pH Dependence of the Stress Regulator DksA

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Abstract

DksA controls transcription of genes associated with diverse stress responses, such as amino acid and carbon starvation, oxidative stress, and iron starvation. DksA binds within the secondary channel of RNA polymerase, extending its long coiled-coil domain towards the active site. The cellular expression of DksA remains constant due to a negative feedback autoregulation, raising the question of whether DksA activity is directly modulated during stress. Here, we show that \textit{Escherichia coli} DksA is essential for survival in acidic conditions and that, while its cellular levels do not change significantly, DksA activity and binding to RNA polymerase are increased at lower pH, with a concomitant decrease in its stability. NMR data reveal pH-dependent structural changes centered at the interface of the N and C-terminal regions of DksA. Consistently, we show that a partial deletion of the N-terminal region and substitutions of a histidine 39 residue at the domain interface abolish pH sensitivity in vitro. Together, these data suggest that DksA responds to changes in pH by shifting between alternate conformations, in which competing interactions between the N- and C-terminal regions modify the protein activity.

Introduction

\textit{Escherichia coli} DksA has been shown to play a key role in regulation of transcription of ribosomal RNA and protein genes [1,2] and may also contribute to genome integrity by preventing conflicts between replication and transcription machineries [3]. In addition, DksA, often in synergy with the alarmone ppGpp, controls expression of a large number of genes required for motility [4,5], fimbriae biogenesis [6], pathogenesis [7,8], and stress responses to very diverse cellular signals, ranging from nutrient limitation [2] to oxidative and nitrosative damage [9]. Although ppGpp and DksA frequently function synergistically, examples of differential and even opposite regulation continue to accumulate [4,10,11]. Most strikingly, while both ppGpp and DksA are required for \textit{rrnB} P1 regulation by many cellular signals [1,12], ppGpp is dispensable during phosphate starvation [10].
ppGpp and DksA bind to distant sites on the core RNA polymerase (RNAP) [13,14] and reduce the stability of the promoter complexes, leading to repression or activation of transcription depending on the properties of a target promoter [1,2]. Their most pronounced effect is to shut down synthesis of very abundant rRNAs by the σ70 holoenzyme, thereby potentially making core RNAP available for binding to alternative σ factors. This indirect control of σ factors activities is consistent with the core enzyme being the target of regulation and is supported by observations that reduced levels or affinity of σ70 for the core RNAP mimics the effect of ppGpp accumulation on activation of σ5 [15] and σS4 [16] transcription in vivo. However, recent data paint a more complex picture in which ppGpp and DksA regulate alternative σs directly [17] and differently [10]. While σS and σE transcription was dependent on ppGpp during both phosphate and amino acid starvation, DksA was required for σS under both conditions but dispensable for σE response to phosphate limitation [10].

The broad repertoire of DksA targets is due to its unusual properties. Although the presence of a zinc finger initially suggested that, similarly to most initiation factors, DksA may recognize specific DNA elements in its target promoters [1], the structure of E. coli DksA [13] revealed striking similarities to a family of regulators that control transcription by directly binding to bacterial RNA polymerase (RNAP) [18,19,20]. These regulators have a common two-domain organization; structurally similar coiled-coil (CC) domains extend through the secondary channel towards the active site of RNAP, whereas dissimilar globular domains bind outside the channel. Acidic residues located at the tip of the CC domains in E. coli DksA and Gre factors and Thermus thermophilus Gfh1 approach the active site, enabling their defined regulatory functions [13,19,21,22].

The regulatory specificity of the secondary channel factors is maintained, in part, by their preferential interactions with a subset of transcription complexes. For GreB, a conformational change in RNAP is thought to enable activity on paused, backtracked complexes [23]. Similarly, two reports [24,25] suggest DksA binds to various transcription complexes with different affinities, which could, in principle, direct DksA to specific targets in the cell. However, observations that cellular levels of DksA and Gre factors remain constant throughout cell growth [26] raise a question of whether their activity could also be modulated in response to cellular environment. For example, Gfh1 has been shown to flip between an active and an inactive conformation upon a pH shift [21]. Although neither the physiological role of Gfh1 nor the regulatory role of this transition is known, it could contribute to T. thermophilus adaptation to acidity; the authors speculated that analogous conformational switches may regulate activities of other secondary channel regulators. Consistent with this idea, dksA deletions in Salmonella typhimurium and Shigella flexneri compromise survival at low pH [8,27].

We report that DksA activity and binding to RNAP increase at lower pH. Our structural analysis suggests a pH-induced structural change in DksA that involves small modifications at the interface between the globular and the CC domains. Consistent with this hypothesis, changes at the interface abolish the characteristic pH-mediated regulation of DksA activity. We demonstrate that DksA is essential for E. coli survival under acidic conditions and that its cellular levels do not change under these conditions. Finally, we propose that DksA could serve as a pH sensor in the cell.

Materials and Methods

Reagents

All plasmids are listed in S1 Table. Oligonucleotides were obtained from Integrated DNA Technologies (Coralville, IA) or Sigma (St. Louis, MO), nucleotide triphosphates (NTPs) from GE Healthcare (Piscataway, NJ), 32P-NTPs from Perkin Elmer (Waltham, MA), restriction and
modification enzymes from New England Biolabs (Ipswich, MA), PCR reagents from Roche (Indianapolis, IN), SYPRO Orange and other chemicals from Sigma. Plasmid DNAs and PCR products were purified using spin kits from Qiagen (Valencia, CA) and Promega (Madison, WI).

**Proteins**

DksA variants and RNAP were purified as described previously [13,20]. HMK-tagged DksA was radiolabeled with [$\gamma$-$^{32}$P]-ATP as described in [20]. Uniformly $^{15}$N-labeled and/or $^{13}$C/$^{15}$N-labeled DksA (either WT or N88D) were purified from cells grown in M9 minimal medium supplemented with 1% (v/v) Eagle Basal Vitamin Mix (Life Technologies, Carlsbad, CA), containing 1g/L $^{15}$N ammonium chloride as the sole nitrogen source and, in $^{13}$C/$^{15}$N samples, 3g/L $^{13}$C glucose as the sole carbon source. For WT DksA, we used a His$_{10}$-TEV DksA construct (pRF2), which was obtained by cloning DksA WT from pVS11 into PstI and HindIII sites of pIA884, and for DksA mutant N88D, we used pIA1119, all of which are derivatives of pET28a (EMD Chemicals, Gibbstown, NJ). Protein production was induced with 0.2 mM IPTG for 7 h at 30°C. Cells were lysed by French Press in a disruption buffer (50 mM Tris-HCl pH 6.9, 150 mM NaCl, 0.1 mM EDTA, 1 mM 2-mercaptoethanol). Cleared lysate was loaded on a Ni-NTA agarose (GE Healthcare) column. After four consecutive washes with 10 vol of the disruption buffer containing 0, 50, and 100 mM imidazole (pH 7.5), DksA was eluted with 250 mM imidazole and loaded on a Resource Q column (GE Healthcare) equilibrated with HepA buffer (50 mM Tris-HCl pH 6.9, 0.1 mM EDTA, 1 mM 2-mercaptoethanol, 5% glycerol). Elution was carried out using 50–1500 mM NaCl gradient; DksA eluted around 350 mM NaCl. To remove the tag, purified DksA was incubated with the His-tagged TEV protease overnight at room temperature and loaded again on Resource Q column to separate the cleaved protein from the short His-tag, the TEV protease, and the uncleaved protein. Following TEV cleavage, the proteins have three additional residues (Gly-Leu-Gln) at the N-terminus.

**In vitro transcription assays**

Transcription initiation assays were performed as described previously [28] in 20 mM Tris-HCl buffer pH 8.2 or 7.3 (measured at room temperature). IC$_{50}$ was obtained by fitting the product band intensity $I$ to the Langmuir binding equation: $I = \text{Baseline} + (A^* [\text{DksA}])/([\text{DksA}] + \text{IC}_{50})$, where $A$ is a proportionality constant, using the Scientist software (MicroMath).

**Binding assays**

Localized Fe$^{2+}$ cleavage assays were performed as described in [25] using $^{32}$P-labeled DksA$_{HMK}$ and core RNAP. Titrations were performed using 1 nM DksA and 25, 50, 100, 200 and 400 nM RNAP in 10 mM Na-HEPES at different pH and 20 mM NaCl. Gels were dried and then visualized and quantified by phosphorimaging (ImageQuant). The fraction of cleaved DksA was plotted against RNAP concentration and fitted into a single binding site equation: $K = (\text{RNAP-X})^* (\text{DksA-X})/X$, $X = \text{RNAP-DksA complex}$. Apparent $K_d$ was calculated using the Scientist software for individual assays and averaged from at least three independent experiments.

**Circular dichroism (CD) and thermostability analyses**

For CD analyses, DksA was dialyzed overnight into 20 mM phosphate buffers at pH 8, 7 and 6 with 50 mM NaCl. CD spectra were recorded using the Jasco J-815 Circular Dichroism Spectrometer from 200 μl samples. DksA concentrations were 50–100 μM. For thermostability
analysis DksA concentration was reduced to 50 μM. CD was measured at 220 nm as a function of temperature (25–85°C); with 1°C/min increments. The data were fit to the Gibbs-Helmholtz equation, assuming a ΔCp = 0, and linear baselines before and after the melting transition. That is, the temperature dependence of the CD of the folded and unfolded states was modelled as linearly dependent on temperature, with their own slopes and y-intercepts. The Kaleidagraph equation used was \((m_1+m_5^\prime(M_0))+((m_2+m_6^\prime(M_0)-(m_1+m_5^\prime M_0))^1/((1-(M_0+273)/(m_4+273))/1.987/(M_0+273))+1);\) where \(m_5\) and \(m_6\) are the slopes of the folded and unfolded states, \(m_1\) and \(m_2\) their y-intercepts, \(m_4\) is the Tm and \(m_3\) is ΔH; \(M_0\) is temperature in degrees C. Absorbance at 600 nm was recorded simultaneously to monitor protein aggregation. No changes in \(A_{600}\) were observed for any of the variants. To further exclude this possibility, DksA renaturation following gradual temperature shift from 85 to 25°C was monitored. Consistent with the lack of aggregation, the CD spectra were nearly superimposable before denaturation and after renaturation.

**Differential scanning fluorimetry**

The assay was performed as described in [29]. DksA (50 μM) was mixed with 5X SYPRO Orange (the stock concentration is not disclosed by the manufacturer) in 50 mM HEPES buffer at pH 6, 7 and 8 with 100 mM NaCl. The 25 μl reaction was incubated in a C1000 Thermal Cycler (Bio-Rad, Hercules, CA) for 25 minutes at 25°C followed by a gradual increase (1°C/30 sec) in temperature; with fluorescence intensity recorded every 1°. The unfolding temperature (Tu) is defined as the temperature at which the fluorescence increase reaches the half-maximum, and was estimated separately from three independent experiments and then averaged.

**NMR analysis**

NMR studies as a function of pH were performed on uniformly \(^{15}\)N-labeled WT DksA in 20 mM sodium phosphate, 200 mM NaCl, 0.02% NaN\(_3\), 10% D\(_2\)O at either pH 6 or 8 at 298K on a Bruker DRX-800 equipped with a 5 mm triple-resonance cryoprobe and z-axis gradient. For backbone assignments, we used the DksA\(^{N88D}\) variant, whose \(^{15}\)N-HSQC spectra are nearly identical to WT but provide higher resolution data (S1 Fig.). DksA\(^{N88D}\) NMR samples contained 0.4 mM protein, 20 mM sodium phosphate, 200 mM NaCl, 1 mM DTT, 50 μM ZnCl\(_2\), and 0.02% NaN\(_3\) at pH 6.0. The sequential backbone assignments were obtained from triple-resonance three-dimensional HNCO, HNCA, HNCACB and CBCA(CO)NH NMR spectra recorded on \([U,^{13}\)C/\(^{15}\)N]-DksA-N88D at 298K on a Bruker DRX-800 and a 3D \(^{15}\)N-edited NOESY (200 ms mixing time) recorded on a \(^{15}\)N-labeled sample at 298K on the Bruker DRX-800. Data were processed with NMRPipe and analyzed with NMRViewJ. Chemical shifts were referenced directly (\(^1\)H, \(^{13}\)C) or indirectly (\(^{15}\)N) to the external standard 2,2-dimethyl-2-silapentane-5-sulfonate (DSS). Amide \(^1\)H and \(^{15}\)N chemical shift changes were quantified as: \(\Delta \delta (ppm) = (\Delta \delta _{\text{H}}^2 + \Delta \delta _{\text{N}}^2)/25)^{1/2}\) [30].

**Growth assays**

Acid-survival was assayed as described in [27]. WT and ΔdksA cells (CH458 and CH2294, a gift from Christophe Herman) from an overnight culture grown in LB medium at pH 7.8 were diluted 1:50 into LB at pH 2.5. The medium was titrated to the desired pH using HCl or NaOH followed by filtration through a 0.2 μm filter. At different time points (0, 0.5, 1, 2, 4 and 8 h), aliquots were removed and serial dilutions were plated onto LB plates. Percentage of survival was determined using viable count after 16 h incubation at 37°C from three independent experiments. For pH adaptation experiments, cells from an overnight culture were diluted 1:50 into LB at pH 7.8, grown for 2 hours, diluted 1:50 into LB at pH 6.5, 5.5 or 4.5, grown for additional
2.5 hours, and diluted again 1:50 into LB at pH 3.5. Survival was determined as described above.

Western blotting

WT (CH458) cells and ΔdksA (CH2294) cells were grown overnight in LB media, pH 7.8. The cultures were then diluted (1:25) into LB at pH 2.5 (at time 0) and grown at 37°C. 100 ml samples were collected at ½, 1, 2 and 4 hour time points. The cells were pelleted by centrifugation, washed three times with 500 μl of 50 mM Tris–HCl, pH 7.9, 100 mM NaCl, 5% glycerol, 0.1 mM DTT, 0.1 mM EDTA, protease inhibitors (Roche-complete ULTRA tablets, Mini EDTA-free) to remove traces of acidic media, resuspended in 50 μl of same buffer, and disrupted by sonication. The extracts were cleared by centrifugation, and total protein concentrations were determined by Bradford assay. Each lane was loaded with 5 μg protein. Proteins were separated on a 4–12% SDS Bis–Tris gel (Invitrogen, Carlsbad, CA, USA), transferred onto Trans-Blot nitrocellulose membrane (Bio-Rad Laboratories, Hercules, CA, USA) in Tris–glycine buffer, pH 8.3, containing 20% methanol. The blots were blocked overnight in 1× PBS-T, pH 7.5, 0.025% Tween 20 containing 5% nonfat dry milk. Following incubation with rabbit anti-DksA polyclonal antibodies diluted 1:500 in Phosphate-Buffered Saline-Tween (PBS-T) for 2 h at room temperature, the membrane was washed and probed with rabbit IgG (GE Healthcare) for 1 h (1:5000 dilution in PBS-T), washed again and exposed to ECL detection reagents (GE Healthcare). Imaging was carried out on Chemi-doc XRS+ Molecular imager (Bio-Rad).

Results

DksA is essential during acid stress

A hypothesis that pH-dependent conformational switches in the secondary channel regulators may play a role in adaptation to pH stress, suggested by a conformational change observed in Gfh1 upon pH downshift [21], is consistent with earlier reports that S. typhimurium and S. flexneri ΔdksA mutants are sensitive to low pH [8,27]. However, growth profiling [31] did not reveal pH-dependent phenotypes for the ΔdksA E. coli strain. Therefore, we decided to directly test the role of DksA in response to acid stress in E. coli. We grew E. coli cells overnight at pH 7.8 and then diluted them into LB, pH 2.5. Next, we measured survival at different time points by plating aliquots at pH 7.8. Fig. 1A shows that, in contrast to the WT E. coli cells which persisted even after an 8-hour exposure to acid stress, the ΔdksA cells were extremely sensitive to the pH change, with 98% of cells dying within 30 minutes of exposure to low pH.

To test if DksA affects cell adaptation to acidic conditions, we grew cells at lower pH (6.5, 5.5 or 4.5) before diluting them to pH 3.5 (more surviving cells, as compared to pH 2.5, enable more reliable measurements). We observed increased survival during gradual pH adaptation in both the wild-type and ΔdksA cells (Fig. 1B). Thus, the requirement for DksA is not alleviated by adaptation.

This result might be explained by a rapid and dramatic change in DksA levels in cells exposed to low pH. However, the steady-state levels of DksA remained relatively unchanged even after 8 hours at lower pH (Fig. 1C). Another possibility is that pH-dependent structural changes enhance DksA activity at low pH.

DksA activity is sensitive to pH

Two recently proposed models generated by docking DksA into hybrid electron microscopy/X-ray models of E. coli RNAP suggest that the main binding site of DksA is a helix-hairpin-
The helix domain of the β’ subunit (rim helices, RH) located at the entrance to the secondary channel [24,25]. Both models, though different in some respects, feature a clash between the N-terminal helix of DksA and the RH, suggesting that DksA might undergo conformational changes in order to productively bind to RNAP (Fig. 2A). Consistent with the models, a deletion of the N-terminal 18 amino acids of DksA increases the activity of the protein (26). The crystal structure of DksA reveals a large interface between the N-terminal and C-terminal regions and the

Fig 1. ΔdksA mutants are sensitive to acidic conditions. (A) WT and ΔdksA E. coli strains were grown overnight in rich medium at pH 7.8. Cultures were diluted 1:50 into LB medium at pH 2.5. At selected time points aliquots were taken and the percentage of survival of bacteria was determined using viable count. (B) WT and ΔdksA E. coli strains were grown at pH 7.8, followed by 2.5 hour adaptation at pH 6.5–4.5, and then diluted into LB medium at pH 3.5. Survival was determined using viable counts; the result after 2 hour incubation at pH 3.5 is shown. (C) DksA concentration remains relatively constant at low pH. Samples taken at different time points after a change in pH were analyzed using Western blotting with anti-DksA antibodies. Extract from the ΔdksA strain and purified DksA were loaded as controls.

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CC (Fig. 2B) that contains charged or ionizable residues; one possible explanation is that changes in the protonation state of interface residues may mediate repositioning of the N-terminal helix in a manner that favors productive interaction with RNAP.

To test whether DksA activity is pH dependent, we measured in vitro transcription from the *rrnB* P1 promoter, one of the main cellular targets for DksA [1]. Although DksA may affect several steps during transcription [1,13,32], its best characterized effect occurs during initiation [1]. To eliminate possible post-initiation effects of DksA, we measured the formation of a short transcript. We incubated increasing concentrations of DksA with RNAP, ApC, UTP and [α-32P]-GTP for 15 minutes prior to the addition of a linear *rrnB* P1 template, and monitored the formation of a 4 nucleotide-long RNA product. The relative transcription was plotted against DksA concentration to determine IC₅₀ (Fig. 3A; see Materials and Methods for details). At pH 7.6, DksA inhibited transcription from the *rrnB* P1 with an IC₅₀ of 0.7 μM. Reducing pH increased the inhibitory effect by DksA, lowering the IC₅₀ to 0.11 μM at pH 6.7. Notably, RNAP activity in the absence of DksA did not change significantly in this range of pH values (S2 Fig.), suggesting that the IC₅₀ changes are due to changes in DksA (or its interactions with RNAP) rather than a change in the transcription complex.

We next tested whether the effect of pH on DksA activity can be observed at other promoters. Lyzen et al. showed that DksA (when present at < 1μM) had a 3-fold stimulatory effect at the *λP*_R promoter at pH 8 in *vitro* [33]. We also observed a stimulatory effect of DksA at pH 7.6 at *λP*_R (S3 Fig.); however, this effect was reduced at higher DksA concentrations and was
completely abolished at 5 \mu M. Interestingly, reducing pH reversed the effect of DksA at the \lambda P_R promoter from stimulation to inhibition of transcription with an IC50 of 1.5 \mu M at pH 6.7.

DksA binding to RNAP is sensitive to pH

Changes in the affinity of DksA for RNAP were shown to correlate with changes in its activity [34]. We measured the binding of DksA to core RNAP at different pH values using localized
Fe\textsuperscript{2+}-mediated cleavage (Fig. 3B). In this assay, hydroxyl radicals generated by Fe\textsuperscript{2+} bound in place of the catalytic Mg\textsuperscript{2+} ion in the RNAP active site induce cleavage of macromolecules within a \(~10\) Å radius. In a functionally bound to RNAP DksA which is labeled for visualization purposes, the CC tip is located close to the active site and is cleaved [13]. The fraction of cleaved protein is then plotted against increasing RNAP concentrations and fitted to the Langmuir binding equation, from which an apparent dissociation constant (Kd app) is derived [35]. The maximum cleavage of DksA was around 20% of the total protein and was constant at different pH. In addition, we measured binding affinities using a previously developed [25] fluorescence anisotropy assay (S4 Fig.) that measures functional as well as any non-functional binding of DksA; in the latter mode, DksA may be bound to the RNAP surface without entering the secondary channel. These different binding modes were reported previously for the yeast functional homolog of GreA, TFIIS [36]. Using both approaches, we found that the affinity of DksA for core RNAP increased dramatically upon shift from pH 8.1 to pH 6 (Fig. 3B and S4 Fig.).

DksA stability is reduced at low pH

The increased affinity of DksA for RNAP could result from structural changes in either or both components. However, the analogy to Gfh1 motivated us to investigate whether DksA may exist in a more ‘active’ form at lower pH. To examine possible changes in its secondary structure, we recorded CD spectra of DksA at pH 8, 7 and 6 (Fig. 4A). As expected from the high proportion of helical structure, the CD spectra are dominated by these features (i.e., minima at 208 and 222 nm). We did not observe significant changes in DksA spectra as a function of pH, indicating that under those conditions pH does not significantly perturb the average secondary structure. Henard et al. have recently reported that DksA activity is regulated during oxidative
and nitrosative stress using a thiol switch mechanism, wherein changes in the cysteine residues that comprise the zinc finger alter the zinc on/off rate and consequently DksA secondary structure and activity [9]. Since our data revealed no pH-dependent changes in the CD spectra, we conclude that the pH-induced change in DksA activity is not regulated by the same mechanism and does not involve cysteines modifications.

To investigate a potential pH-dependent change in the tertiary structure of DksA, we measured the protein stability using CD thermal melt assay and a differential scanning fluorimetry assay, which detects the increased fluorescence that accompanies binding of a hydrophobic dye SYPRO Orange to unfolded protein regions [29,37]. Both assays showed reduced DksA stability at pH 6 compared to pH 8 (Fig. 4 and S4 Fig.). In contrast, E. coli σ70 showed a very modest pH-induced shift in its unfolding temperature (S4 Fig.). The differential scanning fluorimetry and the CD melt show large differences in the melting temperature of DksA at both pH 8 and 6; 65 and 55°C as measured by CD vs 54 and 42°C measured by differential scanning fluorimetry. Such differences in melting temperatures are usually indicative of unfolding in two steps [38] and it is tempting to speculate that DksA unfolding involves changes in its tertiary structure followed by an alteration of its secondary structure.

**DksA exhibits structural changes at low pH**

To probe pH-induced structural changes in DksA, we collected two-dimensional $^1$H-$^{15}$N correlated NMR spectra at different pH values (Fig. 5). Upon changing from pH 8 to pH 6, a number of amide signals are significantly shifted. Backbone resonance assignments were obtained through analysis of triple resonance NMR spectra recorded at pH 6; excessive line broadening and resonance overlap allowed only partial assignments (94 of 149 amides could be assigned). Significant perturbations were observed for the backbone amide signals of residues in the

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Fig 5. DksA structure is sensitive to pH. Two-dimensional $^1$H-$^{15}$N HSQC spectra at pH 8 (black) and 6 (red) reveal large chemical shift changes at (A) Tyr23 and (B) many other residues.

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interface between the protein CC and the N- and C-terminal regions (e.g., Tyr23, Gln24, Asn33, Glu34, Gln36, Phe40, Arg41, Ile43, and Leu44; Fig. 5B and S5 Fig.). These localized chemical shift perturbations indicate localized conformational differences between pH 6 and 8.

Histidine 39 plays a role in the pH sensitivity of DksA

To probe the significance of pH-dependent changes at the interface between the C- and N-terminal regions revealed by NMR, we wanted to test whether changes at the interface would abrogate the pH-dependent change in DksA activity. Small C-terminal deletions completely abolish DksA activity [25] and hence cannot be tested. By contrast, a deletion of N-terminal 18 residues increases the protein activity at both rrnB P1 and λPR. Our results demonstrate that this deletion variant is not sensitive to pH (Fig. 6A and S6 Fig.), supporting the hypothesis that the N-terminal region is involved in the pH response of DksA.

The activity of DksA changes within pH range of 7.6 to 6.7, near the pKa values of free histidines. DksA contains two histidine residues, at positions 39 and 61. His61 is located in the middle of the CC domain and does not interact with other DksA regions whereas His39 is positioned at the interface between the two domains (Fig. 2B) and makes contacts to residues in both the N- and C-terminal regions. To determine whether these residues contribute to the pH-dependent changes in DksA, we substituted His39 and His61 individually with alanine. We first monitored transcription from rrnB P1 and λPR promoters at different pH. The DksA H61A variant exhibited a slightly reduced activity (relative to the WT protein) but displayed a similar dependence on pH, whereas the DksA H39A variant was insensitive to pH changes at both promoters (Fig. 6A and S6 Fig.). Changing His39 to the polar amino acids, Asn and Lys, also abolished the pH sensitivity at rrnB P1 (Fig. 6A).

CD analysis of the H39 variants indicated that they share a similar secondary structure as the WT, albeit exhibit dramatic reduction in their stability (S6 Fig.). This result suggests that
the substitution abrogates intramolecular interaction that helps stabilizing the tertiary but not the secondary structure of the protein.

A single Asn for Ile substitution at the 88 position of DksA increases the activity of the protein and its binding to RNAP [34]. We explored a hypothesis that the N88I substitution might lock DksA in an active conformation that can otherwise be achieved by reducing pH. We measured the activity of DksA<sup>N88I</sup>, its affinity for core RNAP, and its stability at different pH. We found that DksA<sup>N88I</sup> responds to pH similarly to the WT protein (S7 Fig.), suggesting that the N88I substitution and pH affect DksA activity through different mechanisms.

**Discussion**

DksA regulates a large set of genes during amino acid, iron, nitrogen, phosphate and carbon starvation, and is required for the proper regulation of gene expression by ppGpp, possibly the most universal stress regulator in bacteria [12]. Interestingly, while expression of some DksA-like proteins may be induced only during the time of need [28], the levels of DksA in *E. coli* are kept constant by a negative feedback mechanism during different growth phases [39]. Consistently, we did not observe dramatic changes in DksA levels upon pH downshift for at least 8 hours (Fig. 1C). Condition-specific regulation by DksA might be explained by at least two mechanisms. First, DksA function may be regulated at the level of activity through a conformational switch, as recently suggested by Henard et al. [9] and as was previously proposed for Gfh1 [21]. Second, the primary function of DksA could be to sensitize RNAP to changes in the levels of ppGpp and NTPs, whose fluctuations account for the rapid responses of DksA-regulated promoters to changes in nutritional conditions. Both mechanisms could be used to control diverse sets of DksA-dependent genes because coordinated activities of DksA and ppGpp are required for response to some stress cues but not others. Indeed, DksA has been shown to act independently of ppGpp at some promoters [4,6,33].

We show that DksA activity is stimulated by a pH downshift and that this change correlates with a stronger affinity for RNAP. We demonstrate that the structure and stability of DksA are sensitive to pH and propose that lower pH favors a more active DksA state. Observations that substitutions of His39 and the deletion of the N-terminal region eliminated DksA activation at low pH suggest that a change in the position of the N-terminal region may account for this effect. Changes in the protonation state of His39 will potentially affect hydrogen bonding between the imidazole side chain and the backbone carbonyl of Glu21 and the side chain carboxylate of Glu127, and between this same carboxylate and the backbone amide of Tyr23. This hypothesis is consistent with a steric clash hypothesized to hinder DksA binding to RNAP and is supported by a large pH-dependent chemical shift change in the 15N-HSQC spectra at Tyr23 located in the N-terminal region.

The His39 mutant is less stable yet as active as the WT protein at pH 8 but, unlike WT, fails to exhibit an increase in activity at pH 6. Both proteins show some reduced stability at lower pH, but in the case of H39A, the apparent Tu approaches the assay temperature, raising a question if the lack or H39A activation under acidic conditions could be due to its destabilization. Although this is possible, the pH-dependent change in apparent stability of the H39A variant is very small, and it is not clear how absolute Tu value measured with an isolated protein is related to the transcription activity. For example, efficient transcription initiation at the lacUV5 promoter is observed at temperatures that exceed Tu for σ<sup>70</sup> measured by this assay [40].

Although our data argue that His39 is involved in the conformational change, other changes may also occur at lower pH. For example, zinc coordination may be jeopardized at lower pH, destabilizing the DksA Zn finger. However, several lines of evidence are inconsistent with this scenario: (i) the pKa of cysteines (8.4; [41]) is above the pH range tested; (ii) we did not observe
chemical shift in C114 and in the areas surrounding the cysteines; (iii) contrary to what we would expect, lower pH increases the WT protein activity; and (iv) we did not detect pH-dependent changes in the CD spectra reported to be altered by changes in the cysteines [9]. At present, we cannot exclude a possibility that residues other than H39 contribute to pH-induced conformational changes that lead to changes in DksA activity and apparent stability. Additional experiments and higher resolution structural information, which at present seems limited by the intrinsic flexibility of the protein, will be required to fully develop the relevant mechanistic hypothesis.

Acidic pH may mimic effects of ppGpp on DksA binding to RNAP

The mechanisms that govern the synergistic effects of ppGpp and DksA are presently unclear. One suggestion is that ppGpp and DksA allosterically increase each other’s affinity to RNAP, potentiating their respective effects on gene expression [34,42]. Consistent with this idea, two dksA alleles can suppress the requirement for ppGpp during growth on a minimal medium without amino acids [34]. The encoded changes were found to increase the binding of DksA to RNAP, suggesting that an increased affinity of DksA for RNAP is sufficient to suppress some of the effects of relA/spoT (ppGpp0) deletion in the cell. Likewise, overexpression of DksA can partially suppress the growth defects of ppGpp0 mutants [43]. Locations of the DksA- and ppGpp-binding sites on RNAP [14,24,44,45] exclude the possibility of their direct interaction and point towards a model in which one regulator favors binding of the other to RNAP allosterically. Our data suggest that a similar outcome (increase in DksA binding to RNAP) could be achieved by a conformational change in DksA which occurs during acid stress. Hence, DksA may serve as a direct sensor that is “turned on” by, and allows RNAP to respond to, changes in the cellular conditions rapidly.

DksA role during acid stress

E. coli grows over a wide range of pH values and its own metabolism shifts the internal pH away from either extreme, depending on available nutrients and electron acceptors [46]. E. coli internal pH can shift significantly, from 4.7 to 7.8, when external pH changes from 2.5 to 6.9 [47]. The ability of some E. coli strains to survive exposure to strong acidic conditions is relevant for pathogenicity since bacteria have to overcome the acidic barrier of the stomach. Four overlapping systems are known to be involved in regulation of pH homeostasis in E. coli: a glucose-repressed system and three amino acid decarboxylase-dependent systems [46]. Each of these are regulated by σ5, an alternative σ factor required for gene expression during the stationary phase, thereby making the acid stress response a growth phase-dependent process [48].

Our data demonstrate that E. coli DksA is essential for survival under acidic conditions and suggest that DksA activity is stimulated by a conformational change upon reduction of the cellular pH, as observed in vitro. Our results raise several key questions that need to be addressed. First, is DksA effect direct and what genes are involved? DksA has many documented effects on the σS regulon and could potentiate expression of known pH homeostasis systems. DksA also elevates amino acids expression, of which some, such as arginine and glutamate, promote the resistance of E. coli to acid stress [47]. Second, what is the mechanism underlying the increased DksA affinity for RNAP? Third, does DksA act alone or in concert with ppGpp? DksA, but not ppGpp, is required for σ70 response to phosphate starvation [10] and ppGpp levels do not increase under mild (pH 5) extracellular acidic conditions [49]. It is possible that DksA also functions as a stand-alone stress regulator during (presumably σ5-dependent) adaptation to acid stress.
Conclusions

We show that DksA is essential for *E. coli* survival in acidic conditions and that DksA activity and affinity for RNAP are increased at lower pH. NMR data reveal pH-dependent structural changes centered at the interface of the N and C-terminal regions of DksA and changes near this interface abolish pH-dependent changes in DksA activity *in vitro*. Our results suggest that conformational switches in response to pH and other cellular cues could be common among the secondary channel regulators.

Supporting Information

**S1 Table. Plasmids.** Details for the plasmids constructed in earlier studies can be found in the cited works [13,25,28,50–52].

**S1 Fig. DksA mutant N88D facilitated backbone assignments via NMR.** Overlayed two-dimensional $^1H-^{15}N$ HSQC spectra of DksA WT (black) and N88D (orange) reveal that while the two variants of the protein yield nearly identical spectra, N88D provides higher resolution data, especially in overlapped regions of the spectra, allowing backbone assignments to be determined.

**S2 Fig. The overall transcription yield remains relatively constant at different pH.** Transcription was carried out for 15 minutes at 37°C in the absence of DksA at different pHs from the *rrnB* P1 promoter. Samples were separated by electrophoresis on 8% polyacrylamide, 7 M urea gels, and dried gels were visualized and quantified by phosphorimaging. The average signal at pH 7.6 was referred to as 1 and was used to normalize the overall signal at pH 6.7 for each experiment. The average fraction of transcription at pH 6.7 relative to pH 7.6 (normalized transcription) was calculated from 5 independent repeats.

**S3 Fig. DksA activity at the $\lambda PR$ promoter.** Increasing concentrations of DksA were added to holo RNAP (30 nM), ApU dinucleotide and [α-$^{32}$P]-GTP followed by incubation for 15 minutes. A linear DNA fragment containing the $\lambda PR$ promoter was added to initiate transcription and the formation of a 3 nucleotide RNA product was monitored on a denaturing 8% acrylamide gel. A dotted line marks the inhibition of 50% of transcription and is denoted as IC$_{50}$. The IC$_{50}$ values (calculated using a single-site binding equation from three independent repeats combined in a best-fit curve, in μM) were: pH 7.6 – >7.5, pH 6.7 – 1.5 ± 0.45.

**S4 Fig. DksA has an increased affinity to RNAP at lower pH.** DksA$^{A35C}$ (10 nM) was labeled using Atto 488 as described previously (18) and incubated with increasing concentrations of RNAP in HEPES buffer, pH 7.9 and pH 6.9, at 30°C for 10 minutes prior to anisotropy measurements. K$_{d_{app}}$ was calculated from three independent measurements. K$_{d_{app}}$ values at different pHs were: pH 7.9 – 160 nM ± 28, pH 6.9 – 48 nM ± 10. (B) Representative scan of the fluorescence emission of SYPRO Orange binding to DksA as a function of temperature. (C) Unfolding temperature (Tu) of DksA at each pH. Tu is the temperature at which fluorescence emission is half-maximal, implying that 50% of DksA is unfolded.

**S5 Fig. Chemical shift perturbations in DksA between pH 6 and 8.** Per residue weighted average amide $^1H$ and $^{15}N$ chemical shift perturbations were calculated as $\Delta \delta$ (ppm) = ($\Delta \delta_1^2 + \Delta \delta_2^2/25$)$^{1/2}$ for all residues for which assignments are available. Red line indicates the mean
value plus one standard deviation.

S6 Fig. Properties of DksA variants. (A) Transcription inhibition by DksA variants at the $\lambda_{PR}$ promoter at different pH was measured as in S3 Fig. The IC$_{50}$ values were calculated based on a single exponential fit from three independent repeats. (B) Overlaid CD spectra of DksA$^{WT}$ and DksA$^{H39N}$ recorded at room temperature using 100 $\mu$M DksA in phosphate buffer pH 8 show no difference in their secondary structure. Similar spectra were observed for other H39 variants. (C) Comparison of thermostability of DksA$^{WT}$ and DksA$^{H39}$ variants measured at 220 nm as described for Fig. 4. $\Delta$Ellipticity values were used at the Y axis to better align the different spectra and denote the change in ellipticity at increasing temperatures for each variant. Ellipticity was recorded at 220 nm wavelength. Each sample contained 50 $\mu$M DksA in phosphate buffer pH 8.

S7 Fig. The effect of pH on DksA$^{N88I}$ activity and stability. (A) The IC$_{50}$ values for DksA inhibition at the $rrnB$ P1 promoter were calculated as described for Fig. 5. (B) DksA$^{N88I}$ affinity to core RNAP was calculated using the localized Fe-mediated cleavage, as described for Fig. 3C. (C) Thermostability of the DksA$^{N88I}$ variant was determined using the differential scanning fluorimetry, as described for S4 Fig.

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Author Contributions

Conceived and designed the experiments: RF EMD MPF IA. Performed the experiments: RF EMD MN IA. Analyzed the data: RF EMD MN CY MPF IA. Contributed reagents/materials/analysis tools: RF EMD MN IA. Wrote the paper: RF EMD MPF IA.

References

1. Paul BJ, Barker MM, Ross W, Schneider DA, Webb C, Foster JW, et al. DksA: A critical component of the transcription initiation machinery that potentiates the regulation of rRNA promoters by ppGpp and the initiating NTP. Cell. 2004; 118(3):311–22. PMID:15294157
2. Paul BJ, Berkmen MB, Gourse RL. DksA potentiates direct activation of amino acid promoters by ppGpp. Proc Natl Acad Sci USA. 2005; 102(22):7823–8. PMID:15899978
3. Tehranchi AK, Blankschien MD, Zhang Y, Halliday JA, Srivatsan A, Peng J, et al. The transcription factor DksA prevents conflicts between DNA replication and transcription machinery. Cell. 2010; 141(4):595–606. doi:10.1016/j.cell.2010.03.036 PMID: 20478253
4. Aberg A, Fernandez-Vazquez J, Cabrera-Panes JD, Sanchez A, Balsalobre C. Similar and divergent effects of ppGpp and DksA deficiencies on transcription in Escherichia coli. J Bacteriol. 2009; 191(10):3226–36. doi:10.1128/JB.01410-08 PMID: 19251846
5. Lemke JJ, Durfee T, Gourse RL. DksA and ppGpp directly regulate transcription of the Escherichia coli flagellar cascade. Mol Microbiol. 2009; 74(6):1368–79. doi: 10.1111/j.1365-2958.2009.06939.x PMID: 19889089
6. Aberg A, Shingler V, Balsalobre C. Regulation of the fimB promoter: a case of differential regulation by ppGpp and DksA in vivo. Mol Microbiol. 2008; 67(6):1223–41. doi: 10.1111/j.1365-2958.2008.06115.x PMID: 18284577
7. Jude F, Kohler T, Branny P, Perron K, Mayer MP, Comte R, et al. Posttranscriptional control of quorum-sensing-dependent virulence genes by DksA in Pseudomonas aeruginosa. J Bacteriol. 2003; 185(12):3558–66. PMID: 12775693
8. Webb C, Moreno M, Wilmes-Riesenbg M, Curtiss R, Foster JW. Effects of DksA and ClpP protease on sigma S production and virulence in Salmonella typhimurium. Mol Microbiol. 1999; 34(1):112–23. PMID: 10540290

9. Henard CA, Tapscott T, Crawford MA, Husain M, Doulias PT, Porwollik S, et al. The 4-cysteine zinc-finger motif of the RNA polymerase regulator DksA serves as a thiol switch for sensing oxidative and nitrosative stress. Mol Microbiol. 2014; 91(4):790–804. doi: 10.1111/mmi.12498 PMID: 24354846

10. Gopalkrishnan S, Nicollot H, Ades SE. Co-ordinated regulation of the extracytoplasmic stress factor, sigmaE, with other Escherichia coli sigma factors by (pppGpp and DksA may be achieved by specific regulation of individual holoenzymes. Mol Microbiol. 2014; 93(3):479–86. doi: 10.1111/mmi.12674 PMID: 24946009

11. Magnusson LU, Gummesson B, Joksimovic P, Farewell A, Nystrom T. Identical, independent, and opposing roles of ppGpp and DksA in Escherichia coli. J Bacteriol. 2007; 189(14):5193–202. PMID: 17496080

12. Potrykus K, Cashel M. (p)pGpp: still magical? Ann Rev Microbiol. 2008; 62:35–51. doi: 10.1146/annurev.micro.62.081307.162903 PMID: 18454629

13. Perederina A, Svetlov V, Vassilyeva MN, Tahirov TH, Yokoyama S, Artsimovitch I, et al. Regulation through the secondary channel-structural framework for ppGpp-DksA synergism during transcription. Cell. 2004; 118(3):297–309. PMID: 15294156

14. Zuo Y, Wang Y, Steitz TA. The mechanism of E. coli RNA polymerase regulation by ppGpp is suggested by the structure of their complex. Mol Cell. 2013; 50(3):430–6. doi: 10.1016/j.molcel.2013.03.020 PMID: 23623685

15. Jishage M, Kvint K, Shingler V, Nystrom T. Regulation of sigma factor competition by the alarmone ppGpp. Genes & development. 2002; 16(10):1260–70.

16. Bernardo LM, Johansson LU, Solera D, Skarfstad E, Shingler V. The guanosine tetraphosphate (pppGpp) alarmone, DksA and promoter affinity for RNA polymerase in regulation of sigma-dependent transcription. Mol Microbiol. 2006; 60(3):749–64. PMID: 16629675

17. Costanzo A, Nicoloff H, Barchinger SE, Banta AB, Gourse RL, Ades SE. ppGpp and DksA likely regulate the activity of the extracytoplasmic stress factor sigmaE in Escherichia coli by both direct and indirect mechanisms. Mol Microbiol. 2008; 67(3):619–32. PMID: 18086212

18. Lamour V, Hogan BP, Erie DA, Darst SA. Crystal structure of Thermus aquaticus Gfh1 a Gre-factor paralog that inhibits rather than stimulates transcript cleavage. J Mol Biol. 2006; 356(1):179–88. PMID: 16337964

19. Laptenko O, Lee J, Lomakin I, Borukhov S. Transcript cleavage factors GreA and GreB act as transient catalytic components of RNA polymerase. EMBO J. 2003; 22(23):322–34. PMID: 14633991

20. Vassilyeva MN, Svetlov V, Dearborn AD, Klyuyev S, Artsimovitch I, Vassilyev DG. The carboxyl-terminal coiled-coil of the RNA polymerase beta'-subunit is the main binding site for Gre factors. EMBO Rep. 2007; 8(11):1038–43. PMID: 17917675

21. Laptenko O, Kim SS, Lee J, Starodubtseva M, Cava F, Berenguer J, et al. pH-dependent conformational switch activates the inhibitor of transcription elongation. EMBO J. 2006; 25(10):2131–41. PMID: 16628221

22. Lee JH, Lennon CW, Ross W, Gourse RL. Role of the coiled-coil tip of Escherichia coli DksA in promoter control. J Mol Biol. 2012; 416(4):503–17. doi: 10.1016/j.jmb.2011.12.028 PMID: 22200485

23. Roghanian M, Yuzenkovs Y, Zhenkin N. Controlled interplay between trigger loop and Gre factor in the RNA polymerase active centre. Nucleic Acids Res. 2011; 39(10):4352–9. doi: 10.1093/nar/gkq1359 PMID: 21266474

24. Lennon CW, Ross W, Martin-Tumasz S, Touloukhonov I, Vrentas CE, Rutherford ST, et al. Direct interactions between the coiled-coil tip of DksA and the trigger loop of RNA polymerase mediate transcriptional regulation. Genes Dev. 2012; 26:2634–46. doi: 10.1101/gad.204693.112 PMID: 23207918

25. Furman R, Tsdkiov OV, Wolf YJ, Artsimovitch I. An insertion in the catalytic trigger loop gates the secondary channel of RNA polymerase. Journal of Molecular Biology. 2013; 425(1):82–93. doi: 10.1016/j.jmb.2012.11.008 PMID: 23147217

26. Rutherford ST, Lemke JJ, Vrentas CE, Gaal T, Ross W, Gourse RL. Effects of DksA, GreA, and GreB on transcription initiation: Insights into the mechanisms of factors that bind in the secondary channel of RNA polymerase. J Mol Biol. 2007; 366(4):1243–57. PMID: 17207814

27. Mogull SA, Runyen-Janecky LJ, Hong M, Payne SM. dksA is required for intercellular spread of Shigella flexneri via an RpoS-independent mechanism. Infect Immun. 2001; 69(9):5742–51. PMID: 11500451

28. Blaby-Haas CE, Furman R, Rodionov DA, Artsimovitch I, de Crey-Lagard V. Role of a Zn-independent DksA in Zn homeostasis and stringent response. Mol Microbiol. 2011; 79(3):700–15. doi: 10.1111/j.1365-2958.2010.07475.x PMID: 21255113
29. Lavinder JJ, Hari SB, Sullivan BJ, Magliery TJ. High-throughput thermal scanning: a general, rapid dye-binding thermal shift screen for protein engineering. J Am Chem Soc. 2009; 131(11):3794–5. doi:10.1021/ja8049063 PMID:19292479

30. Garrett DS, Seok YJ, Peterkofsky A, Clore GM, Gronenborn AM. Identification by NMR of the binding surface for the histidine-containing phosphocarrier protein HPr on the N-terminal domain of enzyme I of the Escherichia coli phosphotransferase system. Biochemistry. 1997; 36(15):4393–8. PMID:9109646

31. Nichols RJ, Sen S, Choo YJ, Beltroo P, Zietek M, Chaba R, et al. Phenotypic landscape of a bacterial cell. Cell. 2011; 144(1):143–56. doi:10.1016/j.cell.2010.11.052 PMID:21185072

32. Furman R, Sevostyanova A, Artsimovitch I. Transcription initiation factor DksA has diverse effects on RNA chain elongation. Nucleic Acids Res. 2012; 40(8):3392–402. doi:10.1093/nar/gkr1273 PMID:22210857

33. Lyzen R, Kochanowska M, Wegrzyn G, Szalewska-Palasz A. Transcription from bacteriophage lambda pR promoter is regulated independently and antagonistically by DksA and ppGpp. Nucleic Acids Res. 2009; 37(20):6655–64. doi:10.1093/nar/gkp676 PMID:19759216

34. Blankschien MD, Lee JH, Grace ED, Lennon CW, Halliday JA, Ross W, et al. Super DksAs: substitutions in DksA enhancing its effects on transcription initiation. EMBO J. 2009; 28(12):1720–31. doi:10.1038/emboj.2009.126 PMID:19424178

35. Lennon CW, Gaal T, Ross W, Gourse RL. Escherichia coli DksA binds to free RNA polymerase with higher affinity than to RNA polymerase in an open complex. J Bacteriol. 2009; 191(18):5854–8. doi:10.1128/JB.00621-09 PMID:19617357

36. Ruan WJ, Lehmann E, Thomm M, Kostrewa D, Cramer P. Evolution of two modes of intrinsic RNA polymerase transcript cleavage. J Biol Chem. 2011; 286(21):18701–7. doi:10.1074/jbc.M111.222273 PMID:21454497

37. Vedadi M, Niesen FH, Allali-Hassani A, Fedorov OY, Finerty PJ, Wasney GA, et al. Chemical screening methods to identify ligands that promote protein stability, protein crystallization, and structure determination. Proc Natl Acad Sci USA. 2006; 103(43):15835–40. PMID:17035505

38. Greenfield NJ. Using circular dichroism collected as a function of temperature to determine the thermodynamics of protein unfolding and binding interactions. Nature protocols. 2006; 1(6):2527–35. PMID:17406506

39. Chandrangsu P, Lemke JJ, Gourse RL. The dksA promoter is negatively feedback regulated by DksA and ppGpp. Mol Microbiol. 2011; 80(5):1337–48. doi:10.1111/j.1365-2958.2011.07649.x PMID:21496125

40. Buc H, McClure WR. Kinetics of open complex formation between Escherichia coli RNA polymerase and the lac UV5 promoter. Evidence for a sequential mechanism involving three steps. Biochemistry. 1985; 24(11):2712–23. PMID:3896304

41. Rich AM, Bombarda E, Schenk AD, Lee PE, Cox EH, Spuches AM, et al. Thermodynamics of Zn2+ binding to Cys2His2 and Cys2HisCys zinc fingers and a Cys4 transcription factor site. J Am Chem Soc. 2012; 134(25):10405–18. doi:10.1021/ja211417g PMID:22591173

42. Magnusson LU, Nystrom T, Farewell A. Underproduction of sigma 70 mimics a stringent response. A proteome approach. J Biol Chem. 2003; 278(2):968–73. PMID:12421813

43. Vinella D, Potrykus K, Murphy H, Cashel M. Effects on growth by changes of the balance between GreA, GreB, and DksA suggest mutual competition and functional redundancy in Escherichia coli. J Bacteriol. 2012; 194(2):261–73. doi:10.1128/JB.06238-11 PMID:22056927

44. Mechold U, Potrykus K, Murphy H, Murakami KS, Cashel M. Differential regulation by ppGpp versus ppGpp in Escherichia coli. Nucleic Acids Res. 2013; 41(12):6175–89. doi:10.1093/nar/gkt302 PMID:23620295

45. Ross W, Vrentas CE, Sanchez-Vazquez P, Gaal T, Gourse RL. The magic spot: a ppGpp binding site on E. coli RNA polymerase responsible for regulation of transcription initiation. Mol Cell. 2013; 50(3):420–9. doi:10.1016/j.molcel.2013.03.021 PMID:23623682

46. Foster JW. Escherichia coli acid resistance: Tales of an amateur acidophile. Nat Rev Microbiol. 2004; 2(11):898–907. PMID:15494746

47. Richard H, Foster JW. Escherichia coli glutamate- and arginine-dependent acid resistance systems increase internal pH and reverse transmembrane potential. J Bacteriol. 2004; 186(18):6032–41. PMID:15342572

48. De Biase D, Tramonti A, Bossa F, Visca P. The response to stationary-phase stress conditions in Escherichia coli: role and regulation of the glutamic acid decarboxylase system. Mol Microbiol. 1999; 32(6):1198–211. PMID:10383761
49. Kanjee U, Gutsche I, Alexopoulos E, Zhao BY, El Bakkouri M, Thibault G, et al. Linkage between the bacterial acid stress and stringent responses: the structure of the inducible lysine decarboxylase. EMBO J. 2011; 30(5):931–44. doi:10.1038/emboj.2011.5 PMID: 21278708

50. Artsimovitch I, Svetlov V, Anthony L, Burgess RR, Landick R. RNA polymerases from Bacillus subtilis and Escherichia coli differ in recognition of regulatory signals in vitro. J Bacteriol. 2000; 182(21):6027–35. PMID: 11029421

51. Furman R, Biswas T, Danhart EM, Foster MP, Tsodikov OV, Artsimovitch I. DksA2, a zinc-independent structural analog of the transcription factor DksA. FEBS letters. 2013; 587(6):614–9. doi: 10.1016/j.febslet.2013.01.073 PMID: 23416301

52. Belogurov GA, Vassylyeva MN, Svetlov V, Klyuyev S, Grishin NV, Vassylyev DG, et al. Structural basis for converting a general transcription factor into an operon-specific virulence regulator. Mol Cell. 2007; 26(1):117–29. PMID: 17434131