

Single-molecule investigations of the stringent response machinery in living bacterial cells

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The RelA-mediated stringent response is at the heart of bacterial adaptation to starvation and stress, playing a major role in the bacterial cell cycle and virulence. RelA integrates several environmental cues and synthesizes the alarmone ppGpp, which globally reprograms transcription, translation, and replication. We have developed and implemented novel single-molecule tracking methodology to characterize the intracellular catalytic cycle of RelA. Our single-molecule experiments show that RelA is on the ribosome under nonstarved conditions and that the individual enzyme molecule stays off the ribosome for an extended period of time after activation. This suggests that the catalytically active part of the RelA cycle is performed off, rather than on, the ribosome, and that rebinding to the ribosome is not necessary to trigger each ppGpp synthesis event. Furthermore, we find fast activation of RelA in response to heat stress followed by RelA rapidly being reset to its inactive state, which makes the system sensitive to new environmental cues and hints at an underlying excitable response mechanism.

cytosolic diffusion | single particle tracking | photoactivated localization microscopy | stroboscopic illumination

Bacterial cells live in a dynamically changing environment and thus need to adapt quickly to changes in the surroundings. Upon amino acid limitation, the RelA-mediated stringent response globally remodels bacterial physiology (for an excellent review see ref. 1). This process of remodeling is mediated by the production of the alarmones guanosine pentaphosphate, pppGpp, and guanosine tetraphosphate, ppGpp. The ribosome acts as an activator of RelA-mediated (p)ppGpp production, with uncharged cognate tRNA in the A site being the primary inducer (2). The (p)ppGpp alarmone exerts its regulatory role chiefly via direct binding to the bacterial RNA polymerase (3), provoking changes in the profile of transcribed messages. (p)ppGpp also regulates bacterial replication via binding to DNA primase (4) and inhibits translation initiation via binding to initiation factor IF2, a translational GTPase (5, 6). Evidently, the stringent response is a core cellular adaptation pathway, acting as a hub in the regulatory network, where it integrates information about the nutritional status of the bacterial cell and regulates cellular metabolism on the transcription, translation, and replication levels.

Recently it became possible to clone and overexpress RelA from several bacterial species (7–9), reigniting interest in the stringent response. Using overexpressed *Escherichia coli* RelA, the most extensive in vitro analysis to date of the mechanism was undertaken in 2002 (10), arriving at the so-called hopping model. In this model, RelA binds to a stalled ribosome, senses the deacylated tRNA in the ribosome A site, becomes catalytically active, and synthesizes one ppGpp molecule. The act of ppGpp formation by RelA leads to dissociation of the factor from the 70S, and consequent rebinding to the next ribosome completes the cycle (see Fig. 14). Successive “hops” from ribosome to ribosome monitor the global state of translational starvation. Because hopping events are unsynchronized within the RelA population in vivo, characterization of such events is beyond the reach of

available in vivo bulk assays. At the very same time, in vitro biochemical methods, however powerful when applied to other ribosome-associated enzymes, have so far failed to characterize these events as well. The original hopping model (10) was inferred within a very general conceptual framework that encompasses a broad range of ribosome-interacting enzymes, be it translational GTPases (11), ribosome-inactivating toxins (12), or ribosome RNA modification enzymes (13). Direct experimental evidence for hopping, for the synthesis of one ppGpp alarmone per one dissociation event, and also for the reaction being performed on the ribosome, is lacking (10).

This prompted the development of an in vivo approach based on single-molecule fluorescence imaging that allows us to directly observe the stringent response machinery, which we use to test the hopping model in vivo. To monitor RelA in its freely diffusing state it was necessary to develop a new method for tracking individual rapidly diffusing proteins in the bacterial cytosol. We demonstrate that free RelA in the bacterial cytosol, and ribosome-bound RelA, have radically different diffusive properties. This allows us to monitor the catalytic cycle of RelA in vivo by determining the diffusion characteristics of individual RelA molecules throughout their binding cycles. From this we infer the time scale within which individual molecules cycle between free and bound states, thus directly testing the hopping model in vivo. Lastly, by determining the fraction of ribosome-bound and freely diffusing RelA molecules we characterize heat shock in individual cells with unprecedented time resolution.

Results

The first two sections are dedicated to two reference experiments: the diffusion of a free cytosolic protein as a reference for free RelA, and the diffusion of individual ribosomes as a reference for ribosome-bound RelA. Neither of these targets have been characterized before, and these investigations are therefore important in their own right because they provide valuable insights into the nature of the bacterial cytosol and into the micro-structure of the bacterial translational machinery. The third section will be devoted to RelA during amino acid starvation and also during heat shock.

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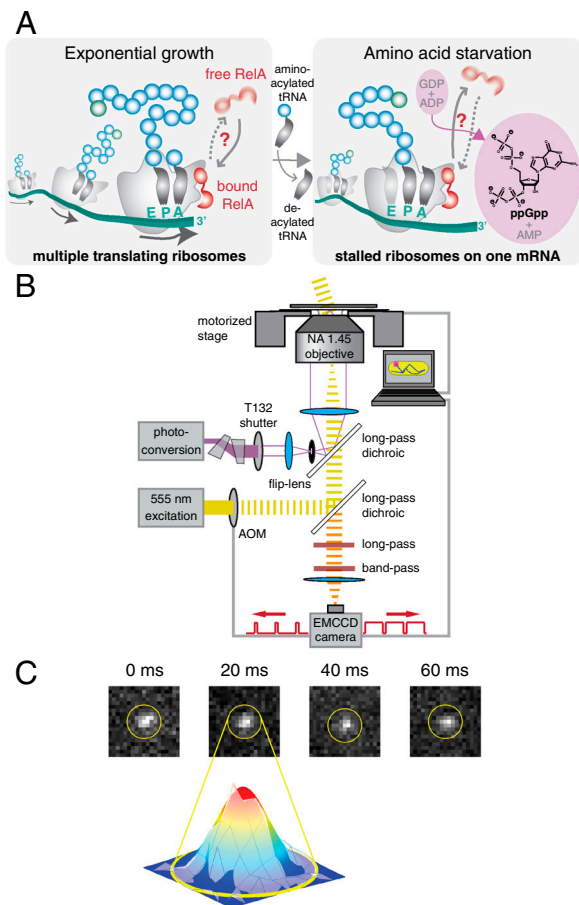


Fig. 1. Single-molecule tracking in living cells. (A) Schematic drawing of RelA interacting with a polysome. During exponential cell growth, RelA could either be bound to the polysome (10) or diffuse rapidly in the cytosol (Left). During amino acid limitation the level of tRNA acylation drops, resulting in accumulation of deacylated tRNA in the ribosomal A site. Under these conditions RelA could either be in tight complex with a polysome (46) or unbind from the ribosome and undergo rapid cytosolic diffusion (Right). (B) Schematic diagram of the optical setup. An acousto-optical modulator is synchronized with an EMCCD camera and shutters a wide-field yellow excitation laser beam to pass short excitation pulses in the middle of each imaging frame. A violet photoconversion laser beam is spatially overlapped and can either be focused onto the back aperture of an Olympus TIRF objective or focused onto the sample plane via a flip-lens. (C) Four consecutive frames of a time-lapse movie of RelA-Dendra2 (in nonstarved cells) with a frame time of 20 ms and an exposure time of 5 ms and a Gaussian fit to frame 2.

The two control experiments presented two challenges for single-molecule tracking. The first one involves the sheer number of ribosomes. An individual *E. coli* cell contains from 7,000 to 50,000 ribosomes (14). Such large copy numbers of targets necessitated a photoconversion approach, in which we only converted and tracked one or a few molecules at a time. These molecules were tracked until they bleached, whereupon the activation cycle was repeated. This approach allowed for the acquisition of good diffusion statistics for individual cells in vivo. The second challenge is the sheer speed at which free proteins move through the cytosol: Current in vivo tracking methodology based on photo-activated localization microscopy (PALM) (15, 16) and stochastic optical reconstruction microscopy (STORM) (17) is limited to the observation of slowly moving molecules, such as proteins bound to membranes (18, 19) or other relatively immobile structures such as DNA (20) or the cytoskeleton (21). These methods have therefore recently been complemented by FCS-based tracking schemes (22) that have excellent temporal resolution but very limited spatial range. To track the cytosolic reference protein

and RelA in its free state we improved in vivo tracking to permit monitoring of fast diffusive processes. This was accomplished by combining superresolution tracking of photoconvertible proteins (18, 21) with a technique borrowed from high-speed photography, stroboscopic time-lapse imaging (20, 23). The key to this is to hardware synchronize short laser excitation pulses with the frame time of the camera such that the fluorophores are effectively immobile during the imaging and do not yield blurred-out views of the diffraction-limited spots, which would be limiting in the presence of the autofluorescent background (see Fig. 1 B and C and Materials and Methods).

Stroboscopic Single-Molecule Tracking of Freely Diffusing Cytosolic Proteins.

As a reference point for free RelA we chose a small cytosolic protein: the GFP variant mEos2. It is a small 26-kDa cytosolic protein whose fluorescence is photoconvertible from green to red (24). We stochastically converted mEos2 to spawn single fluorophores, which allowed for repeated single-molecule tracking within the same living cell. Here we present the first dataset from single-molecule tracking in the cytosol (see SI Materials and Methods in the SI Appendix). The dataset consists of hundreds of trajectories each for eight individual cells, comprising a total of 3,766 single-molecule mEos2 trajectories, which allowed us to characterize how the bacterial cytosol is perceived by a small protein all the way down to the 4-ms time scale. Individual trajectories of mEos2 display several characteristics. First, mEos2 molecules are very fast, covering a considerable part of the bacterial cells between the individual frames even at a frame rate of 250 Hz (Fig. 24). Second, diffusion traces of mEos2 molecules seem to be evenly distributed over the entire volume of the cell, which is especially obvious when individual trajectories are overlaid (Fig. 2B) and sample the whole volume (Fig. 2 A and B).

A quantitative way of analyzing the trajectories in Fig. 2B is by calculating the local apparent diffusion coefficients throughout the cell. These rates are based on how far individual molecules originating from small cellular subregions move within the 4-ms frame time (Fig. 2C, Upper). There appears to be some spatial variation in apparent diffusion coefficients across the cell (8 to 16 $\mu\text{m}^2\text{s}^{-1}$). This variation can be erroneously interpreted as variations in intracellular viscosity. Identical patterns emerge when we simulate normal diffusion using a uniform rate of diffusion throughout the cell and take into account cellular geometry (see Fig. 2C, Lower): Bacterial geometry results in apparent diffusion that is faster in the middle of the cells where molecules can diffuse in a less restricted manner and slower in the quarter positions where molecules have encountered the wall and returned within 4 ms.

Fig. 2D depicts an experimental mean square displacement (MSD) curve for mEos2. We compare the experimental data points to MSD curves calculated from simulated normal diffusion trajectories within the cell geometry. The good fit suggests that cytosolic diffusion is indistinguishable from a Brownian walk. As can be seen in Fig. S1 in the SI Appendix, a single microscopic diffusion coefficient of 13 $\mu\text{m}^2\text{s}^{-1}$ can adequately describe all of our data obtained from all eight cells, because all eight experimental MSD curves fall within their respective confidence intervals. Whereas we present a more detailed treatment of the dataset and of microscopic simulations for mEos2 diffusion in SI Materials and Methods in the SI Appendix, in what now follows, we will use apparent diffusion coefficients throughout. We also complemented our single-molecule analysis with ensemble photo-activation (PA) performed on one of the analyzed cells (see Fig. S2 in the SI Appendix). The obtained diffusion coefficient of 11 $\mu\text{m}^2\text{s}^{-1}$ is in reasonable agreement with our single-molecule analysis, especially when considering the spatial coarseness of PA.

Taken together, our results establish that the small cytosolic protein mEos2 freely diffuses within the entire volume of the bacterial cell, seemingly unperturbed by internal structures such

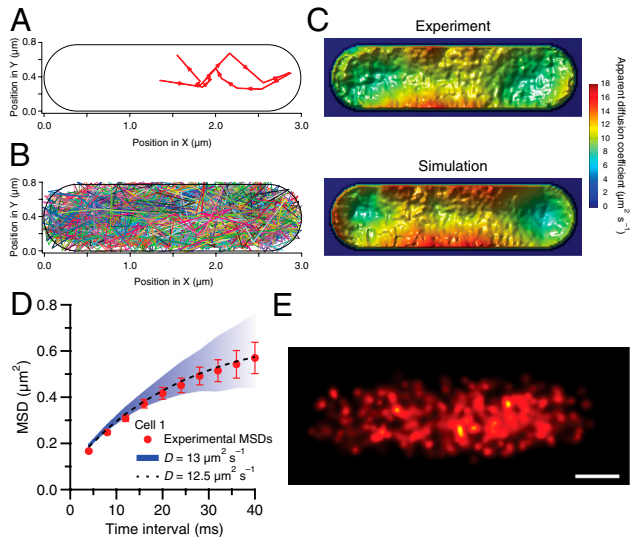


Fig. 2. Normal free diffusion in the cytosol of *E. coli*. (A) A single experimentally obtained single-molecule mEos2 trajectory with a frame time of 4 ms and an exposure time of 1 ms. (B) Overlay of 1,355 single-molecule mEos2 trajectories obtained in an individual *E. coli* cell with a frame time of 4 ms and an exposure time of 1 ms. (C) Local apparent diffusion coefficients in the sample (x-y) plane. The apparent diffusion coefficients are evaluated every 20 by 20 nm in an x-y grid. Each point in the figure is false-colored according to the apparent diffusion coefficient calculated from the mean square displacement over 4 ms for experimental (Upper) and simulated (Lower) displacements originating within 200 nm of this point. The simulations assume normal diffusion at $D = 12.5 \mu\text{m}^2 \text{s}^{-1}$ in the volume defined by the geometry of this cell. There is good agreement between apparent experimental diffusion coefficients and those obtained from simulations. The apparent diffusion coefficients are higher in the middle of the cell, as the molecules are less confined along the long axis. Noise contributions make the apparent diffusion faster close to the cell wall. (D) Mean square displacements (MSDs) in the sample (x-y) plane for different time intervals. Experimental MSDs and error bars representing experimental standard errors of the means are displayed in red. The confidence intervals (blue) are obtained from simulations in the volumes defined by the cell geometry by calculating and sorting MSDs for trajectories using a diffusion coefficient of $13 \mu\text{m}^2 \text{s}^{-1}$. The average MSDs (black, dashed) are also obtained from simulations. Here we vary the diffusion coefficient for each cell to obtain the closest match to the experimental curve. (E) Overlay of 500 positions of single-molecule mEos2 trajectories in one *E. coli* cell with a frame time of 4 ms and an exposure time of 1 ms. Each position is represented by a Gaussian with a standard deviation equal to the localization error. The mean localization error is 44 nm and the scale bar represents 500 nm.

as the nucleoid or the cytoskeleton. This inherent intracellular inertness, which gives rise to the simple diffusion properties, makes mEos2 and similar GFP variants [Dendra2 (25), PA-GFP (26), Dronpa (27), etc.] ideal fusion partners when tracking other proteins for which deviations from simple Brownian diffusion is of biological interest, such as RelA.

In Vivo Single-Molecule Tracking of Bacterial Ribosomes. As a reference point for ribosome-bound RelA we tracked individual fluorescently labeled ribosomes in exponentially growing cells, because reports on the motion of individual ribosomes in the bacterial cell are generally lacking in scientific literature.

Fig. 3A contrasts an MSD curve of 70S diffusion (from 537 individual ribosomal trajectories, comprised of 3,421 positions) with that of mEos2, which we have previously described. A striking difference is immediately obvious; the apparent diffusion coefficients of ribosomes are at least 10 times lower (see Fig. 3A, slopes). Because mEos2 diffusion is extremely rapid, the cytosolic confinement is already noticeable at the 4-ms time scale, which reduces the apparent diffusion coefficients to values that are much lower than the microscopic diffusion coefficient for mEos2

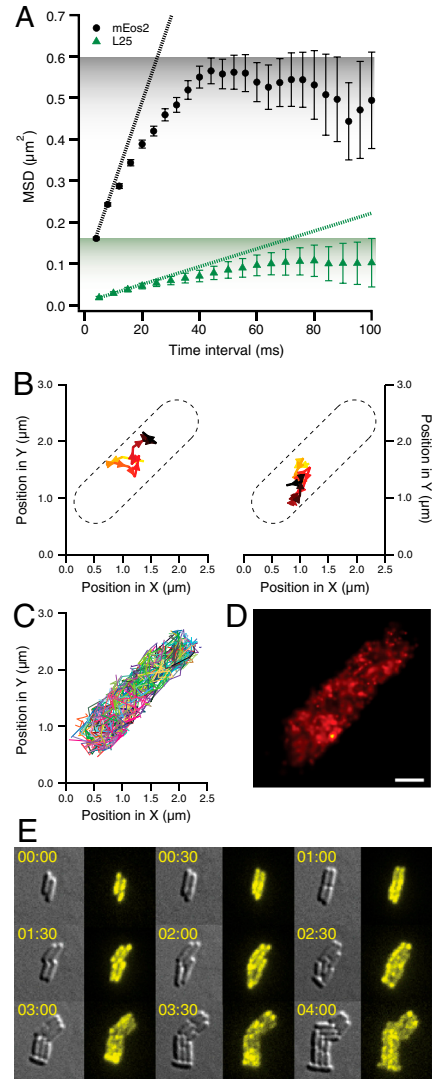


Fig. 3. Single-molecule ribosome tracking and ensemble time-lapse imaging in individual living cells. (A) Mean square displacements (MSDs) in the sample (x-y) plane for mEos2 (in black, from 3,766 trajectories) and ribosome (in green, from 537 trajectories) tracking for different time intervals with error bars representing standard errors of the means. For ribosome tracking, we labeled the ribosomal protein L25 with the photoconvertible GFP variant Dendra2 (25) (see *Materials and Methods*). The dotted lines are calculated from the slope of the first two time points (corresponding to D_{app} of 5.1 and $0.5 \mu\text{m}^2 \text{s}^{-1}$, respectively). As estimated from the initial slopes of the MSD curves, mEos2 has a 10-fold higher apparent diffusion coefficient than Dendra2-labeled ribosomes. The shading represents the different plateaus for mEos2 and L25 MSDs. (B) Two experimentally obtained single-molecule ribosome trajectories with a frame time of 50 ms. The individual ribosome trajectories are recorded for 0.65 and 1.15 s, respectively. The ribosome is tagged via an N-terminally Dendra2 labeled L25 ribosomal protein. (C) Overlay of all 224 single-molecule ribosome trajectories in one *E. coli* cell with a frame time of 50 ms. (D) Overlay of 1,000 positions of single-molecule ribosome trajectories in one *E. coli* cell with a frame time of 50 ms. Each position is represented by a Gaussian with a standard deviation equal to the localization error. The mean localization error is 43 nm and the scale bar represents 500 nm. (E) Time-lapse image acquisitions (differential interference contrast and fluorescence imaging) of ribosome distributions in dividing *E. coli* cells. The fluorescence of Dendra2 from chromosomally labeled C-terminal ribosomal protein S2 is activated at time zero. Subsequent time-lapse imaging followed the initial distribution of the photoconverted ribosomes as they are passed between the cells upon repeated cell division. The cellular distribution of this photoconverted Dendra2 is recorded every five minutes. We present nine snapshots over a period of four hours. Experiments with ribosomal protein L19 and L31-labeled strains resulted in similar data.

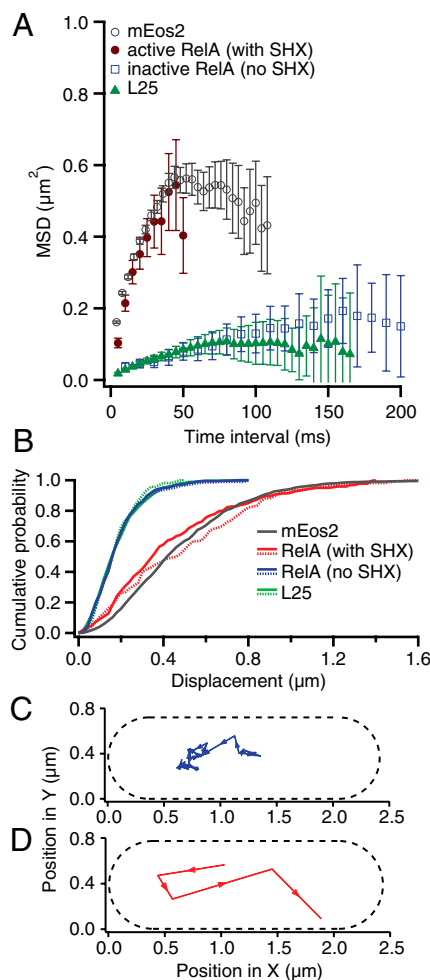


Fig. 4. Diffusion comparison of the ribosome to RelA in exponentially growing *E. coli* cells and to RelA during stringent response. (A) Mean square displacements (MSDs) in the sample plane of *E. coli* cells over different time intervals. The error bars represent the experimental standard errors of the means. MSDs from the ribosomal protein L25 are displayed in green (5-ms frame time, 537 single-molecule trajectories, 3,421 positions). When *E. coli* cells are growing exponentially the MSDs of RelA diffusion (in blue, 10-ms frame time, 90 trajectories, 527 positions) are indistinguishable from L25 diffusion. When cells are starved by addition of 2.5 mM of L-Serine Hydroxamate (brown MSD curve, 5-ms frame time, 216 trajectories, 977 positions), the diffusion of RelA changes dramatically, and is similar to that of mEos2 (average MSDs obtained from all 3,766 single-molecule mEos2 trajectories is displayed in gray). (B) Cumulative distribution functions (CDFs) of displacements over 20 ms in the sample plane of *E. coli* cells. Two CDFs for trajectories of RelA in its inactive unstarved state (in solid-blue with 20-ms frame time, 129 trajectories, 1,325 positions, and dashed-blue with 10-ms frame time, 90 trajectories, 527 positions) and that of L25 (in dashed-green with 20-ms frame time, 49 trajectories, 478 positions, and solid-green with 5-ms frame time, 537 trajectories, 3,421 positions) are indistinguishable. The apparent diffusion coefficient of RelA when cells are starved increases more than eightfold (red and dashed-red curves) and is very similar to the CDF of mEos2 (in gray, average CDF obtained from eight cells). (C) One experimentally obtained single-molecule RelA trajectory with a frame time of 20 ms and an exposure time of 2 ms when cells are exponentially dividing. (D) One experimentally obtained single-molecule RelA trajectory with a frame time of 20 ms and an exposure time of 2 ms when cells are starved using 2.5 mM L-SHX.

RelA molecules are expected to be engaged in prolonged active catalysis.

During the SHX treatment we observe that RelA diffusion undergoes a radical change (Fig. 4D) and becomes very similar to that of mEos2. RelA now diffuses much more rapidly and no longer displays subdiffusive behavior, instead sampling the entire volume of the bacterial cell (compare Figs. 2A and 4D). As we do

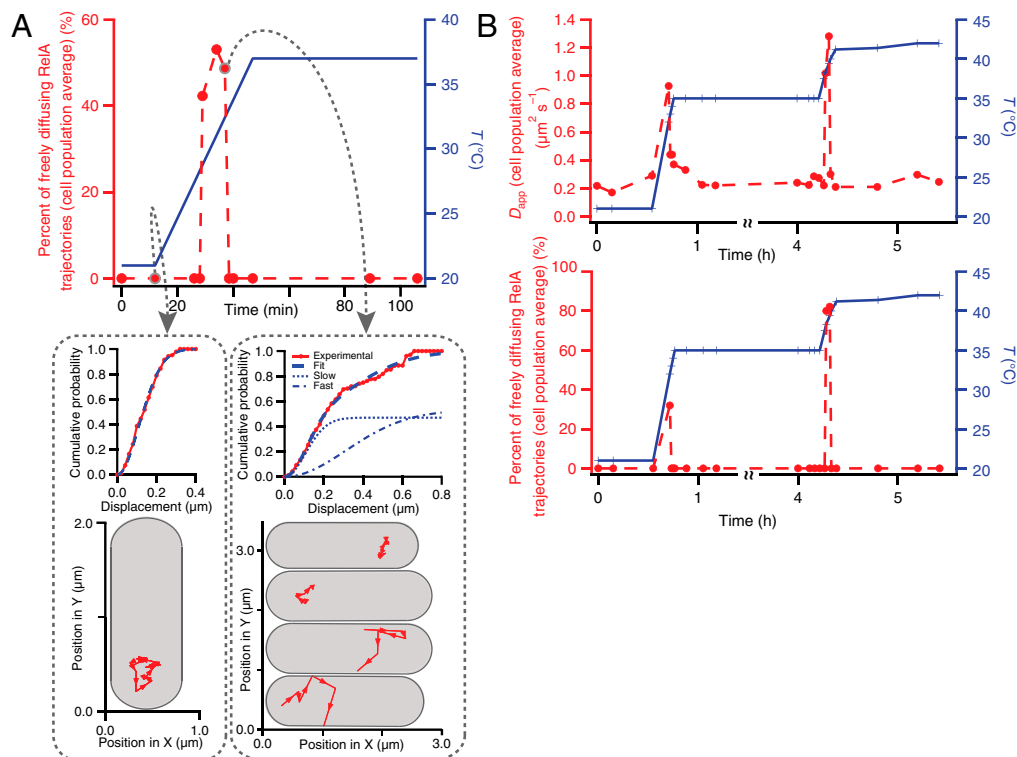
not observe any 70S rebinding events, RelA appears to spend the major portion of its time off, rather than on the ribosome. The close similarity between mEos2 and active RelA trajectories is confirmed when we compare their MSD curves (Fig. 4A). The brown curve in Fig. 4A displays an MSD curve of a representative dataset of 216 individual single-molecule RelA trajectories (composed of 977 positions) obtained from cells plated on an SHX-containing agarose pad. Fig. 4B depicts the CDFs of two datasets each for RelA under active and inactive conditions, reconfirming the striking difference between the two states. In the active state, the CDF curves for RelA dynamics closely resemble those of mEos2 (compare gray and red CDFs, respectively), while in the inactive state, the CDF curves for RelA dynamics are indistinguishable from those of the 70S (compare green and blue CDFs, respectively). These experiments clearly show that one prediction of the “hopping” model holds true: RelA does indeed dissociate from the ribosome under starvation conditions. However, we do not observe RelA “hopping” in the sense that it is immediately rebinding to another ribosome. On the contrary, it stays off the ribosome for hundreds of milliseconds, which suggests an extended hopping model (see Discussion).

Having characterized the two extreme states of the RelA cycle (i.e., perpetually active and inactive RelA), we next applied our methodology to a more physiologically common and a more transient cellular stress response—namely, a temperature upshift (40–42). We took advantage of RelA behaving as a logical switch, changing its diffusion characteristics between two distinct states (active and inactive—i.e., fast or slowly diffusing, respectively) and we utilized this as a readout of cellular stress on the single cell level.

We heat-shocked *E. coli* cells by rapidly increasing the temperature from 21 to 37°C, from 21 to 35°C and from 35 to 42°C (Fig. 5). In all of these cases, within several minutes after the temperature upshift, RelA dissociates from the ribosome and, as in the case of SHX-induced stringent response, diffuses freely. The striking difference to the SHX-treated cells is that after the temperature upshift the initial inactive state is restored within 10 min, as all RelA molecules observed rebound to ribosomes.

To estimate the ratio of the enzymatically active versus inactive RelA molecules in each time point during the temperature upshift, we fit the CDF curve for all recorded trajectories with a linear combination of two CDFs: a fast one, representing free RelA, and a slow one, representing the 70S-bound RelA from data also presented in Fig. 4B. In doing so, we have a straightforward way of deconvoluting the entire dataset and estimating the percentage of the two RelA states at any given time. This is only possible due to an order of magnitude difference in the apparent diffusion coefficients for the two RelA states (see Figs. 3A and 4A), which allows us to use the apparent diffusion coefficients as a logical operator for sorting the particles. As a control for the automated sorting procedure we also sort our trajectories manually into two categories (see Fig. 5A, Bottom), and this approach gives the same ratio as our global deconvolution method described above.

Looking at the individual trajectories at any given time during the very fast temperature response, we observe rapid RelA diffusion in about half of the cells (Fig. 5A and B). When cells overcome the temperature shock, the entire RelA population shifts to the inactive state. All cells will have responded within approximately 10 min, and all cells return to the slow inactive state even when shocked to a final temperature of 42°C. Our observations corroborate nicely with the transient ppGpp level increase documented in refs. 40 and 42. Just as in the case of our SHX-induced amino acid starvation tracking experiments, the observed timing of our responses follows the same pattern of high ppGpp levels reported in microbiological experiments (39, 40, 42).



gested that RelA strongly binds to the ribosome under starvation conditions (46).

We confirm *in vivo* that under nonstarved conditions RelA is indeed ribosome-bound and that under amino acid starvation it does dissociate from the ribosome. The hopping model predicts that RelA subsequently hops from ribosome to ribosome producing exactly one ppGpp molecule per dissociation event. Hence RelA is predicted to toggle between its ribosome-bound and free states with a corresponding frequency. Reported *in vitro* turnover rates of RelA are about 70–75 per second (7), and *in vivo* estimates of ppGpp production under starvation conditions are of order 50,000 molecules per cell per second (47), which taking into account RelA cellular copy number of about 100 (35), gives similar turnover estimates. Such rapid alterations in diffusion characteristics of RelA would easily have been detected in our single-molecule tracking assay, as discussed below.

We, however, observe all RelA molecules remaining dissociated from the ribosome for hundreds of milliseconds. The temporal resolution of our assay is 4 ms due to the limitations in the EMCCD camera readout time. At the same time, the dynamic range spans from 4 ms (the fastest achievable frame time, see above) to seconds (limited by trajectory lengths). In the scenario where the catalytic cycle of RelA is much, much too fast to be captured by our assay, and we therefore would miss all of the thousands of binding events during the course of one trajectory, the two substrates of RelA (ATP and the G nucleotide, GTP or GDP), and RelA itself would have to bind to a stalled ribosome, RelA would have to conclude its catalytic event, and dissociate from the ribosome all in well under 1 ms.

This is orders of magnitude faster than ribosome-binding kinetics determined for other translational components and is much faster than even ricin catalyzed rRNA cleavage (48), which is, to our knowledge, the fastest enzyme “hopping” between the ribosomes, and is probably beyond the diffusion limit for a protein the size of RelA (49).

In the other scenario where the RelA catalytic cycle happens far too infrequently to be captured by our assay (i.e., much longer than seconds), it would take hours for RelA to produce enough ppGpp to cause the stringent response: The cellular copy number of RelA is not over 100 (35), and ppGpp concentrations of 0.5 mM are reached *in vivo* (50), which corresponds to 300,000 molecules of ppGpp. Hence even in the fastest scenario in the absence of cellular ppGpp degradation, it would take about an hour to build up adequate ppGpp levels at a synthesis rate of 1 per second, which is far too long to be compatible with the available *in vivo* data. Both our heat-shock experiments (where cellular adaptation occurs within minutes (and not hours), as well as the documented rise of ppGpp in about a minute after SHX shock (39) are consistent with RelA turnover rates of 10s to 100s per second, not less than 1 per second.

This is why we argue that multiple ppGpp molecules have to be synthesized every time RelA dissociates from the ribosome. We therefore propose an “extended hopping” model—a modification of the original hopping model where many ppGpp molecules are produced upon the dissociation of enzymatically active RelA from the ribosome, and with RelA active off, rather than on the ribosome. Our new model can be rationalized in the framework of existing biochemical data for RelA, which show that RelA has an intrinsic ability to produce ppGpp, which it can auto-inhibit through the use of a dedicated inhibition domain (51). When this domain is removed, RelA switches into a perpetually active state (51). Possibly, activation by the ribosome is in fact a suppression of auto-inhibition. Within this framework it seems only natural that following dissociation from the ribosome, RelA remains active for some time before its auto-inhibition is engaged. The severe and lasting starvation induced by SHX was used to keep RelA in its active mode for an extended time in order to probe its catalytic cycle. We also studied RelA dynamics during heat-shock

response, which is a more natural and transient form of stress. Furthermore, RelA's direct involvement in heat shock has not been possible to study *in vitro*, since the primary heat-shock sensor is unknown. Our *in vivo* enzymatic assay reveals that in this case, as well, RelA dissociates from the ribosome at the onset of stress, but in this case its excursion into a fast active state is more transient in nature: Within less than 10 min, the RelA sensory system adapts to the new temperature and resets, ready to respond to further changes in temperature. There are at least two possible topologies for the heat sensing network, which is known to be mediated by RelA and SpoT (40), a close homologue to RelA. First, both RelA and SpoT sense the heat change independently. Second, SpoT acts as the primary sensor: Because temperature changes are unlikely to be sensed by increased levels of deacylated tRNA, it is conceivable that SpoT triggers RelA activation via an unidentified intracellular messenger. Both of the proposed topologies are consistent with a much smaller but still detectable increase in ppGpp levels even in RelA knockout strains, where ppGpp-mediated heat shock is still functional, and where SpoT acts as the sole source of intracellular ppGpp (40). In the latter topology, RelA, while being transiently unbound from the ribosome, acts as an amplifier of the heat response by significantly boosting ppGpp production after the initial rise in intracellular ppGpp synthesized by SpoT, the initiator of the heat response.

On our path to the *in vivo* mechanism of RelA, we have characterized the intracellular diffusion of both ribosomes and free proteins. We have shown that the movement of individual ribosomes is very slow and highly subdiffusive in nature. The subdiffusive ribosomal movement is most likely due to mRNA tethering, the diffusion of which displays very similar properties (32). On a much longer time scale, time-lapse movies of fluorescently labeled ribosomal proteins have not revealed preferential partitioning of ribosomes during cell division.

While more thorough investigations would benefit from a 3D superresolution approach, be it 3D SPT (52) or 3D PALM (15), these investigations lay the groundwork for a more detailed single particle tracking investigation of the translation process in living cells, as well as into ribosomal protein dynamics during ribosome assembly (53) and into ribosomal aging and repair by exchange of ribosomal proteins (54). It also lays the foundation for forthcoming studies with a more detailed analysis of ribosomal distributions during the exponential phase (55), stationary phase (56–58), and during cold shock (59).

Our *in vivo* single-molecule investigations into the stringent response were made possible due to the development of an assay that can detect also the very fast freely diffusing protein states that would otherwise be overlooked. We found that the diffusion characteristics of the inert freely diffusing mEos2 protein are indistinguishable from Brownian motion. This inherent intracellular inertness, which gives rise to the simple diffusion properties, makes mEos2 and similar GFP variants ideal fusion partners when tracking other proteins for which deviations from simple Brownian diffusion is of biological interest. In addition, the time resolution of this stroboscopic tracking assay makes it possible to monitor binding events in the millisecond regime. This time resolution is on par with even the fastest cellular protein dynamics, which makes the assay a general tool for investigation of intracellular kinetics in living cells.

Materials and Methods

Laser Microscopy. A schematic diagram of the optical setup is shown in Fig. 1B. An acousto-optical modulator (AOM, IntraAction, 40 MHz) shutters a wide-field yellow excitation laser beam into an Olympus IX81 inverted microscope. The excitation laser light from a 555-nm DPSS laser (CrystaLaser) is first filtered (Z 550/20×, Chroma Technology) and then focused onto the back aperture of an Olympus TIRF objective (NA = 1.45). The objective collimates the light and excites either a 4-μm- (for mEos2 tracking) or a 20-μm-wide area (for ribosomal and RelA tracking) at the sample plane with laser power densities

of 50 to 200 kW cm⁻². The AOM is synchronized with a PhotonMax EMCCD camera (Princeton Instruments) by a NI-DAQ M-series data acquisition card (PCI-6259, National Instruments), and is controlled via LabVIEW 8.5 (National Instruments) to pass short 0.3- to 5-ms excitation pulses in the middle of each imaging frame. Our stroboscopic illumination time is optimal in the range of 1 ms (for mEos2 tracking) to 5 ms (for L25 tracking). The short duration of the excitation pulse is a critical parameter for obtaining distinct fluorescence snapshots from our rapidly moving single molecules. This makes it possible to track them at superresolution throughout all frames (see Fig. 1C for four consecutive raw image frames of a RelA–Dendra2 trajectory). Another equally important advantage of stroboscopic illumination is that it allows for a direct comparison between fast and slow molecules under identical illumination conditions since snapshots can be spaced out with varying time intervals.

A violet photoconversion laser beam (405 nm Radius, Coherent) is spatially overlapped using a long-pass dichroic filter (Z405RDC, Chroma) and is focused at the sample plane. An anamorphic prism pair (06 GPA 204, Melles Griot) and a 50- μ m pinhole (Thorlabs) correct the asymmetry of this laser beam to produce a TEM₀₀ mode. The photoconversion laser is independently shuttered using a UNIBLITZ T132 shutter controller that delivers 3-ms activation pulses at regular time intervals at power densities of order 0.1 kW cm⁻².

A long-pass dichroic filter (Z555RDC, Chroma) is used to separate excitation and photoconversion laser light from the fluorescence emission of mEos2. Emission light is further filtered using both band-pass (HQ630/140m-2p, Chroma) and long-pass (HQ 565LP, Chroma) emission filters for mEos2 fluorescence, or a band-pass filter (HQ605/75m, Chroma) for Dendra2 fluorescence.

Individual *E. coli* cells are imaged for 10,000 to 60,000 frames at 250 Hz using up to 28 pixel lines, 200 Hz using up to 40 pixel lines, and 20 Hz using up to 128 pixel lines on a PhotonMax EMCCD camera (Camera shutter: always open; digitizer: 10 MHz EM gain). Metamorph 7.5 (Molecular Devices) is used to control the microscope as well as the camera.

Data Processing. Sample frames of RelA–Dendra2 are shown in Fig. 1C. The time-lapse movies are first edited by hand to remove frames with photoconversion pulses and frames where we observe multiple molecules in one frame. Next, we track the fluorophores in our hand-edited movies using the particle tracking software Diatrack (v3.03, Semasopht), which identifies and fits the intensity spots of our fluorescent particles with symmetric 2D Gaussian functions (see Fig. 1C). All routines for trajectory analysis are written in IGOR Pro 6.12A. For each of the analyzed cells all trajectories with three or more points are first overlaid to determine the precise cellular geometry. This geometry is then approximated as a cylinder with two hemispherical end caps. The final geometry of the cell is determined by subtracting twice the mean localization error (40 nm) from all sides of the initial geometry

of the cell. This corrects for apparent cell broadening due to the experimental fitting noise. The localization errors are calculated using equation 6 in ref. 60.

Sample Preparation for SPT Experiments. *E. coli* cells expressing RelA–Dendra2 are imaged on a M9-glucose agarose pad in a FCS2 flow chamber (Bioptechs). The cells are grown overnight to stationary phase and are diluted 1,000-fold in M9 buffer containing amino acids (M5550, Sigma-Aldrich) on the day of the experiments. The cells are grown to an optical density at 600 nm of 0.05 and placed on a 2.5% agarose pad (SeaPlaque GTG Agarose, Lonza) containing the same M9 medium.

For exponential growth experiments, cells are imaged only after having undergone several rounds of cell division until the individual cells have formed small microcolonies on the agarose pad to ensure that they have adjusted to the agarose pad. The temperature is held constant at 30 °C. The temperature is controlled via an electrical cell heater and an objective heater (both Bioptechs).

Individual RelA trajectories are recorded after red Dendra2 is generated using short pulses of the photoconversion laser beam. To obtain adequate statistics, we calculate CDFs and MSDs from all obtained single-molecule RelA trajectories from the entire microcolony for each of the experimental conditions and for each frame time.

For SHX experiments, the agarose pad is presoaked with L-SHX to a final concentration of 2.5 mM L-SHX. The cells are grown as above. When the cells reach an optical density at 600 nm of 0.05, L-SHX is added to the cells to a final concentration of 2.5 mM, and they are placed on the SHX-containing agarose pad. The cells undergo stringent response and immediately stop dividing on the agarose pad.

For the purpose of single-ribosome tracking we constructed a plasmid expressing an N-terminal Dendra2 fusion with the ribosomal protein L25. Low levels of leakage Dendra2–L25 expression provided hundreds of fluorescently labeled ribosomes per individual cell. For the ensemble time-lapse studies we C-terminally labeled three ribosomal proteins with Dendra2: S2, L19, and L31 in three separate strains using the genome integration approach based on the λ Red system (34). Please see *SI Materials and Methods* in the *SI Appendix* for details on strain preparation, validation, and a more detailed description of experimental procedures.

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