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Insight, innovation, integration

When bacteria encounter certain environmental stresses, the EnvZ/OmpR two-component system alters gene transcription patterns which enables adaptation. We constructed chromosomally-encoded, active, fluorescent fusion proteins and used super-resolution microscopy to visualize the EnvZ/OmpR two-component regulatory system for the first time. Technical innovation enabled us to serially image the chromosome, OmpR and the membrane and then overlay these in a combined image. During environmental stresses such as low osmolality and acid pH, OmpR was juxtaposed next to the membrane and the bacterial chromosome was condensed. In axial cross-sectional images, a linkage stretching from the DNA to the membrane was observed, suggesting a mechanism for mechanical coupling.

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Single cell super-resolution imaging of *E. coli* OmpR during environmental stress[†]

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Two-component signaling systems are a major strategy employed by bacteria, and to some extent, yeast and plants, to respond to environmental stress. The EnvZ/OmpR system in E. coli responds to osmotic and acid stress and is responsible for regulating the protein composition of the outer membrane. EnvZ is a histidine kinase located in the inner membrane. Upon activation, it is autophosphorylated by ATP and subsequently, it activates OmpR. Phosphorylated OmpR binds with high affinity to the regulatory regions of the ompF and ompC porin genes to regulate their transcription. We set out to visualize these two-components in single bacterial cells during different environmental stress conditions and to examine the subsequent modifications to the bacterial nucleoid as a result. We created a chromosomally-encoded, active, fluorescent OmpR-PAmCherry fusion protein and compared its expression levels with RNA polymerase. Quantitative western blotting had indicated that these two proteins were expressed at similar levels. From our images, it is evident that OmpR is significantly less abundant compared to RNA polymerase. In cross-sectional axial images, we observed OmpR molecules closely juxtaposed near the inner membrane during acidic and hyposomotic growth. In acidic conditions, the chromosome was compacted. Surprisingly, under acidic conditions, we also observed evidence of a spatial correlation between the DNA and the inner membrane, suggesting a mechanical link through an active DNA-OmpR-EnvZ complex. This work represents the first direct visualization of a response regulator with respect to the bacterial chromosome.

Introduction

All cells need to sense and respond to their environment in order to survive and bacteria use two-component signaling systems to perform this function. Environmental changes are sensed by histidine kinases (HKs) that in gram-negative bacteria are most often embedded in the inner membrane. Upon activation, HKs are autophosphorylated by intracellular ATP. The phosphoryl group is then transferred to an aspartic acid residue of the second component, the response regulator (RR). Most RRs are DNA binding proteins and phosphorylation enhances or represses transcription.

In the archetype EnvZ/OmpR system, bacteria respond to increasing extracellular osmolality by concentrating their cytoplasm.

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This intracellular signal drives a disordered region around the active site histidine of EnvZ to become ordered, activating phosphorylation.^{1,2} Phospho-OmpR binds to the regulatory regions of the porin genes, activating *ompC* transcription at high osmolality and repressing *ompF*. The two porins are reciprocally regulated³ and OmpF has a larger pore and a faster flow rate to allow for enhanced nutrient exchange in dilute environments.⁴ In *Salmonella enterica*, the EnvZ/OmpR system is essential to activate a second two-component system, SsrA/B, which drives expression of virulence factors that enable *Salmonella* to replicate in the acidic environment of the macrophage vacuole.^{5,6}

In the EnvZ/OmpR system, we recently established that signals that were once believed to be extracellular, were actually intracellular.^{1,2,7} For example, at high external osmolality, the cytoplasm of *E. coli* becomes concentrated, up to 1.8 osmoles,⁸ altering EnvZ conformation.¹ Likewise, when Salmonella is in a macrophage vacuole, its cytoplasm acidifies to pH 5.6.⁷ This acidification is dependent on the cytoplasmic domain of EnvZ (EnvZc), which contains both kinase and ATP-binding subdomains. Acidification works through EnvZ-OmpR, as OmpR represses the cadaverine transport system cadC/BA to prevent neutralization of the cytosol after acid stress. Furthermore, when E. coli and Salmonella are exposed to external osmolytes, they respond by acidifying the cytoplasm.9 EnvZ responds to these intracellular signals by increasing its autophosphorylation and activating OmpR via phosphotransfer.¹ This may be in part a response to mechanical cues that arise as E. coli undergoes substantial volume changes during large alterations in osmolality. Recent studies used ChIP-seq to identify OmpR binding sites in response to external acid stress.¹⁰ Those studies employed a FLAG-tag fused to the C-terminus of OmpR. However, previous studies have shown that alterations to the OmpR C-terminal DNA binding region are deleterious.¹¹ Our goal was to establish a photoactivatable construct of OmpR that was chromosomally-expressed under its native promoter and capable of porin gene regulation so that we could examine the OmpR distribution in single cells in response to various environmental stresses.

To investigate the EnvZ/OmpR system at the single-cell level, a fluorescent label can be attached to the protein of interest, which facilitates the microscopic study of its intracellular localization. Due to the small size of bacterial cells and the diffraction limit of light, conventional fluorescence microscopy is of limited use. Super-resolution microscopy allows visualization of biological structures at scales far below the diffraction limit of light.¹² We applied single-molecule localization microscopy (SMLM)¹³ to visualize protein and chromosome structures at the nanometer scale. In SMLM, the fluorescence signal is separated in time using photoactivatable or photoswitchable fluorescent probes. By activating only a subset of fluorophores, single molecules can be detected and their position determined with a precision of a few nanometers.¹⁴ Next to imaging cells with subdiffraction resolution, photoactivated localization microscopy (PALM)¹⁵ can provide copy numbers of the target protein in single cells, an approach that was recently demonstrated for RNA polymerase in bacteria.¹⁶ Point accumulation for imaging in nanoscale topography (PAINT)^{17,18} and direct stochastic optical reconstruction

microscopy (dSTORM)¹⁹ can be used to visualize the bacterial envelope or chromosomal DNA, respectively. Combining these techniques sequentially enabled us to investigate the localization of OmpR under different stress conditions, to determine its relative position within the cell cylinder, and its spatial relationship with the chromosome. In this study, we labeled OmpR with the photomodulatable proteins mEos²⁰ and PAmCherry,²¹ while conserving its functionality.

In the case of two-component signaling systems, the HK is often not an abundant protein and it is usually embedded in the inner membrane, making it difficult to study. In the case of envZ, it is in an operon with ompR (the ompB operon), with the promoter upstream of ompR.²² The stop signal for ompR overlaps the start signal for envZ, resulting in substantially fewer copies of EnvZ compared to OmpR. This provides a significant imaging challenge, as the low copy prevents the generation of a suitable super-resolution image. We overcame this difficulty by over-expression of EnvZc-mEos2 using an arabinose inducibleplasmid and induction with 0.02% arabinose.² This produced a level of EnvZ that was as functional as the wildtype and the protein was widely distributed throughout the cytoplasm. However, at copy numbers approaching in vivo levels, not enough molecules could be collected to produce a superresolution image. To overcome these obstacles and examine the interaction of EnvZc with OmpR as a function of environmental stress, we turned to fluorescence correlation spectroscopy (FCS) to examine the EnvZc/OmpR interactions in vitro, in solution.

Imaging the distribution of OmpR within single cells using PALM indicated that OmpR was largely localized near the plasma membrane during growth in acidic conditions and in low osmolality medium. We also provide evidence of chromosomal compaction during acid stress. Our results suggest an *in vivo* linkage with EnvZ in the inner membrane, to OmpR to the nucleoid²³ and explains why some dominant mutations in *ompR* require *envZ* for expression of their phenotype.²⁴ Furthermore, the visible linkages observed stretching from the nucleoid to the inner membrane provide evidence of structural interactions between the chromosome and the membrane and suggest mechanical stress could be easily transmitted from the membrane to the DNA.

Methods

Construction of an active, chromosomally-expressed OmpR-photoactivatable fusion protein

In constructing an OmpR fluorescent fusion protein, we placed a flexible linker of GGSG repeats (encoded by 5'-ggtggttctggt-3') in between *ompR* and the fluorophore. Plasmid pCAH63-PompRompR-GGSGx4-PAmCherry was constructed in two steps. First, a fragment with the *ompR* promoter and the *ompR* gene (PompRompR) was amplified from *E. coli* genomic DNA with primers PompR-EcoR1-F and FP-GGSGx4-ompR-R, and PAmCherry was amplified with primers *ompR*-GGSGx4-PAmC-F and PAmC-BamH1-R. These two fragments were equally mixed and used

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as a template to amplify PompR-ompR-GGSGx4-PAmCherry, which was flanked with EcoR1 and BamH1 restriction sites. The resulting fragment PompR-ompR-GGSGx4-PAmCherry was digested with EcoR1 and BamH1, and then inserted into pCAH63 that had been treated with the same restriction enzymes, generating pCAH63-PompR-ompR-GGSGx4-PAmCherry. The mEos2 plasmid (pCAH63-PompR-ompR-GGSGx4-mEos2) was created using a similar method except that mEos2 was amplified with primers ompR-GGSGx4-mEos2-F and mEos2-BamH1-R. PompRompR and mEos2 was fused together with primers PompR-EcoR1-F and mEos2-BamH1-R, yielding a PompR-ompR-GGSGx4-mEos2 fragment. PompR-ompR-GGSGx4-mEos2 and pCAH63 were digested with EcoR1 and BamH1, and ligated to generate pCAH63-PompRompR-GGSGx4-mEos2. Plasmid pCAH63-PompR-ompR-GGSGx10mEos2 was randomly generated during the construction of pCAH63-PompR-ompR-GGSGx4-mEos2.

Genome integration was conducted following the method of Haldimann and Wanner.²⁵ Briefly, strains MH225.101 and MH513.101 that were ompR null²⁶ were transformed with plasmid pINT-ts by electroporation. After that, the recombinant strains MH225.101/pINT-ts and MH513.101/pINT-ts were made competent and transformed with pCAH63-PompR-ompR-GGSGx4-PAmCherry, pCAH63-PompR-ompR-GGSGx4-mEos2 and pCAH63-PompR-ompR-GGSGx10-mEos2 plasmids, and colonies were then recovered on plates using chloramphenicol selection. A single colony was re-streaked and the correct integration was confirmed by colony PCR with primers P1, P2, P3 and P4. The pINT-ts plasmid was eliminated by incubation at the restrictive temperature, as it possesses a temperaturesensitive origin of replication. Finally, we obtained the following strains MH225.101 attB:PompR-ompR-GGSGx4-PAmCherry, MH225.101 attB:PompR-ompR-GGSGx4-mEos2, MH225.101 attB:PompR-ompR-GGSGx10-mEos2, MH513.101 attB:PompR-ompR-GGSGx4-PAmCherry, MH513.101 attB:PompR-ompR-GGSGx4mEos2 and MH513.101 attB:PompR-ompR-GGSGx10-mEos2.

Primer	Sequence $(5'-3')$
PompR-EcoR1-F	ccggaattcggtttgagttcatcttttcgg
FP-GGSGx4-ompR-R	accagaaccaccaccagaaccaccacc
	agaaccaccaccagaaccacctgcttta
	gagccgtccggtac
ompR-GGSGx4-mEos-F	ggtggttctggtggtggttctggtggtggttc
	tggtatgtcagcaatcaaaccggatatg
ompR-GGSGx4-PAmC-F	ggtggttctggtggtggttctggtggtggttc
	tggtatggtgagcaagggcgaggaggataac
PAmC-BamH1-R	cgcggatccctacttgtacagctcgtccatgccgcc
mEos2-BamH1-R	cgcggatccttagcgacgggcgttatccggcag
P1	GGCATCACGGCAATATAC
P2	ACTTAACGGCTGACATGG
P3	ACGAGTATCGAGATGGCA
P4	TCTGGTCTGGTAGCAATG

β-galactosidase assays

β-Galactosidase assays were performed as previously described.²⁷

Protein purification and fluorescent-labeling

EnvZc and EnvZc^M were sub-cloned into the pET28b vector with an N-terminal His-tag, over-expressed and purified as described previously.¹ OmpR was over-expressed on a pET15b vector and purified using a similar method as EnvZc. The proteins were stored in 20 mM phosphate (pH 7.4), 200 mM NaCl (500 mM for OmpR), 5% glycerol (v/v) and 0.05% Tween-20. EnvZc purified as a dimer based on the elution time on the FPLC. Alexa dyes were added in five-fold molar excess for labeling of EnvZc, EnvZc^M and OmpR. The reaction was allowed to proceed at 4 °C overnight with gentle rocking. The labeled protein was separated from the excess dye using two PD-10 desalting columns in tandem (GE Healthcare).

Native PAGE

EnvZc, EnvZc^M and OmpR were incubated at final concentrations of 7 μ M in a 10 μ l volume reaction in 50 mM Tris-HCl, pH 8.0, 50 mM KCl and 5% glycerol buffer. The samples were loaded onto a native gel with 2× loading buffer (40 mM Tris-HCl, pH 7.5, 80 mM β -mercaptoethanol, 0.08% bromophenol blue, 8% glycerol). The composition of the stacking gel was 5% acrylamide/bis (37.5:1) in 63 mM Tris-HCl (pH 7.4) and the composition of the separation gel was 10% acrylamide/bis (37.5:1) in 188 mM Tris-HCl (pH 8.8), while the running buffer was 83 mM Tris-HCl (pH 8.9), 33 mM glycine. The gel was run at a constant current of 0.02A at 4 °C. The native gel was then imaged using a UV imager (BioRad, ChemiDoc MP) before staining with Coomassie based InstantBlueTM (Expedeon).

Fluorescence correlation spectroscopy (FCS) and fluorescence cross-correlation spectroscopy (FCCS) measurements

FCS and FCCS were performed on the LSM710-ConfoCor3 (Carl Zeiss) system with a 40×, NA 1.2 C-apochromat objective. Alexa Fluor 488 and 568 were excited using the 488 nm and 561 nm laser line, respectively. The protein samples were incubated in 20 mM phosphate (pH 7.4), 50 mM NaCl, and 0.05% Tween-20 at room temperature for 30 min before spotting (20 μ l) onto a clean coverslip for measurements. The autocorrelations functions (ACFs) from the EnvZc^M titrations FCS were fitted with either a single component (for the lower EnvZc^M concentrations) or two-component 3D diffusion model (for the higher EnvZc^M concentrations where there are two distinct diffusion times):²⁸

$$G(\tau) = G(0) \left[1 + \frac{F_{\text{trip}}}{1 - F_{\text{trip}}} \exp\left(-\frac{\tau}{\tau_{\text{trip}}}\right) \right] D(\tau) + G(\infty)$$

where $G(\infty)$ is a convergence value for the ACF at long (infinite) delay times. F_{trip} is the fraction of the particles that reside in the triplet state and τ_{trip} is the triplet state relaxation time. G(0) is the amplitude of the ACF and is related to the average number of molecules *N* detected in the observation volume by:

$$G(0) = \frac{1}{N}$$

$$D_{3\mathrm{D}}(\tau) = \left(1 + \frac{\tau}{\tau_{\mathrm{D}}}\right)^{-1} \left(1 + \frac{\tau}{K^2 \tau_{\mathrm{D}}}\right)^{-1/2}$$

and $D(\tau)$ for a two-component 3D diffusion model is given by:

$$\begin{split} D_{\rm 3D,2C}(\tau) &= F_{\rm bound} \\ & \cdot \left(1 + \frac{\tau}{\tau_{\rm D(bound)}}\right)^{-1} \left(1 + \frac{\tau}{K^2 \tau_{\rm D(bound)}}\right)^{-1/2} + (1 - F_{\rm bound}) \\ & \cdot \left(1 + \frac{\tau}{\tau_{\rm D(free)}}\right)^{-1} \left(1 + \frac{\tau}{K^2 \tau_{\rm D(free)}}\right)^{-1/2} \end{split}$$

where F_{bound} is the fraction of bound molecules with a higher diffusion time of $\tau_{\text{D(bound)}}$. $\tau_{\text{D(free)}}$ is the diffusion time for the faster free molecule. *K* is the structural parameter defining a confocal laser spot obtained from calibrating the microscope with either Alexa 488 or 568 dyes.

In the FCCS measurements, as the concentration of EnvZc labeled with Alexa 568 is always in excess, cross-talk of Alexa 488 into the Alexa 568 detection channel can be neglected. Therefore, the fraction of Alexa 488-labeled OmpR (OmpR-A488) bound can be simply obtained from the ratio of the amplitude of the cross-correlation function (CCF) to the amplitude of the ACF of Alexa 568 labeled protein (cross-correlation ratio):

$$\frac{G_{\rm CCF}(0)}{G_{\rm ACF}(0)} = \frac{N_{\rm complex}}{N_{\rm Tot,OmpR}}$$

where N_{complex} is the number of molecules in complex and $N_{\text{Tot,OmpR}}$ is the total number of molecules of OmpR-A488 (bound and free). The microscope was calibrated using known controls to obtain the range of cross-correlation ratio values obtainable (ESI,† Fig. S1A and B). The range of the cross-correlation ratio for the microscope system was determined to be between 0.02 and 0.99.

Super-resolution microscopy

Cell culture. The OmpR–PAmCherry strain was inoculated from glycerol stocks and grown over night (ON) in the respective media, supplemented with 10 μ M chloramphenicol, at 32 °C or 37 °C and 200 rpm shaking. Working cultures were inoculated 1:200 from ON cultures and OD₆₀₀ was determined every 30 minutes to obtain the doubling time for each culture. For media composition and respective doubling times see ESI,† Table S1. EdU (Baseclick) was added at 10 μ M concentration for 40 minutes to the LB culture at an OD₆₀₀ of ~0.25. Due to overlapping replication cycles, EdU-incorporation should be incorporated throughout the entire chromosome. The massdoubling times were significantly longer in the minimal media, therefore EdU was added for 1.5 mass-doubling times.

Fixation and immobilization. Cultures were fixed in solution by adding sodium-phosphate-buffer pH 7.5 and methanol-free formaldehyde (Sigma) to a final concentration of 15 mM and 1%, respectively. After 30 minutes, the cells were pelleted for 3 minutes at 5000 \times *g* and washed once with 100 mM sodiumphosphate-buffer pH 7.5 containing 50 mM ammonium-chloride to reduce excess formaldehyde. Resuspended bacteria were then immobilized on KOH-cleaned (3 M, 30 min), poly-L-lysinecoated 8-well-chamberslides (Sarstedt) and washed twice with phosphate buffer. The buffer was then removed and the sample was placed on ice 5 min before proceeding with gel matrix coating.

Coating and post-labeling. To maintain the positions of immobilized bacteria and prohibit detachment from the poly-L-lysine surface, we added a hydrophilic gel matrix to the sample as described previously, but with shorter gel polymerization time (15 min).²⁹ During the sequential workflow, we also followed the provided protocol for DNA post-labeling. In brief, immobilized bacteria were permeabilized with 0.5% Triton-X 100 in PBS (the osmolality was adjusted for each growth condition) for 60 min. After two washing steps, 300 µl click-reaction buffer (100 mM Tris pH 8.0, 1 mM CuSO₄ 10 µM Alexa Fluor 647 azide, 100 mM ascorbic acid, osmolalityadjustment with NaCl if required) was added to the sample for 60 minutes. Afterwards, the sample was washed 3 times for 5 min and further 2 times for 30 min with PBS (osmolality adjusted). The sample was then transferred onto the microscope for dSTORM imaging.

Super-resolution imaging. Samples were imaged on a custombuilt microscope.²⁹ A mechanical stage (TI-S-ER, Nikon) and an adjustable TIRF mirror were used to relocate bacteria during the sequential imaging workflow and adjust the angle of the inclined laser for proper HILO imaging. µManager³⁰ was used to control the microscope and camera. PALM imaging of RNAP-PAmCherry in fixed cells was performed using 568 nm for excitation (1.5 kW cm^{-2}) and 404 nm for activation (gradually increased up to 12 W cm⁻²). Typically, between 5000 and 7000 frames were recorded at a camera frame-rate of 10 Hz in HILO mode until all PAmCherry-molecules were photoconverted and bleached. Afterwards, 200 pM Nile red in phosphate buffer was added to facilitate PAINT-imaging (10 000 frames at 33 Hz camera frame-rate) using 568 nm excitation light (0.7 kW cm⁻²). dSTORM-imaging of Alexa Fluor 647 was performed in PBS containing 100 mM cysteamin (MEA, Sigma) pH 8.5 using excitation at 647 nm (2 kW cm⁻²) and 488 nm for reactivation (up to 30 W cm⁻²). Appropriate dichroic mirrors and filters (AHF, Germany) were used for each spectral channel. The data was analyzed with rapidSTORM³¹ and resulting images were post-processed with Fiji.32

Image processing. For visualization, a Gaussian blur of the mean localization precision was applied to the SMLM images using Fiji. For the averaging analysis of OmpR distribution, cells were angular reorientated and normalized for cell length and width using custom-written ImageJ macros. In order to prevent orientation artifacts, each bacteria was mirrored along the length axis, the cross axis and both axes with subsequent averaging of the resulting four images. 13–20 images per condition were averaged to obtain 'heat maps' of OmpR localization. Cells of similar size were chosen in order to average bacteria in comparable cell cycle stages and to prevent normalization artifacts (LB: $3.75-4.25 \mu$ m, Tris: 2.0–2.5 μ m, MES: $1.75-2.25 \mu$ m, $0.5 \times$ M9: 1.5 to 2.0 μ m).

Results

A chromosomally-expressed fluorescent fusion protein of OmpR is active

Initially, we constructed photoactivatable fusion proteins based on a previously published strategy³³ that consisted of a four amino acid linker between the end of the ompR gene and the gene encoding the fluorophore. That study did not examine the functionality of the fusions that were constructed, and more recently, it was reported that the fluorescent fusion protein had significant effects on its distribution.³⁴ To avoid this problem, we used a functional assay and examined the ability of our OmpR construct to activate its target genes, using ompF and ompC transcriptional fusions.^{26,35} These fusions contain the ompF/C promoter fused to the *lacZ* gene, encoding β -galactosidase. The original fusion was \sim 20% active (data not shown), similar to other OmpR chimeric fluorescent protein fusions³⁶ that we have also measured. OmpR does not tolerate substitutions at its C-terminus, which disrupt its ability to bind to DNA.¹¹ Increasing the linker length in four amino acid increments eventually led to an active chimera at sixteen amino acids (Fig. 1A and B, see also ref. 37). A linker length of 16 amino acids was sufficient for PAmCherry, but a longer linker length was required for mEos (Fig. 1A and B), and it was not as active as the PAmCherry chimera (79% compared to 93% at ompF and 60% compared to 71% at ompC). As we have also previously observed, ompC transcription was more sensitive than ompF to substitutions in OmpR that alter function.³⁸ Our OmpR-PAmCherry was expressed from its native promoter and was integrated at the attB site of the E. coli chromosome, to avoid disrupting its downstream gene *envZ* (see Methods).

Correlating OmpR distribution with the bacterial nucleoid

We were next interested in visualizing the distribution of OmpR in single bacterial cells using super-resolution microscopy. In order to correlate the distribution of OmpR near the membrane or the bacterial chromosome, we applied a sequential workflow²⁹ (Fig. 2). In medium at high osmolality (LB, 460 mOsm per kg), OmpR molecules were located close to the edge of the nucleoid, while the chromosomal core was not decorated by OmpR (Fig. 2A). This behavior was not readily apparent in low salt media (100 mM Tris pH 7.2, 210 mOsm per kg) and the nucleoid appeared to be more condensed (Fig. 2B, compare green circles outlining the chromosome in the third row panels). When cells were grown in acidic conditions (100 mM MES medium pH 5.6, 205 mOsm per kg) (Fig. 2C) or $0.5 \times$ M9 medium (115 mOsm per kg) (Fig. 2D), OmpR localized near the plasma membrane and appeared to be more clustered than in rich media at higher osmolality (LB, Fig. 2A). In order to quantify this distribution, we constructed average localization maps of OmpR, each consisting of 13-20 bacteria (bottom panel, Fig. 2). While OmpR shows a more uniform distribution in LB and Tris, the preferential localization near the membrane is evident for acidic and low osmolality $0.5 \times$ M9 medium. Furthermore, the bacterial nucleoids appeared to be more condensed during acidic growth compared to cells grown at neutral pH (Tris) (ESI,† Fig. S2).



Fig. 1 Comparison of OmpR-PAmCherry with OmpR-mEos and wildtype OmpR. MH513 and MH225 are bacterial strains containing a chromosomal ompF-lacZ (A) or ompC-lacZ (B) transcriptional fusion, respectively. Wildtype OmpR (column 1) represented 100% and the other backgrounds were normalized to it. In the ompR null strain (ompR101), there was no activation of ompF or ompC. The PAmCherry fusion was 93% as active as the wildtype, when a 16 amino acid linker, GGSGx4, was placed between the 3' end of ompR and the beginning of PAmCherry. Its activity was higher than the mEos2 fusion that contained the same linker length (striped column 4, 37%), but a longer linker (40 amino acids, GGSGx10) improved activity to 79% of the wildtype (column 5). Similarly, at ompC, the PAmCherry fusion was 71% of the wildtype activity and better than the mEos2 fusion (17% with a 16 amino acid linker and 60% with a 40 amino acid linker). The photoactivatable fusion affected ompC activity more than ompF as we have observed with other ompR mutants³⁸ as well as with envZ photoactivatable fusions.²

In preparing bacteria samples for microscopy, the cells were spatially fixed in a gel matrix. We found that some bacterial cells were oriented axially, which allowed cross-sectional imaging (Fig. 3). The membrane association and positioning around the chromosome became more obvious for cells visualized in this orientation, and the preferred localization of OmpR juxtaposed next to the cell membrane was clearly evident in both $0.5 \times$ M9 (Fig. 3A) and pH 5.6 (Fig. 3B). For cells grown in acidic conditions (MES, pH 5.6), we also observed a hollow-tube structure for DNA (Fig. 3B), from which thin DNA fibers appeared to reach towards the membrane (see arrows).

We next investigated qualitative levels of OmpR expression in single bacterial cells. We recorded PALM images of bacterial cells expressing RNAP-PAmCherry endogenously (Fig. 4A) to



Fig. 2 Sequential super-resolution imaging of OmpR–PAmCherry (red hot), bacterial cell membrane (green) and DNA (cyan) in various growth media. LB (A), 100 mM Tris pH 7.2 (B), 100 mM MES pH 5.6 (C) and $0.5 \times$ M9 buffer (D). Evidence for DNA compaction is noticeable in the third row panels, where the green outlines the nucleoid edges. OmpR was distributed around the nucleoid edges (outlined in green) in LB, whereas it was more uniformly distributed in Tris buffer. In acidic (MES, pH 5.6) and hypotonic ($0.5 \times$ M9) conditions, OmpR was recruited to the plasma membrane (scale bar 1 µm). Images in the bottom row show the averaged distribution of the bacterial membrane (green) and OmpR (red hot). These average images indicate a localization of OmpR near the membrane in acid and low osmolality. The recruitment of OmpR to DNA in LB is averaged out because of the nucleoid heterogeneity and asymmetric segregation.⁷⁴ Cells grown in Tris and $0.5 \times$ M9 medium exhibited a smaller cell diameter (~0.8 µm), resulting in projection effects for the membrane average image. The number of images used for averaging was: 19, cell length $3.75-4.25 \mu$ m (A), 20 cell length $2.0-2.5 \mu$ m (B), 15 cell length $1.75-2.25 \mu$ m (C) and 13 cell length $1.5-2.0 \mu$ m (D).

compare with OmpR–PAmCherry (Fig. 4B). Quantitative western blotting reported that OmpR was present at 3500 copies per cell at high osmolality and 2000 copies at low osmolality.³⁹ Copy numbers of RNAP were determined in a previous study using quantitative PALM.¹⁶ In rich media such as LB, an average of 3600 RNAP per cell were counted, *i.e.*, similar to the copy number reported for OmpR by quantitative western blots. Compared to the PALM images of OmpR–PAmCherry recorded using the same experimental settings (Fig. 4B), it is immediately apparent that OmpR expression levels were significantly lower than those of RNAP. These observations question the accuracy of quantitative western blots in determining protein copy number (see Discussion).

FCS and FCCS measurements of EnvZ/OmpR interaction

Because OmpR appears to be more closely associated with the membrane where its cognate HK EnvZ is located (Fig. 3A and B),

we attempted to determine whether OmpR interaction with EnvZ might be dependent on osmolality, pH or other activating signals. Unfortunately, the EnvZ copy number is too low to generate super-resolution images using photoactivatable fusions when *envZ* is expressed from its native chromosomal locus.² Thus, we turned to *in vitro* methods to examine their interaction in solution in response to signals that are known to activate porin gene expression.

Fluorescence correlation spectroscopy $(FCS)^{40,41}$ and its two-color variant, fluorescence cross-correlation spectroscopy $(FCCS)^{42}$ are useful for determining protein–protein interactions at low concentrations (~ nanomolar), which is typically the physiological concentrations of many proteins. FCS is capable of determining the stoichiometry of ligand–receptor binding,⁴³ while FCCS reports the strength of protein–protein interactions.⁴⁴ For these experiments, we purified OmpR and the cytoplasmic domain of EnvZ (EnvZc), which is the



Fig. 3 Cross-section of bacteria in $0.5 \times M9$ (A) and MES pH 5.6 (B). Covering the bacterial cells with a hydrophilic gel matrix enabled some bacteria to orient axially, these provide cross-sectional images of the cells under various growth conditions. (i), (ii) and (iii) show the PALM, dSTORM and PAINT images of OmpR (red hot), DNA (cyan) and membrane (green), respectively. OmpR was distributed around the chromosome (iv) and along the plasma membrane (v), likely enabling recruitment to membrane-embedded EnvZ molecules during acidic and hypotonic shock. Interestingly, the chromosome seems to form a tube in the middle of the cell cylinder (vi) under acidic conditions, which exhibits small DNA fibers stretching from the tube towards the membrane (yellow arrows). (vii) Represents the composite image of all three SMLM channels (scale bar 0.5 μ m).

osmolality-sensing domain.^{1,2} EnvZc consists of a four-helix bundle formed by a dimer that contains the phosphorylated histidine and an ATP-binding domain. EnvZc and OmpR were both labeled with Alexa 568 and Alexa 488 at their lone cysteines, respectively.

EnvZc was labeled at its lone cysteine C277 using the maleimide functional group attached to Alexa Fluor 568. We used FCCS to

measure the EnvZ/OmpR interaction. OmpR was labeled with Alexa 488 at its lone cysteine at C67 (OmpR-A488, 90 nM), incubated with excess EnvZc-A568 (750 nM), and then FCCS measurements were performed (Fig. 5A). The cross-correlation ratio (the relative amplitude of the cross-correlation function to the "red" autocorrelation function of EnvZc-A568) is an indication of the fraction of OmpR-A488 molecules that were bound to EnvZc-A568. This value was 0.14 ± 0.02 , which was very low, given that the range of the cross-correlation ratio for the microscope system is between 0.02 and 0.99 (ESI,† Fig. S1). The low level of complex formation determined by FCCS was also confirmed by native PAGE (ESI,† Fig. S3). This weak interaction was presumably due to interference from the label, as labeling at OmpRC67 is known to decrease phosphorylation.²³

In contrast, when the native cysteine of EnvZ was substituted with methionine⁴⁵ and a lone cysteine was introduced in a new location between the four-helix bundle and the ATP binding subdomains (C277M/M294C), the labeled EnvZc^M (EnvZc^M-A568) was able to interact with OmpR. The FCCS measurement indicated that the fraction of OmpR that was bound to EnvZc^M-A568 was three times higher (0.42 ± 0.08) than that of the EnvZc-A568/ OmpR interaction (Fig. 5B). This result suggests that the fourhelix bundle where C277 resides is important for interaction with OmpR, since labelling at C277 prevented OmpR interaction with EnvZc. This finding is consistent with previous studies analyzing conserved specificity determinants,⁴⁶ co-variance,⁴⁷ NMR titration experiments,⁴⁸ and hydrogen : deuterium exchange mass spectrometry,¹ which predicted that the four-helix bundle, and the second helix in particular, is the site of interaction for EnvZ with OmpR. Again, we confirmed our FCCS measurements by native PAGE (ESI,† Fig. S3).

Environmental conditions such as high osmolality that activate the EnvZ/OmpR system may do so by stimulating formation of an EnvZ-OmpR complex. We therefore examined the effect of high osmolality on the EnvZc interaction with OmpR (Fig. 5C). Similar FCCS measurements were performed in 20% sucrose and produced a fraction of OmpR bound to EnvZ that was 0.37 \pm 0.03, *i.e.*, similar to the result in low osmolality (Fig. 5B). Thus, the interaction of OmpR with EnvZ was not sensitive to increased osmolality and complex formation does not appear to play a role in activation during osmotic stress.



Fig. 4 Representative super-resolution fluorescence images of OmpR (right panel, red hot) and RNAP (left panel, red hot) in *E. coli* cells grown in LB medium. The cell membrane was labeled with Nile red and visualized *via* PAINT imaging (green color). OmpR is clearly less abundant compared to RNAP (scale bar 1 μm).



Fig. 5 (A) FCCS measurement of EnvZc-A568 + OmpR-A488. Inset shows the close up of the relative amplitude of the CCF to the ACF of EnvZc-A568 (cross-correlation ratio). The fraction of OmpR-A488 bound to EnvZc-A568 is 0.14 ± 0.02 . (B) FCCS measurement of EnvZc^M-A568 + OmpR-A488. The fraction of OmpR-A488 bound to EnvZc^M-A568 is 0.42 ± 0.08 . (C) FCCS measurement of EnvZc^M-A568 + OmpR-A488 in the presence of 20% sucrose. The fraction of OmpR-A488 bound to EnvZc-A568 is 0.37 ± 0.03 .

Two OmpR molecules bind to an EnvZ dimer

In order to determine the stoichiometry of the EnvZ/OmpR interaction, we performed an FCS experiment by titrating a fixed amount of OmpR-A488 (280 nM) with increasing amounts of unlabeled EnvZc^M. The auto-correlation functions of OmpR-A488 provide information such as the average number of OmpR-A488 molecules and the change in diffusion time upon binding to the larger EnvZc^M dimer (ESI,† Fig. S4). The number of OmpR-A488 molecules obtained from FCS at low $EnvZc^{M}$ concentrations was ~ 20 (Fig. 6A). At higher concentrations of EnvZc^M (>100 nM), the number of OmpR-A488 molecules decreased by nearly 50% to ~ 11 (Fig. 6A). This observation indicates that two OmpR molecules bind to a single EnvZc^M dimer. The reduction in the number of OmpR-A488 molecules was a result of detection of an EnvZc/OmpR complex as a single molecule instead of two OmpR individual molecules, halving N. As expected, the fluorescence count rate per molecule (cpm), obtained



Fig. 6 Titration of $EnvZc^{M}$ with 280 nM OmpR-A488. (A) The average number of OmpR-A488 (*N*). The dotted line indicates the value of *N* for the sample with only OmpR-A488. (B) Count rate per molecule (cpm) values were obtained by normalizing the fluorescence intensity by *N*. The lower dotted line indicates the cpm for the sample with only OmpR-A488. The upper dotted line indicates the expected cpm value (*i.e.*, twice that of a monomer) of an OmpR-A488 dimer.

from normalizing the total fluorescence intensity by the number of molecules obtained from FCS at higher $EnvZc^{M}$ concentrations increased ~ two fold compared to that obtained at low $EnvZc^{M}$ (Fig. 6B). The doubling of cpm values further supports the premise that two OmpR molecules are present in a single EnvZ/OmpR complex, resulting in a stoichiometry of 2EnvZ:2OmpR. This configuration is presumably 1 OmpR bound to each EnvZ molecule of the dimer (see Discussion).

Discussion

In the present study, we constructed a photoactivatable fusion of OmpR that was expressed under its native promoter on the chromosome. The PAmCherry fusion was nearly as active as the wildtype protein, especially at *ompF* (93%), and it was ~70% as active as wildtype at *ompC* (Fig. 1). For reasons that were not obvious, the mEos2 construct containing identical linkers was not as active as the PAmCherry fusion (37% and 17% at *ompF* and *ompC*, respectively). Increasing the linker length to 40 amino acids improved the activity, but it was still lower compared to OmpR–PAmCherry. The active PAmCherry fusion enabled us to image OmpR in single cells under different environmental stress conditions, which provided new insights into the EnvZ/OmpR two-component system.

OmpR localizes to the edges of the nucleoid and near the inner membrane

Super-resolution microscopy allowed us to visualize OmpR, the bacterial chromosome, and the cell membrane in single cells. PALM imaging revealed that OmpR localized preferentially at the nucleoid edges (Fig. 2A) and close to the cell membrane under low salt and acidic conditions (Fig. 2C and D). We anticipate that OmpR would be bound to EnvZ under these conditions, but we were unable to generate a super-resolution image of EnvZ due to its low copy number. Visualizing EnvZc by super-resolution required over-expression.²

Interestingly, bacterial nucleoids appeared to be more condensed under acidic conditions (ESI,† Fig. S2). Nucleoid compaction was also evident in Fig. 2C, by comparing the green outline of the chromosome between samples (compare panels in the third row).

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This compaction could be an acid stress response, reducing acidinduced damage by reducing the exposed chromosome surface. Nucleoid-associated proteins such as Dps, contribute to acid stress tolerance in *E. coli* O157:H7.^{49,50} Further experiments are needed to provide more insight into the underlying mechanisms of acid-induced DNA compaction and the role of nucleoidassociated proteins in this process.

Chromosomal linkages to the inner membrane

One of the elements of the replicon model was that the chromosome was attached to the inner membrane.⁵¹ The attachment was postulated to play a role in the initiation of replication and to enable effective chromosome partitioning.⁵² Other studies reported an attachment at 20-30 membrane sites in addition to oriC.53 Recently, dynamic interactions of DNA with membrane proteins or associated pathways have been identified.54,55 Alternately, this linkage might result from an interaction with DNA coupling transcription, translation and insertion of membrane proteins^{56,57} i.e. a DNA-RNAP-mRNA-membrane interaction (transertion). We observed linkages stretching from the chromosome to the membrane that were evident in axial cross-sections (Fig. 3B). Understanding the molecular basis of these linkages requires further experimentation (e.g. a means of generating a high number of axially orientated bacteria) and the ability to identify EnvZ and other candidate molecules at low copy number.

OmpR localization near the membrane

It was interesting that in the cross-sectional images (Fig. 3), OmpR was located very close to the membrane. We attempted to determine whether it was in a complex with EnvZ, which would be expected. Unfortunately, we could not generate a super-resolution image because of the low in vivo levels of EnvZ. Other transcription factors are known to localize to the inner membrane as a regulatory strategy, including PutA,⁵⁸ Mlc,⁵⁹ MalT,⁶⁰ TraR⁶¹ and NifL,⁶² but this strategy has not been shown to be involved in the EnvZ/OmpR signaling system. Experiments with artificially tethered BglG and LacI proteins have demonstrated that membrane tethering (or interaction) doesn't necessarily lead to inactivation,^{63,64} as these tethered chimeras were still capable of binding to DNA. There are also examples of response regulators that lack the receiver domain, *i.e.* ToxR⁶⁵ and CadC,^{66,67} but instead they are embedded in the inner membrane and they are clearly capable of binding to DNA. This brings us to the question of whether a complex of DNA-OmpR-EnvZ exists in vivo?²³ Because of the difficulties of two-color PALM imaging in bacteria, addressing this question will require the development of new fluorescent proteins, a task that is underway currently in our laboratory.

The discrepancy between molecule counting by PALM and quantitative western blots

Comparing PALM images of OmpR with those of RNAP allowed us to assess the relative expression level of OmpR. In rich medium such as LB, RNAP copy numbers were determined to be ~ 3600 copies per cell.¹⁶ OmpR copy numbers were significantly lower compared to RNA polymerase (Fig. 4). This result raises doubts as to the levels of OmpR that were previously determined by quantitative western blotting to be 2000–3500 copies per cell.³⁹ This discrepancy between copy numbers determined from super-resolution imaging and western blots has recently been reported for numerous other proteins.^{34,39,68} If OmpR levels are indeed substantially lower than previously reported, this explains why OmpR ~ P levels were barely detectible using Phos-tag labeling, except in the presence of certain *envZ* mutations.⁶⁹

Numbers for the detection efficiency of photoactivatable proteins vary widely. Copy numbers obtained by PALM-imaging compared to quantitative western blots yielded only 3.6% detection efficiency for PAmCherry,³⁴ while the use of a nanotemplate resulted in 45–50% detectable proteins.⁶⁸ Our previous study determined RNAP copy numbers using quantitative PALM resulted in ~3600 RNAP molecules per cell, in good agreement with quantitative western blots (~2600 molecules per cell).⁷⁰ Even if a fraction of PAmCherry molecules does not photoactivate or fold properly, the number of active molecules can be compared, since both OmpR and RNAP fusions carry the same fluorescent protein with similar biological and photophysical properties, further emphasizing that the number of OmpR molecules are substantially lower than the 3500 copies reported from quantitative western blotting.³⁹

OmpR binding to EnvZ is cooperative

When the EnvZc concentration exceeded 280 nM, i.e., when the stoichiometry of EnvZc to OmpR was more than 1:1, our data showed that the number of OmpR molecules in the EnvZc/ OmpR complex remained at ~2, at least up to ~2 μ M EnvZc^M (Fig. 6). This would be expected if OmpR binds to EnvZ cooperatively. At an excess of EnvZc^M (>280 nM), a noncooperative reaction would ensure that most EnvZc^M dimers would bind only one OmpR molecule. This would result in a decrease of cpm in the FCCS experiment back to the value of a OmpR monomer (*i.e.* cpm of \sim 17). If the reaction is cooperative, the first OmpR molecule that binds to an EnvZc^M dimer will recruit the second OmpR molecule due to the higher affinity for OmpR compared to the EnvZc^M dimer that does not have OmpR bound to it. Our results are consistent with similar findings using native PAGE.⁷¹ It clearly is not possible for OmpR to already be a dimer in solution before binding an EnvZc^M dimer, as that result would not halve the number of OmpR molecules during the $EnvZc^{M}$ titration. This finding is also consistent with reports of monomeric OmpR in solution⁷² and our finding that $OmpR \sim P$ (which does form dimers in solution) has a lower affinity for EnvZ compared to OmpR.73

In summary, we created a chromosomally-encoded, active OmpR–PAmCherry fusion protein. Using super-resolution microscopy, we showed that OmpR was localized near the membrane in low salt and acidic conditions and that its expression level was substantially lower than RNA polymerase, even though quantitative western blots had suggested they were expressed at similar levels. A linkage from the DNA to the inner membrane was observed, suggesting the possibility of a mechanical link from membrane-bound EnvZ to OmpR to the chromosome.

Abbreviations

FCS	Fluorescence correlation spectroscopy
FCCS	Fluorescence cross-correlation spectroscopy
PAmCherry	Photoactivatable mCherry
Min	Minutes
RT	Room temperature
PCR	Polymerase chain reaction
ACF	Autocorrelation function
CCF	Cross-correlation function
PAGE	Polyacrylamide gel electrophoresis
RNAP	RNA polymerase

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