Transcription factor YcjW controls the emergency H$_2$S production in *E. coli*

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Prokaryotes and eukaryotes alike endogenously generate the gaseous molecule hydrogen sulfide (H$_2$S). Bacterial H$_2$S acts as a cytoprotectant against antibiotics-induced stress and promotes redox homeostasis. In *E. coli*, endogenous H$_2$S production is primarily dependent on 3-mercaptopyruvate sulfurtransferase (3MST), encoded by *mstA*. Here, we show that cells lacking 3MST acquire a phenotypic suppressor mutation resulting in compensatory H$_2$S production and tolerance to antibiotics and oxidative stress. Using whole genome sequencing, we identified a non-synonymous mutation within an uncharacterized LacI-type transcription factor, *ycjW*. We then mapped regulatory targets of YcjW and discovered it controls the expression of carbohydrate metabolic genes and thiosulfate sulfurtransferase PspE. Induction of *pspE* expression in the suppressor strain provides an alternative mechanism for H$_2$S biosynthesis. Our results reveal a complex interaction between carbohydrate metabolism and H$_2$S production in bacteria and the role, a hitherto uncharacterized transcription factor, YcjW, plays in linking the two.
The ubiquitous gaseous molecule H$_2$S can be generated in many enzymatic pathways and from various substrates, among which L-cysteine is usually the predominant one$^{1,2}$. The main pathway by which E. coli generates H$_2$S when grown aerobically in nutrient-rich LB is via 3-mercaptopyruvate sulfurtransferase, 3MST, encoded by mstA, formerly known as sscA$^{3,4}$. Phenotypic consequences of decreased bacterial H$_2$S production include greater susceptibility to multiple classes of antibiotics and oxidative stress$^{5-8}$. The H$_2$S-mediated resistance against H$_2$O$_2$ can, in part, be explained by sequestration of ferric iron to diminish damaging Fenton chemistry$^7$. Moreover, enzymatic production of H$_2$S consumes the excess of intracellular cysteine$^8$, which otherwise could behave as a pro-oxidant that fuels the Fenton reaction$^7$. Bacterial H$_2$S has also been demonstrated recently to be important in protecting pathogens against host innate immune response$^7$. Given the overall importance of endogenous H$_2$S production for bacterial defense against stress, it has been considered as a promising target for antimicrobial therapy$^8,9$. However, to design specific inhibitors of H$_2$S production, one has to be aware of alternative endogenous sources of H$_2$S, which can be induced in response to inactivation of the main H$_2$S enzymatic pathway(s). Here, we describe one such new alternative source of H$_2$S production in E. coli and the mechanism of its regulation.

**Results and discussion**

**Phenotypic suppression of ΔmstA depends on alternative mechanism for H$_2$S synthesis.** In the course of our work, the antibiotics-sensitive strain ΔmstA often reverted to the resistant phenotype of the isogenic parent when challenged with different antibiotics. The ΔmstA variant, referred to as ΔmstA-sup in this study, was indistinguishable from wild-type in time-kill assay analysis and growth curves of cells exposed to gentamicin, nalidixic acid, and carbenicillin (Fig. 1a; Supplementary Fig. 1). Furthermore, this strain also had increased tolerance to hydrogen peroxide (H$_2$O$_2$), compared with its still sensitive parent ΔmstA strain (Fig. 1b). Using both the classic lead acetate reactivity test for H$_2$S detection and a fluorescent-based probe, WSP5$^{10}$, we confirmed that this phenotypic reversion was concurrent with increased H$_2$S production, comparable with wild-type (Fig. 1c). In contrast, significant levels of H$_2$S remained undetectable in ΔmstA till OD$_{600}$ 1.5.

S258N substitution in the transcription factor YcjW restores both H$_2$S production and antibiotic tolerance in ΔmstA-sup. We utilized whole-genome sequencing to identify possible SNPs coding regions that could be responsible for the observed phenotypic suppression. We mapped and validated by PCR a missense mutation unique to ΔmstA-sup to an uncharacterized transcription factor, ycjW. The nucleotide substitution, G to A on the coding strand, results in an amino acid change from serine to cysteine at residue 258 (Fig. 2a). Sanger sequencing confirmed the presence of the same SNP in a second, independent, ΔmstA-sup isolate as well.

YcjW is annotated as a putative member of the LacI/GalR family of repressors that are largely responsible for carbohydrate metabolism. Common features of the family include an N-terminal helix-turn-helix DNA-binding domain, a linker domain, and a C-terminal ligand-binding domain$^{11}$. To investigate SNP functionality, we constructed two strains, bearing a plasmid expressing either wild-type YcjW (pLLY1) or S258N YcjW (pLLSN1), in the background of ΔmstA/ΔycjW. Figure 2b shows that only plasmid-expressed mutated YcjW is able to restore H$_2$S production, quantitated by utilizing the WSP5 probe, and qualitatively shown by lead acetate assay. Furthermore, only ΔmstA/ΔycjW;P$_{LL}$-ycjW (S258N) has an increased survival rate when challenged with gentamicin, H$_2$O$_2$, and nalidixic acid (Fig. 2c; d; Supplementary Fig. 2). Thus, we confirm that S258N YcjW in ΔmstA-sup is responsible for the increased hydrogen sulfide production and antibiotics and oxidative stress tolerance relative to ΔmstA.

**Genome-wide mapping of the YcjW regulon in vivo.** To identify transcriptional targets of YcjW, we performed ChIP-seq using an antibody against chromosomal 3xFLAG-tagged YcjW from wild-type, and ΔmstA cells, and 3xFLAG-tagged YcjW S258N from ΔmstA-sup cells. Figure 3a shows representative peaks identified by MACS2$^{12}$ from aligned sequence reads. The most enriched regions, for all three strains, are at two sites near ycjM, the first site is before the translation start site of ycjM but after a predicted transcription start site, and the second lies between ycjT and ycjU. The binding motifs for LacI type family transcription factors are typically palindromes with a conserved central CG pair$^{13}$. Recently, Zuo and Stormo experimentally tested the predicted binding motif for YcjW$^{14}$. Combined with our analysis of peak summits, we found the same sequence in our data. Using the putative binding sequence, we further restricted peaks to ones containing the conserved 14-bp motif, allowing up to three mismatches and with a fold enrichment greater than five. With those criteria, we identified two additional peaks specifically in ΔmstA that are not enriched in wild-type or ΔmstA-sup. The two sites are near the promoter of narP, encoding a two-component nitrate/nitrite response system and the other is located within the coding sequence of cyaA, encoding adenylate cyclase (Supplementary Fig. 3).

We then validated transcription factor binding through electrophoretic mobility shift assay (EMSA). We designed 50-bp DNA probes containing the predicted binding sequence in the center. The YcjW protein reduced the mobility of the upstream ycjM DNA probe at about a 1:0.5 DNA:protein ratio. Increasing amounts of protein corresponded to an increase in YcjW–DNA complex (Fig. 3b). YcjW (S258N) also reduced DNA probe mobility at the same DNA:protein ratio, using narP probe (Fig. 3c). Titration of the normal protein and S258N YcjW showed that they both bound DNA probe starting at a DNA:protein ratio of 1:0.5. At a ratio of 1:2, no free DNA probe could be detected.

YcjW is a repressor and S258N derepresses the YcjW regulon. The region downstream of ycjM contains a predicted operon consisting of ten genes-ycjMNOPQRSTUV and ompG. To test functionality of transcription factor-DNA binding to gene expression, we determined the amount of relative mRNA fold change using qRT-PCR. In ΔycjW, representative genes, ycjM, ycjT, ycjU, and ompG are significantly upregulated, confirming that YcjW is a repressor (Fig. 3d). The absence of YcjW results in constitutive derepression of its regulatory transcriptional targets. The same genes exhibit a similar pattern, increased expression, in ΔycjW;P$_{LL}$-ycjW S258N relative to wild-type MG1655 but not to the extent of its isogenic parent, ΔycjW. Consistent with ΔycjW;P$_{LL}$-ycjW S258N, those genes are also upregulated in ΔmstA-sup, but not ΔmstA, suggesting that S258N YcjW affects DNA occupancy in vivo but not necessarily in vitro. Mutational analyses of other LacI-TFs have demonstrated that a single amino acid change in the C-terminal can alter effector or co-repressor binding and therefore DNA affinity at target sites$^{15,16}$. The SNP is located in the C-terminal effector pocket of the protein, thus raising the possibility that it broadens specificity of inducer recognition, co-repressor binding affinity, or oligomerization. qRT-PCR of narP and cyaA showed no significant change in...
A small subset of genes regulated by NarP was tested as downstream targets (Supplementary Fig. 4). Two genes, *nrfA* and *ydhU* did have a modest increase, while two others did not. NarP regulation, however, is complex and involves multiple regulators. Therefore, it is difficult to assess if YcjW-binding upstream of *nrfA* and *cyA* is functional.

Sugar kojibiose allosterically regulates the DNA-binding activity of YcjW in vitro. Because many LacI-type repressors act locally in response to some specific effector, we sought to identify the inducer for YcjW by considering its targets. YcjT is homologous to kojibiose phosphorylase from *Thermoanaerobacter brockii* and *Pyrococcus* sp. Strain ST0417,18. Kojibiose phosphorylase can reversibly catabolize kojibiose to D-glucose and beta-D-glucose 1 phosphate. The downstream gene, *ycjU*, has experimentally been shown to encode a beta-phosphoglucomutase19. Again, utilizing EMSA, we tested to see if kojibiose is the effector molecule for YcjW. The addition of kojibiose at 1 mM disrupts the YcjW–DNA complex (Fig. 3e). Other disaccharides tested in excess of up to ten times, trehalose and sucrose, did not affect binding. However, attempts to grow *E. coli* K-12 MG1655 on minimal media with kojibiose as the sole carbon source were unsuccessful20. Growth on EZ Rich Defined media supplemented with kojibiose as the carbon source did grow but had a rather pronounced defect. Deletion of *ycjW* did not improve growth rates either (Supplementary Fig. 5). However, the concentration of kojibiose added to media is limited by its low solubility. It is possible that a higher concentration of kojibiose supplied would support enhanced growth. Taken together, our results indicate that kojibiose might not be the natural inducer of YcjW, but perhaps some derivative of kojibiose. Recently, the substrate for YcjM was identified as glucosylglycerate, alongside kojibiose for YcjT. Glucosylglycerate is an osmoprotectant in bacteria and archaea, and accumulates under salt stress and limited nitrogen availability21,22. However, most of our experiments were conducted in LB with amino acids constituting the main carbon source. We find it unlikely that synthesis of either the glycoside or disaccharide could occur without the appropriate substrate, and therefore is not likely involved in ΔmstA phenotypic suppression.

**Fig. 1** *E. coli* MG1655 lacking 3MSTA acquires phenotypic suppression and has increased H₂S levels and tolerance to Gm and H₂O₂. **a** ΔmstA-sup has increased survival rate compared with ΔmstA when treated with 2 μg ml⁻¹ gentamicin in a time-kill assay. Values correspond to colony-forming units (c.f.u). **b** ΔmstA-sup also has increased tolerance after exposure to 5 mM H₂O₂ for 30 min. **c** H₂S production as measured with fluorescent probe, WSP5. Relative fluorescent units are normalized to OD₆₀₀ and minus the background fluorescent of PBS buffer + 100 μM L-cysteine and WSP5. H₂S reacts with lead acetate, leading to staining of strips (Sigma-Aldrich). Values are means ± SD (n = 3). *p < 0.05 as determined by the Student’s t test. Source data are provided as a Source Data file.
in ∆mstA-sup/∆pseE (Fig. 4b). However, at late logarithmic phase, H2S levels are now detectable to the same degree as MG1655 and ∆mstA-sup. In addition, overnight incubation with lead acetate strips shows no discernable difference in H2S extracellular production between the three strains (Supplementary Fig. 6). We conclude from the significant delay of H2S generation in ∆mstA-sup/∆pseE that PseE is capable of generating H2S in early growth phases as observed in ∆mstA-sup. However, at later growth stages, another pathway for H2S production is activated and/or PseE is no longer sufficient. Moreover, ∆mstA-sup/∆pseE also has increased sensitivity to gentamicin treatment compared with ∆mstA-sup and E. coli MG1655, but not quite as sensitive as ∆mstA. Overexpression of PseE in ∆mstA increases survival rate but only to the extent of ∆mstA-sup/∆pseE, not ∆mstA-sup or wild-type (Fig. 4c). Altogether, we conclude that the SNP in ycjW resulted in increased expression of pseE in ∆mstA-sup. This is sufficient but not wholly responsible for increased H2S biosynthesis and in turn, the phenotypic suppression observed in ∆mstA-sup. We propose a model wherein, E. coli cells lacking 3MST acquire a SNP in transcription factor YcjW. The SNP imparts moderate constitutive expression of both YcjW targets and of pseE. Thiosulfate sulfurtransferase PseE is then able to increase H2S production in ∆mstA, and subsequently protect the cells from antibiotics and H2O2 induced stress (Fig. 4d).

The region upstream of pseE neither contains a strong binding motif for YcjW nor do the regions flanking regulators PspA and PspF. This is not entirely unexpected since none of the other...
Fig. 3  YcjW shows binding enrichment near ycjM and ycjU and regulates expression of operon ycjMNOPQRSTUV-ompG. a Represented on Integrative Genomics Viewer (IGV), are sorted, aligned sequences containing pileup data to reference genome NC_000913.3. MACS2 was used for peak calling. Enriched peaks are upstream of ycjM and ycjU. A 14 nucleotide sequence identified as the binding motif for YcjW. b YcjW protein was titrated to DNA: protein ratios of 1:0.5, 1:1, and 1:2. Unlabeled ycjM probe was added to the reaction in excess to compete for binding. Unbound (free) probe and YcjW-probe complexes are denoted as U and B, respectively. c YcjW and S258N YcjW proteins were titrated to DNA:protein ratios of 1:0.125, 1:0.25, 1:0.5, 1:1, and 1:2. with narP probe. Unbound (free) probe and YcjW-probe complexes are denoted as U and B, respectively. d qRT-PCR of a subset of genes in the ycjM-V and ompG operon. The absence of ycjW results in massive upregulation. mstA-sup and ycjW;P_{LL758}ycjW both showed moderate and significant increased expression, while mRNA levels are repressed in mstA. Values are means ± SD (n = 3). *p < 0.05 as determined by the Student’s t test. e YcjW protein was pre-incubated with Kojibiose, trehalose, or sucrose before radiolabeled DNA probes were added to the mixture. Only kojibiose prevented increased expression, while mRNA levels are repressed in ΔycjM-ΔmstA-sup or ΔycjW (Supplementary Fig. 7). In addition, the moderate but significant increase of pspE mRNA in ΔycjW, in comparison with ycjMNOPQRSTUompG, suggests a more complicated interaction than direct DNA binding. It may be indicative of “leaky” expression driven by the proximity of the 3′ end of pspE to predicted transcription start sites for ycjM and the binding site of YcjW.

Activation of pspE in the ΔmstA-sup with mutated ycjW mimics a natural physiological condition where the entire YcjW regulon is derepressed. Such condition is likely to be a change in carbohydrate availability, as most members of the YcjW regulon are predicted to function in carbohydrate metabolic pathways. Therefore, our results point to a link between changes in carbohydrate availability and PspE-dependent H₂S production. The SNP in YcjW, regulating metabolism to at least two rare sugars, not only presents an interesting link between H₂S and cysteine metabolism to carbon availability but also highlights the difficulty in studying a system in isolation. Pleiotropic phenotypes, especially should be considered within global cellular context. Moreover, a SNP in YcjW reflects the striking genetic plasticity employed by bacteria to promptly adapt to...
environmental changes and stimuli, and highlights the survival advantage imparted by endogenous H$_2$S. Finally, both MstA and rhodanese enzymes are found in the mitochondrion of human cells where they play an important role in sulfide oxidation pathways. Therefore, it is possible that the same interplay of these enzymes described here in a bacterial model, is highly evolutionary conserved and has implications in human health and disease.$^{26,27}$.

**Methods**

**General growth conditions.** For the general cultivation of E. coli, strains were grown in LB broth supplemented with 50 µg ml$^{-1}$ kanamycin (cat. number 60615, Sigma), or 30 µg ml$^{-1}$ chloramphenicol (cat. number C0378, Sigma) as appropriate. Growth on solid medium contained 1.5% agar added to LB. Where noted, MOPS EZ Rich Deoxycholate broth (cat. number 1593024, Cayman) followed a modified protocol from Peng et al.$^{10}$. Brieﬂy, primers pLL14 and pLL15 were used to PCR amplify CmR cassette from pKD4. PCR product was transformed into appropriate electrocompetent strains.

**Construction of strains and plasmids.** For a list of all strains used throughout this work, refer to Table 1. BW25112 and its derivatives are from the E. coli Keio Knockout Collection (Thermo Scientiﬁc). Introduction of new mutations into E. coli MG1655 were achieved through P1 transduction as previously described.$^{30}$ Temperature-sensitive FLP recombinase plasmid pCP20 was used for the excision of selective markers, as needed.$^{31}$ All constructs were veriﬁed with PCR and sequencing. Primers used throughout this study are listed in Table S1.

To generate pLLY1, ycjW was PCR ampliﬁed from E. coli MG1655 using primers LL10 and LL11 and cloned into pACYC184 plasmid (NEB) using the Gibson Assembly Mastermix, according to the manufacturer’s protocol (NEB). Plasmid pLSN3 was generated as above, except ycjW was PCR ampliﬁed from mstA-sup. The Q5 Site-Directed Mutagenesis Kit (NEB) was used to generate pLLSN1 from pLLY1, according to the manufacturer’s protocol.

Transformations were performed using the CaCl$_2$ competent cell protocol.$^{32}$ All plasmids were sequenced for veriﬁcation.

**Addition of 3xFLAG tag to ycjW** at its chromosomal locus was achieved as previously described, with slight modiﬁcations.$^{31}$ Brieﬂy, primers pLL14 and pLL15 were used to PCR ampliﬁy CmR cassette from pKD4. PCR product was transformed into appropriate electrocompetent strains.

**H$_2$S detection.** End-point detection of H$_2$S production by lead acetate strips (cat. number WHA2602501A, Sigma) were performed, as previously described.$^{3}$ Monitoring H$_2$S generation with the WSP5 ﬂuorescent probe (cat. number 1593024–78–2, Cayman chemicals) followed a modiﬁed protocol from Peng et al.$^{10}$. Brieﬂy, cells were grown in LB at 37°C to desired OD$_{600}$ and aliquots of $4\times10^8$ cells were taken. The extinction coeﬃcient used for calculations is OD$_{600}$ of 1.0 is equal to $8\times10^8$ cells. A working solution of WSP5 was made immediately before use, and added to cells for a ﬁnal concentration of 10 µM. Samples were incubated at 37°C for 30 min and then washed in PBS buffer, pH 7.4.
to remove excess probe. Cells were resuspended in PBS buffer and incubated at room temperature for 30 min. Cytation3 (Biotek) was used to take fluorescent readings at excitation 488 nm and emission 533 nm. All experiments were repeated for a total of three times. Background subtracted values were normalized to cell number during analysis.

Time-kill assay and growth curves. Overnight cultures of E. coli were diluted 1:300 into fresh media and grown to an OD_{600} of ~0.2. A 1-ml aliquot was serially diluted and plated onto LB agar plates to determine initial colony-forming units per ml (c.f.u. ml^{-1}) after overnight incubation at 37°C. Antibiotics were added to the cultures at indicated concentrations. Aliquots of 1 ml were collected at specified time intervals, serially diluted and plated. The results from three independent experiments were plotted in GraphPad version 5.0.

Growth curves were generated from Bioscreen C automated growth analysis system as previously described. Antibiotics were purchased from Sigma-Aldrich or Gold Biotechnology.

Whole-genome sequencing. Overnight cultures of E. coli cells were used for genomic DNA isolation. The MasterPure Complete DNA Purification Kit (Epicentre) was used to purify DNA according to the manufacturer’s protocol. DNA samples were quantified using the Quant-IT PicoGreen dsDNA assay kit (Thermo Fisher) according to the manufacturer’s protocol. DNA was sheared to appropriate size with Covaris, followed by adaptor ligation. Sequencing was performed at New York University School of Medicine’s Genome Technology Center.

Quantitative RT-PCR. Cells were grown until appropriate OD_{600} and aliquots were collected and treated with RNAProtect Bacteria Reagent (Qiagen). After 5 min, cells were harvested and resuspended in lysis buffer (RNase-free TE buffer, 10 mg ml^{-1} 1% SDS, 100 µg ml^{-1}). Trizol LS (Thermo Scientific) was used according to the manufacturer’s protocol to extract the total RNA. Samples were treated with DNase (Invitrogen) and purified using a spin column (Innogenetics). Superscript III reverse transcriptase (Invitrogen) was used to synthesize cDNA. qPCR reactions were amplified using Power SYBR Green PCR Master Mix (Applied Biosystems) with appropriate primer sets and the cDNA template.

ChiP-seq. ChiP was carried out as previously described with the following modifications. Briefly, cells were grown at 37°C to OD_{600} ~0.4 and a final concentration of 1% formaldehyde was added for in vivo cross-linking of nucleoprotein. A final concentration of 0.5 M glycine was added to the culture to quench the reaction after a 20-min incubation. Cells were collected by centrifugation and washing twice with 1x cold Tris-buffered saline then frozen in liquid nitrogen and stored at −80°C. Cells were resuspended in lysis buffer (50 mM Tris [pH 7.5], 100 mM NaCl, 1 mM EDTA, protease inhibitor [Roche], 10 mg ml^{-1} 1% SDS). After incubation at 37°C, IP buffer (50 mM HEPES-KOH, 150 mM NaCl, 1 mM EDTA, 1% Triton X100, 0.1% sodium deoxycholate, 0.1% SDS, protease inhibitor [Roche]) was added at a 1:3 ratio. DNA was sheared using ultrasonicator Covaris M220 on a 10 s on and 10 s off cycle for a total of 50 cycles. The supernatant was incubated with 3xFLAG antibody (Biolegend) and Dynabeads Protein G (Thermo Scientific) overnight at 4°C. Samples were then washed twice with IP buffer, once with IP buffer + 500 mM NaCl, once with wash buffer (10 mM Tris, 250 mM LiCl, 1 mM EDTA, 0.5% NP-40, 0.5% sodium deoxycholate), and a final wash with TE. Immunoprecipitated complexes were eluted in ligation buffer (50 mM Tris, 10 mM EDTA, 1% SDS) at 65°C for 20 min. Samples were treated with RNase A (Qiagen), at 42°C and then uncross-linked with elution buffer + pronase for 2 h at 42°C, followed by 6 h at 65°C. DNA was purified using ChiP Clean and Concentrate (Zymo Research). Prior to sequencing, DNA was checked on TapeStation 2200 for appropriate size (Agilent). ChiP experiments were repeated for a total of three replicates.

For sequencing, sample libraries were prepared by using the NEBNext ChiP-seq Library (Illumina), according to the manufacturer’s protocol. Samples were sequenced on NextSeq500 (Illumina). Bowtie and MACS2 were used for aligning and peak calling, respectively.

Electrophoretic mobility shift assay. Protein purification: Yeast and S25RN Yeast were cloned into plasmid p287 SUMO using the Gibson Assembly Mastermix kit, according to the manufacturer’s protocol (NEB). Auto-induction media was used for maximizing protein yield. Cells were harvested and resuspended in lysis buffer (1 M NaCl, 5 mM imidazole, 5% glycerol, protease inhibitor cocktail [Roche]) and sonicated. AKTA Start system was used for chromatography with HisTrapHP columns (GE Healthcare Life Sciences). Columns were washed in wash buffer (50 mM Tris-Cl [pH 8.0], 10 mM imidazole, 5% glycerol, 500 mM NaCl), followed by gradient elution with ligation buffer (50 mM Tris-Cl [pH 8.0], 250 mM imidazole, 5% glycerol, 250 mM NaCl). The SUMO tag was cleaved with SUMO protease in dialysis buffer (200 mM NaCl, 50 mM Tris-Cl [pH 8.0], 5% glycerol, 1 mM DTT). Samples were applied to HiTrap Heparin HP column (GE). Columns were washed with buffer (20 mM Tris [pH 8.0]). In all, 50 mM NaCl, 5% glycerol), and eluted in ligation buffer (20 mM Tris [pH 8.0], 1.5 M NaCl, 5% glycerol). The sample was concentrated to 5 ml and injected onto a Superdex 200 column with GF buffer (20 mM Tris-Cl [pH 8.0], 50 mM NaCl, 1 mM DTT).

EMSA: DNA probes containing the binding sequence were radiolabeled with gamma-^{32}P rATP using T4 polynucleotide kinase (NEB). Labeled probes were purified by passage through size-exclusion columns (Bio-Rad). Binding reactions were done as previously described. The gel was then exposed to a phosphor screen and visualized on Storm 820 Phosphorimager (GE Healthcare). Experiments with various disaccharides were done in a similar fashion, except purified protein was incubated with appropriate sugar for 20 min at room temperature before addition of radiolabeled probe.

Data availability. Data underlying Figs 1, 2, 3, 4, and Supplementary Figs 4 and 7 are provided as Source Data files. All other data are available from the corresponding author upon reasonable request. All sequencing data that support the findings of this study have been deposited in NCBI SRA with the accession code PRJNA524143.
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Author contributions
L.L. and A.R. performed experiments and analyzed the data. I.S. performed NGS. L.L., A.R. and E.N. wrote the paper. E.N. supervised the project.

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