Title

Vast heterogeneity in cytoplasmic diffusion rates revealed by nanorheology and Doppelgänger simulations

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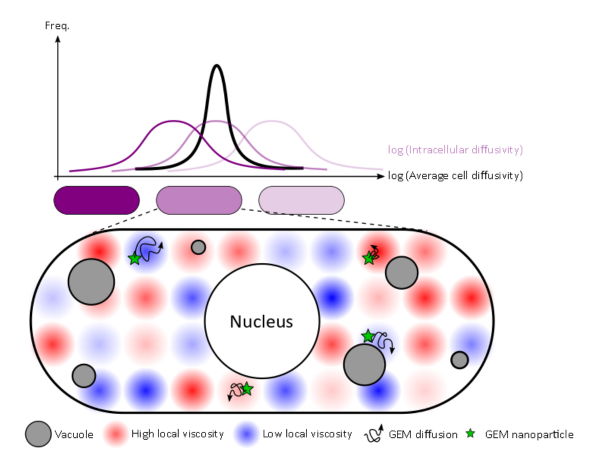
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Graphical abstract



Abstract

The cytoplasm is a complex, crowded, actively-driven environment whose biophysical characteristics modulate critical cellular processes such as cytoskeletal dynamics, phase separation, and stem-cell fate. Little is known about the variance in these cytoplasmic properties. Here, we employed particle-tracking nano-rheology on genetically encoded multimeric 40-nm nanoparticles (GEMs) to measure diffusion within the cytoplasm of the fission yeast Schizosaccharomyces pombe. We found that the apparent diffusion coefficients of individual GEM particles varied over a 400-fold range, while the average particle diffusivity for each individual cell spanned a 10-fold range. To determine the origin of this heterogeneity, we developed a Doppelgänger Simulation approach that uses stochastic simulations of GEM diffusion that replicate the experimental statistics on a particle-by-particle basis, such that each experimental track and cell had a one-to-one correspondence with their simulated counterpart. These simulations showed that the large intra- and inter-cellular variations in diffusivity could not be explained by experimental variability but could only be reproduced with stochastic models that assume an equally wide intra- and inter-cellular variation in cytoplasmic viscosity. To probe the origin of this variation, we found that the variance in GEM diffusivity was largely independent of factors such as temperature, cytoskeletal effects, cell cycle stage and spatial locations, but was magnified by hyperosmotic shocks. Taken together, our results provide a striking demonstration that the cytoplasm is not "well-mixed" but represents a highly heterogeneous environment in which subcellular components at the 40-nm size-scale experience dramatically different effective viscosities within an individual cell, as well as in different cells in the population. These findings carry significant implications for the origins and regulation of biological noise at cellular and subcellular levels.

Main Text

Introduction

Life at the molecular scale is stochastic, with macromolecules continually being jostled by Brownian motion. This emergence of "biological noise" at the molecular level permeates all aspects of cell biology, inducing stochastic fluctuations in subcellular processes and driving natural variation among cells in a population. Critical roles for biological noise in signaling (Huang et al., 2016), cell size control (Facchetti et al., 2019; Patterson et al., 2019; Schmoller, 2017), organelle size scaling (Chang and Marshall, 2017; Gray et al., 2019; Mohapatra et al., 2016), and gene expression (Battich et al., 2015; Oates, 2011; Raj and van Oudenaarden, 2008; Raser and O'Shea, 2005). In general, biological noise presents a challenge to cellular homeostasis and signaling mechanisms, and is often suppressed in order for biological functions to be robust. For example, signaling frequently depends on strong amplification of initially weak signals, which can erroneously amplify noise unless proofreading mechanisms are in place (Huang et al., 2016). However, biological noise can also confer a selective advantage. In a fluctuating and unpredictable environment, biological variation between cells in an isogenic population can ensure population-level survival (Levien et al., 2021; Suderman et al., 2017).

One potentially significant source of biological noise that has been largely ignored is that of heterogeneity in the cell cytoplasm. The cytoplasm is composed of a highly diverse and actively-mixed assembly of resident macromolecules of various size (Milo and Phillips, 2015; Requião et al., 2017), charge (Requião et al., 2017), and hydrophobicity (White and Jacobs, 1990). The complexity of the cytoplasmic milieu could influence molecules' behavior locally. Indeed, spatiotemporal heterogeneity in the diffusion of particles has been observed in multiple contexts such as *E. coli*, fungi, mammalian cells and even *Xenopus* egg extract using methods ranging from fluorescence correlation spectroscopy (FCS) to particle tracking (Bakshi et al., 2011; Baum et al., 2014; Dross et al., 2009; Huang et al., 2021; Manley et al., 2008; McLaughlin et al., 2020; Scipioni et al., 2018; Xiang et al., 2020).

Heterogeneity of cytoplasmic properties have potentially far-reaching effects in cell biology, as the cytoplasm hosts a wide variety of critical molecular processes ranging from protein synthesis and turnover, to cytoskeletal transport and force production, metabolism, and beyond (Cadart et al., 2019; Neurohr and Amon, 2020). Further, changes to physical cytoplasmic properties such as the macromolecular density, viscosity, and degree of crowding have been shown to impart widespread effects within the cell—including sudden and significant impacts on growth and viability (Knapp et al., 2019; Tsai et al., 2019). For example, altering cytoplasmic crowding by changing the concentration of ribosomes has strong effects on phase separation (Delarue et al., 2018), and high osmotic shocks can completely halt microtubule dynamics (Molines et al., 2022). Additionally, alterations in cytoplasmic density have been implicated in cellular aging and senescence (Neurohr et al., 2019) and differentiation (Guo et al., 2017).

Here we establish a combined experimental and computational approach to examine cytoplasmic heterogeneity through the lens of diffusion. Single particle motion-tracking allows for a robust quantification and statistical analysis of particle behavior, revealing variations between particles which would otherwise be averaged out in bulk measurements obtained in photobleaching (e.g. FRAP) and FCS (Charras et al., 2009; Neurohr and Amon, 2020) experiments. Further, this kind of "passive" rheology approach requires minimal perturbations to the cell.

Previously, particle tracking rheology on fluorescent proteins has proven difficult due to their fast diffusion rates and tendency to photobleach. The development of GEMs (Genetically Encoded Multimeric nanoparticles)(Delarue et al., 2018)), has enabled large improvements on this front. These

bright and photostable protein spheres are expressed as fluorescently-tagged monomers which self-assemble into hollow shells of nearly-uniform size and shape (Delarue et al., 2018; Szórádi et al., 2021). Because each particle contains tens of fluorescent proteins, they can be tracked for relatively long periods of time without photobleaching. Additionally, the near-diffusive movements of GEMs suggest they do not interact strongly with eukaryotic cellular components, making them ideal reagents for rheological studies (Delarue et al., 2018; McLaughlin et al., 2020). Critically, their relatively large size and slow diffusion rates - comparable to large protein complexes such as ribosomes - allow GEMs to be tracked using modern high-speed cameras (which is still not attainable for individual fluorescent proteins). Initial studies have established their utility in quantitatively probing diffusion and crowing in the cytoplasm and nucleoplasm in various cell types including yeast and mammalian cells (Alric et al., 2021; Carlini et al., 2020; Delarue et al., 2018; Lemière et al., 2021; McLaughlin et al., 2020; Molines et al., 2022; Szórádi et al., 2021).

The fission yeast *Schizosaccharomyces pombe* provides an excellent model system for the study of cytoplasmic heterogeneity because of their uniformity in many other aspects of their cell biology. In standard laboratory conditions, these rod-shaped cells exhibit very tight distributions in their cell size at division ($CV \sim 6\%$;(Facchetti et al., 2019; Patterson et al., 2019)) and cell shape (Abenza et al., 2015; Davì et al., 2018; Saunders et al., 2012), as well as cell cycle progression and intracellular density ($CV \sim 10\%$;(Odermatt et al., 2021)). The relatively low phenotypic variability within and between fission yeast cells permits the study of cytoplasmic heterogeneity in a well-controlled system in the presence of minimal confounding factors.

Using live cell high-speed imaging and quantitative tracking of 40 nm-diameter GEMs in *Schizosaccharomyces pombe*, we measured cytoplasmic diffusivity for thousands of individual particles. These data revealed large heterogeneity in diffusion coefficients both within single cells as well as between cells in the population. To analyze this variability, we developed an automated pipeline, which we call the Doppelgänger Simulation approach, to reproduce our experimental results computationally using simulations of diffusion, and assay heterogeneity using statistical techniques for analysis of variance. Using these methods, we showed that orders of magnitude of variability in GEM cytoplasmic diffusivity within and between cells arose from an equally wide distribution of cytoplasmic viscosity. This variance was not affected by temperature, the cytoskeleton, or cell size, but was increased by hyperosmotic shock. Our studies support a growing body of evidence that the cytoplasm is not physically well-mixed (Bakshi et al., 2011; Baum et al., 2014; Dross et al., 2009; Manley et al., 2008; McLaughlin et al., 2020; Weiss and Korge, 2001; Xiang et al., 2020) and reveal this heterogeneity in diffusion as an important potential source of biological noise.

Results

Statistical characterization of cytoplasmic GEM particle diffusion in fission yeast

To assay cytoplasmic diffusion in fission yeast, we expressed 40 nm-diameter GEM nanoparticles in wildtype *S. pombe* from a multicopy plasmid on an inducible promoter (Molines et al., 2022). Tuning the expression of the GEMs construct allowed us to titrate particle formation to small number of particles (<10) per cell. To reduce environmental variability, these unicellular cells were grown at 30 °C under optimal conditions in shaking liquid cultures to exponential growth phase and mounted in imaging chambers with fixed dimensions under constant temperature and imaged acutely. Using variable angle epifluorescence microscopy (VAEM) (Delarue et al., 2018; Konopka and Bednarek, 2008; Lemière et al., 2021; Molines et al., 2022) we tracked GEM particle motion at 100 Hz for 10 s, as described previously (Delarue et al., 2018; Lemière et al., 2021; Molines et al., 2022) (Fig. 1a-b, *Methods*). Each field of view (FOV) contained multiple cells that were individualized post-acquisition. Images were manually curated to eliminate from the data set a small

subset of cells that had died, exhibited grossly abnormal morphologies, or contained a single bright aggregate of GEM particles. From a dataset of 145 cells, 3681 tracks were analyzed, with an average of 25 ± 10 (AVG \pm SD) tracks per cell, a mean step size of 104 ± 72 nm (AVG \pm SD), and a mean track length of 273 ± 268 ms (AVG \pm SD) (Fig. 1c).

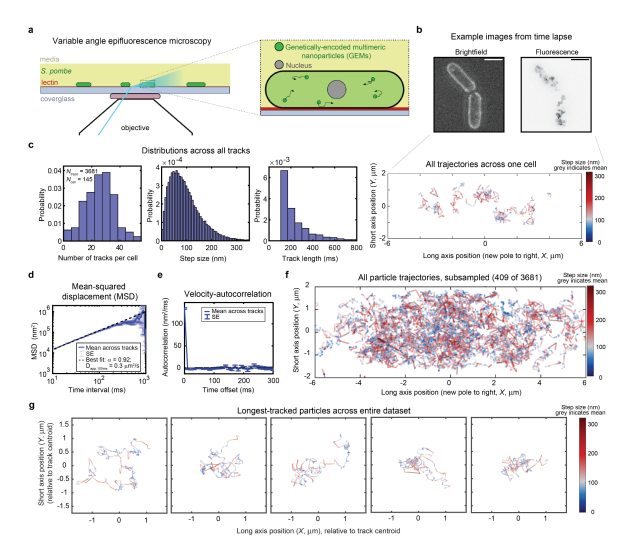


Figure 1: High-speed particle-tracking nano-rheology of GEMs allows detailed statistical analysis of cytoplasmic diffusion. (a) Schematic of the experimental imaging set-up. (b) Example brightfield image (top left) and maximum intensity projection through time of the GEM particle fluorescence (top right) for one representative field of view, alongside the measured nanoparticle trajectories (bottom) for the upper cell in the image. Trajectories are colored by the step size of the particle in nanometers between each time frame of the movie. Grav indicates the mean step size across all tracks in the dataset. Scale bar is 5 µm. (c) Histograms of the number of tracks per cell (left), the step-size for all time-points (middle), and the duration of time that each particle was tracked (right). Note that tracks shorter than 10 time-points were not included in the analysis. (d) The mean-squared displacement (MSD) of the particle tracks. The time-averaged MSD was first calculated individually for each track, and then a second averaging was performed to find the (ensemble averaged) MSD across all tracks. Note the logarithmic scale along the x- and y-axes. (e) The average velocity autocorrelation across all article tracks. Averaging was performed in the same order as the MSD. (d-e) Error bars represent the standard error. (f) Plots of particle trajectories drawn from many experiments and cells, randomly subsampled for better visibility of individual particle behaviors. Subsampled trajectories include at least one track from 141 of the 145 cells in the dataset. Gray indicates the mean step size across all tracks in the dataset. (c-f) Dataset includes 3681 tracks among 145 cells, recorded from 5 different samples and over 3 different days. (g) Individual trajectory plots for five of the longest-tracked particles (in time), excluding stationary particles. Color scaling of the step size was identical in all panels included in f-g (using the mean and standard deviation of the step size across the entire dataset).

From these trajectories we computed the time-averaged, ensemble-averaged (i.e., track-averaged) Mean Squared Displacement (MSD) as a function of time offset, and fitted the resulting MSD curve to a power law (Fig. 1d, Methods). MSD analysis showed that GEM particle motion in the cytoplasm was largely diffusive (MSD \approx D τ), following a robust power law with apparent diffusivity $D_{app,100ms}$ = 0.3 \pm 2*10 $^{\!\!\!\!\!-3}$ $\mu m^2/sec$ (AVG \pm CI) and anomalous diffusion exponent α = $0.92 \pm 5*10^{-3}$ (AVG \pm CI), which were similar to previously published measurements in fission yeast (Lemière et al., 2021; Molines et al., 2022). The diffusivity of the 40 nm GEMs in the cytoplasm was roughly 40 times slower than the theoretical prediction for simple Stokes-Einstein diffusion in water -and corresponded to the particle's expected diffusion rate in a 75% glycerol solution in water. We note that diffusion along the long and short axes of the cell were comparable by our measurements (Fig. S1a), and we found that the MSD plots do not plateau, indicating that diffusion of the GEMs was not confined on timescales less than a second (e.g., most particles do not run into the cell wall within the measured time window). The time-averaged, ensemble-averaged (i.e., track-averaged) velocity autocorrelation of particle trajectories was also consistent with simple unconstrained diffusion (Fig. 1e). Notably, the autocorrelation plot lacked the characteristic negative peak associated with subdiffusive motion and viscoelastic response seen in other systems (Fig. S1b-c) (Guigas et al., 2007; Tolić-Nørrelykke et al., 2004; Weber et al., 2010, 2012a). Therefore, at least with this approach at this 40-nm size scale, we detected no elastic response in the yeast cytoplasm.

Cytoplasmic diffusivity spans orders of magnitude

We next analyzed individual particle tracks, which revealed a rich phenotypic variability (Fig. 1f-g) that was obscured by the ensemble averaging-based analysis described above (e.g., MSD - Fig. 1d). Notably, even within a single trace, individual particles exhibited large fluctuations in their step size (Fig. 1g). To investigate the variety of comportment displayed by individual particles, we calculated and fit the time-averaged MSD individually for each track (Fig. 2a) and the time-averaged, ensemble-averaged MSD over all tracks in each cell (Fig. 2b). These data showed that variability in particle motion ranged over orders of magnitude; fits of particle and cell MSDs (Fig. 2c-d) showed that diffusivity follows a long-tailed, log-scale distribution, consistent with Brownian motion in a heterogeneous environment (Santos et al., 2020; Ślęzak and Burov, 2021). The distribution of apparent diffusivities exhibited a single peak (Fig. 2c-f), which appeared more normally-distributed in log space (Fig. 2f) than in real space (Fig. 2e). Therefore, we performed all further statistics and visualization on the log₁₀ of the fitted apparent diffusivities. The median of the diffusivity distribution in log space, which we then converted to real space (see Methods), corresponded to a diffusivity of 0.29 µm²/sec for the track-wise distribution and 0.33 µm²/sec for the cell-wise distribution, both similar to the bulk estimate. The standard deviation of the diffusivity distribution in log space (representing the number of orders of magnitude spanned by the dataset) can be converted to real space as a fold-range at 2.5 standard deviations away from the median (see Methods), giving a 392-fold range across tracks and 11-fold range across cells. We chose 2.5σ as our cutoff as it gave a range consistent with our outlier estimation algorithm (Fig. 2c, see caption). Overall, we showed that diffusivities vary by over 2.5 orders of magnitude among individual GEMs and one order of magnitude among cells. Hereafter we use the terms *inter*cellular variation to refer to the spread of the cell-wise diffusivity (Fig. 2c, right) and intracellular variation to indicate the spread in the track-wise diffusivity (Fig. 2c, left).

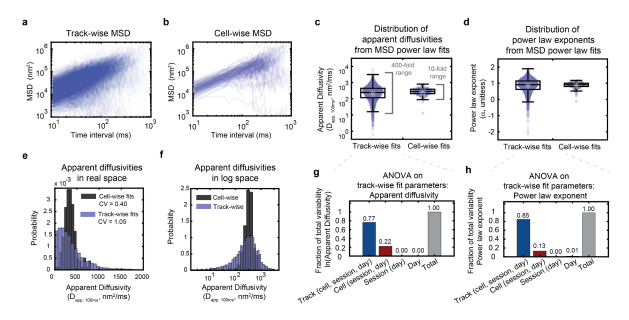


Figure 2: GEM diffusivity varies over 400-fold across tracks and 10-fold across cells. (a-b) Mean-squared displacements averaged either (a) by track (averaged over time for each track), or (b) by cell (averaged over time for each track and then averaged across all tracks in each cell). Note the logarithmic scale along the x- and y-axes. (c-d) Apparent diffusivities (c) and power law exponents (d) calculated from fits of the track-wise and cell-wise MSDs to a power law. Note the logarithmic scale along the y-axis. Boxplots: Central line, median; grey dot, mean; boxes, 25th and 75th percentiles; whiskers, furthest data point that is not an outlier; outliers, any point that is more than 1.5 times the interquartile-range past the 25th and 75th percentiles. (e-f) The same distributions of the fitted apparent diffusivities plotted in (c), now plotted as a histogram either in real space (e) or log space (f). Probabilities represent the probability density per histogram bin width, such that the sum of the bin heights multiplied by the bin width equals 1. (g-h) Results from a nested ANOVA performed on track-wise fits of diffusivities (g) and power law exponents (h). The amount of the experimentally-observed variance that can be explained by track-to-track, cell-to-cell, imaging session-to-session, and day-to-day variability is plotted as a fraction of the total variance. (a-h) Dataset is identical to that shown in Fig. 1 c-f, including 3681 tracks among 145 cells, recorded from 5 different samples and over 3 different days.

To understand whether variation arose from cell-to-cell variability, from different microenvironments within a single cell, or from experimental day-to-day variation, we performed an Analysis of Variance (ANOVA) on the track-wise diffusivity measurements (Fig. 2g). The ANOVA revealed that the vast majority (~80%) of the measured spread in diffusivity came from intracellular variation (i.e., *Track* in Fig. 2g), but there was also a significant amount of variance (~20%) explained by cell-to-cell variability (i.e., *Cell* in Fig. 2g). Only < 1% could be attributed to experiment-to-experiment variability. Similar results were observed for the fitted anomalous diffusion exponent (Fig. 2d, h), which was not surprising given the strong correlation between the fitted apparent diffusivities and power law exponents in our dataset (Supp. Fig. 2a).

Another common way to differentiate sources of noise in biological data is to separate the observed spread into intrinsic (uncorrelated within cells) and extrinsic (correlated within cells) components (Bauer et al., 2021; Raj and van Oudenaarden, 2008). Our ANOVA results suggested that noise in this system was almost entirely intrinsic, as $\sim 80\%$ of the variation was maintained after controlling for cell-to-cell variability. Indeed, by plotting the apparent diffusivities of random pairs of GEM particles, where each pair is randomly chosen from particles within a single cell, we found that the noise had only a very weak correlation between particles within the same cell (Spearman correlation: r = 0.21, $p = 5*10^{-35}$, Supp. Fig. 2b). We noted that the large intercellular and intracellular

variation observed in our data cannot be explained by differences in GEM particle expression levels, as the mean apparent diffusivity among track-wise diffusivity fits within a cell was not significantly correlated with the number of tracks in the cell (Spearman correlation: r = -0.003, p = 0.97, Supp. Fig. 2c), and the coefficient of variation among track-wise diffusivity values within a cell was only very weakly correlated with the number of tracks in the cell (Spearman correlation: r = 0.25, p = 0.004, Supp. Fig. 2d). In addition, the coefficient of variation of particle diffusivities within each cell was ~ 1 , and was uncorrelated with the mean particle diffusivity across all particles in the cell, consistent with Poisson statistics (Spearman correlation: r = -0.1, p = 0.15, Supp. Fig. 2e).

Stochastic simulations demonstrate that spread is not due solely to statistical measurement noise

As diffusion is an inherently stochastic process, we next explored whether the measured variation in particle mobility was due to statistical properties of our measurements. It is known, for example, that datasets with shorter track lengths will produce wider distributions of measured diffusivities (Weber et al., 2010). We therefore developed what we called the Doppelgänger Simulation (DS) approach, employing a custom algorithm to automatically read in and replicate the experimental measurement statistics *in silico* cell-for-cell and track-for-track (Fig. 3a). With DS, simulated cells have the exact same cell length and number of tracks as their experimentally-measured counterparts, and each simulated track has an identical length (i.e., number of time points tracked) to the associated experimental trajectory. This straightforward and powerful approach allowed us to produce simulated data that could be directly compared to the experimental tracks and analyzed using identical statistical methods.

Using DS, particle motion was then recapitulated using stochastic Brownian dynamics simulations of diffusion inside a box representing the exterior cell boundary (Fig. 3). We opted for a simple diffusion model because GEM particle motion is observed experimentally to be largely diffusive (Fig. 1d) and did not display characteristics of constrained or viscoelastic behavior (Fig. 1d-e). The model assumes an average cytoplasm viscosity forty times that of water, giving a mean diffusivity of 0.35 µm²/s that closely matches that of the experimental data. The simplest iteration of the model (Fig. 3b-f, Model #1: uniform viscosity), which we will hereafter refer to as the uniform viscosity model (due to its assumption of constant viscosity within and among cells), accounted for only a fraction of the experimentally-measured spread in GEM particle mobility (Fig. 3d-f) – including ~50% of the track-to-track variability in diffusivity, and <10% of the cell-to-cell variability, as measured by ANOVA (Fig. 3f). We therefore concluded that neither the stochastic nature of diffusion nor the statistical properties of our experimental measurement statistics were the major source of heterogeneity in GEM particle diffusion.

Simulations suggest that heterogeneity in diffusion must arise from an equally vast spread in cytoplasmic viscosity.

As the data set of experimentally-measured GEM particle motion fitted well to a model of simple diffusion, there were only a finite number of sources in this simple model from which heterogeneity in mobility could arise. The major parameter defining diffusion is the diffusivity, D, which theoretically (by the Stokes-Einstein equation) is simply equal to the ratio of the thermal energy, k_BT , to the viscous drag on the particle, γ . For a spherical particle, $\gamma = 6\pi\eta R$, where η is the viscosity of the cytoplasm and R is the radius of the particle (Einstein, 1905). Of these parameters, viscosity is the only parameter that could be varying within and between cells, as temperature is held constant and the radii of GEM nanoparticles have been shown by electron microscopy to be fairly uniform (CV \approx 0.1) when expressed in mammalian and budding yeast cells (Delarue et al., 2018).

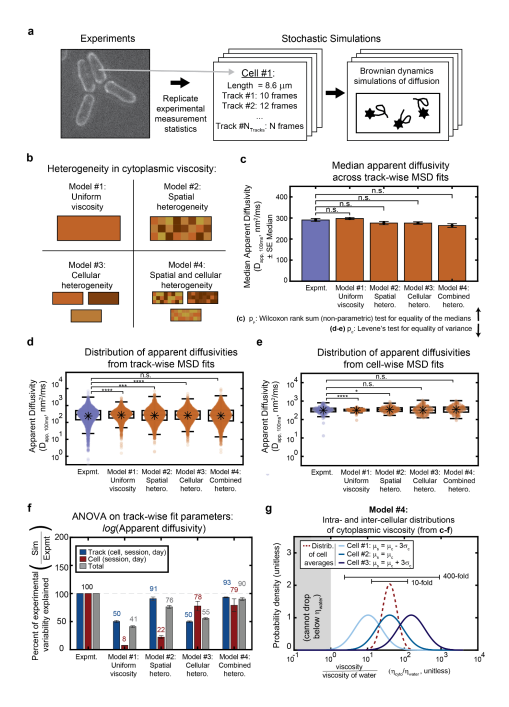


Figure 3: Stochastic simulations reveal both spatial and cellular heterogeneity in viscosity are required to reproduce experimentally observed variation. (a) Schematic of the Mirror Image Simulation approach. Each experimentally-measured cell and particle were reproduced one-to-one in the simulated dataset, with every simulated cell having the same long-axis length as its experimentally-measured counterpart, and each particle being tracked for the same amount of time. (b) Schematic demonstrating different types of heterogeneity in cytoplasmic viscosity included in each of the four models. (c) Median apparent diffusivity (averaged across all tracks) plotted for the experimental dataset as well as each model. Error bars represent the standard error of the median. Significance stars represent the result of the Wilcoxon rank sum test for equality of the medians. (d-e) Distributions of apparent diffusivities calculated from fits of the track-wise (d) or cell-wise (e) MSD curves displayed for the experimental data as well as each of the models. Note the logarithmic scale along the y-axis. Boxplots are drawn as in Figure 2. Significance stars represent the result of Levene's test for equality of variance. (c-e) * p<0.05. **p<0.01, *** p<0.001, **** p<0.0001. (f) Results from a nested ANOVA performed on track-wise fits of diffusivities (d). The percent of the experimentally-observed cell-to-cell and track-to-track variability that can be explained by each of the models. (g) The distribution of cytoplasmic viscosities, shown relative to the viscosity of water, needed to

most closely reproduce the experimental data (i.e., simulations from Model #4, using the same parameters used to generate (**c-f**)). Histograms are shown for the distribution of average cell viscosities (intercellular heterogeneity, red dashed line) and the distribution of intracellular viscosities for three example cells (blue lines of varying darkness). The examples include a cell whose average viscosity equals that of the cell-wide average (medium blue line), a cell with an average viscosity three standard deviations above the cell-wide average (dark blue line), and a cell with an average viscosity three standard deviations below the cell-wide average (light blue line). Note the logarithmic scale along the x-axis. The simulation did not allow viscosities below that of water.

We therefore generated three other versions of our model incorporating viscosity variation (Fig. 3b), while keeping the mean viscosity (and thus mean particle diffusivity) constant (Fig. 3c). In one version, which we refer to as the spatial heterogeneity model (Fig. 3, Model #2: spatial heterogeneity), we aimed to explore whether intracellular spatial variations in viscosity could account for the experimentally measured spread in diffusivity. A spatially varying viscosity was consistent with the observation that individual particles can display significant variations in step size within a single track (Fig. 1g). This model assumed that viscosity varies across the cell with a fixed domain size, approximated by a grid of discrete viscosity domains where each region was randomly assigned a distinct viscosity value. The average cellular viscosity was held constant. In another variation of the model, termed the cellular heterogeneity model, viscosity was uniform within each cell, but the uniform viscosity value varied between cells (Fig. 3, Model #3: cellular heterogeneity). Finally, we developed a fourth model combining both intracellular and intercellular heterogeneity, which we called the combined heterogeneity model (Fig. 3, Model #4: combined heterogeneity). In all three variations on the original model, viscosity values were chosen from a log-normal distribution, mimicking the distribution of the experimentally-measured step sizes (Fig. 1c, middle) and diffusivities (Fig. 2f).

We then ran each model multiple times to account for their stochastic nature and assayed whether each model could reproduce the experimentally-observed spread in diffusivity (1) as measured by ANOVA (Fig. 3f), and (2) such that the variance was not statistically significantly different from the experiments according to Levene's test for equality of variances (Fig. 3d-e)(Levene, 1960). While the spatial heterogeneity model could only account for the track-to-track variation in experimentally-measured diffusivity (but not the cell-to-cell variation), and the cellular heterogeneity model could only reproduce the cell-to-cell variation (but not the track-to-track variation), only the model combining both spatial and cellular heterogeneity could fully reproduce the amount of spread observed in the experimental data (Fig. 3d-f). Further, only a viscosity variation spanning orders of magnitude (Fig. 3g) could quantitatively recapitulate the experimentally-measured spread. In particular, the viscosity was required to vary 10-fold among cells, 100-fold within any individual cell, and 400-fold across the dataset in order to best match the experiments. Overall, our simulations showed that our data is best explained by a model in which effective viscosity experienced by cytoplasmic GEMs particles varies drastically within and between cells.

Heterogeneity in diffusion does not arise from density fluctuations related to the cell cycle or cell tip growth

We next tested what factors might be responsible for such a large heterogeneity in cytoplasmic viscosity. Within an asynchronous population, fission yeast cells exhibit an approximately two-fold range in cell size, which corresponds to the cell cycle stage (Mitchison, 1957). A recent study used quantitative phase imaging (QPI) to show that the overall intracellular dry-mass density of fission yeast cells fluctuates over the cell cycle, with density decreasing during interphase and increasing during mitosis and cytokinesis (Odermatt et al., 2021). To test whether GEM diffusion also varies over the cell cycle, we examined the relationship of GEM diffusion with

cell length as a proxy for cell cycle stage (Supp. Fig. 3a-b). We detected no significant correlation of diffusivity with cell length, making it unlikely that the cell to cell variability in GEM diffusion is cell cycle dependent.

We next tested whether spatial variations of density could explain the variability of GEM diffusion. QPI experiments demonstrated a subtle gradient of intracellular density in a subset of fission yeast cells, in which growing cell tips generally appear to be less dense than the rest of the cell (Odermatt et al., 2021). Regional cytoplasmic differences have also been shown in *Ashbya gossypii*, in which GEMs have decreased diffusivity in the perinuclear region (McLaughlin et al., 2020). To test for spatial variations in fission yeast, we mapped the GEM tracks relative to their positions in the cell (Supp. Fig. 3c-d). This analysis yielded no obvious regional differences in diffusivity within the fission yeast cell; specifically, we noted no strong differences in diffusion at growing cell tips or at the perinuclear regions. Therefore, it is unlikely that systematic regional differences in intracellular density are responsible for the variance in diffusivity.

Variance in diffusion is impacted by osmotic shock but not by cytoskeletal or temperature perturbations.

We then probed what factors could affect the variance by submitting the cells to different perturbations. For each perturbation, we measured the distribution of track-wise and cell-wise fits of GEM diffusivities, and performed the Wilcoxon rank sum non-parametric test for equality of medians (Mann and Whitney, 1947) and Levene's test for equality of variances (Levene, 1960) to establish whether changes to the median and variance were statistically significant (Methods).

One cytoplasmic constituent implicated in the rheological properties of the cytoplasm is the cytoskeleton. A rigid and interconnected cytoskeleton network can act as a barrier (Potma et al., 2001), or elastically resist particle motion – properties which can be described by poroelastic models (Charras et al., 2009; Moeendarbary et al., 2013). In addition, the cytoskeleton is responsible for transporting and positioning organelles and "actively mixes" the cytoplasm (Brangwynne et al., 2008). The cytoskeleton may also create structured intracellular regions with distinct biophysical properties (Fletcher and Mullins, 2010). We used a combination of latrunculin A (LatA) and methyl benzimidazol-2-yl-carbamate (MBC) to depolymerize actin and microtubules in interphase fission yeast cells (Fig. 4a). This treatment however had only subtle effects on GEM diffusivity; we detected a small, statistically insignificant increase in the median diffusivity (Fig. 4d, Wilcoxon t-test, Track-wise fits: 6% increase, p-value = 0.17; Cell-wise fits: 18% increase, p-value = 0.3), and a small, statistically insignificant increase in the variance (Fig. 4g,j, Levene test, Track-wise fits: 71% increase, p-value = 0.08; Cell-wise fits: 27% increase, p-value = 0.18)). We therefore concluded that the cytoskeleton is not the main determinant of cytoplasmic viscosity or variance at the 40-nm size scale in fission yeast.

Another main determinant of diffusivity D is temperature T. In addition to purely physical effects of temperature of diffusion as defined by the Stokes-Einstein equation where $D \propto T$. temperature can also have also a multitude of biological effects. For instance, temperature shifts may alter active mixing of the cytoplasm (Weber et al., 2012b), and trigger viscosity adaptation mechanisms via production of viscogens (Persson et al., 2020). There are also reports of regional differences in effective temperature within single cells (Chrétien et al., 2018; Hayashi et al., 2015; Okabe et al., 2012). To assay the effects of temperature on GEMs diffusivity, we grew fission yeast cells overnight at 30 °C, and then imaged them ~ 5 min after shifting cells down to 20 °C (Fig. 4b). This 10 °C decrease in temperature corresponds to $\approx 3\%$ decrease in the absolute temperature (in Kelvin), and thus the Stokes-Einstein equation predicted a similar decrease in diffusivity. We observed a slightly larger than predicted drop in the median track-wise diffusivity of GEMs (Fig. 4e, Wilcoxon

test, Track-wise fits: 11% decrease, p-value = 0.03; Cell-wise fits: 6% decrease, p-value = 0.54). The track-wise variance exhibited a statistically significant decrease, but

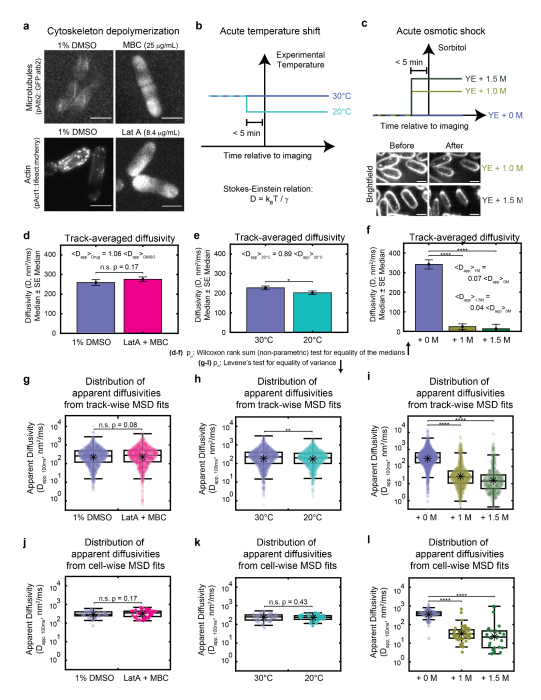


Figure 4: Heterogeneity in cytoplasmic diffusion has varied responsiveness to experimental perturbations. (a) Fluorescence images of fluorescent tubulin (top) and actin (bottom) in the context of the DMSO control (left) and addition of cytoskeleton depolymerizing drugs (right). Scale bar is 5 μ m. (b) Schematic of experiments varying the experimental temperature (top) and prediction of the relationships between the diffusivity, D, and the experimental temperature, T, as well as the Boltzmann constant, k_B , and the viscous drag coefficient, T (bottom). (c) Schematic of experiments varying osmotic shock with sorbitol (top) and example brightfield images of osmotically-shocked cells showing a reduction in cell volume (bottom). Scale bar is 5 μ m. (d-f) The median diffusivity is plotted for each experimental condition. Significance stars represent the result of the Wilcoxon rank sum test for equality of the medians. (g-l) Distributions of apparent diffusivities calculated from fits of the track-wise (g-i) or cell-wise (j-l) MSD curves displayed for each condition. Note the logarithmic scale along the y-axis. Boxplots are drawn as in

Figure 2. Significance stars represent the result of Levene's test for equality of variance. (**d-l**) * p<0.05. **p<0.01, *** p<0.001, **** p<0.0001.

the cell-wise variance did not change significantly (Levene test, Track-wise fits: 49% decrease, p-value = 0.006; Cell-wise fits: 28% decrease, p-value = 0.43). Overall, increasing the temperature had no effect on cell-to-cell variation but slightly increased intracellular heterogeneity.

Finally, we tested the effects of osmotic shocks. Osmotic shocks acutely alter the concentration of molecules in the cytoplasm by removal or addition of water (Knapp et al., 2019; Lemière et al., 2021; Molines et al., 2022). We performed hyperosmotic shocks with 1 M and 1.5 M sorbitol (which roughly doubles the concentration of the cytoplasm; Fig. 4c). Consistent with previous reports (Lemière et al., 2021; Molines et al., 2022), these hyper-osmotic shocks induced a striking decrease in the median track-averaged diffusivity of GEM particles compared to control experiments (Wilconox t-test, Track-wise fits: 93% and 96% decreases, p-values = $5*10^{-273}$, $3*10^{-153}$ for 1 M and 1.5 M shocks, respectively; Cell-wise fits: 92% and 94% decreases, p-values = $3*10^{-21}$, $4*10^{-10}$ for 1 M and 1.5 M, respectively). Interestingly, it also induced a sizable increase in both the track-wise and cell-wise variance in measured diffusivity (Fig. 4 i,l, Levene test, Track-wise fits: 275% and 420% increases, p-values = $1*10^{-9}$, $2*10^{-10}$ for 1 M and 1.5 M shocks, respectively; Cell-wise fits: 530% and 16,083% increases, p-values = $4*10^{-4}$, $3*10^{-9}$ for 1 M and 1.5 M, respectively). Thus we found that increasing the concentration of the cytoplasm slowed diffusion but also drastically increased both intracellular and intercellular cytoplasmic heterogeneity. These results suggest that hyperosmotic shocks may make the cytoplasm even more heterogeneous.

Discussion

Here we used a combined experimental and theoretical analysis to reveal a high degree of cytoplasmic heterogeneity experienced by objects on the scale of large protein complexes. In particular, our results indicated the effective cytoplasmic viscosity in fission yeast varies more than 10-fold among cells, and 100-fold within cells. Although the source of this heterogeneity is not yet understood, our analyses showed that viscosity variation is independent of the cytoskeleton, cell cycle stage, and temperature – but increases under hyperosmotic shock.

Generalizability of cytoplasmic heterogeneity

It is highly likely that the large diffusive heterogeneity we observed in fission yeast is generalizable to most, if not all, cell types. In fact, because fission yeast exhibit strikingly regular cell shape and growth properties, they may be expected to have much less cytoplasmic variability than many other systems. Although most previous work has not explicitly focused on variability, studies of GEM particle diffusion in the cytoplasm or nucleoplasm of budding yeast, the filamentous fungus *Ashbya gossypii*, *Xenopus* egg extract, and several mammalian cell types (Delarue et al., 2018; Huang et al., 2021; Szórádi et al., 2021), as well as other studies of diffusion in *E. coli* (Parry et al., 2014), show that comparable variability in diffusion exists in these diverse contexts. In particular, McLaughlin et al. reported sizeable variation in both inter- and intra-cellular heterogeneity of GEM diffusivity in *Ashbya (McLaughlin et al., 2020)*. Beyond measurements of diffusion, a study directly probing viscosity also revealed substantial variability (Wang et al., 2019). Thus, large variability in cytoplasmic properties may be a fundamental, conserved property of cells.

Hints from the literature suggest that heterogeneous cytoplasmic diffusion is also not limited to large protein complexes. Both larger objects such as lipid droplets (Tolić-Nørrelykke et al., 2004) and smaller particles such as individual fluorescent proteins (Bakshi et al., 2011; Baum et al., 2014;

Dross et al., 2009; Manley et al., 2008; Scipioni et al., 2018; Xiang et al., 2020; Yan et al., 2020) seem to exhibit substantial amounts of diffusive heterogeneity.

Sources of cytoplasmic heterogeneity

What might be the origin of this variability in cytoplasmic properties? Heterogeneity may originate from multiple non-exclusive sources. At the micron-scale, obstruction by organelles (Gu et al., 2007; Parkinson et al., 2008) and other cytoplasmic structures such as condensates, as well as localized active mixing, could contribute to cytoplasmic variability (Chaubet et al., 2020). At the nanometer-scale there are some enticing sources of heterogeneity that remain unexplored, notably those intrinsic to the macromolecular milieu: crowder density, size, charge, and hydrophobicity. Indeed, the fact that diffusion varies strongly with probe size and molecular species (Arrio-Dupont et al., 1996, 2000; Banks and Fradin, 2005; Luby-Phelps et al., 1986, 1987; Verkman, 2002), suggests that the local molecular structure of the cytoplasm plays a large role in the diffusion of macromolecules. Similarly, all-atom molecular dynamics simulations of the cytoplasm show that thermal fluctuations in the local cytoplasmic composition can lead to significant variability in diffusion rates (Yu et al., 2016). Therefore, the molecular and cellular features contributing to viscosity may themselves be highly dynamic and transient. Future studies of diffusive heterogeneity across different species, cell types, and physiological states will be invaluable for dissecting the biophysical determinants of cytoplasmic variation.

Consequences of cytoplasmic heterogeneity

The heterogeneity of the cytoplasm may act as a highly significant source of biological noise for any diffusion-limited process. For example, spatial heterogeneity in diffusivity could lead to differences in diffusion-limited reaction rates across the cell. In particular, if the regions of high viscosity (low diffusivity) are long lived, they could act as "traps", locally increasing the concentration of larger protein complexes or organelles, potentially influencing the speed and localization of certain reactions. The effects of stochasticity should be particularly strong for complexes which exist at low copy number or whose biological function depends on rare binding events. At the cell population level, having a wide range of diffusivities might be advantageous, allowing different cells to react to changes in the environment at different rates, permitting strategies such as bet-hedging to take place.

In fact, it is hard to imagine a biological process that would not be affected by such a large variation in the effective viscosity. For example, many reactions driving gene expression, biosynthesis and metabolism are considered to be diffusion limited. For example, cytoplasmic viscosity has been demonstrated to have strong effects on microtubule dynamics in vivo (Molines et al., 2022). Interestingly, the dynamics of individual microtubules were much less variable than those of the GEMs, suggesting that cellular systems may employ compensatory mechanisms that buffer the effects of heterogeneity in viscosity. Cellular control of viscosity and other aspects of the cytoplasm such as intracellular density represents a potential global mode of regulation.

Generalization of the Doppelgänger simulation approach

Our analyses of biological noise were made possible by using our Doppelgänger simulation approach. This approach explicitly reproduces the experimental measurement statistics in silico, which allowed us to definitively distinguish between statistical noise and biological heterogeneity. This simulation approach may be generalizable to many other systems (Supp. Fig. 4), and could be useful for instance in the analysis of noise suppression. Overall, we believe this powerful approach combining experiment and theory will provide needed clarity for studies of stochastic processes in biology, such as cytoskeletal dynamics, signaling, and gene expression.

Methods

Table 1: Reagents and Resources

REAGENT AND RESOURCE	SOURCE	IDENTIFIER		
S. pombe strains				
h- [pREP41X-PfV-Sapphire] ade+ his+ leu+ ura+	Chang lab collection (Molines et al., 2022)	FC3287		
h+ GFP-atb2:kanMX ade- leu- ura- his+	Chang lab collection	FC2861		
h+ pAct1-Lifeact-mCherry::leu+ ade- leu- ura- his+	Chang lab collection (Huang et al., 2012)	FC2781		
Chemicals				
D-sorbitol	Sigma	S1876		
Carbendazim	Sigma	378674		
Latrunculin A	Abcam	ab144290		
Edinburgh Minimal Media (EMM)	MP Biomedicals	4110-032		
YES 225	Sunrise Science Products	2011-500		
Lectin (glycine max)	Sigma	L1395		
Dimethyl sulfoxide	Sigma	472301		
Supplies				
μ-Slide VI 0.4 ibiTreat	IBIDI	80606		
Software				
FIJI	Schindelin et al., 2012	https://imagej.net /contribute/fiji		
ImageJ	Schneider et al., 2012	https://imagej.nih. gov/ij/		
MOSAIC for ImageJ	Sbalzarini et al., 2005	https://imagej.net /plugins/mosaicsui te		
Matlab	Mathworks	https://www.mathw orks.com/		

Micromanager	Edelstein et al., 2010	https://micro-man ager.org/Citing_Mi cro-Manager
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Yeast strains and culture conditions

Standard methods for growing and genetically manipulating *Schizosaccharomyces pombe* were used (Moreno et al., 1991). The constructions of the GEMs expressing strains were described previously (Lemière et al., 2021; Molines et al., 2022). In brief, the encapsulin-mSapphire chimera was expressed under the control of the inducible nmt1 promoter (Maundrell, 1990) on a multcopy pREP41X plasmid containing a leucine selection cassette. Cells were grown overnight in Edinburgh minimal medium (EMM) containing adenine, histidine and, uracil at 0.25 g per liter (here called EMM LEU-) and 0.1 μg/mL thiamine with shaking at 30 °C to exponential phase (OD600 between 0.2 - 0.8). See Table 1 for reagents and strain list. Expression of the *Pyrococcus furiosus* encapsulin-mSapphire construct produces particles of 40 nm in diameter, with the encapsulin proteins facing the inside of the particle and the fluorescent proteins facing the cytoplasm (Delarue et al., 2018; Szórádi et al., 2021).

Microscopy

S. pombe cells were imaged in commercial microchannels (Ibidi μ-slide VI 0.4 slides; Ibidi 80606, Ibiditreat #1.5). Channels were pre-treated with 50 μl of 100 μg/ml lectin solution for 5 min. The lectin solution was removed by pipetting and 50 μl of cell culture were introduced then incubated for 5 to 10 minutes to allow adhesion to the lectin then cells were washed with EMM LEU-. For the 20°C condition the Ibidi slide and the buffer were equilibrated at 20°C before cells were added. For the 30°C condition, slides and buffers were equilibrated at 30°C before cells were added to it. For hyper-osmotic shocks, the medium was manually removed from the channel via pipetting and quickly replaced with pre-warmed (30°C) hyper-osmotic media. Cells were imaged immediately and for no longer than 5 minutes after the medium was exchanged to minimize adaptation. For cytoskeleton depolymerization cells were introduced in the Ibidi slide as described previously then the buffer was exchanged for pre-warmed (30°C) EMM LEU- containing Latrunculin A (8.4 μg/mL or 20 μM) and methyl benzimidazol-2-yl-carbamate (MBC) (25 μg/mL or 131 μM). Cells were incubated at 30°C with the drug cocktail for 5 minutes prior to imaging. We confirmed that this treatment caused depolymerization of the microtubule and actin cytoskeletons in < 5 min by imaging cells expressing Lifeact-mCherry or GFP-Atb2.

For imaging GEMs, yeast cells were imaged with a Nikon TI-2 equipped with a Diskovery Multi-modal imaging system from Andor and a SCMOS camera (Andor, Ixon Ultra 888) using a 60x TIRF objective (Nikon, MRD01691). Cells were imaged sequentially, first a brightfield (BF) image then 1,000 fluorescence images at 100 Hz (for ~ 10 s) with a 488 nm excitation laser and a GFP emission filter 525 +/- 25 nm. Variable angle epifluorescence microscopy (VAEM) (Konopka and Bednarek, 2008) was used to reduce background fluorescence and allow for the high imaging frequency required. Cells were selected for sparse numbers of labeled motile nanoparticles (< 10 GEMs per cell) to ensure proper particle tracking. Note that each GEM can be imaged multiple times during the acquisition giving more tracks per cell than the number of visible nanoparticles.

Particle tracking

Cells were individualized from the field of view by cropping the images. Images of individual

cells were rotated so that cell length (long axis) was horizontal. From the brightfield image cell length was measured by tracing a straight line joining each pole and passing through the center of the cell. Cell contours were drawn manually from the brightfield image and used to determine cell centroid. Cell length and centroid were used to plot GEMs tracks in real and normalized space (Figure 1). GEMs nanoparticles in each cell were tracked using the MOSAIC plugin (Fiji ImageJ)(Sbalzarini and Koumoutsakos, 2005; Schindelin et al., 2012) with the following parameters for the 2D Brownian dynamics tracking in MOSAIC: radius = 3, cutoff = 0, per/abs = 0.2-0.3, link = 1, and displacement = 6. Tracks shorter than 10 timepoints were removed from further analysis.

Diffusivity Analysis

Mean Square Displacement (MSD) Analysis: The $MSD = \langle (x(t+\tau)-x(t))^2 \rangle$ was computed using non-overlapping intervals and plotted versus time offset, τ . A linear fit of ln(MSD) vs $ln(\tau/\tau_0)$ for the first 7 time offsets (~70 ms) was used to determine the values of the anomalous exponent and the apparent diffusivity (see our rationale for choice of τ_0 below in this paragraph). As the length of the trajectories is an exponentially decaying distribution (Fig. 1d - histograms), the statistical error grows with time (Fig. 1e - MSD) -- hence, we fit the only first part of the MSD function. The fitting resulted in two fit parameters corresponding to the equation $MSD = A(\tau/\tau_0)^{\alpha}$, where A has units of nm² (representing the MSD when $\tau = \tau_0$) and α is unitless. We can convert these values to an apparent diffusivity by assuming $MSD_{\tau=\tau_0} = A = 2nD_{app,\tau_0}\tau_0$, where n is the number of spatial dimensions (in this case, n=2). Solving for the apparent diffusivity D_{app} in nm²ms¹¹, we find the following conversion: $D_{app,\tau_0} = A/(2n\tau_0)$ (representing the apparent diffusivity specifically at τ_0). We choose $\tau_0 = 100$ ms, to represent the intermediate regime measured in our dataset. For track-wise fits, the time-averaged MSD was calculated and fit separately for each trajectory. For cell-wise and condition-wise MSD calculations, the time-averaged MSDs for each track were then ensemble-averaged over all tracks in each cell or condition, respectively, and subsequently fit.

Doppelganger Simulations

Simulations of particle diffusion were implemented using fixed time step Brownian dynamics, according to the Stokes-Einstein relation for diffusion of a spherical particle in a viscous medium ($D = k_B T/\gamma$, where D is the diffusion coefficient such that the mean-squared displacement MSD = 2nDt is linear with time t and the number of dimensions n, k_B is the Boltzmann constant, T is the temperature, and $\gamma = 6\pi\eta R$, where η is the viscosity of the cytoplasm and R is the radius of the particle). See Table 2 for a list of the parameters used. All code was written in custom MATLAB scripts. Cells were implemented as 2D rectangular boxes with reflecting boundary conditions at the edges of each box. All simulated cells had a short-axis width of 3 μ m, and a long-axis width equal to that of it's experimentally-measured doppelgänger. (Note that the short-axis width was chosen to be 3 μ m, rather than the known 4 μ m diameter of fission yeast cells, to best represent the imaging conditions in the experimental data. VAEM imaging only captures the lower portion of the cell near the coverslip, where the cross-section is smaller than at the equatorial plane.)

Each simulated cell had the same number of particles as its experimental doppelgänger. Each particle was initialized randomly within the rectangular cell wall boundary. After initialization, particle positions were updated using fixed time step Brownian dynamics, where the fixed time step, Δt , was equal to the acquisition frame rate of the experimental measurements. In each time step, a random number generator (randn, seeded randomly at the beginning of each set of simulations with rng('shuffle')) selected each particle's step size and direction from a normal distribution with a mean of zero and a standard deviation of $\xi = \operatorname{sqrt}(2*k_{\rm B}T*\gamma/\Delta t)$. If a particle left the cell boundary during a timestep, the particle's position was reflected across the cell boundary (or boundaries) that the particle crossed, in order to keep the particle inside the cell (i.e., reflective boundary conditions). For ease of

implementation, all particle tracks were simulated for the longest length of time any particle in the experimental dataset was tracked; then after simulations were complete, each simulated particle's data were pruned to match their experimental doppelgänger -- all other timepoints that were not tracked for the experimental doppelgänger were deleted from the simulated dataset.

For simulations with cell-cell variations in viscosity (Models #3 and #4), viscosity values for each cell were chosen from a random log-normal distribution ($\eta = \mu e^{(\sigma/\mu)^* randn(t)}$), with a mean viscosity equal to 40 times that of water, and a standard deviation of 45% of the mean. For simulations with spatially-varying viscosity (Models #2 and #4), each rectangular cell was broken up into spatial domains of equally-sized squares with 1 μ m side-length. As all cells were 3 μ m in width but variable in length, simulated spatial domains within cells were arranged in a 3xm grid, where m is the number of domains along the long axis. If the cell length along the long dimension was not an integer multiple of 1 μ m, then the remainder was placed in its own spatial domain of smaller size. Viscosity values in each domain were chosen from a random log-normal distribution ($\eta = \mu e^{(\sigma/\mu)^* randn(t)}$), with a mean equal to the mean viscosity of that cell, and a standard deviation equal to 85% of the mean.

Statistical analysis

Velocity autocorrelation analysis: Velocity autocorrelations were defined as $VAC(\tau) = \langle (v(t+\tau)v(t)) \rangle_t$ and were performed using non-overlapping intervals.

ANOVA: A nested, n-way analysis of variance was performed using MATLAB's anovan() function. Track identity was nested under cell, session, and day identities, cell identity was nested under session and day identities, and the session identity was nested under the day identity. ANOVA was performed separately on the power law exponents and the natural logarithm of the diffusivities. ANOVA was performed identically on the experimental and simulated datasets. Because the Doppelgänger simulation approach computationally reproduces the exact experimental distribution of tracks, cells, sessions, and days, the exact magnitudes of the variance attributed to each category can be directly quantitatively compared (e.g. Figure 3f).

Comparison of median diffusivity values between conditions: A Wilcoxson rank sum non-parametric test for equality of medians (Mann and Whitney, 1947) was performed to determine whether differences in the medians between conditions were statistically significant. We chose a non-parametric test, and compared the medians instead of the means, so that our analysis would be less sensitive to the fact that the distributions were long-tailed and not perfectly Gaussian (even in log space). Statistical tests were performed on the logarithm (base 10) of the apparent diffusivities.

Comparison of variance in diffusivity values between conditions: Levene's test for equality of variance (Levene, 1960) was performed to determine whether differences in the variances between conditions were statistically significant. While Levene's test is not a non-parametric test, it is less sensitive to non-normality than many other parametric tests, and is MATLAB's recommended test for equality of variance for non-normal distributions. Statistical tests were performed on the logarithm (base 10) of the apparent diffusivities.

Converting summary statistics from log space to real space: Because the diffusivities were more normally distributed in log space than in real space, all of the summary statistics (medians, standard deviations, etc.) were calculated on the distribution in log space. For a distribution that is log-normally distributed, medians and standard deviations calculated in log space are not the same as those calculated in real space and so are not interchangeable (i.e. $10^{<a>} =/= <10^a>$, and $10^{sqrt(<(a-<a>)^{\sim}2>)} =/= sqrt(<(10^a-<10^a>)^2>)$ and have different interpretations. The median of the log-scale diffusivity distribution ($\mu_{log} = <log_{10}(D_{app,100ms})>$ represents the median order of magnitude of diffusivities in the dataset. The medians reported in this work were first calculated from the distribution in log space, and then converted to real space as $\mu_{real} = 10^{\mu_{-}log}$, and also represents the median order of magnitude (but now presented in real space). The standard deviation of the log-scale diffusivity distribution ($\sigma =$

sqrt(<(log₁₀(D_{app,100ms})-<log₁₀(D_{app,100ms})>)^2>) represents the number of orders of magnitude spanned by the dataset). In real space, the associated number which best captures the data's span in order of magnitudes is the fold-range of the distribution measured at some specified number of standard deviations away from the mean. In our dataset, a 2.5 σ threshold best matched the outlier exclusion algorithm used in our box-plotting software (1.5 times the interquartile range past the 25th and 75th percentiles, Fig. 2c, see caption). To determine the fold-range, the standard deviation was calculated for the diffusivity distribution in log space, then the ratio of the diffusivities at $10^{(\mu_{-1}log+2.5*\sigma_{-1}log)}$ (i.e., the fold-range) was evaluated as $10^{(\mu+2.5*\sigma)}/10^{(\mu-2.5*\sigma)} = 10^{5*\sigma}$. In perturbation conditions (Fig. 4), the reported percent change in the median and fold-range were de

Table 2: Model Parameters - Input parameters

Variable	Meaning	Value	Source
variable	Meaning	value	Source
k _B	Boltzmann's constant	0.0138pN nm / K	-
Τ	Cytoplasm temperature	303.15 K (30 °C)	This work
R	Particle radius	20 nm	(Delarue et al., 2018)
L	Cell length	That of the experimentally- measured counterpart	This work
W	Cell width	3 μm	This work
$\eta_{\scriptscriptstyle w}$	Dynamic viscosity of water	2.414×10 ⁻⁸ ×10 ^{(247.8/(T-140))}	-
		(8×10 ⁻⁷ pN ms / nm² at 30 °C)	
$\langle \eta_{c,s} angle$	Dynamic viscosity of the cytoplasm (mean across all cells and subcellular spatial domains)	40 η _w (320×10 ⁻⁷ pN ms / nm² at 30 °C)	Approximated to match experimentally-measured average diffusivity assuming the Stokes-Einstein relationship D = k_BT/γ
$\sigma_{\eta,c}$	Standard deviation of the average cellular viscosity across all cells in the population	$0.45\langle\eta_{c,s} angle$	Best fit to this work
$\sigma_{\eta,s}$	Standard deviation of the viscosity among all spatial domains in a cell	$0.85\langle\eta_c\rangle$ where $\langle\eta_c\rangle$ is the average across all spatial domains in a particular cell	Best fit to this work
λ	Spatial domain size within a single cell	1 μm	Best fit to this work

Table 3: Model Parameters - Derived parameters

Variable Meaning Value

k _B T	Thermal energy	4.18 pN nm
⟨γ⟩	Viscous drag coefficient given Stokes' law (for the average particle) $\gamma = 6\pi \langle \eta_c \rangle R$	0.012 pN ms / nm (at 30 °C)
$\langle D_{c,s} \rangle$	Diffusivity (averaged across all cells and spatial domains) $D = k_B T/\gamma$	350 nm²/ms

Data and code availability

All raw imaging data are available upon request. All tracking data and code are freely available on Gitlab:

https://gitlab.com/theriot_lab/vast-heterogeneity-in-cytoplasmic-diffusion-rates-revealed-by-nanorheology-and-doppelgaenger-simulations.git

Author Contributions

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Investigation, A.T.M., and R.M.G;

Resources, A.T.M., and R.M.G.;

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Writing - Original Draft, A.T.M., R.M.G..;

Writing – Review & Editing, A.T.M., R.M.G., J.A.T., and F.C.;

Visualization, R.M.G. and A.T.M.;

Supervision, A.T.M. and R.M.G.;

Project Administration, A.T.M. and R.M.G.;

Funding Acquisition, J.A.T. and F.C.;

R.M.G. and A.T.M. contributed equally and have the right to list their name first on their C.V.

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Supplementary Figures

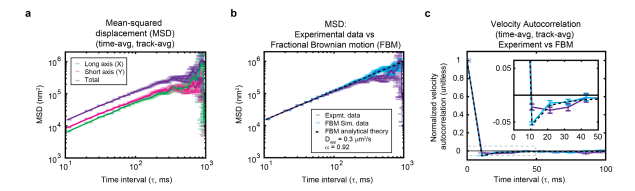


Figure S1: Experimental data is consistent with nearly unconstrained diffusion. (a) Mean-squared displacement (MSD) along the long and short axes of the cell, plotted alongside the total MSD. (b) The predicted MSD for Fractional Brownian motion (FBM), including both analytical theory and results from simulated data, using the experimentally-measured values of D and α . The experimental data is also plotted for comparison, showing good agreement with the theory. (a-b) Note the logarithmic scale along the x- and y-axes. (c) The predicted velocity autocorrelation for FBM, showing the characteristic negative peak which then decays to zero. Experimental data shows a wide and very shallow negative basin, which does not match the shape or depth of the peak predicted by FBM.

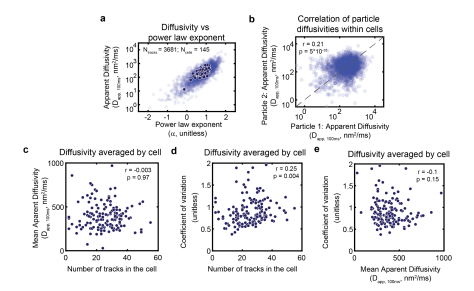


Figure S2: Additional evidence for intrinsic and extrinsic sources of noise. (a) The relationship between the apparent diffusivity and power law exponent. (b) The apparent diffusivity of each particle in the dataset plotted against a randomly chosen particle from the same cell. Each particle is represented exactly once in the plot. For cells with an odd number of particles, one particle would not be represented for that cell. (c) Mean diffusivity across tracks in each cell plotted vs the number of tracks in each cell. (d) Coefficient of variation across tracks in each cell plotted vs the number of tracks in each cell. (e) Coefficient of variation vs mean diffusivity calculated by averaging across all tracks for each cell. (a-e) Fits of track-wise MSD data are shown in light blue, with cell-wise fits overlaid in dark blue. (a-b) Note the logarithmic scale along the y-axis. (b) Note the logarithmic scale along the x-axis. (b-d) r- and p-values determined by a Spearman correlation algorithm.

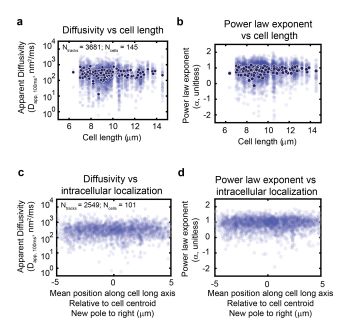


Figure S3: The large heterogeneity in diffusivity cannot be explained by the cell cycle or subcellular GEM particle localization. (a-b) Fitted values for diffusivity (a) and power law exponent (b) plotted as a function of cell length. (c-d) Track-wise fit values for diffusivity (c) and power law exponent (d) plotted against the mean (time-averaged) particle position along the long axis of the cell. There are fewer cells and tracks represented in (c-e) compared to (a-b) because the new pole could be distinguished from the old pole for only a subset of cells. (a-d) Fits of track-wise MSD data are shown in light blue, with cell-wise fits overlaid in dark blue. (a, c) Note the logarithmic scale along the y-axis.

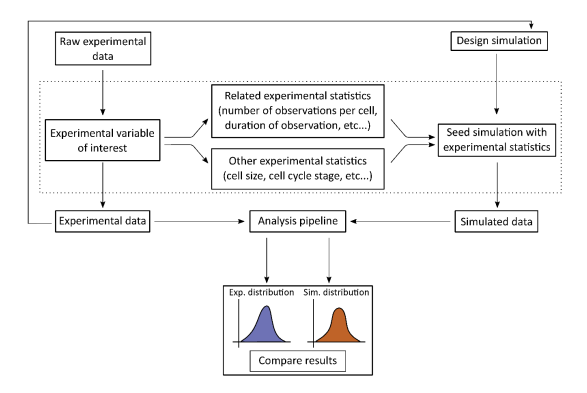


Figure S4: Schematic of the generalized Doppelgänger approach.