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Mechanical slowing-down of cytoplasmic diffusion allows *in vivo* counting of proteins in individual cells

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Many key regulatory proteins in bacteria are present in too low numbers to be detected with conventional methods, which poses a particular challenge for single-cell analyses because such proteins can contribute greatly to phenotypic heterogeneity. Here we develop a microfluidics-based platform that enables single-molecule counting of low-abundance proteins by mechanically slowing-down their diffusion within the cytoplasm of live *Escherichia coli* (*E. coli*) cells. Our technique also allows for automated microscopy at high throughput with minimal perturbation to native physiology, as well as viable enrichment/retrieval. We illustrate the method by analysing the control of the master regulator of the *E. coli* stress response, RpoS, by its adapter protein, SprE (RssB). Quantification of SprE numbers shows that though SprE is necessary for RpoS degradation, it is expressed at levels as low as 3–4 molecules per average cell cycle, and fluctuations in SprE are approximately Poisson distributed during exponential phase with no sign of bursting.
A surge of single-cell fluorescence studies has shown that genetically identical cells residing within the same environment can display extensive cell-to-cell variability in the expression levels of various proteins\(^1^–^3\). A substantial challenge when analysing these phenomena is that the heterogeneity typically originates in reactions involving low-abundance components, while only the high-abundance components that indirectly respond to the heterogeneity are relatively straightforward to measure. For example, many of the key regulatory proteins in Escherichia coli (E. coli) are present in such low numbers—recent studies suggest that at least 10% of the proteins in E. coli are present in <10 copies per cell\(^4^–^8\)—that fluorescent protein (FP) fusions are difficult to detect over the cellular auto-fluorescence\(^9\). Furthermore, when fluorescent levels are detectable, they are typically quantified in terms of total fluorescence and reported in arbitrary units\(^7\). Quantifying the total fluorescence intensity rather than counting separate copies can also introduce measurement errors, as problems with, for example, uneven excitation or detection becomes hard to separate from actual cell heterogeneity. Finally, fluctuations in protein abundances are easier to analyse mathematically when absolute numbers are known\(^7,^8\). The ability to count low-abundance proteins in individual cells would thus substantially help analyse single-cell dynamics.

A recent study\(^4\) quantified levels of low-abundance FPs by deconvoluting the cellular autofluorescence distribution from that of the total fluorescence, which was measured separately. Though the variation in autofluorescence makes it impossible to infer the FP fluorescence in any particular single cell, that approach can still estimate the distribution over the population of cells, at least in arbitrary units of fluorescence. The challenge is that for low copy proteins, where the FP signal is a small fraction of the total, this procedure essentially infers a small quantity by taking the difference between two relatively large quantities, and is thus exceedingly sensitive to measurement errors due to imaging, growth conditions or differences in cell size.

A potentially less error-prone approach is to directly count spatially separate molecules. One early technique used single-cell capture and lysis, followed by downstream binding to antibodies to detect single protein copies\(^9\). Fine tuning allowed ~60% of the molecules to be detected, but only for high-abundance proteins: the lowest abundance detected was ~600 proteins per cell, and it was estimated that any protein present in <10 copies would fall entirely under the detection limit\(^9\). Quantifying protein abundances in vivo by microscopy could help improve detection, but the challenge is that individual proteins diffuse rapidly and appear smeared for typical exposure times. Several approaches have been used to address this problem. Chemical fixation can be used to immobilize and detect single proteins via standard total internal reflection fluorescence (TIRF) microscopy or super-resolution methods\(^12\) but at the expense of substantial denaturation of FPs\(^8\) and an increase in the cellular auto-fluorescence\(^13\). Although super-resolution methods can be used to infer stoichiometries\(^14\), an accurate enumeration of the protein-of-interest (POI) remains challenging because the FPs used for super-resolution imaging exhibit complicated photophysics and suffer from a low yield of conversion into the fluorescently detectable state\(^15\). Otherwise cytoplasmic FPs have also been targeted to the cell membrane\(^16\) to slow down the diffusion, at the cost of disrupting the function of the POI. To address this issue a cotranslationally cleavable linker was added between the membrane-targeted FP and the POI\(^7\), but even if that could be made to work with high accuracy, the method is limited to counting proteins produced within a certain time window. All these different methods further face the challenges that the shallow depth of focus of high numerical aperture objectives is typically smaller than the height of even E. coli cells, making it difficult to detect all copies of the POI in a cell, and that the fluorescence of a single FP can still be difficult to separate from the cellular auto-fluorescence.

Statistical throughput can also be almost as important as resolution in single-cell studies. Interrogating large numbers of cells is not only necessary to ensure that observed differences are statistically significant\(^18\), but is useful for determining distributions more accurately\(^19\) as well as for detecting rare phenotypes\(^20\). High sampling further permits binning of data, where cells are grouped for instance according to their size or gene expression levels before analysing other properties. Such analysis can greatly facilitate interpretations, but the number of distinct bins increases exponentially with the number of properties measured and thus sample sizes quickly become limiting. The ability to count single molecules in single cells with a significant throughput is therefore a key requirement for analysing low-copy number protein fluctuations and the resultant cell-to-cell heterogeneity.

Here we use a simple microfluidic platform (MACS: microfluidics-assisted cell screening) to mechanically compress cells in a controlled manner. This causes diffusional slowing-down of cytoplasmic molecules without loss of fluorescence in E. coli, and thus enables detection of single molecules on a standard TIRF microscope set-up. The resultant flattening of the pressed cells also reduces local autofluorescence, separates the molecules spatially, and makes it easier to keep all copies within the objective depth of focus. Moreover, MACS provides automation with high throughput while growing cells in conventional liquid culture until just before the moment of imaging, and makes it possible to retrieve/enrich rare cells. To illustrate the capabilities of this technique, we applied MACS to study the control of RpoS, the master regulator of stress response in E. coli, by the low abundance adapter protein, SprE (RssB). Though RpoS is one of the most important and well-studied proteins in E. coli\(^21\), and SprE plays an important role in controlling RpoS levels\(^21\), little is known about the dynamics of this circuit because SprE is present in too low numbers\(^21\) to be reliably detected with conventional methods.

Results

Description of the MACS set-up. MACS uses poly-dimethylsiloxane (PDMS)-based microfluidic on-chip valves\(^22\) (Fig. 1a–c), with pressure-driven flow (Fig. 1d) instead of syringe pumps to allow for easy streamlining and fast response times. Although MACS essentially exploits valve actuation to immobilize cells between a glass coverslip and a PDMS membrane similar to what was described earlier\(^23,^24\), simply collapsing the valve (that is, going directly from open to closed state) yields extremely poor trapping efficiency due to the rapid displacement of liquid. Instead, MACS relies on three distinct valve states, achieved by controlling both the pressure of the valve \(P_{\text{valve}}\) and the pressure driving the flow \(P_{\text{flow}}\) of the cell suspension. First, the valve is closed at a certain pressure \(P_{\text{valve}}>0\), while the flow is off \(P_{\text{flow}}=0\) corresponding to the closed state. The pressure driving the flow is then adjusted to a level \(P_{\text{flow}}>0\) that breaks the seal between PDMS and the coverslip, where cells start slipping through as a monolayer, corresponding to the half-open state (Fig. 1e). New cells are introduced, trapped and imaged by sequential cycling between half-open, closed and open valve states. Since each cycle typically takes ~5–15 s (corresponding to ~240–720 frames per hour), this allows automated imaging of E. coli cells with high throughput. As a proof-of-concept for the stability and
Throughout capabilities of MACS, we acquired unattended snapshots of approximately one million stationary phase E. coli cells in 4 h at a single valving intersection. We also imaged large numbers of cells every few minutes along the growth curve from exponential to stationary phase, revealing subtle but reproducible features of growth (Supplementary Fig. 2).

**Mechanical slowing-down of diffusion in E. coli cytoplasm.** We expressed various FPs or translational fusions to FPs in E. coli and monitored them using HILO imaging typically with a 30-ms exposure time (see Methods). Comparing the area of cells imaged on an agar pad versus MACS revealed that cells were compressed and flattened via MACS, increasing the cell area under typically applied pressures ($P_{\text{valve}} = 20$ p.s.i.) on average by 72% under our conditions (Fig. 2a,b). When imaged on agar pads (without any applied pressure), cells displayed a uniform cytoplasmic signal due to the rapid diffusion of molecules (Fig. 2c and Supplementary Movie 2). In contrast, when cells were squeezed with the MACS chips with $P_{\text{valve}} = 20$ p.s.i., individual molecules appeared as diffraction-limited spots due to slowing-down of diffusion (Fig. 2c and Supplementary Movie 3). We speculate that this phenomenon is due to water being expelled from the cell, increasing the density of the E. coli cytoplasm26. Figure 2c shows results for a SprE-mNeonGreen translational fusion expressed from its native chromosomal locus, but similar effects were observed for other FPs tested (Supplementary Fig. 3). To further characterize this mechanical slowing-down of cytoplasmic diffusion, we carried out fluorescence recovery after photobleaching (FRAP) measurements (Fig. 2d, Supplementary Fig. 4 and Methods). For cells imaged on agar, the diffusion coefficient ($D$) of RFP mKate2 was $14 \pm 4 \mu m^2 s^{-1} \ (\pm \text{s.d., } n = 21 \text{ cells})$ (Fig. 2d), in agreement with the previously reported values27. Increasing $P_{\text{valve}}$ from 5 to 20 p.s.i. decreased $D$ below $1 \mu m^2 s^{-1}$ (Fig. 2d). The average displacement of molecules within 30 ms is then $\approx 250$ nm and single molecules of FPs should appear punctate, which is consistent with the discrete spots that we observe.

These results indicate that MACS could be used to image single cytoplasmic proteins in individual E. coli. It was previously suggested that proteins deform under increased molecular crowding, which could lead to denaturation of the FPs28. We therefore confirmed that, in contrast to other fixation methods13, there was no significant loss of signal as seen by comparing the total fluorescence distributions between agar pad versus MACS (Fig. 2e). We confirmed low levels of false positive detection with cells expressing no fluorescent marker, typically showing less than one spot per average cell, and under some conditions as low as 0.3 spots per average cell (Supplementary Fig. 5). Occasional complete immobilization further allowed us to detect single-step photobleaching traces (Fig. 2f), a hallmark of single-molecule
Figure 2 | Characterization of mechanical slowing-down and single-molecule counting on MACS. (a–d) MACS induces cell deformation and mechanical slowing-down of cytoplasmic proteins. (a) Cartoon depicting cell flattening and appearance of discrete spots (that is, single molecules, see below) for MACS versus agar pad imaging. (b) In comparison with agar, cells are flattened when imaged under MACS. (c) SprE tagged with mNeonGreen results in discrete spots on MACS as opposed to a diffuse signal on agar. (d) FRAP measurements quantify the extent of mechanical slowing-down on MACS as a function of \( P_{\text{value}} \) (Supplementary Fig. 4 and Supplementary Movies 4 and 5). Cells were treated with cephalixin and were thus elongated to enable FRAP measurements on agar, since FRAP occurred too rapidly to be measured otherwise. (e–g) Single-molecule counting is feasible with MACS. (e) Comparison of total intensities between agar and MACS imaging for the highly expressed segmentation marker (CFP) of the SprE-mNeonGreen strain suggesting that MACS does not cause signal loss. Comparing images of strains for FP-tagged versus wild-type SprE implies that the spots are specific to the mNeonGreen tagging of SprE. (f) Representative time traces of the SprE spots from one cell exhibiting single-step photobleaching (Supplementary Movie 6). (g) Undercounting due to spatial spot overlap in the cells was quantified using two independent computer simulations. First, a numerical simulation was carried out considering that the Euclidian distance between the spots be smaller than the diffraction-limited resolution (Supplementary Note 1), compared with unpressed cells, facilitates detection of the whole cytoplasmic volume, by ensuring all molecules to be within the depth of focus. Second, the spreading out of cytoplasmic volume also causes a reduction in background autofluorescence density, which improves the signal-to-noise ratio for the detected FPs. Finally, the flattening spreads the diffraction-limited spots over a larger area, reducing the probability of two spots to overlap. With any method, spot overlap can have counterintuitive effects. Specifically, cells that by chance have more molecules will have a higher fraction of overlapping spots. As opposed to most experimental errors, this artefact will narrow the observed distributions, at least the right tail (discussed more below). Our analysis suggests that for cell detection, as expected since we use monomeric FPs and do not fuse them to oligomeric native proteins\(^{10}\).

As a final control, we compared the results from single and dual FP labelling, to determine the rate of false negatives, for example, due to incomplete FP maturation, which is particularly problematic during fast growth. Under the most challenging conditions of 25 min generation times we observed that single-labelled proteins still captured about 80% of the double-labelled proteins, perfectly consistent with the expected maturation time of 5–10 min\(^{29}\) (Supplementary Fig. 9). We further found that with dual labelling the fluorescent spots were almost twice as bright after background subtraction. Thus though maturation is a problem in all FP-based studies, the effects appear small even in the worst-case scenario for mNeonGreen and can be reduced further yet by double labelling.

The flattening of cells under MACS further helps counting in several ways. First, the ~50% decrease in cell height (Supplementary Note 1), compared with unpressed cells, facilitates detection of the whole cytoplasmic volume, by ensuring all molecules to be within the depth of focus. Second, the spreading out of cytoplasmic volume also causes a reduction in background autofluorescence density, which improves the signal-to-noise ratio for the detected FPs. Finally, the flattening spreads the diffraction-limited spots over a larger area, reducing the probability of two spots to overlap. With any method, spot overlap can have counterintuitive effects. Specifically, cells that by chance have more molecules will have a higher fraction of overlapping spots. As opposed to most experimental errors, this artefact will narrow the observed distributions, at least the right tail (discussed more below). Our analysis suggests that for cell
sizes typical of rapidly growing *E. coli*, spot overlap does not significantly interfere until levels reach above 7–8 spots per cell (Fig. 2g), and a substantial fraction of the proteome is present in lower abundances than that. However, our method can also be used to infer numbers with reasonable precision even for higher abundances; and for statistical metrics such as averages (Supplementary Fig. 10) or even distributions and variances as the data can in principle be corrected for overlap.

**Analysing the control of stress response in *E. coli***. To illustrate the approach, we used MACS to analyse fluctuations in the control of the master regulator of stress response in *E. coli*—the alternative sigma factor RpoS (also known as σS or s38). In the presence of various stress factors such as oxidative stress, low pH, high osmolarity or nutrient limitations, RpoS replaces its vegetative counterpart RpoD to regulate the transcriptional program of *E. coli* by redirecting RNA polymerase to transcribe ~500 genes (Fig. 3). To prevent this from occurring under non-stressful conditions, RpoS is delivered to the ClpXP protease by the adapter protein SprE, rendering RpoS one of the most short-lived proteins in *E. coli* during exponential growth. SprE has been reported to be rate-limiting for RpoS degradation in exponential phase but levels are so low as to be almost undetectable using either western blots or standard fluorescence imaging. Although the activity of SprE is regulated

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**Figure 3 | Using MACS for studying the general stress response in *E. coli***. In the presence of stress, RNAP preferentially binds to RpoS (σS or σ38) instead of RpoD (σ70) to direct the transcriptional program towards the expression of stress-response genes. SprE (RssB) is an adapter protein, which is involved in controlling the RpoS levels by binding to RpoS, and delivering it to the ClpXP protease for ATP-dependent degradation. (a) Cell-length dependence of the RpoS-Venus strain measured along the growth curve suggest that the balanced growth (where the average cell length remains steady) is sustained for a very limited period (OD600 ≤ 0.07). Grey-shaded area is the s.d., and black line is smoothing via moving-window average. (b) RpoS-Venus intensity distribution for balanced growth. Brightest cells display only twice as much intensity compared with that of the auto-fluorescence (inset). (c) Absolute number distribution of SprE in balanced growth (extreme cell sizes were excluded). Inset shows the normalized histogram (black) overlaid with a Poisson distribution of the same mean (red). (d) SprE counting at various OD600 allows monitoring (i) number of SprE molecules per cell, (ii) number of SprE molecules per cell normalized by the cell area as a metric for SprE concentration, (iii) fraction of cells with zero SprE molecules and (iv) Fano factor (which is equal to σp^2/µp, where σp is the s.d. and µp is the mean of SprE number distributions) along the growth curve of *E. coli*. Fano factor after length conditioning appears to deviate from that of Poisson distribution (Fano factor = 1) after OD600 ~ 1.2 most likely due to undercounting (Supplementary Fig. 13). Inset shows coefficient of variation (CV = σp/µp). Grey-shaded areas represent the s.d., and blue lines are line segments connecting data points.
in various ways, this raises the question of whether heterogeneity in the abundance of SprE could create heterogeneity in RpoS. Specifically, if SprE levels are low, a substantial fraction of cells may contain zero SprE copies—particularly if production is burst-like as reported for many other proteins. Because RpoS is so short-lived and its levels are sensitive to SprE (Supplementary Fig. 11), it could quickly accumulate during those time windows and ensure that a subpopulation of cells is ready for the stress before it occurs.

To test whether RpoS is randomly activated in the absence of external cues, we considered the balanced growth regime where cells have fully adjusted to the growth conditions but have not yet started affecting each other by depleting available nutrients or accumulating metabolic waste. In keeping with previously reported results, we observed that this regime is only sustained at extremely low cell density, before OD_{600} can be reliably measured (Fig. 3a). As demonstrated above, MACS can be used to image on the order of >10^3 cells per hour in dense culture. Here we instead exploited the automation and ability to perform imaging under native conditions to image about 10^3 cells per hour in the very dilute balanced growth regime without concentrating cells, since the latter could trigger an RpoS response. We measured SprE levels in a strain where a truncated version of RpoS fused with a rapidly maturing YFP (RpoS750-Venus) is present as an additional chromosomal copy. The RpoS750 truncation has been widely used to report protein abundances, unless the low protein abundances are caused by lower mRNA numbers, which is not the case here because sprE is strongly transcribed.

In balanced growth we did not observe a single cell with zero SprE-mNeonGreen molecules, and based on the apparent Poisson statistics the probability of such events should be on the order of 10^-4 (or less given that a few immature FPs are missed). Because SprE-mNeonGreen is not substantially degraded, the results further suggest that cells produce ∼4–8 molecules per average cell cycle, while the Poisson statistics and absence of bursts suggest that those production events are effectively independent. Thus the few cells that temporarily have zero SprE molecules should only remain in that state for a few minutes on average, providing little time to boost RpoS levels.

We next measured SprE-mNeonGreen levels at various OD_{600} (Fig. 3d) to decipher SprE dynamics along the growth curve. Starting at an observed average of 8.3 ± 3.2 copies per cell (± s.d., n = 458 cells), SprE-mNeonGreen levels went through a minimum of 3.3 ± 1.9 copies per cell (± s.d., n = 2928 cells) in mid-exponential phase (OD_{600} ∼ 1.2), and then went up again to 6.7 ± 2.7 copies per cell (± s.d., n = 4326 cells) in early-stationary phase (OD_{600} ∼ 1.9). As discussed earlier, the values in early exponential phase are likely to be slightly underestimated due to incomplete FP maturation, but this problem decreases with increased cell division time. The reported dip, which has not been previously observed because even average SprE levels were experimentally unobservable, should thus be slightly more pronounced when correcting for mNeonGreen maturation time. Because the cells become substantially smaller during this interval, the SprE concentration increases about five-fold (Fig. 3d). The mechanisms causing the dip are unknown, but could reflect the fact that SprE is expressed from two promoters or that competition for the gene expression machinery is reduced after ribosomal genes are no longer expressed.

After conditioning on cell size (Supplementary Fig. 12), the distributions were again close to Poisson for most of the growth curve (Fig. 3d). In late exponential phase, the raw distributions appear even narrower than Poisson. This could in principle be explained by the fact that RpoS and SprE are thought to form a negative feedback loop in mid-exponential phase. However, a more likely explanation is that the method approaches its limits since the cell size of the MC4100 strain shrinks greatly in late exponential phase. The diffraction-limited spots then inevitably overlap and the observed distributions are in fact expected if the actual distribution is Poisson (Supplementary Fig. 13). However, spot overlap has a marginal impact on the average abundance and on the left tail of the distribution, which is particularly interesting in this context. We observe virtually no cells with zero SprE copies, except close to the minimum average abundance at OD_{600} ∼ 1.2, where this fraction reaches as high as a few percent (Fig. 3d). However, even in that regime, the RpoS distribution does not show any substantial outliers (Supplementary Fig. 14), perhaps because the RpoS half-life then is much longer making RpoS much less responsive to brief periods with zero SprE molecules. Thus despite the fact that bet-hedging has been suggested for RpoS, and the dedicated adapter protein SprE necessary for its degradation is present in such extremely low numbers, we see no evidence for bet-hedging at the frequencies we can measure. Instead we observe that, given the low abundances, the SprE distribution is quite narrow and with less signs of bursts than observed for most other proteins in E. coli. We further used our set-up to show that SprE production ceases quickly on exit from stationary phase: the total numbers of SprE-mNeonGreen per cell remain virtually unchanged until cells become large enough to divide at which time they become diluted between multiple cells (Supplementary Fig. 15).
Discussion

We show here that applying pressure to cells in a controlled manner makes it possible to count low-abundance proteins. We speculate that the diffusional slowing-down reflects increased cellular crowding effects (Supplementary Note 2) where objects as small as individual proteins experience the cytoplasm as a glassy medium. Indeed the bacterial cytoplasm has been shown to display properties of a colloidal glass for molecules larger than 30 nm (ref. 42) under normal conditions, and chemically induced osmotic compression can lead to reduction of cytoplasmic diffusion43 and hindrance of intracellular signalling due to overcrowding44. However, regardless of the underlying physical explanation, the insignificant loss of fluorescence on compression as opposed to chemical fixation (Supplementary Fig. 16) appears somewhat blurred due to their constant movement within the exposure time. Scale bar (white), 20 μm. (B) RFP-expressing cells, spiked in with a dilution factor of 1:100,000 (redgreen), could be captured within the field of view typically in ~3 min. After the red cell was immobilized (circled), snapshots in RFP and GFP fluorescence channels were taken (shown here as overlaid). Scale bar (white), 5 μm. (C) Minor modification of MACS enables cell retrieval (Methods, Supplementary Movie 10). (D) Bright-field and RFP fluorescence images are overlaid to show the captured cell of interest (circled) within the trapped volume, which is outlined by the red dashed line (control valve is open, valves 1–4 are closed). Scale bar (white) is 40 μm. (E) When the trapped volume was collected, grown overnight and imaged on the agar pad; the RFP-expressing cells were enriched. Counting red versus green cells suggested an enrichment factor of 10^2 (n = 4 runs). At low cell densities this allows for the immediate retrieval of cells, and at high densities a second round is necessary to achieve 100% purity. Scale bar (white), 2 μm.

Methods

Chip fabrication and MACS properties. MACS chips were produced via soft lithography using PDMS. The base (part A) and the curing agent (part B) of a two-part silicone elastomer kit (Sylgard 184, Dow Corning) were mixed at particular ratios (part A:part B) in weight to produce PDMS. The master mould for the flow channel was produced by spin-coating positive photoreist (PR) (AZ1010 (AZ, Electronic Materials) to a height of 10 μm on a silicon wafer. After ultra violet patterning the PR using a transparency mask (Output city), the wafer was heated for rounding the features to achieve dome-shaped channels. After rounding, the channel height becomes 8 μm. The wafer was then baked on a hotplate overnight to stabilize the positive PR. The control layer master was made by spin-coating the negative PR SU-8 2025 (Microchem) to yield a height of 25 μm, and ultra violet patterning it using a transparency mask defining the channels. To produce the soft MACS chip, 20% PDMS was spin-coated on the flow channel master at 1,250 r.p.m. for 45 s to yield an ~65-μm-thick PDMS membrane. For this condition, the minimum pressure required for closing the valve is ~5 p.s.i. For even gentler handling of cells, if required, a thinner membrane can be made to achieve valve closing at lower pressures. For control channels, PDMS with a 5:1 ratio was poured onto the control layer master. After both masters were partially cured at 65 °C for 30 min, they were aligned and cured for another 6 h at 65 °C to achieve thermal bonding. Finally, the two-layer PDMS chip was plasma-bonded permanently against glass coverslips. Since the freshly bonded chips did not work due to altered surface properties following plasma treatment, they were kept at room temperature for at least 1 day to regain the native surface properties. For single-molecule counting experiments, the chips were kept at the 65 °C for a total of 3 days after cover glass bonding since ‘cytoplasmic slowing-down’ works better with stiffer PDMS.

Closing properties of the valve depend on multiple parameters48. To achieve the half-open state for the 200-μm-wide control and flow channels, we typically used ~20 p.s.i., both for P_{valve} and P_{flow}, though different combinations of P_{valve} and P_{flow} also work. Compared with the full foot-print of the valve (200 μm x 200 μm), cell trapping happens within a subregion (approximately, 100 μm x 50 μm), which can be varied by slight modifications of P_{valve} and P_{flow}. The number of cells captured per field-of-view (FOV) also depends on the relative values of P_{valve} and P_{flow} as well as durations of the valve states. Since Quake valves can be actuated millions of times without signs of fatigue22, the bottleneck for long-term stability of MACS is the accumulation of debris within the valving intersection. This is problematic only during actuation of the valve and the presence of sample flow: debris does not get stuck permanently unless pressed against the surface during valve actuation and eventually gets washed away otherwise. Therefore, the intersections that remain passive do not collect debris and a neighbouring intersection can be used on demand if the actively used intersection becomes clogged. To minimize debris, we filtered all buffers and media using 0.22-μm-pore-size filters (Corning). Cells were grown in plastic tubes (BD Falcon, round-bottom) instead of glass vials to prevent crumbled glass. Sonicating the PDMS chips in isopropanol for 30 min, followed by 4 h of drying at 65 °C before bonding them to the cover glass removes PDMS crumbs that form at the inlets during hole punching45. In addition, prior to using the chips, flow channels were extensively rinsed with PBSA buffer (1 x PBS with 4 mg/ml BSA) to wash away debris that was stuck on the walls of the chip, as well as for passivating the channel surfaces to minimize cell sticking. We were able to keep the same chip on the microscope and use it for multiple days, everyday using a fresh flow channel.

Figure 4 | MACS allows for rapid screening and recovery/enrichment of rare cells. (a) Single frame of a movie (Methods, Supplementary Movie 9) of GFP-expressing E. coli cells flowing during the half-open state. Cells appear somewhat blurred due to their constant movement within the exposure time. Scale bar (white), 20 μm. (b) RFP-expressing cells, spiked in with a dilution factor of 1:100,000 (red-green), could be captured within the field of view typically in ~3 min. After the red cell was immobilized (circled), snapshots in RFP and GFP fluorescence channels were taken (shown here as overlaid). Scale bar (white), 5 μm. (c) Minor modification of MACS enables cell retrieval (Methods, Supplementary Movie 10). (d) Bright-field and RFP fluorescence images are overlaid to show the captured cell of interest (circled) within the trapped volume, which is outlined by the red dashed line (control valve is open, valves 1–4 are closed). Scale bar (white) is 40 μm. (e) When the trapped volume was collected, grown overnight and imaged on the agar pad, the RFP-expressing cells were enriched. Counting red versus green cells suggested an enrichment factor of 10^2 (n = 4 runs). At low cell densities this allows for the immediate retrieval of cells, and at high densities a second round is necessary to achieve 100% purity. Scale bar (white), 2 μm.
Microscopy. Epi-fluorescence and HILO (highly inclined and laminated optical sheet) microscopy was carried out similar to what was described previously15. In brief, images were obtained using an Electronic Trinocular (ECM CCD camera (EMCCD), iXon3-897, Andor), and OPLS lasers (Coherent) were used for HILO and FRAP measurements. The EM gain was set to 300 for single-molecule imaging. The HILO angle was adjusted using a custom-built stepper motor system. A manual flip mirror (Newfocus) allowed for switching between two different laser paths set-up for HILO and FRAP mode, respectively. For HILO experiments, a full mirror was replaced with an 80:20 beam-splitter to allow for easy switching between a focused laser pulse and epi-fluorescence illumination. Applying a 100-ms photo-bleaching laser pulse focused on one pole of an E. coli cell using a mechanical shutter (Uniblitz), the fluorescence recovery was monitored via epi-fluorescence imaging. The microscope was controlled by Micro-Manager (http://www.micro-manager.org/) and custom-written MATLAB scripts. Fluorescence imaging was performed with an LED system (SOLA light engine, Lumenkor) and appropriate filter cubes (Semrock): for cyan fluorescence, CFP-2432A; green fluorescence, GFP-3035B; yellow fluorescence, YFP-2427A; and red fluorescence, mcCherry-A. Unless otherwise stated, we used a ×100 TIRF objective (Nikon, TIRF; numerical aperture = 1.45) in combination with a ×2.5 relay lens in front of the EMCCD camera. At the expense of smaller FOV (hence lower throughput), this magnification provides a near-optimal effective pixel size (64 nm) to resolve spots and segment cell boundaries. Solid-state lasers (Coherent) in combination with proper filters (Semrock) are used for detecting single molecules (dichroic: D01-R488, emission filters: FF01-550/88 and LP02-514RS, laser: Genesis MX 514-1000 STM OPS/Laser-Diode System for mNeonGreen; dichroic: D01-R532, emission filters: FF01-607/70 and LP03-532RS, laser: Genesis MX 532-1000 STM OPS/Laser-System for mEos2).

Construction of E. coli strains. Strain construction is described in Supplementary Note 4. All E. coli strains, plasmids and primers used are listed in Supplementary Tables 2–4, respectively.

Sprl and RpoS measurements. Overnight cultures were grown in 1 × M9 salts supplemented with 10% (v/v) LB (M9 + 10% LB), and kept at stationary phase for a defined duration (~16 h). After diluting the overnight cultures typically in 30 ml of M9 + 10% LB by a factor of 10³, we diluted the 30-ml culture into 10 falcon tubes of 3 ml each and grew cultures at 37°C with shaking at 200 rpm. We used separate tubes for different time-points along the growth curve to ensure constant growth conditions for each sampling to ensure reproducibility since the RpoS levels are sensitive to aeration. Once the cells were manually transferred from the shaking flask into an airtight pressure tube (Supplementary Fig. 17), we took images for 15–30 min (TShaking = 25 min) in an automated fashion. At the end of each set of data acquisitions, we were able to clear the system off the cells completely. Ensuring that there is no carryover between different samples is vital since any cell that sticks around from earlier samplings may encounter stress, which is particularly important for studying the components of the stress response (namely, RpoS and Sprl) (Supplementary Note 5). To gather reproducible statistics, we pooled data that sticks around from earlier samplings may encounter stress, which is particularly important for studying the components of the stress response (namely, RpoS and Sprl) (Supplementary Note 5). To gather reproducible statistics, we pooled data.

Numerical simulations for single-molecule counting. A computer simulation was developed to assess the degree of potential protein undercounting due to apparent spatial clustering of molecules caused by the ‘large’ point-spread function (that is, a typical full-width at half-maximum is ~250 nm) and the relative small size of the bacterial cell (2–6 μm). Different cell sizes and geometries were compared. In short, N molecules were randomly placed in a virtual bacterial cell and the number of spatially resolved molecules and non-resolved clusters of molecules were calculated using Euclidian geometry and a spatial resolution of 250 nm (Supplementary Note 2 and Supplementary Figure 19). The computer code was developed and executed in MATLAB, which is available on request.

Spot-finding analysis and simulation of EMCCD images. Determining the number of spots in single cells was achieved in two steps. First, a spot-finding software was used to detect single molecules in the entire FOV. The next step involved specifying those spots to specific cells. Therefore, we used fluorescence images of a cytoplasmic CFP as a segmentation marker to obtain an outline of the individual cells such that spots could be assigned to each cell according to their x-y coordinates. The software used for spot-finding was modified based on a previously published single-particle tracking software12,13. To summarize, each image was first computationally filtered prior to spot localization using a band-pass filter to remove high-frequency noise and low-frequency features like cellular autofluorescence signal. This process results in a smooth zero-background based image. We detected local maxima with pixel level accuracy in the image via a user-defined intensity threshold. Sub-pixel localization of the spots was then estimated from the centroids of the spots calculated using a 7 × 7 pixel square centred on the local maxima. The box size, intensity threshold, and parameters for the band-pass filter were empirically optimized to minimize false positives and false negatives in the spot detection.

As described in the main text, the small confinement volume of the E. coli cytoplasm imposes severe limitations for counting performance using a diffusion-limited imaging system. Since the capabilities of MACS or agar pad-based imaging may also depend on the image analysis software used to detect the molecules, we have therefore quantified the limits of the counting performance using simulated EMCCD images that closely mimic our actual microscopy data in addition to the numerical simulations (Supplementary Note 3).

Capturing rare phenotypes and their retrieval with MACS. For the spiking-in experiments, overnight cultures of GFP- and RFP-expressing strains inoculated from fresh bacterial re-streaks on plates were used. We found that cells grown from older-than-a-week plates display higher tendency to stick to the PDMS chips surfaces. After mixing the RFP-expressing cells with GFP-expressing cells in the pressure tube using the dilution factor of 10³, cells were sent through the MACS chip in the half-open valve state. Detection of the RFP-expressing cell-of-interest was achieved manually. Two inlets and two outlets on the modified design can all be controlled via on-chip valves (1–4), and allow for cell collection. Screening is carried out while valves 3 and 4 are closed, and valves 1 and 2 are open. When a cell-of-interest is captured within the FOV, cell flow is stopped. After taking detailed images, valves 1–4 are closed, and the control valve is opened to release the pressure on the cells. Subsequently, the trapped volume is sent out to collection by opening valves 3 and 4, and flowing in oil. Using an oil phase for cell collection provides precise control of the volume that is retrieved. To facilitate the collection using the oil phase, chips were treated with a commercial water repellent (Aquapel) after plasma bonding and kept at room temperature until use.

Data availability. The data that support the findings of this study are available from the corresponding authors on request.

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Corrigendum: Mechanical slowing-down of cytoplasmic diffusion allows in vivo counting of proteins in individual cells

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The original version of this Article contained an error in the spelling of the author Somenath Bakshi, which was incorrectly given as Somenath Bakhsi. This has now been corrected in both the PDF and HTML versions of the Article.