Regulation of global transcription in *E. coli* by Rsd and 6S RNA

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ABSTRACT

In *Escherichia coli*, the sigma factor $\sigma^{70}$ directs RNA polymerase to transcribe growth-related genes, while $\sigma^{38}$ directs transcription of stress response genes during stationary phase. Two molecules hypothesized to regulate RNA polymerase are the protein Rsd, which binds to $\sigma^{70}$, and the non-coding 6S RNA which binds to the RNA polymerase-$\sigma^{70}$ holoenzyme. Despite multiple studies, the functions of Rsd and 6S RNA remain controversial. Here we use RNA-Seq in five phases of growth to elucidate their function on a genome-wide scale. We show for the first time that Rsd and 6S RNA facilitate $\sigma^{38}$ activity throughout bacterial growth, while 6S RNA also regulates widely different genes depending upon growth phase. We discover novel interactions between 6S RNA and Rsd and show widespread expression changes in a strain lacking both regulators. Finally, we present a mathematical model of transcription which highlights the crosstalk between Rsd and 6S RNA as a crucial factor in controlling sigma factor competition and global gene expression.

INTRODUCTION

In bacteria, the core RNA polymerase ($\alpha_2\beta\beta'\omega$, referred to as E) transcribes all cellular genes. Proteins called sigma ($\sigma$) factors bind to E forming $E\sigma$ holoenzymes, and direct it to transcribe from specific promoters. *Escherichia coli* has seven sigma factors
(Ishihama 2000) – $\sigma^{70}$ (RpoD), $\sigma^{38}$ (RpoS), $\sigma^{32}$, $\sigma^{54}$, $\sigma^{28}$, $\sigma^{24}$ and $\sigma^{19}$. Though it was previously reported that the cellular concentration of E exceeded that of sigma factors (Jishage and Ishihama 1995; Maeda et al. 2000), recent quantitation has shown that sigma factors exceed E under common culture conditions (Grigorova et al. 2006; Piper et al. 2009), and may compete to bind to E. This is supported both by *in vitro* assays demonstrating competition between sigma factors under limiting E (Maeda et al. 2000; Jishage et al. 2002; Laurie et al. 2003; Bernardo et al. 2006) and *in vivo* studies where increasing activity of one sigma factor decreased the activity of others and vice-versa (Farewell et al. 1998; Jishage et al. 2002; Laurie et al. 2003).

$\sigma^{70}$ (RpoD) is the housekeeping sigma factor, directing transcription of genes essential for growth. The major alternative sigma factor, $\sigma^{38}$ (RpoS), is present in low concentration during exponential growth, but regulates several hundred genes (Rahman et al. 2006; Dong et al. 2008). Upon entry into stationary phase, while the concentrations of E and $\sigma^{70}$ show little change, $\sigma^{38}$ accumulates (Gentry et al. 1993; Tanaka et al. 1993; Piper et al. 2009; Table 1) and directs transcription of genes for multiple stress tolerance (Lange and Hengge-Aronis 1991; Hegge-Aronis et al. 1991; Hengge-Aronis and Fischer 1992, Weber et al. 2005). However, $\sigma^{70}$ remains the most abundant sigma factor, and has the highest affinity for E (Maeda et al. 2000; Colland et al. 2002; Piper et al. 2009; Ganguly and Chatterji 2012), implying that additional regulators are needed for $\sigma^{38}$ to compete effectively with $\sigma^{70}$. 

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Table 1: Cellular concentrations of RNA polymerase, sigma factors, Rsd and 6S RNA.

| Molecule        | Exponential | Stationary   | Reference                  |
|-----------------|-------------|--------------|----------------------------|
| Core RNA Polymerase | ~4.3 μM     | ~4.3 μM     | Piper et al. 2009           |
| σ70             | ~12.1 μM    | ~12.0 μM    |                            |
| σ38             | Not Detected | ~2.7 μM     |                            |
| Rsd             | ~5.5 μM     | ~10.4 μM    |                            |
| 6S RNA          | ~1000 molecules/cell | ~10,000 molecules/cell | Wassarman and Storz 2000 |

An example of such a regulator is the Crl protein, which binds to σ38 and increases its affinity for E, and promotes transcription by the Eσ38 holoenzyme at some promoters (Bougdour et al. 2004; Typas et al. 2007; England et al. 2008). The nucleotide ppGpp also increases the ability of alternative sigma factors to compete with σ70 (Jishage et al. 2002). However, ppGpp is produced transiently on entry into stationary phase (Cavanagh et al. 2010), and Crl has been shown to decrease in extended stationary phase (Bougdour et al. 2004).

On the other hand, two regulators - the protein Rsd and the non-coding 6S RNA - act on σ70. Rsd binds to σ70, sequestering it from E, and inhibits Eσ70-dependent transcription at several promoters in vitro (Jishage and Ishihama 1998). An Rsd null strain showed
increased transcription from a $\sigma^{70}$ dependent promoter, and reduced transcription from a $\sigma^{38}$ dependent promoter, in stationary phase, while Rsd overexpression had the opposite effect (Jishage and Ishihama 1999). It was hypothesized that in stationary phase, Rsd reduces $\mathrm{E}\sigma^{70}$ formation, and, by freeing $E$ to bind $\sigma^{38}$, increases $\mathrm{E}\sigma^{38}$ formation. However, a microarray experiment found no significant change in gene expression on knocking out Rsd, and even comparison of the knockout with an Rsd overexpressing strain found changed expression of only a few $\sigma^{38}$-dependent genes (Mitchell et al. 2007). Also, effects of Rsd were observed only during stationary phase, supposedly due to low Rsd levels in exponential phase. However, Rsd was later observed to be present at $\sim 50\%$ of the level of $\sigma^{70}$ in exponential phase and $\sim 90\%$ in stationary phase (Piper et al. 2009) – raising the question of why no change in expression was seen in its knockout.

6S RNA is a non-coding RNA expressed from the ssrS gene, which binds to the $\mathrm{E}\sigma^{70}$ holoenzyme (Wassarman and Storz 2000). It has been shown to block $\mathrm{E}\sigma^{70}$ binding to a target promoter (Wassarman and Saecker 2006) and inhibit transcription from several promoters in vitro and in vivo (Wassarman and Storz 2000; Trotochaud and Wassarman 2004; Gildehaus et al. 2007; Cavanagh et al. 2008; Cavanagh et al. 2010). A 6S RNA knockout showed increased expression from some $\mathrm{E}\sigma^{70}$ promoters containing extended -10 elements, and reduced expression from a few $\mathrm{E}\sigma^{38}$ promoters, in stationary phase (Trotochaud and Wassarman 2004). It was suggested that sequestration of $\mathrm{E}\sigma^{70}$ by 6S
RNA allows $\sigma^{38}$ to compete more effectively for E, increasing transcription by $E\sigma^{38}$. An alternative hypothesis was that 6S RNA regulates a trans-acting factor important for $E\sigma^{38}$ activity. There have been two microarray-based genome-wide studies on 6S RNA knockouts. While the first (Cavanagh et al. 2008) showed that 6S RNA regulated hundreds of genes in stationary phase, and an extended $-10$ element and weak $-35$ element could make a promoter sensitive to 6S RNA, the second (Neusser et al. 2010) found no correlation of 6S RNA sensitivity with promoter sequence or sigma factor preference. There was also little overlap between the regulated genes found in the two experiments.

Rsd and 6S RNA are present at high concentrations during growth, increase in stationary phase, and are thereafter at high levels (Jishage and Ishihama 1998; Wassarman and Storz 2000; Piper et al. 2009). Yet their regulatory function remains controversial and studies have been contradictory. Both are hypothesized to reduce $E\sigma^{70}$ and increase $E\sigma^{38}$ formation in stationary phase. However, it is not clear whether Rsd affects the expression of $\sigma^{70}$ target genes, nor whether it has any effect during growth – and if not, why not? Similarly, with 6S RNA, while studies concur that it reduces transcription from several $\sigma^{70}$ promoters, one study found that it regulated $\sigma^{38}$ target genes during stationary phase whereas another did not. Does it have such an effect? If so, is it a direct effect of $E\sigma^{70}$ sequestration or mediated by a trans-acting factor? What is that trans-acting factor? Finally, although both Rsd and 6S RNA have
been proposed to help $\sigma^{38}$ compete with $\sigma^{70}$, one sequesters $\sigma^{70}$ while the other sequesters the $\mathrm{E}\sigma^{70}$ holoenzyme. What impact does this have on their regulatory effects? Why does the cell require two molecules to sequester $\sigma^{70}$?

Here, we present a genome-wide investigation of the functions of Rsd and 6S RNA in *E. coli*. We used RNA-seq to identify genes regulated by Rsd and 6S RNA in five phases of growth and demonstrated that both regulate global transcription during exponential as well as stationary phase. We showed that both increase expression of $\sigma^{38}$ target genes, with 6S RNA also regulating hundreds of $\sigma^{70}$ targets, varying greatly with growth phase. We found evidence of substantial crosstalk between Rsd and 6S RNA, with each regulating the other's expression and non-additive effects on over a thousand genes. Finally, we developed a mathematical model of sigma factor competition in *E. coli*, which explains our experimental results and highlights crosstalk between Rsd and 6S RNA as a vital factor controlling sigma factor competition.

**MATERIALS AND METHODS**

**Growth conditions:** Luria-Bertani broth and agar (20 g/L) were used for routine growth. M9 defined medium (0.6% Na$_2$HPO$_4$, 0.3% KH$_2$PO$_4$, 0.05% NaCl, 0.01% NH$_4$Cl, 0.1 mM CaCl$_2$, 1 mM MgSO$_4$, 5 x 10$^{-4}$% Thiamin) supplemented with 0.5% glucose and 0.1% casamino acids was used for RNA-seq and validation. During strain construction,
ampicillin or kanamycin were used at final concentrations of 100 μg/ml and 50 μg/ml respectively.

**Strain construction:** Single gene deletions were achieved by λ Red recombination (Datsenko and Wanner 2000), using plasmids pKD46 and pKD13 and specific primers. This introduced a kanamycin resistance cassette into the chromosome. Knockout strains were selected on LB Kanamycin plates. In the rsd knockout, the resistance cassette was removed by FLP-mediated site-specific recombination using plasmid pCP20. The ΔrsdΔssrS double knockout was generated by P1 transduction from single knockouts (Thomason et al. 2007). The 3x-FLAG epitope was added to the C-terminus of Rsd by PCR using plasmid pSUB11 as template (Uzzau et al. 2001) and introduced onto the MG1655 chromosome by λ Red recombination using specific primers. The ssrS knockout was moved into this strain using P1 transduction. Strain constructions were verified by PCR using specific primers and Sanger sequencing. Primer sequences are given in Table S4.

**RNA extraction and mRNA enrichment:** Overnight cultures in M9 glucose were inoculated in 100 mL fresh M9 glucose to a final OD$_{600}$ of 0.02 and incubated at 37 °C with shaking. Two biological replicates were performed for each strain. Cells were collected by centrifugation at the early exponential (OD$_{600}$ ~0.3), mid-exponential (OD$_{600}$ ~0.8), transition to stationary (OD$_{600}$ ~1.6), stationary (16 hrs, OD$_{600}$ ~2), and late
stationary (48 hrs, OD$_{600}$ ~1.6) phases of growth. RNA was extracted using TRizol (Invitrogen), following the manufacturer's protocol. Total RNA was treated with DNase I (Invitrogen, 18068-015) according to the manufacturer's protocol. Further precipitation of RNA and ribosomal RNA cleanup was achieved using the MICROBExpress bacterial mRNA purification Kit (Ambion, AM1905) according to the manufacturer's protocol. RNA concentration was determined using a Nanodrop 2000 (Thermo Scientific) and quality was checked by visualization on agarose gels.

**RNA-Seq:** Sequencing libraries were prepared using TruSeq RNA sample preparation kit v2 (Illumina, RS-122-2001) according to the manufacturer's guidelines, checked for quality on an Agilent 2100 Bioanalyzer, and sequenced for 50 cycles from one end on an Illumina HiSeq1000 platform at the Centre for Cellular and Molecular Platforms, Bangalore. RNA-Seq data is summarized in Table S2.

**qRT-PCR for RNA-Seq validation:** qRT–PCR was carried out using specific primers to selected mRNA targets (primers in Table S4, results in Table S3). 5 ng RNA was used for each RT-PCR reaction. TAKARA One-step SYBR PrimeScript RT-PCR kit II (RR086A) was used according to the manufacturer’s protocol, on an Applied Biosystems ViiA 7 Real-Time PCR system.
Western Blotting: Cells were grown as for RNA-seq. For stationary phase samples, 5 ml culture was harvested by centrifugation. For mid-exponential phase, 10 ml was harvested. Lysates were prepared and protein concentration was estimated using BCA assay (Thermo Fisher Scientific, 23227). Lysates containing equal amounts of protein were loaded onto an SDS-PAGE gel. Proteins were elecroblotted onto a nitrocellulose membrane and probed with mouse primary antibody against the protein of interest followed by horseradish peroxidase-conjugated anti-mouse secondary antibody. The primary antibodies were: Mouse monoclonal antibody to RpoB (Neoclon, WP023), Mouse monoclonal antibody to σ^70 (Neoclon, WP004), Mouse monoclonal antibody to σ^38 (Neoclon, WP009), Mouse monoclonal anti-FLAG antibody (Sigma-Aldrich, F3165), Mouse monoclonal antibody to GroEL (Abcam, ab82592). Bands were visualized using SuperSignal West Dura Chemiluminescent Substrate (Thermo Fisher Scientific, 34076) and imaged using an ImageQuant LAS 4000 system (GE Healthcare Life Sciences). Band intensities were quantified using ImageJ (http://imagej.nih.gov/ij).

Data sources: The E. coli K-12 MG1655 genome was downloaded from NCBI (NC_000913.2). Gene coordinates were taken from RegulonDB v8.0 (Salgado et al. 2013). Lists of σ^38 target genes and ppGpp regulated genes were obtained from (Weber et al. 2005) and (Traxler et al. 2008) respectively. Lists of genes regulated by 6S RNA were obtained from (Cavanagh et al. 2008) and (Neusser et al. 2010). Coordinates of RNA polymerase binding regions and their occupancy were obtained from (Cho et al.
A list of 501 genes under constitutive $\sigma^{70}$ promoters was obtained from (Shimada et al. 2014). Of these, we selected 270 genes which were not regulated by $\sigma^{38}$ or other sigma factors according to RegulonDB (Salgado et al. 2013), (Weber et al. 2005), or our RNA-Seq from the $\sigma^{38}$ knockout.

**Data Analysis**: RNA-Seq reads were mapped to the *E. coli* K-12 MG1655 genome using BWA$^{52}$, and reads mapping uniquely were used for further analysis. DESeq (Anders and Huber 2010) was used for differential expression analysis. Fold changes and p-values for all genes are given in the Supplementary Dataset. Genes with $\leq 10$ reads mapping to them under all conditions were excluded from all analyses and plots. All statistical analyses were performed in R version 3.0.1.

**Mathematical model**: The reactions in Figure 6A are represented by a set of differential equations that determine how the dynamical variables (levels of sigma factors, holoenzymes, etc.) change with time. These equations are given in File S2. Since the timescales on which these reactions occur is much faster than typical timescales of cell division, or processes such as stress responses in stationary phase, we assume that the levels of sigma factors, holoenzymes, etc., are in quasi steady-state in the cell. Parameters that determine the specific steady-state include the dissociation constants of the various complexes, total levels of Rsd, 6S RNA, E, sigma factors, and some
others, listed in Table 2 with their default values. These values are altered in specific simulations as described in the results.

The model of holoenzyme formation (shaded in Figure 6A) describes the following reactions:

\[ E + \sigma^{70} \rightleftharpoons E\sigma^{70} \]
\[ E + \sigma^{38} \rightleftharpoons E\sigma^{38} \]
\[ Rsd + \sigma^{70} \rightleftharpoons Rsd - \sigma^{70} \]
\[ 6SRNA + E\sigma^{70} \rightleftharpoons 6SRNA - E\sigma^{70} \]

In steady state, the following equations must be fulfilled:

\[ \frac{[E][\sigma^{70}]}{[E\sigma^{70}]} = K_{E\sigma^{70}} \]
\[ \frac{[E][\sigma^{38}]}{[E\sigma^{38}]} = K_{E\sigma^{38}} \]
\[ \frac{[Rsd][\sigma^{70}]}{[Rsd - \sigma^{70}]} = K_{Rsd} \]
\[ \frac{[6SRNA][E\sigma^{70}]}{[6SRNA - E\sigma^{70}]} = K_{6S} \]
\[ E_{total} = [E] + [E\sigma^{70}] + [E\sigma^{38}] + [6SRNA - E\sigma^{70}] \]
\[ \sigma^{70}_{total} = [\sigma^{70}] + [E\sigma^{70}] + [Rsd - \sigma^{70}] + [6SRNA - E\sigma^{70}] \]
\[ \sigma^{38}_{total} = [\sigma^{38}] + [E\sigma^{38}] \]
\[ Rsd_{total} = [Rsd] + [Rsd - \sigma^{70}] \]
\[ 6SRNA_{total} = [6SRNA] + [6SRNA - E\sigma^{70}] \]
Steady-state levels of the dynamical variables were obtained from the above equations both by solving them numerically and by integrating the corresponding differential equations until they reached steady-state.

The model with DNA (represented by the full schematic in Figure 6A) includes, in addition, the following reactions:

\[ E + DNA \rightleftharpoons EDNA \]
\[ E\sigma^{70} + DNA \rightleftharpoons E\sigma^{70}DNA \]
\[ E\sigma^{38} + DNA \rightleftharpoons E\sigma^{38}DNA \]
\[ E\sigma^{70} + P_{70} \rightleftharpoons E\sigma^{70}P_{70} \rightarrow E_{e70} + \sigma^{70} + P_{70} \]
\[ E\sigma^{38} + P_{38} \rightleftharpoons E\sigma^{38}P_{38} \rightarrow E_{e38} + \sigma^{38} + P_{38} \]
\[ E_{e70} \rightarrow e E + mRNA_{70} \]
\[ E_{e38} \rightarrow e E + mRNA_{38} \]

c represents the rate of promoter clearance and \( e \) represents the rate of transcript elongation. At steady state these fulfil the following equations, in addition to equations (1) - (4) and (8) – (9). \( K_{NS} \) represents the dissociation constant for non-specific binding of RNA polymerase to DNA, which we assume is equal for \( E, E\sigma^{70} \) and \( E\sigma^{38} \). We also assume the specific binding constants of \( E\sigma^{70} \) and \( E\sigma^{38} \) to their target promoters to be equal in stationary phase.
Here, steady-state levels were obtained by integrating the corresponding differential equations until they reached steady-state.

**Data availability:** RNA-Seq data are deposited with NCBI GEO under the accession number GSE74809. Supplementary material has been uploaded to Figshare. These include Figures S1-S12, Tables S1-S4, and Files S1-S3.

**RESULTS**
RNA-seq to identify genes regulated by Rsd and 6S RNA: Figure 1A shows a schematic of the binding activity of 6S RNA and Rsd. Rsd sequesters $\sigma^{70}$ and prevents it from binding to core RNA polymerase (E), while 6S RNA binds to the E$\sigma^{70}$ holoenzyme and prevents it from binding promoters. To identify their effects on gene expression, we carried out RNA-seq to identify genes regulated by Rsd and / or 6S RNA in five growth phases.

Figure 1

Figure 1. Rsd regulates RpoS target genes throughout growth. (A) Schematic showing the binding activity of Rsd and 6S RNA. (B) Boxplot of log2 fold change in gene expression ($\Delta rsd$/wild-type, EE phase) for 313 genes whose expression is reduced at least twofold in $\Delta rpoS$/wild-type in EE phase, compared to all other genes. (C) Boxplot of log2 fold change in gene expression ($\Delta rsd$/wild-type, stationary phase) for 634 genes whose expression is reduced at least twofold in $\Delta rpoS$/wild-type in stationary phase, compared to all other genes. $p$-values are for Wilcoxon Test.
Table S1 lists strains and plasmids used. Five strains: E. coli K-12 MG1655 (Wild-type), Rsd knockout (Δrsd), 6S RNA knockout (ΔssrS), 6S RNA-Rsd double knockout (ΔrsdΔssrS), and σ^{38}/RpoS knockout (ΔrpoS) were used for RNA-seq. These had similar growth rates in M9 glucose (Figure S1). RNA-seq was performed in five growth phases: early exponential (EE), mid-exponential (ME), transition to stationary (TS), stationary (S), and late stationary (LS; time points in Methods).

* Rsd increases σ^{38}/RpoS activity throughout growth: We defined differentially expressed genes as genes whose expression changed >=2-fold in a mutant strain relative to wild-type, with FDR-adjusted p-value < 0.05. Using these criteria, the Δrsd strain showed only 16 differentially expressed genes. These included several non-coding RNAs (ryfD, sokA, oxyS, sroH, sibD) which were increased two- to six-fold in Δrsd during stationary phase. Notably, 6S RNA expression was altered in Δrsd; 6S RNA was 2.3 times its wild-type level in stationary phase, but in ME phase was reduced to about half its wild-type level.

As very few genes were differentially expressed in Δrsd, we looked for smaller changes. We found that in all growth phases, genes whose expression was significantly reduced (>=2-fold, p < 0.05) in ΔrpoS also displayed reduced expression in Δrsd. The box plots in Figure 1B-C show the distribution of log_{2} fold change in gene expression in Δrsd relative to wild-type. ~75% of genes whose expression was significantly reduced in
ΔrpoS showed reduced expression (log₂ fold change < 0) in Δrsd. Similar plots for other growth phases are in Figure S2A. This reduced expression was less than twofold in magnitude and so was not seen when searching for differentially expressed genes.

Though these fold changes are small (the median decrease in expression is 10%), we consider them important for several reasons. First, they represent a consistent and highly significant (Wilcoxon test p < 10⁻¹⁵ in all growth phases except LS) decrease in expression across hundreds of rpoS regulated genes, in all five growth phases. Second, the average overlap between the set of rpoS regulated genes in successive growth phases is only 54% - so it is not a single set of genes whose expression is reduced in Δrsd, but a different set in each growth phase. Third, the Δrsd samples showed high inter-replicate correlation in all growth phases (Table S2). Fourth, this trend of decreased expression held true when only previously reported σ³⁸ targets (Weber et al. 2005) were considered (Figure S2B). Fifth, genes whose expression was increased >=2-fold in ΔrpoS also showed increased expression in Δrsd, during EE, ME and stationary phases (Figure S2C). On the other hand, expression of genes controlled by constitutive σ⁷⁰ target promoters (Shimada et al. 2014) was not substantially altered in any phase (Figure S3).

Thus, the Rsd knockout behaved like a σ³⁸ knockout, only with smaller changes in gene expression. Western blots in stationary phase showed that σ³⁸ protein levels were
similar in Δrsd and wild-type (Figure S4; σ^{38} level in Δrsd is 94% of wild-type); however, it is extremely difficult to obtain reliable measures of small differences in protein level using western blots. We cannot be certain based on this experiment whether the reduced expression of σ^{38} targets in Δrsd is due to reduced σ^{38} protein, reduced E-σ^{38} association, or both. We attempt to address this question theoretically later.

**6S RNA regulates distinct sets of genes in all phases of growth:** What role does the Eσ^{70} - sequestering 6S RNA play in gene regulation? Is its function similar to that of the σ^{70}-sequestering Rsd? Our RNA-seq showed that the 6S RNA knockout (ΔssrS) was very different from the Rsd knockout. To begin with, it showed ≥2-fold differential expression of 447 genes in total.

Figures 2A and B show that there was very little overlap between genes differentially expressed in ΔssrS in successive growth phases, even in the two stationary phase time points. Only a few genes were differentially expressed in ΔssrS throughout growth, particularly *fau* (*ygfA*), which is downstream of 6S RNA in the same operon. *fau* expression was increased when 6S RNA was overexpressed in a wild-type background (Figure S5), indicating that this is at least partly a regulatory effect of 6S RNA and not merely a polar effect.
Figure 2. The 6S RNA regulon is highly growth-phase dependent. (A) Venn diagram showing genes whose expression is increased at least twofold in ΔssrS, in different phases of growth. (B) Venn diagram showing genes whose expression is reduced at least twofold in ΔssrS, in different phases of growth.

During exponential phase, expression of several genes for amino acid transporters (artM, artI, hisP, hisQ, hisJ, hisM, tdcC) and amino acid biosynthesis (argH, argB, thrA, thrB, thrC, asnB, glyA) was increased in ΔssrS, while expression of genes involved in stress responses (rmf, appY, yadC, ybcM, yciF, gadW, ydel, yodD, dps, hdeA, hslV, oppA, osmE, dosC) was reduced. On the other hand, in LS phase, genes involved in global transcriptional regulation (crp, hha) and the TCA cycle (sdhD, sdhC, gltA, aceB, ppc, sucA) were upregulated. Thus, the physiological function of 6S RNA appears largely growth phase-dependent; nevertheless, there are certain patterns throughout growth.
6S RNA increases $\sigma^{38}$/RpoS activity throughout growth: Like $\Delta rsd$, the $\Delta ssrS$ strain showed reduced expression of $\sigma^{38}$ target genes (Figures 3A and B, Figure S6, Wilcoxon test $p < 10^{-15}$ in all growth phases except LS). This was not due to reduced $\sigma^{38}$ protein, as western blots in stationary phase showed that the $\Delta ssrS$ strain had higher $\sigma^{38}$ protein than the wild-type (Figure S4). On the other hand, expression of genes under constitutive $\sigma^{70}$ promoters was slightly increased in the ME, TS and LS phases (Figure S7).

Figure 3

Figure 3. 6S RNA regulates RpoS target genes throughout growth. (A) Boxplot of log2 fold change in gene expression ($\Delta ssrS$/wild-type, EE phase) for 313 genes whose expression is reduced at least twofold in $\Delta rpoS$/wild-type in EE phase, compared to all other genes. (B) Boxplot of log2 fold change in gene expression ($\Delta ssrS$/wild-type, stationary phase) for 634 genes whose expression is reduced at least...
twofold in ΔrpoS/wild-type in stationary phase, compared to all other genes. p-values are for Wilcoxon Test.

6S RNA regulates RpoB, Crl and Rsd: Since the wild-type expression of 6S RNA increased in successive growth phases (Figure 4A), its effect should increase with growth phase. Indeed, with the exception of the TS phase, the number of 6S RNA regulated genes increased with growth phase. A previous study (Cavanagh et al. 2010) showed increased ppGpp in a 6S RNA knockout during the TS phase, and we observed consistent changes in gene expression (Figure S8). Since ppGpp also favors competition of alternative sigma factors with σ70 (Jishage et al. 2002), increased ppGpp may reduce the effects of ΔssrS deletion during this phase.

We found 36 genes where 6S RNA had a dose-dependent effect throughout growth, increasing in each successive growth phase (Figure S9). For instance, expression of rpoB (encoding the β subunit of core RNA polymerase) was reduced in ΔssrS, and the magnitude of reduction increased with growth phase (Figure 4B). Reduced RpoB was validated in stationary phase by qRT-PCR (Table S3) and western blotting (Figure S10). Since excess Eσ70 inhibits rpoB transcription (Fukuda et al. 1978), we suggest that deleting 6S RNA leads to higher free Eσ70, which proportionally represses rpoB. Since RpoB is limiting for the formation of core RNA polymerase (Piper et al. 2009), this implies that the cell compensates for 6S RNA loss by reducing RNA polymerase synthesis. Similarly, we observed decreased expression of genes encoding ribosomal
proteins (Figure 4C), consistent with previous reports in stationary phase (Neusser et al. 2010).

**Figure 4**

(A) log2 fold increase in 6S RNA levels in wild-type *E. coli* over successive growth phases (relative to its expression in EE phase), based on RNA-Seq. (B) log2 fold change in *rpoB* expression in ΔssrS/wild-type, in successive growth phases. (C) log2 fold change in the expression of 46 ribosomal protein genes in ΔssrS/wild-type, in successive growth phases. (D) log2 fold change in *crl* expression in ΔssrS/wild-type, in successive growth phases. (E) Western blot for 3xFLAG-tagged Rsd in wild-type and ΔssrS backgrounds, in mid-exponential and stationary phases. (F)
Boxplots showing the stationary phase wild-type expression level of genes that are reduced >=2-fold in ΔssrS/wild-type, genes that are not differentially expressed in ΔssrS, and genes that are increased >=2-fold in ΔssrS/wild-type. (G) Boxplots showing the stationary phase RNA polymerase occupancy (by ChIP-chip) of promoters belonging to genes that are reduced >=2-fold in ΔssrS/wild-type, genes that are not differentially expressed in ΔssrS, and genes that are increased >=2-fold in ΔssrS/wild-type. Only genes that were first in their transcription unit and were associated with a single RNA polymerase binding site, were included in (G). All p-values are for Wilcoxon Test.

6S RNA also represses some genes in a dose-dependent manner. Figure 4D shows that crl expression is increased in ΔssrS, and the magnitude of this effect increases with growth phase. As with the effect of reducing RNA polymerase synthesis to compensate for the loss of 6S RNA, increasing σ^38 protein and Crl may be a means to compensate for reduced Eσ^38 activity in ΔssrS bacteria.

Since we observed that Rsd regulated 6S RNA expression, we checked whether 6S RNA in turn regulated Rsd. Our RNA-Seq showed that rsd was not differentially expressed in ΔssrS. To check if 6S RNA regulated Rsd post-transcriptionally, we added a 3xFLAG tag to the C-terminal of the Rsd protein, as in previous studies (Piper et al. 2009). Indeed, western blots showed that 3xFLAG-tagged Rsd was reduced in the ΔssrS background, in both ME and stationary phases (Figure 4E).

Finally, though 6S RNA binds to Eσ^70, its effects are highly promoter-specific (Trotochaud and Wassarman 2004; Cavanagh et al. 2008; Cavanagh et al. 2010;
Neusser et al. 2010). We did not observe a link between 6S RNA sensitivity and an extended -10 or weak -35 promoter sequence as reported in (Cavanagh et al. 2008) (Figure S11), nor with any other sequence features. Instead, during stationary phase, 6S RNA sensitivity was associated with low wild-type expression (Figure 4F) and low promoter occupancy by RNA polymerase (Figure 4G, based on ChIP-chip data from (Cho et al. 2014)). Thus, our data support a model in which sequestration of RNA polymerase by 6S RNA primarily represses promoters that bind weakly to RNA polymerase.

The Rsd/6S RNA double knockout shows differential expression of a distinct set of genes: We have discovered several instances of crosstalk between 6S RNA and Rsd; apart from the fact that both sequester σ^{70} in different forms, each regulates the other's expression, and both favor the activity of σ^{38}. We therefore asked whether the double knockout of Rsd and 6S RNA showed effects distinct from the single knockouts.

We observed that in every growth phase, hundreds of genes were differentially expressed in the double knockout relative to the wild-type. This far exceeded the number of differentially expressed genes in both single knockouts, demonstrating significant crosstalk between the two regulators. Figures 5A and B highlight genes that showed differential expression in the double knockout, but less than twofold change in both single knockouts added together; there were 1780 such genes in total. These
included genes encoding DNA Gyrase and Topoisomerase I, which maintain DNA supercoiling and regulate gene expression (Peter et al. 2004), the nucleoid-associated proteins HU, H-NS and StpA, the global transcriptional regulators ArcA, LRP and IHF, and the small RNA chaperone Hfq. This suggests that Rsd and 6S RNA act together, in a partially redundant manner, to control global gene expression and nucleoid structure.

Figure 5.

Figure 5. Rsd and 6S RNA have widespread non-additive effects on gene expression. Scatterplots of log2 fold change in gene expression in the ΔrsdΔssrS double knockout versus the sum of log2 fold changes in the Δrsd and ΔssrS single knockouts, for each gene, in (A) Early exponential and (B) Stationary phase. Blue points represent genes that show differential expression (>=2-fold increase or decrease, FDR-adjusted p-value<0.05) in the double knockout, but less than twofold change in expression in both single knockouts added together.

A theoretical model suggests an explanation for the behavior of Rsd and 6S RNA:

Our RNA-Seq demonstrated that Rsd and 6S RNA regulate sigma factor competition
and gene expression at a global scale; however, there are two major results that appear counter-intuitive. First, how is it that Rsd, which sequesters $\sigma^{70}$, increases transcription of $\sigma^{38}$ target genes but has very little effect on $\sigma^{70}$ target genes? Second, how does 6S RNA, which sequesters not only $\sigma^{70}$, but also core RNA polymerase which is required for transcription by all sigma factors, nevertheless increase transcription by $\sigma^{38}$?

To answer these questions, we constructed a mathematical model of transcription during stationary phase, using parameters from literature (Table 2). We focused on stationary phase as that is when $\sigma^{38}$, 6S RNA and Rsd are at high concentrations. Our model is similar in structure to previous studies (Grigorova et al. 2004; Mauri and Klumpp 2014); however, it is the first to model stationary phase and to include both 6S RNA and Rsd. Consequently, our results differ from previous models.

Table 2: Default values of parameters used for simulations. Wherever possible, values are specific to stationary phase. These values are discussed in File S1.

| Parameter | Meaning                                      | Value    | Reference                  |
|-----------|----------------------------------------------|----------|----------------------------|
| Volume    | Average cell volume                          | $10^{-15}$ L | (Ali Azam et al. 1999)     |
| $E_{total}$ | Total cellular concentration of RNA Polymerase | 4.3 $\mu$M | (Piper et al. 2009)        |
| $\sigma^{70}_{total}$ | Total cellular concentration of $\sigma^{70}$ | 12.0 $\mu$M |                           |
| $\sigma^{38}_{total}$ | Total cellular concentration of $\sigma^{38}$ | 2.7 $\mu$M |                           |
| Symbol | Description | Value/Unit | Source |
|--------|-------------|------------|--------|
| Rsd<sub>total</sub> | Total cellular concentration of Rsd | 10.4 μM | |
| 6S RNA<sub>total</sub> | Total cellular concentration of 6S RNA | 13 μM | (Wassarman and Storz 2000) |
| DNA<sub>total</sub> | Nonspecific binding sites per cell | 4.6 x 10<sup>6</sup> | MG1655 genome size |
| P<sub>70</sub>, P<sub>38</sub> | σ-cognate promoters per cell | 200 | (Mauri and Klumpp 2014) |
| K<sub>Eo70</sub> | Dissociation constant for E - σ<sup>70</sup> binding | 3.3 nM | (Colland et al. 2002) |
| K<sub>Eo38</sub> | Dissociation constant for E - σ<sup>38</sup> binding | 15.2 nM | |
| K<sub>NS</sub> | Dissociation constant for nonspecific binding of RNA Polymerase and DNA | 10<sup>-4</sup> M | (deHaseth et al. 1978) |
| K<sub>Rsd</sub> | Dissociation constant for Rsd - σ<sup>70</sup> binding | 32 nM | (Sharma and Chatterji 2008) |
| K<sub>6S</sub> | Dissociation constant for 6S RNA - Eσ<sup>70</sup> binding | 131 nM | (Beckmann et al. 2011) |
| K<sub>EoP</sub> | Dissociation constant for holoenzyme - promoter binding | 10<sup>-7</sup> M | (deHaseth et al. 1998); (Bintu et al. 2005) |
| Operon length | Average operon length | 1000 nt | RegulonDB (Salgado et al. 2013) |
| c | Rate of promoter clearance | 0.005 s<sup>-1</sup> | (Smith and Sauer 1996); (Mauri and Klumpp 2014) |
| E | Rate of escape from elongation | 0.021 s<sup>-1</sup> | (Proshkin et al. 2010) |
Figure 6A shows a schematic of the model. Core RNA polymerase (E) binds to sigma factors ($\sigma^{70}$ and $\sigma^{38}$) forming holoenzymes (E$\sigma^{70}$ and E$\sigma^{38}$). Holoenzymes initiate transcription from target promoters (P$_{70}$ and P$_{38}$), releasing the sigma factor. The elongating RNA polymerase (E$_{e70}$ and E$_{e38}$) transcribes until released. Holoenzymes and E also bind DNA non-specifically. We focus on the steady state of this model, determined by equations (1) - (4) and (8) - (17) in Methods.

**Figure 6**

(A) Schematic of the model. Shaded area represents reactions involved in holoenzyme formation (without DNA) (B) (i) Steady-state levels of $E\sigma^{38}$ and $E\sigma^{70}$ (ii) + 6S RNA (iii) + 6S RNA (iv) + 6S RNA

**Figure 6. A mathematical model of Rsd and 6S RNA activity.** (A) Schematic of the model. Shaded area represents reactions involved in holoenzyme formation (without DNA) (B) (i) Steady-state levels of $E\sigma^{38}$ and $E\sigma^{70}$ (ii) + 6S RNA (iii) + 6S RNA (iv) + 6S RNA

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$E\sigma^{70}$ (black) and $E\sigma^{38}$ (gray), computed from equations (1) - (9), as a function of total Rsd, when 6S RNA = 0. (ii) Same when total 6S RNA = 13 μM. (iii) Steady-state levels of the 6S RNA-$E\sigma^{70}$ complex, as a function of total Rsd. (iv) Steady-state rate of transcription from $E\sigma^{70}$ dependent promoters (black) and $E\sigma^{38}$ dependent promoters (gray), as a function of total Rsd. Vertical dashed lines represent wild-type cellular concentrations in stationary phase. (v) Steady-state rate of transcription from $E\sigma^{70}$ dependent promoters (black bars) and $E\sigma^{38}$ dependent promoters (gray bars) in the wild-type, absence of Rsd, and simulated Rsd knockout. (C) (i) Steady-state rate of transcription from $E\sigma^{70}$ dependent promoters (black) and $E\sigma^{38}$ dependent promoters (gray), as a function of total 6S RNA. Vertical dashed lines represent wild-type cellular concentrations in stationary phase. (ii) Steady-state rate of transcription from $E\sigma^{70}$ dependent promoters (black bars) and $E\sigma^{38}$ dependent promoters (gray bars) in the wild-type, absence of 6S RNA, and simulated 6S RNA knockout.

To understand how Rsd and 6S RNA regulate competition between sigma factors, we initially modeled the formation of $E\sigma^{70}$ and $E\sigma^{38}$ in the absence of DNA. This is represented by the shaded area in Figure 6A and steady-state equations (1) - (9) in Methods. Figure 6B(i) depicts what happens when Rsd is added to a system containing only E, $\sigma^{70}$ and $\sigma^{38}$. Each value of Rsd on the x-axis corresponds to a separate ‘run’ of the model where we compute the steady-state for those parameter values, before moving on to the next Rsd value. We predict that as Rsd is increased, it sequesters $\sigma^{70}$, reducing $E\sigma^{70}$ formation. This allows more E to bind $\sigma^{38}$, increasing the formation of $E\sigma^{38}$, consistent with previous predictions of Rsd function (Jishage and Ishihama 1999).
However, Figure 6B(ii) predicts that when 6S RNA is also present, Rsd increases Eσ\textsuperscript{38} with little effect on Eσ\textsuperscript{70}. How? This paradoxical result can be understood with Figure 6B(iii), which shows that the concentration of the 6S RNA–Eσ\textsuperscript{70} complex decreases as Rsd is increased. When 6S RNA is present, increasing Rsd still reduces E-σ\textsuperscript{70} association; however, the reduction in Eσ\textsuperscript{70} is compensated for by the release of Eσ\textsuperscript{70} from its complex with 6S RNA, and so there is little change in the overall Eσ\textsuperscript{70} level. This prediction remains when we include DNA in the model (represented by the complete schematic in Figure 6A). Figure 6B(iv) shows that simulating increased Rsd increases transcription by Eσ\textsuperscript{38} with less effect on Eσ\textsuperscript{70} transcription.

Our Δrsd strain also displayed a ~2.3-fold increase in 6S RNA during stationary phase. Figure 6B(v) shows the predicted rate of transcription from Eσ\textsuperscript{70} and Eσ\textsuperscript{38} target promoters in the wild-type and when Rsd = 0. The third pair of bars is an approximation of conditions in the Δrsd strain, where 6S RNA is increased 2.3-fold. We see that increased expression of 6S RNA could reduce Eσ\textsuperscript{70} dependent transcription in the Δrsd strain to almost the wild-type level, so that the main observable effect of knocking out Rsd would be reduced transcription by Eσ\textsuperscript{38}.

This explains why, in both our RNA-seq and a previous study\textsuperscript{27}, Rsd increases transcription of Eσ\textsuperscript{38} target genes, with little to no effect on Eσ\textsuperscript{70}. We hypothesize that
high levels of 6S RNA cause Rsd to act as a $\sigma^{38}$ regulator and not a $\sigma^{70}$ regulator, and, moreover, that Rsd controls the level of 6S RNA to modulate its own function.

We also observed a 2.4-fold increase in oxyS (a repressor of rpoS translation) in the Rsd knockout. We did not include this in the model due to lack of quantitative data on oxyS; however, it is possible that a small reduction in $\sigma^{38}$ protein levels, perhaps due to increased oxyS, may contribute to reduced $\sigma^{38}$ activity in the $\Delta$rsd strain. However, it is difficult to confirm this.

What is the effect of 6S RNA? Our model predicts that as 6S RNA levels increase, it sequesters E$\sigma^{70}$, reducing the amount of E available to bind to both sigma factors and thus inhibiting formation of both holoenzymes. This decreases the rate of transcription by both holoenzymes (Figure 6C(i)). However, our RNA-seq showed that deleting 6S RNA actually results in reduced transcription of E$\sigma^{38}$ target genes, i.e. 6S RNA increases transcription by E$\sigma^{38}$. This is also supported by single-promoter studies in (Trotochaud and Wassarman 2004). How is this possible? A 10-fold increase or decrease in any of the parameters was not sufficient to reproduce this observation.

Apart from losing 6S RNA, our $\Delta$ssrS strain also displayed reduced Rsd and RpoB, and increased $\sigma^{38}$. Figure 6C(ii) shows the predicted rate of transcription from E$\sigma^{70}$ and E$\sigma^{38}$ target promoters in the wild-type and when 6S RNA = 0. The third pair of bars is an
approximation of conditions in the ΔssrS strain; Rsd and E are reduced by 50% and σ^{38} is increased by 50%. Eσ^{38} transcription is now reduced to less than the wild-type level; in fact, within the default parameters of our model, reducing Rsd alone from 10.4 μM to 8 μM is sufficient to lower Eσ^{38} transcription in the 6S RNA knockout below its wild-type level.

We also observed increased crl mRNA in ΔssrS. We have not modeled this due to lack of quantitative data on Crl. However, if the change in mRNA corresponds to increased Crl protein, it could partially mitigate the effect of reduced Rsd in the ΔssrS strain.

Thus, from our model we can make the strong claim that the binding reactions of Rsd and 6S RNA shown in Figure 6A cannot explain the reduced transcription of σ^{38} target genes in the ΔssrS strain. However, adding reduced Rsd levels to the model is sufficient. We therefore hypothesize that the reduced Eσ^{38} transcription in the 6S RNA knockout is due to its indirect effects primarily via Rsd.

The quantitative predictions of this model would ideally be validated in detail by in vitro transcription experiments, measuring the change in transcription of σ^{38} and σ^{70} dependent promoters with Rsd and 6S RNA in varying concentrations. We predict that the effect of Rsd on σ^{70} dependent transcription should decrease as 6S RNA is increased. We also predict that increasing 6S RNA alone should reduce both σ^{38} and...
σ^{70} dependent transcription, but σ^{38} dependent transcription can be increased on increasing Rsd. To our knowledge, such studies have not been carried out. However, such experiments are necessarily limited to a few promoters. Some predictions may also be testable by looking at distributions of gene expression in vivo; for instance, increasing Rsd expression in ΔssrS back to the wild-type level should largely mitigate the reduced expression of σ^{38} target genes.

**DISCUSSION**

Rsd and 6S RNA have long been known to regulate RNA polymerase in *E. coli*. Here we report for the first time that Rsd regulates gene expression from early exponential to stationary phase. Though it sequesters σ^{70}, its primary function is reducing transcription by the alternative sigma factor σ^{38}. Based on theoretical modeling, we suggest that this is due to 6S RNA, which minimizes the effect of Rsd on Eσ^{70} levels. We show that Rsd regulates 6S RNA expression, thereby minimizing its own effect on Eσ^{70}. Since Rsd overexpression increases transcription by σ^{24} and σ^{54} in a ppGpp^0 background (Laurie *et al.* 2003; Costanzo *et al.* 2008), Rsd may generally assist alternative sigma factors to associate with RNA polymerase, under suitable conditions.

6S RNA is more complex, regulating expression of >400 genes; however, we did not observe associations between promoter sequence and 6S RNA susceptibility as
previously reported (Cavanagh et al. 2008). Our data is more similar to that of (Neusser et al. 2010), who reported reduced expression of rpoB and ribosomal genes in ΔssrS in stationary phase, and increased ppGpp without increase in relA (spoT levels were found unchanged in all studies). However, there is still relatively low overlap in the list of 6S RNA regulated genes (46 genes). This is likely because of differences in time points and media. Even within our dataset, there is little overlap between genes regulated by 6S RNA at different time points. It seems that the 6S RNA regulon varies greatly under different conditions. This supports a model where 6S RNA acts as a background-level regulator operating on RNA polymerase, with gene-level outcomes depending strongly upon cellular environment; this could involve DNA topology, transcription factors, and other RNA polymerase-binding factors, all of which vary with growth phase (Dorman 2013). Regulators that interact with 6S RNA can be experimentally identified, for example by transposon insertion sequencing to identify genes whose disruption leads to a larger survival defect in a 6S RNA knockout than in a wild-type background, or by in vitro transcription experiments testing whether the ability of 6S RNA to regulate known targets is affected by the presence of other regulators in the medium.

Previous work using individual promoters (Trotochaud and Wassarman 2004) had suggested that 6S RNA might increase transcription by Eσ38 during stationary phase. It was unknown whether this was a direct effect of 6S RNA, allowing σ38 to compete more effectively for E, or indirect, via a trans-acting factor important for σ38 activity. However,
another study (Neusser et al. 2010) found no evidence of this link. We show that 6S RNA increases transcription by Eσ³⁸ globally, from early exponential to stationary phase, and show mathematically that this is extremely unlikely to be a direct effect of sequestration. We show that 6S RNA increases Rsd protein level and suggest that Rsd is primarily responsible for 6S RNA's effect on Eσ³⁸ activity.

It has been asked, given 6S RNA's function as a global regulator, why its deletion does not cause a growth defect. Our data, along with others (Cavanagh et al. 2010; Neusser et al. 2010), shows multiple feedback effects in the ΔssrS strain, where the cell reduces RNA polymerase expression to compensate for 6S RNA loss, and increases σ³⁸ and possibly Crl to compensate for reduced Eσ³⁸ activity. Apart from these two, increased ppGpp may also have such a compensatory role during TS phase, while in LS phase, the 6S RNA knockout has higher expression of ppk, responsible for synthesis of polyphosphate - which activates transcription by Eσ³⁸ (Kusano and Ishihama 1997). In stationary phase, the Rsd/6S RNA double knockout shows reduced expression of DNA supercoiling enzymes; supercoiling regulates promoter binding by Eσ⁷₀ and Eσ³⁸ (Kusano et al. 1996; Bordes et al. 2003); in ME phase, it shows reduced expression of Hfq, which increases σ³⁸ expression (Zhang et al. 1998). These relationships, illustrated in Figure S12, demonstrate that competition between sigma factors is very tightly controlled, and 6S RNA is connected to multiple pathways involved in this process.
We see that 6S RNA and Rsd regulate each other’s expression, and control each other’s effect on gene expression; finally, we report that 1780 genes across 5 growth phases – almost 40% of the genes in the cell - are differentially expressed in the 6S RNA-Rsd double knockout but not in the single knockouts added together. This provides direct evidence that 6S RNA and Rsd have non-additive effects on gene expression, since the single knockouts cannot explain the widespread gene expression changes observed in the double knockout. Previous studies have only examined single knockouts; this underscores the importance of studying 6S RNA and Rsd as a unit that controls global RNA polymerase activity.

Given that 6S RNA homologs are widespread in bacteria, co-occurring with Rsd and with other RNA polymerase regulators such as the actinobacterial RbpA, and many species have two or three 6S RNA homologs with different expression patterns and potentially different binding partners (Barrick et al. 2005; Wehner et al. 2014), we suggest that studying the relationships between RNA polymerase regulators will give greater insights into transcriptional control across the bacterial kingdom.

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