene expression,
as a recent publication demonstrates the ability of *δ* to selectively affect individual
promoters (7, 8) via binding to specific promoter elements (50). As the disruption of
*rpoE* results in slower bacterial growth in a variety of organisms (e.g., an extended lag
phase [8, 51, 52]), removal of this subunit from RNAP may actually present a survival
advantage during stress, redirecting resources away from division and toward repair
and cellular maintenance. In agreement with this is a recent study with *Streptococcus
pneumoniae* biofilms, where there appears to be selective pressure for *rpoE* mutations
leading to small-colony variant (SCV) phenotypes (53). SCV formation is a common
mechanism used by many different bacteria which would initially appear to present a
defect in viability but in actuality results in enhanced survival (54) and antibiotic
resistance and the development of persister cells (55). Interestingly, our group has
observed similar effects in *S. aureus*, where *δ*-deficient mutants display variable, and
decreased, colony size after recovery from murine models of infection (unpublished
observation). Thus, it would appear that the significant compositional rearrangement
within *ω*-less RNAP is far from random. Rather, these changes in binding of accessory
factors are seemingly directed to specifically adjust promoter selectivity and preference
to facilitate survival. Furthermore, these findings regarding subunit swapping within
the transcription complex appear to confirm long-made contentions regarding a level
of cooperativity for *δ* and *σ* factor binding within RNAP (56–58).

![FIG 7 Model for the role of *ω* in the Gram-positive RNA polymerase.](http://jb.asm.org/)

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In spite of these changes in subunit abundance, which we suggest are utilized by the cell as a survival strategy, rpoZ deletion itself results in wide-reaching cellular stress that is associated with an impaired transcriptional process. This contention is supported by our RNA sequencing data, which demonstrate the upregulation of various genes coding for transcriptional and translational factors, putatively in an attempt to overcome diminished or inefficient RNAP function. Furthermore, we identified increased expression of genes involved in the general stress response, sugar uptake pathways, and energy metabolism, all of which may be designed to overcome unfavorable conditions (59). The latter group emphasizes that intracellular stress resulting from ω deletion, and the subsequent countermeasures it elicits (e.g., production of chaperones, RNAP subunits, and ribosomal subunits), consumes energy and therefore has the potential to deplete cellular energy pools. This in turn leads to the upregulation of sugar uptake and energy metabolism, which in our study resulted in a minor increase in ATP pools (data not shown). In accordance with this, studies investigating S. aureus heat stress demonstrate that following exposure to elevated temperatures, cellular energy levels are in fact not depleted, but rather a steady (60) or transient (61) increase in ATP pools is observed, presumably due to the upregulation of energy-generating processes. It is perhaps not surprising that the transcriptional and physiological changes observed from our in vitro experiments also translate into the impairment of virulence-related processes, as shown by the impaired biofilm formation in the rpoZ mutant.

We suggest that, collectively, our findings extend the existing knowledge regarding ω and its role within RNAP, provide a unique insight into structural rearrangements within the transcription complex in response to stress, and demonstrate the necessity of this important subunit for the cellular homeostasis.

MATERIALS AND METHODS

Bacterial strains and growth conditions. All bacterial strains used in this study are listed in Table 1. Unless otherwise indicated, cultures were grown as described by Shaw et al. (62). Briefly, S. aureus was grown at 37°C with shaking (250 rpm) in tryptic soy broth (TSB) or chemically defined medium (CDM) (45), while E. coli was cultivated in lysogeny broth (LB) with agitation (250 rpm) at 37°C. Overnight cultures of S. aureus were seeded at a 1:100 dilution into fresh medium. After 3 h of growth, cultures were used to inoculate fresh TSB to an optical density at 600 nm (OD₆₀₀) of 0.05 and grown to the desired density in a Synergy2 plate reader (BioTek) in 96-well plates.

Mutant construction. In order to investigate the role of the ω subunit, a strain carrying a markerless deletion of the rpoZ gene was created, using a method based on the plasmid pJ138 as outlined by Bose and coworkers (63). Briefly, the majority of the rpoZ gene was removed via allelic replacement, as initially described elsewhere (64). To do this, both the 5' and 3' ends of the rpoZ gene, together with ~1 kb of up- and downstream DNA, were PCR amplified using primers OL2889 to OL2892 and fused together using MluI sites contained in the amplification primers. This was then cloned into pJ138 using SacI and KpnI sites in primers OL2889 and OL2892. Following several selection and counterselection steps in S. aureus as described previously (63), rpoZ deletion was confirmed using internal (OL2991 and OL2992) and external (OL3008 and OL3009) primer pairs relative to the deleted region within the gene.

Complement construction. In order to exclude that secondary mutations were causative for phenotypic changes observed in the ΔrpoZ strain, a plasmid was constructed to complement the rpoZ gene in trans. Here, the complete rpoZ operon, including its native promoter, was PCR amplified using primers OL3753 and OL3807. After BamHI digestion of the amplicon and the pMK4 shuttle vector, the PCR fragment and plasmid were ligated together and transformed into electrocompetent E. coli DH5α. The plasmid isolated from ampicillin-resistant clones was confirmed via Sanger sequencing (using standard M13 primers). Correct plasmids were transformed into electrocompetent S. aureus RN4220 (65) and finally transduced into the S. aureus USA300 rpoZ mutant using φ11 (66).

RNA isolation and Northern blots. The detection of RNA transcripts was performed using a Northern blotting protocol outlined by Caswell et al. (67). Briefly, RNA from cultures grown for 3 h was isolated using an RNaseasy kit (Qiagen), with DNA removed using a Turbo DNA-free kit (Ambion). RNA concentrations were determined using an Agilent 2100 Bioanalyzer and an RNA 6000 Nano kit (Agilent). RNA (15 μg) was separated by gel electrophoresis (10% polyacrylamide containing 7 M urea and 1× Tris-borate-EDTA [TBE]) before being transferred to an Amersham Hybond N+ membrane (GE Healthcare) by electroblotting. In order to cross-link samples, membranes were exposed to UV radiation. Following this, membranes were prehybridized (1 h, 43°C) in ULTRAhyb-Oligo buffer (Ambion) and incubated (16 h, 43°C) with a 32P-labeled oligonucleotide (OL3377) specific to the rpoZ target sequence. Labeling was carried out using T4 polynucleotide kinase (Thermo Scientific) according to the manufacturer’s protocol. After overnight incubation, each membrane was washed (30 min, 43°C) with decreasing (2×, 1×, and 0.5×) concentrations of SSC buffer (300 mM sodium chloride, 30 mM sodium...
Finally, membranes were exposed to X-ray film in order to detect the size and abundance of the rpoZ transcript.

RNA sequencing and data analyses. RNA sequencing was performed for rpoZ and sigB mutants and their respective parental strains as previously outlined by our group (8, 68–70). Briefly, RNA was isolated and the quality determined as described for Northern blot experiments. Following this, RNA from three biological replicates was pooled in equimolar amounts and rRNA removed by consecutive treatment with the MICROBExpress (Ambion) and RiboZero (Epicentre) kits. Removal of rRNA was confirmed using an Agilent 2100 Bioanalyzer (RNA 6000 Nano kit; Agilent). The rRNA-depleted RNA samples were prepared for sequencing with an Ion Personal Genome Machine (PGM) system as described previously (68). First, cDNA libraries were constructed with an Ion Total RNA-seq kit, v2 (Ion Torrent). The prepared libraries were then used to generate template-positive ion sphere particles (ISPs) using an Ion PGM Template OT2 200 kit (Ion Torrent) in combination with an Ion OneTouch 2 system (Ion Torrent). The template-positive ISPs were loaded onto Ion 318 v2 chips (Ion Torrent) and sequencing runs performed with an Ion PGM Sequencing 200 kit, v2 (Ion Torrent). Raw data files in fastq format were exported and analyzed using the CLC Genomics Workbench software (Qiagen) and the USA300_FPR3757 and NCTC 8325 reference genomes (GenBank accession numbers CP000255 and NC_007795, respectively). Expression values for each gene were calculated as reads per kilobase per million mapped reads (RPKM), and a quantile normalization approach was applied (71) with a lower limit of 10 RPKM. Genes that displayed fold changes of $\geq 2$ when comparing expression in the mutant to that in the wild-type strain were included in further analyses.

### TABLE 1 Strains, plasmids, and primers

<table>
<thead>
<tr>
<th>Strain, plasmid, or primer</th>
<th>Description or sequence</th>
<th>Reference or source</th>
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<tbody>
<tr>
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<td>S. aureus</td>
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<td>Restriction-deficient transformation recipient</td>
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<td>USA300 HOU</td>
<td>MRSA isolate cured of pUSA300-HOU-MRSA</td>
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<td>B325-4 derivate with functional rsbU; rsbU +</td>
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<td>USA300 HOU ΔrpoZ mutant</td>
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MRSA, methicillin-resistant S. aureus. Underlining denotes restriction sites.
RT-qPCR. In order to confirm expression values that were generated by RNA sequencing, a subset of genes were chosen to confirm transcriptional changes via quantitative reverse transcription-PCR (RT-qPCR), as described by us previously (8, 69, 72, 73). PCR amplification was performed using primers for each gene (Table 1) (qpfL, OL1297/OL1298; asp23, OL2129/OL2130; SAUSA300_0174, OL3136/OL3137; aur, OL3747/OL3748; rpoC, OL4099/OL4100; and SAUSA300_0777, OL4109/OL4110), alongside 16S rRNA–specific primers as standard controls (OL1184/OL1185) (74).

Investigation of RNA polymerase composition. In order to determine the effects of ω loss on the stability and composition of RNAP, we adopted an approach first outlined by Gunnelius et al. (an overview of this procedure can be found in Fig. S1 in the supplemental material) (25). Wild-type and ΔrhoZ strains were grown to exponential phase, and their cytoplasmic protein fractions were isolated as outlined previously (75). In order to differentiate between RNAP subunits unbound or within the RNAP Δ subcomplexes, specific primers as standard controls (OL1184/OL1185) (74). Peptides were resuspended in H2O–0.1% formic acid for LC-MS/MS processing in MaxQuant (www.maxquant.org) and searched against the UniProtKB [ACN] with 0.1% formic acid). Fractions were analyzed on a linear ion trap Orbitrap instrument (Orbitrap XL; Thermo Fisher Scientific), with full MS survey scans acquired at 60,000 resolution. The top 10 most abundant ions were selected for tandem MS (WS/MS) analysis in the linear ion trap. Raw data files were processed in MaxQuant (www.maxquant.org) and searched against the UniProtKB S. aureus USA300 protein sequence database. Search parameters included constant modification of cysteine by carbamidomethylation and the variable modification methionine oxidation. Proteins were identified using filtering criteria of 1% protein and peptide false-discovery rates. Protein levels were normalized to the overall protein content in each of the investigated fractions.

Characterization of protein content using mass spectrometry. Protein extracts isolated via the size selection protocol outlined above were processed by filter-aided sample preparation (FASP), as described by us previously (76). Proteins were reduced with dithiothreitol (DTT), alkylated with iodoacetamide (IAA), and digested with trypsin–Lys-C (Promega) overnight at 37°C. Peptides were collected by centrifugation, desalted using Pierce SPE C18 columns with a Supelco vacuum manifold, and dried in a vacuum concentrator (Labconco). Peptides were resuspended in H2O–0.1% formic acid and separated using a 75-μm by 10-cm C18 reversed-phase high-pressure liquid chromatography (HPLC) column (New Objective) on a NanoLC Ultra (Eksigent) with a 120-min gradient (4 to 40% acetonitrile [ACN] with 0.1% formic acid). Fractions were analyzed in parallel on an ion trap Orbitrap mass spectrometer (Orbitrap XR; Thermo Fisher Scientific), with full MS survey scans acquired at 60,000 resolution. The top 10 most abundant ions were selected for tandem MS (WS/MS) analysis in the linear ion trap. Raw data files were processed in MaxQuant (www.maxquant.org) and searched against the UniProtKB S. aureus USA300 protein sequence database. Search parameters included constant modification of cysteine by carbamidomethylation and the variable modification methionine oxidation. Proteins were identified using filtering criteria of 1% protein and peptide false-discovery rates. Protein levels were normalized to the overall protein content in each of the investigated fractions.

Western blotting. The assessment of β subunit abundance was performed as described by us previously (66). Size-selected protein fractions containing proteins and protein complexes of >100 kDa were prepared as for experiments determining RNAP composition. Samples were separated using 12% SDS-polyacrylamide gel electrophoresis (SDS-PAGE), as described by us previously (76). Proteins were transferred to a polyvinylidene difluoride (PVDF) membrane and detected using a monoclonal mouse RNAP antibody (8RB13; Santa Cruz Biotechnology) and a horseradish peroxidase (HRP)-conjugated secondary antibody. HRP activity was assessed using the SuperSignal West Pico substrate (Thermo Fisher Scientific) and visualized on X-ray film.

MIC assessment assay. To determine whether transcriptional changes observed within the mutant strain resulted in detectable physiological sensitivities, we assessed MICs for various antibacterial agents, as described by our group previously (77). Briefly, overnight cultures were diluted 1:1,000 into fresh plates, 1 ml of these fresh cultures was transferred into each well and incubated for 48 h (static, 37°C). After incubation, supernatants were gently removed from each well and the biofilm washed twice with PBS before fixation with absolute ethanol. Biofilms were stained with crystal violet solution (2% [wt/vol])

Biofilm assay. The ability of strains to form biofilms was assessed as described by us previously (78). The wells of non-tissue-culture treated 12-well polystyrene plates were incubated for 48 h (static, 4°C) with 1 ml of human serum in order to facilitate attachment of cells. Concurrently, bacteria were cultured overnight in biofilm medium (BFM) (TSB supplemented with 0.5% [wt/vol] dextrone and 3% [wt/vol] NaCl) and used to seed fresh BFM to an OD600 of 0.05. After removal of the human serum from the wells of plates, 1 ml of these fresh cultures was transferred into each well and incubated for 48 h (static, 37°C). After incubation, supernatants were gently removed from each well and the biofilm washed twice with PBS before fixation with absolute ethanol. Biofilms were stained with crystal violet solution (2% [wt/vol])

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for 10 min, followed by two more rounds of washing with PBS. Residual supernatants were aspirated, and plates were dried overnight. After this time, 300 μl of absolute ethanol was added to the wells and incubated for 10 min before being removed, and the absorbance was measured at 570 nm using a Synergy2 plate reader (BioTek).

**REFERENCES**


**ACKNOWLEDGMENT**

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**Supplemental Material**

Supplemental material for this article may be found at https://doi.org/10.1128/JB.00459-16.


30. Reference deleted.


38. Reference deleted.


