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The (p)ppGpp-binding GTPase Era promotes rRNA processing and cold adaptation in
*Staphylococcus aureus*

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Abstract

Ribosome assembly cofactors are widely conserved across all domains of life. One such
group, the ribosome-associated GTPases (RA-GTPase), act as molecular switches to coor-
dinate ribosome assembly. We previously identified the *Staphylococcus aureus* RA-
GTPase Era as a target for the stringent response alarmone (p)ppGpp, with binding leading
to inhibition of GTPase activity. Era is highly conserved throughout the bacterial kingdom
and is essential in many species, although the function of Era in ribosome assembly is
unclear. Here we show that Era is not essential in *S. aureus* but is important for 30S ribo-
sonal subunit assembly. Protein interaction studies reveal that Era interacts with the 16S
rRNA endonuclease YbeY and the DEAD-box RNA helicase CshA. We determine that both
Era and CshA are required for growth at suboptimal temperatures and rRNA processing.
Era and CshA also form direct interactions with the (p)ppGpp synthetase Rel*Sau*, with Rel*Sau*
positively impacting the GTase activity of Era but negatively affecting the helicase activity
of CshA. We propose that in its GTP-bound form, Era acts as a hub protein on the ribosome
to direct enzymes involved in rRNA processing/degradation and ribosome subunit assembly
to their site of action. This activity is impeded by multiple components of the stringent
response, contributing to the slowed growth phenotype synonymous with this stress
response pathway.

Author summary

The bacterial ribosome is an essential cellular component and as such is the target for a
number of currently used antimicrobials. Correct assembly of this complex macromolec-
ule requires a number of accessory enzymes, the functions of which are poorly character-
ised. Here we examine the function of Era, a GTase enzyme involved in 30S ribosomal
subunit biogenesis in the important human pathogen *S. aureus*. We uncover that Era is
not an essential enzyme in *S. aureus*, as it is in many other species, but is important for
correct ribosome assembly. In a bid to determine a function for this enzyme in ribosomal
assembly, we identify a number of protein interaction partners with roles in ribosomal
RNA maturation or degradation, supporting the idea that Era acts as a hub protein
facilitating ribosomal biogenesis. We also uncover a link between Era and the (p)pGpp synthetase Rel\textsubscript{Sau}, revealing an additional level of control of rRNA processing by the stringent response. With this study we elaborate on the functions of GTPases in ribosomal assembly, processes that are controlled at multiple points by the stringent response.

Introduction

Ribosomes are macromolecular machines responsible for the synthesis of proteins in all living cells. In bacteria these complexes consist of two subunits, with the large 50S subunit containing 33 large proteins (L1-36) and two rRNAs, and the 30S small subunit containing 21 small proteins (S1-21) and the 16S rRNA. As such, assembly of the ribosome is a tightly regulated process, with correct maturation requiring the help of assembly cofactors, one class of which are the P-loop ribosome-associated GTPases (RA-GTPase). These enzymes are widely conserved across all domains of life and act as molecular switches, cycling between inactive GDP-bound, and active, effector-binding GTP-bound states. Within the P-loop GTPase class lies the Era family (\textit{Escherichia coli} Ras-like protein), which is characterised by the presence of a distinct derivative of a KH domain \[1\]. Era, the protein after which this family is named, is highly conserved throughout the bacterial kingdom, although is missing in \textit{Chlamydia} and mycobacterial species \[1\]. This GTPase is essential in \textit{E. coli} \[2–5\], \textit{Salmonella} Typhimurium \[6\] and in some strains of \textit{Bacillus subtilis}, where deletion mutants of \textit{B. subtilis} strains IS75 and CRK6000 were unobtainable \[7, 8\], although an era knockout was created in strain BR151 \[7\]. The reasons behind this difference are not known. The first indication that Era is involved in ribosomal biogenesis came from the ability of the 16S rRNA methyltransferase KsgA to suppress a cold-sensitive phenotype of an Era E200K mutation \[9\]. While important for ribosomal maturation, additional phenotypic defects associated with depleted cellular levels of Era include cell cycle control and chromosome segregation, as well as carbon and nitrogen metabolism \[10–12\].

Era is composed of two domains, an N-terminal GTPase domain and a C-terminal RNA-binding K-homology domain \[13\]. A cryo-electron micrograph structure of Era in complex with the 30S subunit of the ribosome reveals that Era binds into the same pocket as small subunit protein S1 \[14\]. In the absence of S1, Era interacts with proteins S2, S7, S11 and S18, as well as with a number of helices of the 16S rRNA. In addition, Era interacts with h45 and nearby residues 1530–1539 (GAUCACCUCC) in the 3’ minor domain of the 16S rRNA via its CTD region \[14, 15\]. These residues include the anti-Shine Dalgarno sequence, critical for the formation of the 30S preinitiation complex. Including Era in \textit{in vitro} reconstitutions of the 30S ribosome promotes the incorporation of several late-stage ribosomal proteins for the RNA \[16, 17\]. Consequently, it has been proposed that Era functions as a checkpoint protein, and that by binding to the 16S rRNA the formation of the initiation complex is prevented until the appropriate time \[14\]. In addition to interacting with ribosomal proteins, Era has also been reported to interact with a number of proteins involved in 16S rRNA maturation. One of these, YbeY, is an endonuclease required in \textit{E. coli} and \textit{B. subtilis} for the maturation of the 3’ end of the 16S rRNA \[18, 19\]. It is proposed that the binding of YbeY to Era and S11 guide the endonuclease to its site of action \[18\].

The stringent response is a signalling system used by bacteria to cope with a variety of environmental stresses, the best characterised of which is nutrient deprivation. The opportunistic pathogen \textit{Staphylococcus aureus} contains three enzymes, Rel\textsubscript{Sau}, RelP and RelQ, which upon sensing a stress, synthesise the nucleotides guanosine tetra- and pentaphosphate ((p)pGpp) \[20, 21\]. Once produced, this alarmone controls cellular responses to aid survival. Our
previous work identified Era, as well as three other GTPase enzymes from *S. aureus*, as target proteins for (p)ppGpp and demonstrated that the production of (p)ppGpp has a negative impact on mature 70S assembly [22]. Here, we examine the role of Era as an enzyme required for ribosome biogenesis. Unlike in *E. coli*, this GTPase is not essential for the growth of *S. aureus*, however mutant cells are defective in 30S subunit maturation. We identify the *S. aureus* endonuclease YbeY and the DEAD-box RNA helicase CshA as interaction partners for Era, and show that both Era and CshA are crucial for rRNA homeostasis and growth at low temperatures. We additionally demonstrate that both Era and CshA interact with the (p)ppGpp synthetase/hydrolase RelSau, and that cellular rRNA processing is controlled in a stringent response-dependent manner. With this, we propose that Era is a protein that facilitates the interactions between a number of rRNA processing and degrading enzymes and the 30S subunit/16S rRNA, and show that under stress conditions the stringent response is important for this process.

**Results**

**Era is not essential in *S. aureus* but is required for normal growth and ribosome assembly**

Previous reports show that *era* is an essential gene in multiple bacterial species [3, 6]. In agreement with this, transposon insertions in this gene have not been identified in a number of published *S. aureus* transposon libraries [23–25]. Closer inspection of the transposon insertion hits within the *era* operon obtained in the Nebraska *S. aureus* mutant library [23] reveals insertions in *ybeZ*, a gene of unknown function, and at the 3' end of the diacylglycerol kinase *dgkA* but not within *ybeY*, *cdd*, *era* or *recO*, suggestive of an additional promoter upstream of *era* and *cdd* (Fig 1A). To rule out polar effects and examine whether *era* in isolation is essential, we replaced the entire coding sequence with the Tet-encoding *tetAM* open reading frame in the presence of the anhydrotetracycline (Atet)-inducible covering plasmid pCN55iTET- *era*. Following deletion of the chromosomal copy of *era*, we attempted to phage transduce this mutation into *S. aureus* strains containing the empty vector pCN55iTET or the complementing plasmid pCN55iTET- *era*. Transduction efficiencies were similar with both recipients, albeit smaller colony sizes were observed in the absence of *era*, indicating that this gene is not essential in *S. aureus*. We next transduced the *era* deletion into the community-associated methicillin resistant *S. aureus* strain LAC*<sup>+</sup>* to rule out secondary mutations and used this strain for further studies. Analysis of growth rates revealed that an *era* mutant strain, while viable, does have a significant growth defect, which can be complemented by the expression of Era from pCN55iTET- *era* (Fig 1B).

A role for Era as a ribosomal subunit assembly cofactor has been reported in both *E. coli* and *B. subtilis*, where depletion leads to a decrease in 70S ribosomes and an accumulation of individual 50S and 30S subunits [26, 27]. We reasoned that the growth defect observed in *S. aureus* may be due to defects in ribosome assembly. To investigate this in the context of a complete *era* deletion, we analysed the cellular ribosomal content of wildtype, mutant and complemented strains by sucrose density centrifugation. This revealed that the *era* mutant strain contained fewer polysomes and mature 70S ribosomes, with a concurrent build-up of 50S subunits, a defect that was reversed in the presence of the complementing plasmid (Fig 1C). As cryo-electron microscopy has shown Era from *Thermus thermophilus* interacting with the 30S subunit [14], an excess of free 50S subunits may be, in and of itself, an indication that there is a defect in small subunit biogenesis, leading to a build-up in free 50S. We reasoned that this defect in ribosome assembly might make the *era* mutant strain more susceptible to ribosome-targeting antibiotics. In agreement with this, we observed that the minimum inhibitory
concentration (MIC) for the Δera strain decreased 2-4-fold when exposed to the 30S-targeting antibiotic spectinomycin (S1A Fig). However, the MIC was unaffected by the 50S targeting antibiotic chloramphenicol (S1B Fig). Together this indicates that while Era is not an essential protein in *S. aureus*, it is important for optimal growth and ribosome maturation.

Uncovering Era protein interaction dynamics in a native background

In *S. aureus*, Era is encoded in an operon with five other genes, one of which encodes YbeY (Fig 1A). YbeY is an endoribonuclease implicated in the maturation of the 3' terminus of 16S rRNA in *E. coli*, as well as a quality control checkpoint protein that together with RNase R is involved in eliminating defective 70S ribosomes [19, 28]. Era interacts with YbeY in *E. coli* [18], although these two genes are not encoded in the same operon in that organism (Fig 1A). To determine whether Era from *S. aureus* interacts with YbeY, and/or any other proteins...
encoded in the *S. aureus* *era* operon, we first used a bacterial two-hybrid approach, heterologically expressing Era in combination with YbeZ, YbeY, DgkA, Cdd or RecO in *E. coli*. Using this approach, we observed that Era potentially interacts with YbeY and YbeZ (Fig 2A).

In order to look at these interactions natively in an *S. aureus* background, we adapted a split-luciferase system recently developed to analyse protein-protein interactions in...
Clostridium difficile for use in *S. aureus* (Fig 2B) [29]. These vectors, while designed for use in *C. difficile*, also replicate in *S. aureus* using an oriV origin of replication. When using this system in *S. aureus*, we noted an increase in non-specific luciferase activity from the pAF256 control vector over the pAF257 negative control, regardless of the inserted gene. For this reason, only interactions that are statistically increased over both negative controls will be considered positive. The *era* gene, in combination with itself and each of the operon genes, were cloned into the split-luciferase plasmid pAP118 and introduced into *S. aureus* strain LAC'. Induction of protein expression with A.et revealed a strong positive interaction for Era with itself (Fig 2C), indicating that Era can form dimers. Dimerisation of Era has previously been observed while solving the crystal structure of the *E. coli* protein [30] and also by gel filtration chromatography [31], and so this interaction confirms the functionality of the split-luciferase system in *S. aureus*. In agreement with the bacterial two-hybrid results, no interactions occurred between Era and DgkA, Cdd or RecO. There was a significant interaction between Era and YbeY, but only when compared to the pAF257-ybeY negative control (Fig 2C). This prompted us to investigate the potential interaction between Era-YbeY *in vitro*. Pulldown assays were performed using glutathione beads coupled to either GST or GST-tagged Era, which were incubated with His-tagged YbeY (Fig 2D). Here we observed low-level cross reactivity of YbeY with the glutathione beads that could not be reduced despite increased washes. However, His-YbeY was pulled down significantly more in the presence of Era (Fig 2D). Altogether the bacterial two-hybrid, split luciferase and affinity pulldown assays suggest a weak interaction between Era and the endonuclease YbeY from *S. aureus*, which may form part of a larger complex, as has been suggested for the *E. coli* protein [18]. With this data we have adapted a luciferase system for confirmation of protein-protein interactions in a native *S. aureus* background.

Era interacts with the DEAD-box RNA helicase CshA

We wished to shed further light on the role played by Era in the cell and sought to identify unknown interaction partners using a genome-wide bacterial two-hybrid screen. A library of *S. aureus* genomic DNA fragments cloned into pUT18C was screened against pKNT25-era. Of the 17 hits obtained, 14 contained fragments mapping to *cshA*, a gene encoding a DEAD-box RNA helicase. Of the remaining 3 plasmids, 2 contained sequences not in-frame with the T18 coding sequence, while 1 contained a sequence that also went blue when co-transformed with the pKNT25 empty vector. CshA functions in mRNA protection and RNA decay in *S. aureus* [32, 33], while the homologue CsdA from *E. coli* is involved in the cold shock degradosome [34]. In *B. subtilis* cells, CshA has also been shown to interact with a number of ribosomal proteins from the 50S subunit, while a deletion strain has a reduction in the amount of mature 70S ribosomes [35].

To confirm the interaction between Era and CshA, full-length *cshA* was cloned into both pUT18 and pUT18C so as to perform a bacterial two-hybrid assay with full-length constructs. Upon co-transformation, interactions occurred between CshA-T18 and both T25-Era and Era-T25 (Fig 3A). To assess this interaction *in vitro*, we performed pulldown assays using glutathione beads coupled to either GST or GST-tagged CshA and incubated them with His-tagged Era, revealing that while His-Era did not interact with the control GST protein, it was pulled down in the presence of CshA (Fig 3B). Finally, to analyse and confirm this interaction in the context of the native host, we used the split luciferase assay (Fig 2B). Assaying for luciferase activity revealed a 4-fold increase in luciferase activity upon co-expression of Era and CshA (Fig 3C). Together these data confirm the identification of a novel Era interaction partner and suggest that CshA can also interact with proteins that associate with the 30S ribosomal subunit.
The N-terminal regions of both CshA and Era are crucial for interactions

CshA is a DEAD-box RNA helicase with an N-terminal helicase core containing two RecA-like domains and a disordered C-terminal (CT) region involved in RNA binding (Fig 3D) [33]. This protein is capable of unwinding both double stranded RNA and RNA-DNA hybrids and is required for both the stabilisation and the degradation of mRNA, the latter of which occurs via interactions of the CT region with components of the RNA degradosome [32, 33].

To determine which regions of both Era and CshA are important for interacting, shorter domain constructs comprising only RecA1 (aa 1–221), RecA2 (aa 222–382), the entire core helicase domain (aa 1–382) or just containing the disordered CT-region (aa 383–506) were
cloned, together with Era, into the split luciferase vector pAP118. Interactions with full-length Era were only apparent in the presence of the full core helicase domain (aa 1–382: Fig 3E), indicating that the RNA-binding CT region of CshA is dispensable for this interaction. To determine whether the GTPase domain of Era is required for binding, a shorter construct comprising the N-terminal G domain (aa 1–180) was cloned into pAP118 alongside full-length CshA. This domain is reported to still retain GTPase activity, but can no longer bind to RNA [13, 36]. Luciferase assays reveal that the G domain of Era is sufficient for interacting with CshA (Fig 3F).

**Interactions between Era and CshA do not affect enzymatic activity**

CshA is an RNA helicase, while Era has GTPase activity. To investigate whether the interaction between these two proteins affects the enzymatic activity of either one, we first performed helicase assays. A double stranded RNA oligo was incubated with each protein singly or in combination. While CshA was able to unwind the dsRNA over time, the addition of Era had no effect on its activity (S2A Fig). In addition, the GTPase activity of Era in the presence of 70S ribosomes was unaltered by the addition of either CshA or YbeY (S2B and S2C Fig), indicating that while these proteins do interact, this binding has little effect on their enzymatic functions. As previously proposed, we suggest that these interactions fit with a role for Era as a guide for rRNA/ribosome maturation enzymes to their substrates [14, 18].

**CshA and Era are both required for growth at suboptimal temperatures and rRNA homeostasis**

In *E. coli*, CshA has been linked to survival at low temperatures [37], while strains with depleted levels of Era are sensitive to both cold and heat shock [9, 10]. To examine the importance of these enzymes in *S. aureus*, we utilised deletion strains in the *S. aureus* MRSA strain LAC*. While both the *S. aureus era* and *cshA* null mutants showed significant growth defects at 37˚C (Figs 1B and S3A), the growth of both was severely compromised at 25˚C (Figs 4A and S3B). This defect was enhanced in a double Δera cshA mutant, which failed to grow at 25˚C even after 48 h (Figs 4A and S3B), highlighting that these two proteins are essential for bacteria to grow at suboptimal temperatures.

Era is an RA-GTPase that, along with a second RA-GTPase RsgA, binds to the 30S ribosomal subunit [14, 39]. In *E. coli*, overexpression of Era can complement a deletion of *rsgA* [40]. We first examined whether expression of Era could complement a deletion of *rsgA* in *S. aureus* using cross-complementation. This revealed that Era can complement for the loss of RsgA (S4A Fig), although RsgA could not complement for the loss of Era (S4B Fig), suggesting partial overlapping but also distinct functions. We next used cross-complementation of the Δera and ΔcshA mutations to investigate whether Era and CshA act in the same pathway and can functionally complement each other. Cross-complementation of the Δera and ΔcshA mutations did not improve growth at 25˚C (S4B and S4C Fig). This indicates that these proteins have separate cellular functions, even though they interact and bacteria lacking both cannot survive cold temperatures. We additionally confirmed that no suppressor mutations had arisen in our Δera cshA double deletion by complementing growth of the double mutant back to ΔcshA levels with pCN55iTET-era (S4E and S4F Fig).

To understand why Era and CshA are so important for growth at cold temperatures, we sought to assess the impact of deleting these genes on the cellular rRNA. We observed that RNA extracted from both the single Δera and double Δera cshA strains grown at 37˚C had a processing or degradation intermediate migrating slightly below the 16S band (Fig 4B). At 25˚C we also observed a defect for the ΔcshA mutant strain and multiple additional bands
were evident in the Δera cshA double mutant (Fig 4B), which could represent either processing intermediates from the 16S, or degradation intermediates from both the 23S and the 16S. An accumulation of degradation products in the cshA mutant could be due to its role in the RNA degradosome [33], or could suggest a role for its helicase activity in rRNA processing. To examine whether an accumulation of unprocessed 17S rRNA could be occurring in these mutant strains when grown at low temperatures, we performed qRT-PCR with RNA extracted from cells grown at 25˚C, with probes that either amplified internally to the 16S rRNA sequence (to amplify total 16S rRNA), or flanked either the 5’ or 3’ processing junctions...
required to convert 17S into mature 16S rRNA (Fig 4C). When strains were normalised to total 16S rRNA levels, significant increases in unprocessed 5’ were evident in both the Δera and ΔcshA strains compared to the wildtype (Fig 4C). Increases were also observed for the processing of the 3’ sequence for the Δera mutant, although these did not reach significance (Fig 4C). As Era is known to interact with the 30S ribosomal subunit, these defects could potentially be due to a failure to recruit the endonuclease YbeY, or indeed other processing enzymes, in the absence of Era, while the defects in the ΔcshA strain could be either direct, due to a loss of helicase activity necessary for 16S 5’ processing at 25˚C, or could reflect the role of CshA in the RNA degradosome, with pre-16S rRNA destined for degradation accumulating in its absence.

RelSau interacts with both Era and CshA

Our previous work demonstrated that the GTPase activity of Era is inhibited by the stringent response alarmone ppGpp [22]. To examine whether ppGpp also interacts with CshA we performed DRaCALA assays, revealing that this nucleotide does not interact with CshA (S5A Fig), nor is the affinity of ppGpp for Era altered in the presence of CshA (Kd of 3.1 ± 0.4 μM in the absence versus 3.2 ± 0.3 μM in the presence of CshA) (S5B Fig).

In E. coli, Era, YbeZ and the (p)ppGpp synthetase/hydrolase enzyme SpoT all interact with YbeY, potentially forming a complex [18]. To investigate whether the stringent response has any additional points of interaction with either Era and CshA in S. aureus, we examined whether one of the (p)ppGpp synthetases might directly interact with either protein via bacterial two-hybrid. This revealed that RelSau, but not RelP or RelQ, interacts with CshA and that RelSau interacts with Era (Fig 5A). To confirm this interaction, we first used affinity pulldown assays, showing that statistically significant interactions occurred between both Era or CshA and RelSau (Fig 5B). Following this, we again used the split luciferase assay, which confirmed an interaction between RelSau and CshA (Fig 5C) or Era (Fig 5D), and also established that the GTPase domain of Era was sufficient for this interaction (Fig 5E).

RelSau influences the enzymatic activity of Era and CshA and is important for rRNA processing

We first examined whether the interactions between RelSau and Era/CshA impact the activity of either enzyme. Recombinantly purified full-length RelSau is known to be in the synthetase-off/hydrolase-on conformation in vitro [41]. Using purified RelSau, we determined that Era and CshA have no effect on the (p)ppGpp hydrolase activity of this enzyme (S5C Fig). To look at the enzymatic activities of Era and CshA in the presence of RelSau we firstly examined the GTPase activity of Era, observing a slight but significant increase in activity in the presence of RelSau (Fig 6A). Following this we performed a helicase assay, revealing a decrease in the unwinding activity of CshA in the presence of RelSau, the opposite effect of binding on the enzymatic activity of Era (Fig 6B).

Through RelSau the stringent response appears to interact with RA-GTPases and associated proteins, increasing Era GTPase activity, while decreasing the helicase activity of CshA. To directly analyse the importance of the stringent response for rRNA processing, we grew the wildtype S. aureus LAC at 25˚C, a condition under which we have shown that both Era and CshA are important for viability, and induced the stringent response with mupirocin, an antibiotic that inhibits the isoleucyl-tRNA synthetase mimicking amino acid deprivation [42]. We again performed qRT-PCR with the probes as in Fig 4C that either amplified internally to the 16S rRNA sequence, or either flanked the 5’ or 3’ processing junctions of the 17S rRNA. For the probe internal to the 16S rRNA we observed a significant decrease in 16S rRNA in the presence of mupirocin (Fig 6C). This is expected, as it is known that rRNA levels decrease upon
activation of the stringent response [43]. However, when we compare the amount of unprocessed 5’ or 3’ sequences relative to the level of 16S in each strain, we observed a significant increase in the amount of unprocessed leader 17S rRNA upon induction of (p)pGpp (Fig 6D: \( p = 0.0031 \)). We have previously shown that ppGpp interacts with Era and that mature ribosome formation is inhibited upon induction of the stringent response [22]. Here we propose that the stringent response can also impact RA-GTPase function through direct interactions of RelSau with both Era and CshA. We suggest that Era acts as a hub protein on the 30S ribosome, promoting the association of processing or degradation enzymes like YbeY and CshA.
RA-GTPases, including Era, are reported to associate to the ribosome when bound to GTP, but dissociate once hydrolysis has occurred [44]. The increased GTPase activity of Era observed in the presence of RelSau may therefore lead to increased dissociation of Era, and associated enzymes, from the ribosome, stalling maturation. Additionally, the inhibition of the helicase activity of CshA may also delay 17S rRNA processing or degradation, especially at 25˚C. The dynamics of RA-GTPase associations to the ribosome are complex, with much further in-depth study required to precisely determine if the Era-YbeY, Era-CshA or RelSau interactions occur on or off the ribosome and the direct impact of these on ribosomal maturation.

Fig 6. The stringent response affects Era and CshA enzyme activity and rRNA processing. A) The GTPase activity of 100 nM Era was measured in the presence of an equal amount of ribosomes, 100 nM RelSau and 1 μM GTP. All reactions contained ribosomes. The RelSau only control reaction contained RelSau and ribosomes but no Era. Hydrolysis of 32P-GTP was monitored by TLC and the percentage GDP formed quantified using ImageJ. Experiments were repeated three times with means and standard deviations shown. Statistical analysis was performed using a two-way ANOVA, followed by Dunnett’s multiple comparisons test ($p < 0.005$, $p < 0.05$). B) The RNA helicase activity of 0.5 μM CshA with and without RelSau was determined using a Cy3-labelled double stranded RNA oligomer. Reactions were incubated at 25˚C for up to 10 min before analysis on a native page gel. Experiments were repeated four times with means and standard deviations shown. Statistical analysis was performed using a two-way ANOVA, followed by Sidak’s multiple comparisons test ($p < 0.05$, $p < 0.0001$). C & D) rRNA extracted from LAC− Strains were grown at 25˚C until an OD600 of 0.4. Cells were split and half exposed to 60 μg/ml mupirocin for 1 h (plus), which at 25˚C corresponds to one doubling. RT-PCR of total RNA using the probes as indicated in Fig 4C that bind internally to the 16S (C) or to the unprocessed leader and trailer sequences (D) of the 17S rRNA. rRNA values are expressed as mean relative expression +/- SD normalised to the expression of rho RNA as an internal control. For comparisons between two groups (C—LAC− minus v plus) a two-tailed, unpaired Student’s t-test was performed ($p < 0.0001$). For comparisons between more than two groups (D) a two-way ANOVA, followed by Sidak’s multiple comparisons test was performed ($p < 0.005$). Cycle threshold values were determined for 3 biological repeats in duplicate (N = 3).

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Discussion

The synthesis of ribosomes and proteins consumes approximately 40% of the energy within a growing bacterial cell [45]. Ribosomal assembly cofactors are, therefore, an essential group of enzymes for coordinating biogenesis as efficiently as possible. Members of this group include RNA helicases, rRNA and protein modification enzymes, chaperones and RA-GTPases. *S. aureus* contains over 11 RA-GTPases, each with potentially varying roles in the biogenesis of the 50S and 30S subunits, although the precise functions of each in this process is unclear.

While the RA-GTPase Era has been extensively studied, its precise function in the cell is unknown. Cryo-electron microscopy shows Era interacting with the 30S ribosomal subunit, however this protein has also been implicated in numerous other cellular processes [14]. Era-depleted *B. subtilis* or *E. coli* cells are elongated, with defects in septum formation. These cells also contain diffuse nucleoid material, implicating Era in cell division and chromosome segregation [5, 12]. In *B. subtilis*, Era-depleted cells have defects in spore formation [7], while growth defects in strains containing Era variants with reduced nucleotide binding abilities can be rescued by truncating either rpoN or ptsN [11]. RpoN, which encodes for the alternative sigma factor σ54, is required for nitrogen assimilation and fixation, while PtsN is involved in sugar transport, suggesting that Era is also involved in regulating carbon and nitrogen metabolism. Localisation studies of Era in *E. coli* have indicated that this protein is present at both the membrane and in the cytoplasm [46], and so it has been suggested that this enzyme cycles between the membrane and the ribosome in response to cellular triggers. In addition to ribosomal biogenesis factors, Era has been reported to interact with MazG, Ndk, Pk and YggG [47–49]. Ndk, a nucleoside diphosphate kinase and Pk, a pyruvate kinase, from *Pseudomonas aeruginosa* were both shown to form a complex with Era, further implicating Era in energy metabolism [48]. MazG is a nucleoside triphosphate pyrophosphohydrolase, while YggG is a membrane-associated heat shock protein. The significance of these interactions with Era is unknown.

Era has also been implicated in helping cells cope with cold stress. Conditional cold-sensitive mutants of Era have been constructed in *E. coli* [10] and these mutations could be suppressed by the overexpression of the 16S rRNA methyltransferase KsgA, providing one of the first links between Era and ribosome biogenesis [9]. Era interacts with the universally conserved endonuclease YbeY in *E. coli* [18]. YbeY is dispensable in *E. coli*, but is essential in *B. subtilis* and potentially in *S. aureus*, given the lack of transposon mutants available [19, 25, 28]. This endonuclease is required for the maturation of the 3' end of the 16S rRNA, and in strains depleted of YbeY the 70S ribosomes are targeted for degradation with the help of RNase R [19, 28]. Era and YbeY are fused in a polypeptide in clostridial species, further highlighting a link between these two proteins. Although not encoded in the same operon as *era* in *E. coli*, this prompted us to examine the interactions of all other proteins encoded in the *era* operon in *S. aureus* with Era (Fig 2). As in *E. coli*, YbeY from *S. aureus* interacts with Era, although quite weakly. YbeZ also interacted weakly by bacterial two-hybrid but not significantly using the quantitative luciferase assay (Fig 2). YbeZ has been shown to interact with YbeY in *E. coli*, in addition to the (p)pGpp synthetase/hydrolase enzyme SpoT [18], and ribosomal proteins S7 and L6 [50]. YbeZ has an ATPase domain and an RNA binding motif, and because of its interactions with ribosomal proteins may also bind to the 17S rRNA and in *E. coli* form a complex with Era, YbeY and SpoT to aid in processing the 17S rRNA to the mature 16S form [18]. So it is possible that YbeY, YbeZ, Era and RelS in *S. aureus* all form a complex, and that the interactions between all these proteins may become more apparent upon isolation of the entire complex.
Here we use interaction studies to show that Era also interacts with a protein involved in survival at suboptimal temperatures, the DEAD-box RNA helicase CshA (Fig 3). In *B. subtilis*, CshA is one of the most abundant RNA helicases produced at low temperatures [35] and in *B. subtilis* and *E. coli* it has been implicated in multiple processes, including 50S ribosome biogenesis and interacting with components of the RNA degradosome [33–35, 37]. In *S. aureus*, CshA has been linked to controlling the turnover of mRNA in the cell [33] [32]. Deletion of cshA results in the stabilisation of some mRNA transcripts, such as the spa mRNA, but CshA also protects a number of other mRNA and sRNA transcripts under stress conditions [32]. Our analysis shows that Era and CshA interact and that the CT disordered region of CshA, which is important for binding to mRNA and the degradosome [33], is dispensable for this interaction (Fig 3). Given that the CshA homologue CsdA in *E. coli* associates with the 50S ribosomal subunit and is involved in its biogenesis [37], our results showing an interaction of CshA with the 30S binding protein Era, may indicate a wider role for CshA in both 30S and 50S subunit biogenesis, especially at 25˚C. This is supported by pulldown experiments performed with CshA from *S. aureus*, where both 30S and 50S ribosomal proteins were identified as interaction partners for CshA [33]. The precise dynamics of the interactions between CshA, Era and components of the ribosome are a question for future study.

Using cshA and era deletion strains we showed that both single mutants are important for growth at cold temperatures, with a double mutant unable to survive at 25˚C (Fig 4). By examining the rRNA of these strains we observe that both mutant strains have increased 17S unprocessed rRNA intermediates (Fig 4C). Additionally, a double Δera cshA mutant has multiple processing and degradation defects (Fig 4B). It has been proposed that Era may act to guide processing enzymes to their site of action [18]. In keeping with this, we have observed that Era and CshA have little effect on the enzymatic activity of each other. Therefore, we support a model whereby Era acts as a hub protein, allowing proteins such as the endonuclease YbeY and the RNA helicase CshA, and potentially others, access to their substrate rRNA. In keeping with this it was observed that over-expressing Era in a ybeY mutant *E. coli* strain suppresses a growth defect and improves 16S rRNA processing in an RNase II, Rnase R and RNase PH-dependent manner, potentially caused by over-recruitment of these exoribonucleases by Era in the absence of YbeY [51].

Cellular levels of rRNA are controlled by the stringent response [43]. Upon activation of the stringent response, transcription from rRNA promoters is reduced (Fig 6C), either by direct binding of ppGpp to the RNA polymerase in Gram negatives, or by tight control of cellular GTP levels in Gram positives [52, 53]. Here we see an additional level of regulation by the stringent response on rRNA. We show a direct interaction between the (p)ppGpp synthetase RelSau, and both CshA and Era (Fig 5), and show that RelSau has a relatively small but significant positive effect on the GTPase activity of Era (Fig 6A) and a negative effect on the helicase activity of CshA (Fig 6B). Additionally, we observed that the activation of the stringent response results in increased processing defects in a wildtype strain grown at 25˚C (Fig 6D). Era has increased association to the ribosome in the GTP-bound form [44]. Hydrolysis of GTP then acts as a signal to promote dissociation, presumably following a maturation event [54, 55]. The increased GTPase activity of Era in the presence of RelSau, could promote premature dissociation of Era from the immature 30S ribosomal subunit, leading to processing defects and contributing to the stalled growth phenotype synonymous with the stringent response. This would be in addition to our previous observation that Era has a higher affinity for ppGpp than GTP [22], which could potentially inhibit the association of the GTPase to the ribosome. All of this would be compounded by the inhibition of helicase activity of CshA, although further research is needed to determine whether any or all of these interactions occur on or off the ribosome.
It is well known that RelSau binds to mature 70S ribosomes and synthesises (p)ppGpp in response to uncharged tRNA docking in the A-site [43]. In addition, a recent paper from Trin-guier et al., describes how the accumulation of immature tRNAs with processing defects activates RelB-dependent production of (p)ppGpp in *B. subtilis*, leading to the inhibition of 16S rRNA 3’ maturation [56]. Here this inhibition is ascribed to decreasing GTP levels upon production of (p)ppGpp, however a direct involvement of (p)ppGpp cannot be ruled out. Our results agree with the reported role for Rel enzymes in rRNA maturation and extend this through the identification of interactions between RelSau and Era/CshA. From this we propose a role for RelSau in controlling immature 30S subunit maturation, in addition to its known function as a sensor of amino acid starvation on mature 70S ribosomes. An interesting area for future study would be to investigate the dynamics of RelSau and Era/CshA interactions in the presence of 30S and 50S subunits, as well as mature 70S ribosomes.

Taken together, we demonstrate a cellular function for Era as a protein important for coordinating 30S ribosomal biogenesis in a stringent response-dependent manner. It is evident that defects in ribosomal assembly lead to altered protein translation, which may be the reason for the plethora of other phenotypes associated with depletion of Era. As Era can complement a defect in RsgA (S4A Fig, [40]), as well as RbfA [26], we propose a broader role for RA-GTPases as hub proteins involved in coordinating biogenesis. It is now of interest to determine what processing events are coordinated by the other RA-GTPases in *S. aureus*, as well as more broadly in other prokaryotes.

**Materials and methods**

**Bacterial strains and culture conditions**

*E. coli* strains were grown in Luria Bertani broth (LB) and *S. aureus* strains in tryptic soy broth (TSB) at 37˚C or 25˚C with aeration. Strains and primers used are listed in S1 and S2 Tables. Information on strain and plasmid construction is provided in supplemental S1 protocol.

**Luciferase assays**

Overnight cultures of *S. aureus* strains were diluted to an OD$_{600}$ of 0.05 and grown for 90 min in the presence of 100 ng/ml anhydrotetracycline (Ate). Strains were normalised to an OD$_{600}$ of 0.1 and luciferase activity measured according to the Nano-Glo Luciferase Assay System Protocol (Promega).

**Pull-down experiments and western blotting**

Glutathione beads were washed 5 times in 1 x wash solution (25 mM Tris pH 7.5, 150 mM NaCl, 1 mM EDTA, 0.5% Triton X-100). 1 μM of glutathione-S-transferase (GST), GST-CshA or GST-Era were coupled to the beads by incubating in 1 x wash buffer for 4 h at 4˚C. Unbound protein was removed by washing in 1 x wash solution between 5–10 times. Protein-bound beads were incubated with 1 μM His-tagged Era, His-YbeY or His-RelSau in the presence of 1 x binding buffer (25 mM Tris pH 7.5, 150 mM NaCl, 5 mM MgCl$_2$, 0.5% Triton X-100) for 16 h at 4˚C. After washing, attached proteins were eluted from the beads with 50 μl elution buffer (25 mM Tris pH 8, 150 mM NaCl, 1 mM EDTA, 0.5% Triton X-100, 10 mM reduced glutathione). Samples were mixed 1:1 with 2 x SDS protein sample buffer. Aliquots were separated on 12% SDS-polyacrylamide gels and proteins subsequently transferred to PVDF membranes. Bound His-tagged proteins were detected using HRP-conjugated anti-His antibodies (Sigma) at a 1:500 dilution. Blots were developed by enhanced chemiluminescence and imaged using a ChemiDoc MP imager (Bio-Rad).
Construction of bacterial two-hybrid library
Genomic DNA of *S. aureus* was extracted and partially digested by incubation with Sau3AI at 37 °C for 20 min. Digested DNA was run on a 0.8% agarose gel and fragments of 500 to 1000 bp and 1000 to 3000 bp gel extracted and purified. This was repeated five times from separate genomic preps and DNA fragments pooled. The vector pUT18C was digested overnight at 37 °C with BamHI and dephosphorylated with antarctic phosphatase for 90 min. Genomic DNA fragments were ligated into the pUT18C linearized vector using T4 ligase, transformed into *E. coli* DH5α competent cells (New England Biolabs) and plated onto LB agar with carbenicillin. Plates were scraped and transformant plasmids isolated using the GeneJET plasmid purification kit (Thermo scientific).

Bacterial two-hybrid
Bacterial two hybrid plasmids containing genes of interest were co-transformed into *E. coli* BTH101 cells, plated on LB agar containing 150 μg/ml carbenicillin and 30 μg/ml kanamycin and incubated at 30 °C overnight. Colonies were isolated, grown overnight at 30 °C in LB broth containing 0.5 mM IPTG and 5 μl spotted onto LB agar containing 0.5 mM IPTG and 40 μg/ml X-gal. Plates were incubated at 30 °C for up to 48 h. In some cases, BTH101 transformants were incubated overnight with 0.5 mM IPTG and spotted directly onto LB agar containing IPTG, X-gal and appropriate antibiotics.

RNA extraction
Strains of *S. aureus* were grown overnight at 37 °C and diluted to an OD$_{600}$ of 0.05. Cultures were grown to an OD$_{600}$ of 0.4 at either 37 or 25 °C and harvested. RNA was extracted using the RiboPure RNA Purification Kit (Invitrogen) as per guidelines. RNA was visualised using a modified agarose gel containing 0.7% agarose and 0.9% Synergel (Diversified Biotech) in 0.5 X TBE (44.5 mM Tris, 1 mM EDTA pH 8, 44.5 mM boric acid) as per Wachi *et al.* 1999 [57]. RNA was visualised using EvaGreen fluorescent nucleic acid dye (Biotium) on a ChemiDoc MP imager (Bio-Rad).

RT-PCR
RNA was extracted as described above. Complementary DNA was synthesised from 1.5 μg RNA with transcript reverse transcriptase (Sigma) and random primers. RT-PCR was performed on 100 ng of cDNA in triplicate using SYBR Green Jumpstart Taq readymix (Sigma). Primers RMC379/380 (16S internal), RMC570/571 (leader) and RMC572/548 (trailer) were designed to amplify 200–220 bp target regions in the 17S rRNA. The housekeeping gene rho was amplified with primers RMC573/574. The rho transcript was chosen as it has been shown to be highly stable under nutrient-deficient conditions under which the stringent response would be induced [38]. Cycle threshold values were determined for 3 biological repeats in duplicate. For each reaction, the ratio to rho transcript number was calculated as follows: $2^{(-\Delta C_{t}\text{target} - \Delta C_{t}\text{rho})}$.

Growth curves
*S. aureus* strains were grown overnight in TSB medium containing the appropriate antibiotics. Overnight cultures were diluted to a starting OD$_{600}$ of 0.05 in the presence of 100 ng/ml Atet where appropriate, incubated at 25 °C or 37 °C with aeration and OD$_{600}$ values determined at 2 h intervals. Growth curves were performed three to four times, and averages and standard deviations are plotted.
Minimum Inhibitory Concentrations

Overnight cultures of wildtype LAC⁺ and Δera strains were adjusted to an OD₆₀₀ of 0.05 in Mueller-Hinton broth and 100 µl incubated in 96 well plates with 2-fold dilutions of various antimicrobials at the following starting concentrations: spectinomycin 250 µg/ml and chloramphenicol 64 µg/ml. Plates were incubated at 37°C overnight with shaking. Assays were performed four times, and averages and standard deviations are plotted.

Protein purifications

Proteins were purified from 1–2 L E. coli cultures. Cultures were grown to an OD₆₀₀ 0.5–0.7, expression induced with 1 mM IPTG and incubated for 3 h at 37°C. Protein purifications were performed by either nickel or glutathione affinity chromatography. For nickel purifications, cell pellets were resuspended in 5 ml Buffer A (50 mM Tris pH 7.5, 150 mM NaCl, 5% glycerol, 10 mM imidazole), lysed with 100 µg lysozyme and sonication and the filtered cell lysate loaded onto a 1 ml HisTrap HP Ni²⁺ column (GE Healthcare) before elution using a gradient of Buffer B (50 mM Tris pH 7.5, 150 mM NaCl, 5% glycerol, 500 mM imidazole). GST-tagged proteins were resuspended in 5 ml PBS, lysed and the cell lysate was loaded onto a 1 ml GSTrap HP column (GE Healthcare) before elution using 50 mM Tris pH 8.0, 10 mM reduced glutathione. Protein containing fractions were dialysed in 50 mM Tris pH 7.5, 200 mM NaCl, 5% glycerol before storage at -80°C. Protein concentrations were determined by A₂₈₀ readings.

Synthesis of (p)ppGpp and differential radial capillary action of ligand assay (DRaCALA)

The synthesis of (p)ppGpp and DRaCALA binding assays were performed as described previously [22].

Ribosomal profiles from S. aureus cell extracts

Crude isolations of ribosomes from S. aureus cell extracts were achieved as described by Chin Loh et al. with some modifications [27]. Briefly, 100 ml cultures of the different S. aureus strains were grown to an OD₆₀₀ of 0.4 in TSB 100 ng/ml Atet. 100 µg/ml chloramphenicol was added to each culture and incubated for 3 min, before being cooled to 4°C to produce runoff ribosomes. Pelleted cells were suspended in association buffer (20 mM Tris-HCl pH 7.6, 8 mM MgCl₂, 30 mM NH₄Cl and 2 mM β-mercaptoethanol), normalized to an OD₆₀₀ of 15, lysed by the addition of 0.2 µg/ml lysostaphin and 75 ng/ml DNase and incubated for 60 min at 37°C. The extracts were centrifuged at 17,000 x g for 10 min and subsequently 250 µl were layered onto 10–50% sucrose density gradients made in association buffer. Gradients were centrifuged for 7 h at 192,100 x g. Gradients were fractionated by upwards displacement of 250 µl aliquots, which were analysed for RNA content at an absorbance of 260 nm.

Helicase assay

Reactions were set up as described [58]. Briefly, 250 nM of Cy3-labelled duplex RNA (Cy3-GCUUUACGGUGCUA, AACAAAAACAAAAUAAGCACCGUAAAGC) purchased from IDT, was incubated with 0.5 µM recombinant protein in reaction buffer (20 mM Tris pH 7.5, 50 mM potassium acetate, 5 mM magnesium acetate, 10 mM DTT, 0.1 mg/ml BSA, 10 U RNasin). The 0.5 µM recombinant protein used is at a concentration similar to that used in the affinity pulldown assays (1 µM) that show an Era-CshA interaction. The reaction was initiated with the addition of 1 mM ATP and incubated at 25°C for 10 min. Reactions were terminated by removing an aliquot at the indicated timepoints mixing with an equal volume of stop buffer.
(1% SDS, 0.5 mM EDTA, 20% glycerol) and loaded onto 20% native polyacrylamide gels before being electrophoresed in 0.5 x TBE. Bands were imaged using a ChemiDoc MP imager (Bio-Rad).

**Nucleotide hydrolysis assays**

The ability of proteins to hydrolyse GTP was determined by incubating 100 nM recombinant Era with equimolar YbeY/CshA or RelSau, 100 nM S. *aureus* ribosomes, 1 μM GTP and 2.78 nM α-<sup>32</sup>P-GTP in 40 mM Tris pH 7.5, 100 mM NaCl, 10 mM MgCl<sub>2</sub> at 37˚C for the indicated times. All reactions were set up in the absence of Era as controls. pppGpp hydrolysis assays were performed by incubating 100 nM recombinant RelSau in the presence and absence of equimolar GST, GST-Era or GST-CshA, 1 μM pppGpp and 2.78 nM α-<sup>32</sup>P-pppGpp in 50 mM Tris pH 8.5, 0.1 mM MnCl<sub>2</sub>, 20 mM KCl at 37˚C for the indicated times. All reactions were set up in the absence of RelSau as controls. Hydrolysis reactions were inactivated with the addition of formic acid to a final concentration of 1.2 M. Precipitated proteins were pelleted by centrifugation at 17,000 x g for 10 min. Reaction products were then visualized by spotting 1 μl on PEI-cellulose thin layer chromatography (TLC) plates (Merck Millipore) and separated using 1 M KH<sub>2</sub>PO<sub>4</sub>, pH 3.6 buffer. The radioactive spots were visualised using an LA 7000 Typhoon PhosphorImager and images quantified using ImageJ.

**Data provision**

All raw data for the above methods are provided in the supplemental raw data file.

**Supporting information**

S1 Fig. Antibiotic susceptibility testing. The susceptibility of LAC<sup>+</sup> and Δera to A) spectinomycin and B) chloramphenicol was measured by growing the strains in 96 well plates with the indicated concentration of each antibiotic. OD<sub>600</sub> readings were determined after 24 h of growth and plotted as % survival compared to growth without antibiotic. Experiments were repeated four times and mean and standard deviation plotted.

(TIF)

S2 Fig. Enzyme activity of CshA and Era. A) The RNA helicase activity of 0.5 μM CshA with and without an equal concentration of Era were determined using a Cy3-labelled double stranded RNA oligomer. Reactions were incubated at 25˚C for up to 10 min before analysis on a native page gel. Experiments were repeated three-four times with means and standard deviations shown. Statistical analysis was performed using a two-way ANOVA, followed by Sidak’s multiple comparisons test. B & C) The GTPase activity of 100 nM Era was measured in the presence of an equal amount of ribosomes and 1 μM GTP, plus and minus 100 nM CshA (B) or 100 nM YbeY (C). All reactions contained ribosomes. Reactions lacking Era were included as controls. Hydrolysis of <sup>32</sup>P-GTP was monitored by TLC and the percentage GDP formed quantified using ImageJ. Experiments were repeated five times with means and standard deviations shown. Statistical analysis was performed using a two-way ANOVA, followed by Dunnett’s multiple comparisons test.

(TIF)

S3 Fig. Growth of Δera and ΔcshA strains at 37 and 25˚C. Growth of S. *aureus* strains LAC<sup>+</sup>, LAC<sup>+</sup> Δera, LAC<sup>+</sup> ΔcshA and LAC<sup>+</sup> Δera ΔcshA at A) 37˚C and B) 25˚C. Overnight cultures were diluted to an OD<sub>600</sub> of 0.05 and grown for 24 h. Growth curves were performed in triplicate, with averages and standard deviations shown.

(TIF)
S4 Fig. Cross-complementation of ΔrgsA, Δera and ΔcshA strains. A) Growth of LAC\(^+\) iTET, LAC\(^+\) ΔrgsA iTET, LAC\(^+\) ΔrgsA iTET-rgsA and LAC\(^+\) ΔrgsA iTET-era at 37\(^\circ\)C. B) Growth of LAC\(^+\) iTET, LAC\(^+\) Δera iTET and LAC\(^+\) Δera iTET-rgsA at 37\(^\circ\)C. C) Growth of LAC\(^+\) iTET, LAC\(^+\) Δera iTET, LAC\(^+\) Δera iTET-era and LAC\(^+\) Δera iTET-cshA at 25\(^\circ\)C. D) Growth of LAC\(^+\) iTET, LAC\(^+\) ΔcshA iTET, LAC\(^+\) ΔcshA iTET-cshA and LAC\(^+\) ΔcshA iTET-era at 25\(^\circ\)C. E) Growth of LAC\(^+\) iTET, LAC\(^+\) ΔcshA iTET, LAC\(^+\) ΔcshA era iTET and LAC\(^+\) ΔcshA era iTET-era at 37\(^\circ\)C. F) Growth of LAC\(^+\) iTET, LAC\(^+\) ΔcshA iTET, LAC\(^+\) ΔcshA era iTET and LAC\(^+\) ΔcshA era iTET-era at 25\(^\circ\)C. Overnight cultures were diluted to an OD\(_{600}\) of 0.05 and grown in the presence of 100 ng/ml Atet for 8 h to 24 h at either 37\(^\circ\)C or 25\(^\circ\)C. Growth curves were performed three to four times, with averages and standard deviations shown.

(TIF)

S5 Fig. ppGpp does not interact with CshA. A) DRaCALA binding assays with recombinant GST, GST-CshA and Era-His and \(^{32}\)P-labelled GTP and ppGpp. Quantification was carried out using ImageJ. The average values and standard deviations of triplicate experiments are plotted. B) Binding curves and K\(_d\) determination for \(^{32}\)P-ppGpp and Era in the absence and presence of CshA. C) Hydrolysis activity of Rel\(_{Sau}\) on \(^{32}\)P-pppGpp in the absence and presence of Era and CshA. 100 nM of each protein were incubated with 1 \(\mu\)M pppGpp over the course of 5 min at 37\(^\circ\)C before reactions were quenched. Reactions lacking Rel\(_{Sau}\) or including just GST in place of GST-Era/CshA were included as controls. Experiments were repeated two to four times with means and standard deviations plotted. Statistical analysis was performed using a two-way ANOVA, followed by Dunnett’s multiple comparisons test.

(TIF)

S1 Table. Bacterial strains used in this study.

(DOCX)

S2 Table. Primers used in this study.

(DOCX)

S1 Protocol. Plasmid and strain construction.

(DOCX)

S1 Dataset. Raw data for figures.

(XLSX)

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