Identification of a High-Affinity Pyruvate Receptor in *Escherichia coli*

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Two-component systems are crucial for signal perception and modulation of bacterial behavior. Nevertheless, to date, very few ligands have been identified that directly interact with histidine kinases. The histidine kinase/response regulator system YehU/YehT of *Escherichia coli* is part of a nutrient-sensing network. Here we demonstrate that this system senses the onset of nutrient limitation in amino acid rich media and responds to extracellular pyruvate. Binding of radiolabeled pyruvate was found for full-length YehU in right-side-out membrane vesicles as well as for a truncated, membrane-integrated variant, confirming that YehU is a high-affinity receptor for extracellular pyruvate. Therefore we propose to rename YehU/YehT as BtsS/BtsR, after ”*Brenztraubensäure*”, the name given to pyruvic acid when it was first synthesized. The function of BtsS/BtsR was also assessed in a clinically relevant uropathogenic *E. coli* strain. Quantitative transcriptional analysis revealed BtsS/BtsR importance during acute and chronic urinary-tract infections.

Exponential growth of bacteria in complex, nutrient-rich media usually ends when at least one nutrient has been used up. We recently reported that the histidine kinase/response regulator system YehU/YehT of *E. coli*, belongs to the LytS/LytTR family and presumably plays a role in tuning bacterial exploitation of available carbon sources1. Strikingly, the YehU/YehT system is the most widespread representative of its family found in γ-proteobacteria – and many LytS/LytTR-type systems regulate crucial host-specific mechanisms during infection of human or plant hosts by members of this bacterial clade2. This system is conserved in non-pathogenic as well as pathogenic *E. coli*.

Our previous studies on YehU/YehT in *E. coli* identified yjiY as its sole target gene1 (Fig. 1). This gene codes for the putative carbon starvation transporter YjiY, which is homologous (61.1% identity) to CstA3 and was found to be expressed in cells that were grown in complex media containing a high content of amino acids, such as LB or CAA (casamino acids), as well as in minimal medium supplemented with certain carbon sources, such as gluconic or glucuronic acid. Studies in *E. coli* revealed that YehT-mediated yjiY transcription is also regulated by the cAMP/CRP complex5 (Fig. 1), and down-regulated in the presence of energetically favorable carbon sources like glucose. Furthermore, YjiY is subject to translational control via the Csr regulatory circuit5 (Fig. 1), which synchronizes the output of *E. coli* central carbohydrate metabolism (glycolysis versus gluconeogenesis) with YjiY production1, 6. Finally, yjiY transcription is under positive feedback regulation by a second two-component system, YpdA/YpdB, and its gene product YjxY2 (Fig. 1).

Here, we performed a comprehensive *in vivo* characterization of yjiY expression in order to identify the primary stimulus sensed by the histidine kinase YehU. We found that the YehU/YehT system responds to depletion of nutrients specifically serine and the concomitant presence of extracellular pyruvate. Biochemical studies revealed that pyruvate binds specifically to the extracellular side of the membrane-spanning domain of YehU. We therefore renamed the system BtsS/BtsR, for ”*Brenztraubensäure*”, the original name given by Jöns Jakob 1

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Berzelius to the compound when he first synthesized pyruvic acid in 1835. Finally, we found that the BtsS/BtsR system of uropathogenic E. coli may contribute to acute urinary tract infection.

**Results**

**Elucidation of the stimulus for BtsS/BtsR (YehU/YehT) by in vivo yjiY expression analyses.** It was previously shown that cultivation of E. coli in amino acid-rich media leads to activation of the BtsS/BtsR system and transient expression of the target gene yjiY in the late-exponential growth phase. To identify a potential quorum sensing-like molecule, we used a combination of chemical fractionation of the medium and measurements of reporter strain activity. For this purpose, we cultivated E. coli MG1655 ΔyjiY/pBBR yjiY-lux in M9-minimal medium with gluconic acid as sole carbon source. Shortly before the induction of yjiY, we removed the cells and fractionated the supernatant by high pressure liquid chromatography. All fractions were analyzed for their potential to induce yjiY using reporter strain E. coli KX1468 pBBR yjiY-lux grown in minimal medium and succinate as C-source. After several rounds of fractionation/freeze-drying we found that the fraction with the highest induction potential contained a high concentration of gluconic acid, the initial carbon source (data not shown). This result ruled out the possibility that E. coli produces and senses a quorum sensing-like molecule. Then we quantified yjiY expression as a function of nutrient levels. For this purpose, we cultivated the reporter strain E. coli MG1655/pBBR yjiY-lux in LB medium with decreasing amounts of nutrients (1.0x, 0.5x, 0.4x, 0.3x LB, 0.2x LB and 0.1x LB), keeping the osmolarity of the medium constant. The growth rates (µ) of E. coli cells decreased with the dilution of LB medium, and exponential growth ceased at different time points (Table S1). Strikingly, expression of yjiY always began shortly before the onset of stationary phase (Fig. S1), and E. coli cells grown in 0.1x LB did not express yjiY. These results suggested that BtsS/BtsR somehow responds to nutrient limitation. We reasoned that supplying the relevant nutrient(s) in excess should suppress or postpone yjiY induction. Therefore, the reporter strain was grown in LB media supplemented with an excess of each individual L-amino acid (Fig. 2A). Particularly, the addition of L-serine delayed the expression of yjiY by almost two doubling periods (Fig. 2A). Subsequently, we tested different concentrations of L-serine in the reporter assay and found a concentration-dependent delay in yjiY expression, accompanied by a decrease in peak expression levels (Fig. 2B). The addition of serine does not influence the growth of E. coli and does not delay the onset of stationary phase (Fig. 2C). Although L-serine is not a preferred carbon source for E. coli and high external concentrations are actually toxic to the organism, it is the first amino acid to be consumed when mixtures of amino acids are available. These data suggest that BtsS/BtsR responds to depletion of nutrients, specifically serine.

**Changes in extracellular serine and pyruvate concentrations during growth of E. coli.** When E. coli is grown in amino acid-rich media, 50.7% of L-serine is converted directly to pyruvate, whereas 36.3% is used for glycine synthesis, 6.5% for cell biomass, and the remainder for other metabolites. Its central role in pyruvate supply provides one explanation for the importance of L-serine in growing E. coli. We therefore monitored the changes in extracellular serine and pyruvate concentrations during growth in LB medium, and found that extracellular levels of serine decreased at a constant rate (Fig. 3). The starting concentration of serine in the medium (approximately 200 µM) was completely exhausted after 120 min of growth at the late-exponential growth phase. At the same time, the abundance of extracellular pyruvate peaked (approx. 500 µM) and shortly after yjiY expression reached its maximum level (Fig. 3). It was previously shown that the external pyruvate derives from overflow metabolism.
metabolism in *E. coli* during growth in amino acid-rich media \(^1\), which was confirmed by monitoring the intracellular concentrations of serine and pyruvate (Fig. S2).

These data reveal that induction of BtsS-dependent *yjiY* expression coincides with the decline of serine in the medium and an extracellular accumulation of pyruvate (Fig. 3). Extracellular pyruvate triggers *yjiY* expression under nutrient limitation. In the next experiment we tested the influence of serine, pyruvate and related metabolites on *yjiY* expression in *E. coli* cells growing in low-nutrient environment. Since *E. coli* harbors a second two-component system, YpdA/YpdB, which responds to high concentrations of extracellular pyruvate (the threshold concentration that leads to induction was determined to be 600 \(\mu\)M) and positively regulates the BtsS/BtsR system \(^1\), we modified our reporter strain by deleting *yhhX*, which is sufficient to interrupt the feedback loop \(^1\). The resulting strain was then cultivated in 10-fold diluted (0.1x) LB medium for 1 h. At this time point, cells do not induce expression of *yjiY*, but experience soon carbon limitation (Fig. 4A, Fig. S1). However, expression of *yjiY* was rapidly triggered upon addition of pyruvate, and the induction level increased linearly with increasing pyruvate concentration (Fig. 4A). Addition of L-serine also induced *yjiY* expression, but only after a 20-min delay (Fig. 4B). Under these conditions the growth of *E. coli* did not differ significantly by addition of pyruvate or serine (Fig. S3). Only supplementation of 10 mM serine prolonged the exponential growth phase. Moreover, higher serine concentrations delayed *yjiY* expression for even longer, and decreased the level of induction attained (Fig. 4B). The threshold concentration of pyruvate required for detectable *yjiY* expression was 10 \(\mu\)M, and that for L-serine 50 \(\mu\)M. None of the other tested compounds (each of the other 19 amino acids, phosphoenolpyruvate, lactate, oxaloacetate, \(\alpha\)-ketoglutarate, valerate, propionate, acetate, malate) were able to induce *yjiY* in this context. These results suggest that extracellular pyruvate acts as a direct stimulus for BtsS/BtsR-mediated *yjiY* expression, whereas delayed *yjiY* induction in response to L-serine

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**Figure 2.** Effects of an excess of individual amino acids on the expression of *yjiY*. (A) *E. coli* MG1655/pBBR *yjiY*-lux was grown in LB medium supplemented with one of the indicated amino acids (at 20 mM), and growth and luminescence were monitored over time. Maximal *yjiY* expression values are depicted. The open circle provides a benchmark and indicates *yjiY* expression in *E. coli* grown in LB medium. (B) Expression of *yjiY* in LB (dotted line) supplemented with increasing L-serine concentrations. (C) Corresponding *E. coli* growth curves in LB media supplemented with increasing L-serine concentrations. Experiments were performed at least three times (standard deviation <10%), and results of a representative experiment are shown.
may depend on uptake of the amino acid, its conversion to pyruvate and excretion of the pyruvate into the culture medium.

**The sensor histidine kinase BtsS binds pyruvate with high affinity.** BtsS cannot be autophosphorylated, possibly owing to a defective ATP-binding site within the G1 box

to determine whether BtsS physically interacts with pyruvate as ligand. The technique is based on the ability of proteins that have been immobilized on a nitrocellulose membrane to bind a radiolabeled ligand, whereas unbound ligands undergo radial diffusion. DRaCALA allows rapid detection of both the total ligand and the ligand sequestered by proteins. The fraction of ligand bound to the protein, defined as \( F_b \), is calculated from the signal intensity of the area with protein (inner circle) and the total signal intensity of the area (outer circle)\(^\text{12}\). For the DRaCALA we used unsealed membrane vesicles prepared from *E. coli* cells overproducing BtsS (MV BtsS) and calculated an \( F_b \) value of \( >0.15 \) (Fig. 5A). Control membrane vesicles (MV, lacking overproduced BtsS) were used and the low value of \( F_b \) (0.05) reflected only minor, non-specific binding. Therefore this assay was judged to be suitable for membrane vesicles, and it clearly indicated binding of radiolabeled pyruvate to BtsS. We also tested \(^3\)H-serine binding to BtsS in membrane vesicles using the DRaCALA technique. However, we only observed unspecific binding (data not shown).

*Figure 3.* Determination of changes in extracellular concentrations of serine and pyruvate during growth of *E. coli*. *E. coli* MG1655/pBBR yjiY-lux was cultivated in LB medium, and growth (OD\(_{600}\)) and luminescence were monitored. At the times indicated, cells were harvested, and serine and pyruvate levels were quantified by hydrophilic interaction liquid chromatography. All experiments were performed in triplicate, and the error bars indicate the standard deviation of the means. The growth phases of *E. coli* are marked as following: lag phase (white), exponential growth (light grey) and stationary phase (dark grey).

*Figure 4.* Stimulus-dependent yjiY expression under nutrient-limiting conditions. *E. coli* MG1655 mutant \( \Delta yhjX \) harboring pBBR yjiY-lux was cultivated in 0.1x LB medium to establish low nutrient conditions. After 1 h (time point 0), the indicated concentration of pyruvate (A), or L-serine (B), or the equivalent volume of water was added. Experiments were performed at least three times (standard deviation <10%), and results of a representative experiment are shown.
In addition, we tested MVs harboring a membrane-integrated truncated variant of BtsS (MV BtsS-TM) lacking its soluble domains, and found that this truncated sensor also binds pyruvate ($F_B > 0.15$). Furthermore, we prepared sealed right-side-out vesicles (RSO BtsS), in which membrane proteins retain their native orientation.\(^{13}\) BtsS in these vesicles also showed pyruvate binding, and no significant change in binding was observed when these vesicles were fragmented by sonication and had a random orientation (sonicated RSO BtsS) ($F_B$ values of each 0.18) (Fig. 5A). These data reveal that pyruvate binds to the external side of the membrane-spanning domain of BtsS.

The specificity of pyruvate binding to BtsS was then addressed by using a competition assay (Fig. 5B), in which several competitors were added in excess to the reaction mixture. Only unlabeled (“cold”) pyruvate was able to prevent binding of radiolabeled pyruvate. L-serine, D-serine or glycine did not interfere with pyruvate binding to BtsS, suggesting that pyruvate binds specifically to BtsS.

Next we investigated the effects of varying the length and side-chain charge on ligand binding by BtsS. Using competition assays, we found that acetate, propionate and butyrate all failed to compete with pyruvate for binding, indicating the importance of the C-$\alpha$ keto group of pyruvate (Fig. 5C). The influence of polarity at the C-$\alpha$ position was then tested by addition of lactate and malate, which contain C-$\alpha$ hydroxyl groups. These molecules were able to decrease pyruvate binding by about 50% suggesting that a negative charge seems to be recognized. It should be noted that these compounds were tested in a 104-fold excess over pyruvate. Phosphoenolpyruvate, which harbors a phosphoryl group at the C-$\alpha$ position, also competed with pyruvate, reducing binding by approximately 50%. In contrast, a positive charge at the C-$\alpha$ position in the form of the amino group in alanine had no effect, reducing pyruvate binding by 1% (Fig. 5C). Notably, none of these compounds were able to induce yjiY expression in vivo, which emphasizes the specificity of BtsS for pyruvate. The dissociation constant ($K_D$) for pyruvate binding to the histidine kinase was found to be 58.6 ± 8.8 µM (Fig. 5D).

To our knowledge this is the first application of DRaCALA to measure the interaction between a ligand and a membrane-embedded protein. Moreover, these assays unambiguously demonstrated that pyruvate binds specifically and with high affinity to BtsS.

Figure 5. Analysis of the interaction of BtsS with selected ligands by DRaCALA. (A) A mixture of membrane vesicles (MV) or right-side-out vesicles (RSO) enriched with the corresponding proteins (indicated by graphical representations) and radiolabeled 14C pyruvate (5 µM) is dropped onto a nitrocellulose membrane, and ligand migration via capillary action is analyzed. (B) Competition assays. Binding of radiolabeled pyruvate (5 µM) to BtsS in MVs was analyzed in the presence of various unlabeled competitors (each 50 mM). NC, no competitor. (C) Relative efficiency of competition by various carboxylic acids. Binding of radiolabeled pyruvate (5 µM) to BtsS in MVs was analyzed in the presence of various carboxylic acids (each 50 mM). The efficiency of competition by cold pyruvate was set to 1.00, and the effect of the indicated compounds was calculated accordingly. (D) Determination of the dissociation constant ($K_D$) for pyruvate to BtsS using DRaCALA. For each reaction radiolabeled pyruvate was used at 5 µM. Normalized $F_B$ values [$F_B = (F_{B\NC} - F_{B\text{pyr}})/F_{B\NC}$; see Methods for details] were plotted as function of the pyruvate concentration. The best-fit line was determined by nonlinear regression using the equation $y = B_{\text{max}} * x/(K_D + x)$. 

\[ \text{competition efficiency} \]

\[ \text{no competition} \]

\[ \text{competition} \]

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\[ \text{competition} \]
BtsS/BtsR importance during urinary tract infection. To understand the potential implications of BtsS/BtsR for pathogenesis, we therefore turned our attention to uropathogenic *E. coli* (UPEC) that colonize a nutritionally demanding environment – the mammalian bladder. However, responses to nutritional stress are of the utmost importance for the survival of both commensal and pathogenic bacteria within a given host.

UPEC strains are the primary causative agent of urinary-tract infections (UTIs) worldwide, accounting for over 85% of all reported episodes\(^{14}\). In the bladder, bacteria adhere to the apical surface of the epithelium and are internalized, before replicating to form biofilm-like pods within host bladder cells\(^{15,16}\). Subsequent to this transient intracellular cascade, adverse outcomes such as chronic colonization can ensue as a result of an aberrant host immune response\(^{17}\). Previous studies have demonstrated that successful UTI requires aerobic respiration and amino acid utilization\(^{18–20}\). To investigate the significance of BtsS/BtsR in bladder infection, we first created a Δ*btsS/btsR* mutant in the UPEC cystitis isolate UTI89. Using the same reporter fusion, we demonstrated that this strain, unlike the wild-type parental UTI strain, failed to induce transient expression of *yjiY* (Fig. 6A). Both pyruvate and L-serine are present in human and murine urine, and levels are elevated in diabetic populations\(^{21–23}\). Given that BtsS/BtsR responds directly to extracellular pyruvate levels and indirectly to L-serine, we asked whether BtsS/BtsR is active in the bladder lumen during acute and chronic UTI.

Mice were infected with the cystitis isolate UTI89 and RNA samples were extracted from bladder homogenates at 6 h post infection (coinciding with the acute stage of UTI), and at 2 weeks post infection from mice that went on to develop chronic cystitis. At this stage, the majority of bacteria are found on the bladder epithelial surface in the form of an extracellular biofilm\(^7\). Subsequent TaqMan-based qPCR analysis compared the expression of *yjiY* in the corresponding cDNA samples to the expression of *yjiY* in the starting bacterial inoculum. Our results demonstrated robust expression of *yjiY* in both the acute and chronic stages of infection from at least three separate mice per time-point (Fig. 6B). Taken together, these data suggest that, in the murine UTI model BtsS/BtsR responds to serine/pyruvate fluctuations and could play a role in promoting the infection process.

Discussion

Although numerous studies continue to demonstrate the importance of histidine kinase-mediated signal transduction in bacterial physiology, the natural ligands have been identified for very few histidine kinases\(^{24–26}\). This study presents compelling evidence demonstrating that the histidine kinase BtsS is a high-affinity receptor for extracellular pyruvate. Induction of the BtsS/BtsR target gene *yjiY*, which codes for a “carbon starvation” CstA-like transporter, correlates with the depletion of serine in complex medium, explaining the observation that BtsS/BtsR senses low serine. At the same time, while serine is being depleted from LB medium, *E. coli* excretes a substantial amount of pyruvate from overflow metabolism into the medium. Shortly afterwards, the concentration of extracellular pyruvate returns to its basal level (Fig. 3 and ref. 11). In previous studies, we demonstrated activation of a different system, YpdA/YpdB in response to high levels of extracellular pyruvate\(^{13}\) and showed that induction of YpdA/YpdB enhances BtsS/BtsR activation\(^7\). The current work provides a comprehensive characterization of how BtsS/BtsR directly responds to extracellular pyruvate levels and fine-tunes the metabolic fitness of *E. coli* under low-nutrient conditions. Sequence comparison between YpdA and YehU did not identify putative amino acids involved in pyruvate binding. In non-pathogenic *E. coli*, each of these signaling systems induces the expression of exactly one gene, which codes for a transporter. One (YhjX) belongs to the major facilitator...
superfamily, and is assumed to be a low-affinity carboxylate antiporter. The other one (YjiY) belongs to the CstA-like transporters, which are characterized by an unusually large number of transmembrane helices (17 in the case of YjiY). Neither transporter has been characterized thus far. It is hypothesized that both are involved in nutrient uptake, but with different affinities. The interconnectivity between the high-affinity pyruvate signaling system BtsS/BtsR with the putative low-affinity pyruvate signaling system YpdA/YpdR by a positive and a negative feedback loop would provide E. coli with a network that could tailor pyruvate uptake in each individual cell according to its availability. Switching between low- and high-affinity phosphate transporters based on the needs of the individual cell was recently demonstrated for S. cerevisiae, and seems to be a widely distributed strategy for the maintenance of nutrient homeostasis as stocks of essential nutrients decline.

Although nutrient sensing is crucial for host-microbe and microbe-microbe interactions, the majority of studies in non-pathogenic E. coli strains have focused on metabolic engineering, aiming to understand how processes such as elevated intracellular pyruvate levels affect metabolite distribution or how central mutations trigger/alter metabolic fluxes. In recent years, technological advances have permitted detailed analyses of in vivo metabolism, revealing its complexity and its influence on virulence and pathogenesis. These studies have shown pyruvate to be one of the major factors connecting cellular metabolism to cell division. Moreover, pyruvate levels are thought to reflect the quantitative relationship between carbon and nitrogen availability in the cell, and affect amino acid biosynthesis.

Investigations of how metabolic decisions determine pathogen fitness within host niches are increasingly uncovering opportunities for the development of robust and pathogen-specific drugs. Different E. coli pathotypes cause various clinical syndromes, depending on their genetic makeup and expression patterns. Different E. coli pathotypes cause various clinical syndromes, depending on their genetic makeup and expression patterns. Different E. coli pathotypes cause various clinical syndromes, depending on their genetic makeup and expression patterns. These studies have shown pyruvate to be one of the major factors connecting cellular metabolism to cell division.

In summary, this study has uncovered a signal transduction network that responds with exquisite sensitivity to extracellular pyruvate levels and fine-tunes carbon utilization in E. coli strains. Furthermore, we provide direct evidence for receptor-ligand interactions between BtsS and pyruvate, and demonstrate that BtsS/BtsR mediates yjiY expression during UTI. Future studies will focus on understanding the contribution of this system to pathogenic E. coli intra-host fitness and dissecting molecular determinants that drive the fine-tuning of bacterial fitness in response to external metabolic cues.

**Methods**

**Strains, plasmids, and oligonucleotides.** In this study we used the E. coli strains MG1655, E. coli MG1655 ΔyjiY, E. coli MG1655 ΔyjiY ΔyehU, BL21(DE3), E. coli KX1468 and the cystitis isolate UTI89. Plasmids used in this study include the transcriptional promoter-luciferase fusion construct for P_yehU activity (pBBR yjiY-lux) and for protein purification we used the arabinose-inducible expression vectors pBAD24, pBAD24-yehU and pBAD24-yehU-TM (YehU(BtsS) amino acids 1–205). DNA fragments for plasmid construction were amplified from genomic DNA by PCR. Plasmid and genomic DNAs were isolated using a HiYield plasmid mini kit (Suedlaborbedarf) and a DNeasy blood and tissue kit (Qiagen), respectively. DNA fragments were purified from agarose gels using a HiYield PCR cleanup and gel extraction kit (Suedlaborbedarf). Q5 DNA polymerase (New England BioLabs) was used according to the supplier’s instructions. Restriction enzymes and other DNA-modifying enzymes were also purchased from New England BioLabs and used according to the manufacturer’s directions.

**Growth conditions.** All strains were grown overnight in LB medium. After inoculation, bacteria were grown without shaking at 37 °C for 24 h. This culture was incubated for another 24 h to maximize expression of type 1 pili, as previously described. The growth phases of E. coli marked in graphs were according to the definition in ref. 48.

**In vivo expression studies.** In vivo expression of yjiY was quantified by means of luciferase-based reporter-gene assays, using bacteria that had been transformed with plasmid pBBR yjiY-lux. Cells from an overnight culture were transferred to fresh medium to give a starting optical density of 600 nm (OD<sub>600</sub>) of 0.05, and
incubated under aerobic growth conditions at 37 °C while OD₆₀₀ and luminescence were continuously monitored. The optical density was determined in a microplate reader (Tecan Sunrise) at 600 nm. Luminescence levels were determined in a luminescence reader (Centro LB960; Berthold Technology) for 0.1 s and are reported as relative light units (RLU; counts s⁻¹).

**Identification of quorum sensing-like molecule.** The cells of *E. coli* MG1655 ΔyjiY/pBBR yjiY-lux were cultivated in M9-minimal medium supplemented with 20 mM gluconic acid and OD₆₀₀ and luminescence were constantly monitored. Cells were harvested shortly before, at and after the induction of yjiY. The supernatant was sterile filtrated (Stericup®, Filter Unit Millipore Express® PLUS(PES), 0.22 µm) and separated by high pressure liquid chromatography (HPLC) using column C18-Hypersil Gold, with the gradient 1–100% (v/v) acetonitrile in 40 min. Fractions were collected, concentrated and used in reporter strain *E. coli* KX146845/pBBR yjiY-lux grown in M9-minimal medium supplemented with 20 mM succinate. The fraction with the highest induction potential was then analyzed via UPLC-UHR-ToF-MS using a Waters XBridge Amide column.

**Production of BtsS-6His in membrane vesicles.** *E. coli* BL21(DE3) cells were transformed with the vector pBAD24-yejU⁴⁰, which codes for BtsS-6His, and grown in LB medium at 37 °C to an OD₆₀₀ of 0.5. Recombinant gene expression was induced by addition of 0.2% (w/v) arabinose. After 3 h of further incubation, cells were harvested by centrifugation, disrupted and fractionated. At each step, the pellet was resuspended in buffer consisting of MES/NaOH (pH 6), 10% (v/v) glycerol and 10 mM MgCl₂. To produce BtsS-6His in right-side-out vesicles, formation of spheroplasts and right-side-out vesicles was monitored under the microscope. His-tagged BtsS was detected by Western blot analysis with an anti-His antibody (Qiagen) and an alkaline phosphatase-conjugated anti-mouse antibody as the secondary antibody.

**Protein-ligand interactions.** Protein-ligand interactions were analyzed via DRaCALA, a method established by Roelofs et al.⁵². Membrane vesicles enriched for overproduced BtsS-6His were mixed with 5µM radiolaabeled (3-¹⁴C) pyruvic acid sodium salt (55 mCi mmol⁻¹, Biorend) or radiolaabeled (¹⁴) L-serine (11.0 Ci mmol⁻¹, Hartmann Analytics) and incubated for 5 min at room temperature. Triplicate 5-µl aliquots were transferred to a nitrocellulose membrane, dried and visualized by a PhosphorImager. Quantification of radioactive signal was done with the software ImageQuant. The fraction bound to protein was calculated according to the signal intensities using an equation defined earlier: $F_B = \frac{I_{\text{sample}} - I_{\text{background}}}{I_{\text{sample}}}$. For the evaluation the corresponding cold ligand (50 mM) was added to the reaction mixture and incubated for further 3 min. To estimate the $K_D$ value, increasing concentrations of cold pyruvate (0 µM to 2.5 mM) were used. In this case the $F_B$ value was normalized as follows: $F_B = \frac{F_{\text{RINC}} - F_{\text{BRIPPY}}}{F_{\text{RINC}}}$. Here, $F_{\text{RINC}}$ is calculated from the sample with pure ¹⁴C-labeled pyruvate, and $F_{\text{BRIPPY}}$ are all values for mixtures containing ¹⁴C-labeled pyruvate together with increasing concentrations of unlabeled pyruvate. The amount of ¹⁴C-pyruvate was kept constant in all assays.

**Extraction and determination of extra- and intracellular metabolites.** The reporter strain *E. coli* MG1655/pBBR yjiY-lux was cultivated in LB medium, and OD₆₀₀ and luminescence were constantly monitored. At selected time points, cells were harvested and the supernatants were saved. Cells were washed with 10% (v/v) glycerol, and subsequently cell pellets and supernatants were analyzed via hydrophilic interaction liquid chromatography (HILIC). Acetonitrile (ACN), methanol (MeOH), ammonium acetate and ammonium hydroxide (all of LC-MS grade) were obtained from Sigma-Aldrich (Sigma-Aldrich GmbH). Water was purified using a Merck Millipore Integral water purification system to a resistance of 18 MΩ and TOC < 5 ppb. Sodium pyruvate and serine were also obtained from Sigma-Aldrich and dissolved in water at a concentration of 100 mM and further diluted with ACN. Cell pellets were extracted with 200 µl ice-cold water/MeOH (50/50, v/v). Samples were vortexed vigorously and lysed in an ice-cold sonic bath for 15 min. After centrifugation at 20,000 × g and 4 °C for 10 min, supernatants were transferred to autosampler vials.

Pyruvate and serine were quantified using a modified version of the method published by Yuan et al.⁴⁹. Separation was achieved using a Waters XBridge Amide column (3.5 µm, 100 mm × 4.6 mm ID) and an ACN/water gradient using a Waters Acquity UPLC system coupled to a Bruker maXis UHR-ToF-MS (Bruker Daltonics). Eluent A consisted of 5% (v/v) ACN, 95% (v/v) water, 20 mM ammonium hydroxide, 20 mM ammonium acetate, pH 9.0 and eluent B was pure ACN.

Metabolites were eluted with the following gradient: After 3 min of 85% B, a linear decrease to 2% B over 9 min, isocratic hold for 3 min and return to initial conditions for 1 min and a 7-min re-equilibration time. Sample (5µl) was injected via partial loop injection.

For quantification a calibration curve was generated from different concentrations of standard compounds (0 µM, 0.5 µM, 1.0 µM, 2.5 µM, 5 µM, 10 µM, 25 µM, 50 µM, 100 µM, 250 µM). If the concentration of a sample was beyond the range of the calibration curve, it was appropriately diluted with water/MeOH. Quantification was performed with Bruker Quant Analysis.

**Murine infections.** A cohort (15) of 7- to 8-week old female C3H/HeN mice was infected with strain UTI189 via transurethral catheterization as described in Hung et al.⁵⁰. Five mice were sacrificed at 6 h after infection, marking the acute stage of UTI in this murine model. Bladders were excised for CFU enumeration and RNA extraction. The remaining 10 mice were monitored for chronic infection using longitudinal urinalysis, as previously described.⁵⁰ Mice that consistently shed more than 100,000 CFUs/ml of urine were separated from the
rest of the cohort, and flagged as chronically infected. These mice were sacrificed at 4 weeks post infection and bladders were obtained for CFU enumeration and RNA extraction.

2. To ensure the proper and humane treatment of animals, all animal studies were carried out in strict accordance with the guidelines of the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health, and the Vanderbilt University Medical Center’s Institutional Animal Care and Use Committee (IACUC), who approved all protocols. Statistical analyses were performed using two-tailed Mann-Whitney t-test.

RNA extraction, cDNA synthesis and qPCR. RNA extraction and bacterial RNA enrichment were performed as described by Conover et al. 47. DNase treatment, reverse transcription, and real-time quantitative PCR were done as described by Guckes et al. 53. qPCR analysis was carried out with three concentrations of cDNA (50 ng, 25 ng, 12.5 ng), each in triplicate for each sample, and levels of gyrB (DNA gyrase) were used for normalization. The following primers (Integrated DNA Technologies) were used for amplification: yjiY_FWD (5'-GGCAGACGCGCAAACCT-3'), yjiY_REVP (5'-GCCGTAGCCGATGAACG-3'), gyrB_FWD (5'-GATGGCGCTGAAGGCTTAGT-3'), gyrB_REV (5'-CACGGGCAGCCGACGATC-3'). The following probes (Applied Biosystems) were used for quantification: yjiY (5'FAM-TGGCTATGAAACGAGC-G-MGBNFQ-3'); gyrB (5'VIC-ACGAACTGTG-GCGGA-MGBNFQ-3').

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Author Contributions
S.B., M.W., P.S.-K., M.H. and K.J. designed the experiments. S.B., I.K., M.W., E.B., A.E., C.S. performed the experiments. S.B., I.K., M.H., and K.J. wrote the manuscript.

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