Statistical Analysis of Scanning Fluorescence Correlation Spectroscopy Data Differentiates Free from Hindered Diffusion

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Supporting Information

ABSTRACT: Cells rely on versatile diffusion dynamics in their plasma membrane. Quantification of this often heterogeneous diffusion is essential to the understanding of cell regulation and function. Yet such measurements remain a major challenge in cell biology, usually due to low sampling throughput, a necessity for dedicated equipment, sophisticated fluorescent label strategies, and limited sensitivity. Here, we introduce a robust, broadly applicable statistical analysis pipeline for large scanning fluorescence correlation spectroscopy data sets, which uncovers the nanoscale heterogeneity of the plasma membrane in living cells by differentiating free from hindered diffusion modes of fluorescent lipid and protein analogues.

KEYWORDS: Brownian, scanning FCS, free and trapped diffusion, actin cytoskeleton, hindered diffusion dynamics, diffusion modes

Quantification of molecule diffusion dynamics in the plasma membrane of living cells is essential for the understanding of their function.1−3 Most fluorescence-based methodologies for examining the heterogeneity of lipid and protein diffusion dynamics, such as single-particle tracking,4,5 fluorescence recovery after photobleaching,6,7 or fluorescence correlation spectroscopy (FCS),1,2 and their advancements, especially super-resolution stimulated emission depletion (STED) FCS,8−10 either necessitate specialized hardware, sophisticated fluorescence labeling, or constrain themselves to quantification with limited spatial and temporal sensitivity. Together, these issues remain to pose practical challenges for the quantification of diffusion dynamics in living cells.

Important advancements in fluorescence acquisition and/or analysis pipelines drew attention to the power of simple measurement equipment like total internal reflection setups, which can be used for camera-based FCS variants.11−13 For instance, for the disclosure of large spatial diffusion maps, the methodology of image mean square displacement (iMSD) analysis provides robust detection of free Brownian and non-Brownian hindered diffusion modes.14,15 Unfortunately, these camera-based acquisitions often limit the accessible diffusion time scales.16

Yet, scanning FCS (sFCS) measurements acquired with confocal laser scanning microscopes offer a good compromise in terms of equipment, adaptability (e.g., combination with iMSD), and high spatiotemporal sensitivity with statistical accuracy and flexibility by combining conventional instrumentation and standard fluorescence labeling.14,17−19 Specifically, the large number of simultaneously acquired FCS curves (>50 curves) and well-established fitting and bleaching correction procedures ensure high confidence in the data analysis.20−22 Although sFCS measurements are often performed in biological membranes at the equatorial plane of cells or vesicles,23,24 they can, in principle, flexibly be applied anywhere in the cell volume.19,22,23 From the biological point of view, it is important to differentiate three main parameters for the complete characterization of molecular diffusion dynamics: the total number of diffusion processes, the

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Figure 1. (a) Schematic of the experiment principle. (Left) sFCS data are recorded by scanning the laser (green dots) along a micrometer line in the membrane (lipids with red headgroups and gray tails), thereby creating (through temporal correlations) FCS data (decaying curves $G(\tau)$ from red to blue against correlation time $\tau$) for each pixel along the line (space). (Right) All FCS data are fitted to obtain values of transit times through the observation spot, in which histograms (blue, probability distributions; right, cumulative; top right, logarithmic values; bottom right, linear representation) are fitted by the LogNorm (red line), with weighted residuals in the respective bottom panels. (b–d) Cumulative (top panels) and linear (bottom panels) transit time histograms (blue) and LogNorm fits (red, single- or double-LogNorm as labeled) with weighted residuals (respectively bottom panels) for simulated sFCS data of (b) freely (input: $D = 0.80 \mu m^2/s$, recovered $D = 0.82 \mu m^2/s$) and (c) hindered diffusion (input: $D_{\text{free}} = 0.80 \mu m^2/s$, $D_{\text{trapped}} = 0.1 \times 10^{-9} \mu m^2/s$, and $p_{\text{trap on}} = p_{\text{trap off}} = 0.001$, recovered $D_1 = 0.39 \mu m^2/s$, $D_2 = 0.31 \mu m^2/s$, and $A = 0.73$), and (d) for experimental data of DPPE-Abberior STAR Red in pure DOPC (blue; $\mu = 4.2$ ms, $D = 2.45 \mu m^2/s$) and DOPC/Chol (green; $\mu = 9.7$ ms, $D = 1.1 \mu m^2/s$) SLBs (see Supporting Information Table S1). Inset (b): Fit results $\mu$ of the LogNorm fits (output values) against the transit times implemented in the sFCS simulations of free diffusion, indicating an accurate recovery of values. Insets (d): Relative likelihood (RL) values of Gaussian (Gauss) and single (sLogn) and double (dLogn) LogNorm fits to the experimental data of pure DOPC (left) and DOPC/Chol (right) SLBs, indicating accurate fitting by a single-LogNorm model.
fraction of molecules contributing to each diffusion mode, and their respective diffusion time scales. Such a parametrization allows one to infer whether molecules undergo free Brownian or hindered diffusion, revealing deeper insights into cellular signaling and function. For example, during the activation of immune cells, well-defined fractions of immune receptors are thought to form receptor clusters through transient trapping of their intracellular domains.\(^{29,30}\) In turn, certain lipids such as 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine (DOPE) and the transferrin receptor have been shown to undergo hindered hop diffusion within the dense cortical actin network beneath the cell membrane.\(^{2,4,25}\) Conventional sFCS recordings theoretically contain all the kinetic information on such hindered diffusion of fluorescent particles, including hop and trapped diffusion modes within the observed probe ensemble. However, this information is mostly lost due to averaging during the analysis process because, traditionally, only the average diffusion coefficients and their standard errors are computed, assuming a normal distribution of the sFCS data sets. Consequently, conventional sFCS quantification analysis does not usually allow for the direct disclosure of diffusion modes, such as for differentiating hindered from free Brownian diffusion, precluding perhaps the discovery of lipid or protein clustering, hopping, or trapping.

Here, we introduce a pipeline for the statistical analysis of sFCS data recorded by confocal microscopy, yielding the systematic and robust quantification of diffusion modes in living cells. The protocol allows a systematic mathematical characterization of the distribution of sFCS diffusion data, involving fitting analysis and the calculation of the biological-relevant parameters, and a quantitative evaluation of the results using weighted fitting residuals and maximum likelihood estimations. Applying this framework to the free and non-Brownian diffusion dynamics of lipids and glycosylphosphatidylinositol (GPI)-anchored proteins demonstrates the power of the method and allows a comprehensive characterization of well-known molecular particle diffusion in computer-simulated data sets, supported lipid bilayers (SLBs), and the plasma membrane of living cells. Combining this analysis pipeline with conventional confocal sFCS experiments may offer alternative avenues for systematic studies of the mechanisms by which living cells adjust their molecular membrane dynamics to their physiological needs.

Analysis of sFCS experiments conventionally involves two steps: (i) simultaneous acquisition of a multitude of FCS curves acquired by repeatedly scanning few-micrometer-long lines, and (ii) obtaining the transit times of the diffusing molecules through the observation spot from fitting the respective time-correlated data for each pixel of the line to specific diffusion models, yielding a value of an average transit time and a standard error. To improve sFCS analysis, we aimed to examine the full kinetic information provided by all the correlation curves. Consequently, our strategy comprised a third step, wherein we computed and analyzed the transit time histograms to extract further information on the diffusion modes (Figure 1a), exploiting the statistical power of large ensemble measurements independent of local spatial heterogeneities.

RESULTS AND DISCUSSION

We first set out to statistically evaluate the distribution of transit time values obtained from a large ensemble of computer-simulated sFCS data (see the Methods section and the Supporting Information). Consistent with previous findings, the histograms over a wide range of transit time values displayed non-Gaussian distributions (Supporting Information Figure S1a). The broadness and asymmetry toward longer transit times of the distribution depended on the magnitude of the measured transit time values, because the error of the measurements increased exponentially with the transit times, accompanied by respective changes in the signal-to-noise and fitting standard deviation (Supporting Information Figure S1b,c). Specifically, slow diffusing particles experienced larger errors than faster particles, primarily depending on the finite scanning frequency but also on other factors such as the finite acquisition time. To this end, the total acquisition time determined the number of points for correlation, limiting the convergence of the corresponding FCS curves.\(^{20}\) Computing the logarithm of the corresponding transit time histograms revealed a normal distribution, indicating, together with exponentially distributed signal-to-noise and fitting standard deviation, that sFCS data were log-normally distributed (Figure 1a and Supporting Information Figure S1b,c). The variance of the distributions increased with increasing transit time (Supporting Information Figure S1a).

In the following, we exploited the mathematical properties of a log-normal distribution, hereafter referred to as LogNorm. Statistical analysis of the sFCS data was achieved by fitting the cumulative, logarithmic, and linear representation of the histograms using analytical forms of the LogNorm (Figure 1a and Supporting Information Figure S2a). LogNorm functions are well-defined through their two characteristic parameters, \(\mu\) and \(\sigma\), which can mathematically be related to the mean and standard deviation of a normal distribution in its logarithmic representation. Consequently, the analysis involves the fitting of three representations to obtain two free parameters. Large ensemble acquisitions of sFCS curves (>400 per condition) ensured sufficient bin counts (30–80 bins). The bins of a given transit time histogram reflected the number of fitting points. Empirically, we found that the LogNorm fitting accurately recovered the input diffusion parameters in the computer simulations of free diffusion over a wide range of transit times (inset of Figure 1b and Supporting Information Figure S2b–d).

Next, we introduced hindered trapped diffusion (transient halts in the diffusion path; see stochastic trapping model in the Methods section and Supporting Information Figure S3a,b) as a second process in addition to free diffusion in the computer simulations. Using our statistical analysis, we found deviations from the single-LogNorm behavior as indicated by the weighted residuals. These transit time histograms were accurately fitted with a double-LogNorm fitting model in all three representations (Figure 1c and Supporting Information Figure S3c). Consistently, the statistical analysis indeed revealed two LogNorm transit time histograms reporting on free Brownian and non-Brownian trapped diffusion.

To confirm the findings of the computer simulations, we further measured the diffusion of fluorescently labeled lipids in SLBs. As expected, analysis of the diffusion dynamics of a fluorescent DPPE analogue (1,2-dipalmitoyl-sn-glycero-3-phosphoethanolamine labeled with Abberior STAR Red) in a fluid SLB of DOPC (1,2-dioleoyl-sn-glycero-3-phosphocholine) accurately detected free diffusion (i.e., best representation of the data by a single-LogNorm fit), with an overall decrease in mobility depending on the cholesterol content (Figure 1d). In contrast to the computer simulations, we did not have any prior knowledge of the diffusion modes. We hence complemented our weighted residual evaluation for the fitting and model selection with maximum likelihood estimations, employ-
The Bayesian information criterion (from which we could calculate the relative likelihood (RL) value for any given model; see the Methods section and Supporting Information Figure S4a). Consistent with the weighted residuals, the RL values
confirmed free diffusion for the lipids in the SLB by best representing the data with a single-LogNorm fit (insets in Figure 1d and Supporting Information Figure S4a and Table S1). Note, in the following, the most likely model (RL value equals 1) was selected as the appropriate fitting model. The weighted residuals were employed to evaluate the quality of the single- or double-LogNorm fits in all experiments.

Next, we investigated the well-described nanoscale diffusion dynamics of fluorescent lipid analogues in living Ptk2 cells. Consistent with previous reports using advanced FCS technologies such as super-resolution STED-FCS, the phospholipid analogue PE (Atto647N-DPPE) diffused freely, and the sphingomyelin (SM) analogue (Atto647N-SM) showed hindered trapped diffusion, as revealed by either single- and double-LogNorm fits to the transit time distributions (Figure 2a,b and Supporting Information Figure S4b), respectively. Trapping sites of SM were also illustrated by the occurrence of the heterogeneities in the unprocessed sFCS data (correlation carpets) of SM compared to PE (Figure 2a,b). Consequently, the statistical analysis of large ensemble sFCS data acquired with a conventional confocal microscope yielded the detection of a small fraction of molecules undergoing trapped diffusion.

To further demonstrate the applicability of our approach, we examined the diffusion of GPI-anchored proteins (GPI-AP) fluorescently labeled with green fluorescent protein (GFP) and compared the results of our sFCS analysis to those from the diffusion dynamics of the GPI-APs in giant plasma membrane vesicles (GPMVs). While GPI-AP diffusion appeared to be trapped in the cells, likely due to the interactions with the cortical actin cytoskeleton, it was seemingly free in actin-depleted GPMVs (Figure 2c,d and Supporting Information Figure S5a) as previously demonstrated. The statistical analysis was therefore able to detect free and trapped diffusion of GFP-tagged GPI-APs in the cell membrane.

Finally, we explored the ability of our analysis pipeline to detect hindered hop diffusion. To this end, we produced spatial random mesh maps with an average boundary distance L in the simulations (Supporting Information Figure S5a) mimicking the cortical actin network beneath the membrane. The particles were able to undergo free motion within the boundaries of the individual meshes and random hop transitions for crossing the confinement barriers with the probability $P_{hop}$ (Supporting Information Figure S5a). To fully characterize the motion conditions of the particles, we calculated the phase diagram of those particles experiencing hop diffusion for a range of hopping probabilities and characteristic average length scale of mesh sizes for a given input diffusion coefficient (Supporting Information Figure S5b,c). Consistent with the previous hindered trapped diffusion simulations, the hop diffusion simulations revealed two processes in the transit time histograms, with the second process dominant on short time scales. Statistical analysis involving fitting and the maximum likelihood estimations of the transit time histogram uncovered a combination of a LogNorm and an exponential probability distribution function reporting on the combination of free Brownian diffusion and a random hop diffusion process, respectively (Supporting Information Figure S5d). Note that the same model selection criteria using the RL values were applied (Supporting Information). Consequently, the statistical analysis of large ensemble sFCS data also allowed the detection of a fraction of molecules undergoing hop diffusion.

**CONCLUSIONS**

In this work, we highlighted how different diffusion dynamics of lipids and proteins in the plasma membrane of living cells can be disclosed using conventional confocal microscopy. Revisiting well-established experiments, we demonstrated how this statistical analysis pipeline of scanning confocal microscopy allows the determination of free Brownian diffusion, double-LogNorm or LogNorm and exponential analysis enabled the detection of hindered trapped and hop diffusion dynamics.

In conclusion, the statistical analysis provides a broadly applicable methodology that revealed findings, which could previously only be characterized by dedicated equipment and experimental procedures such as spot variation or STED-FCS. Expanding the statistical analysis further to quantification of full reaction diffusion dynamics including transient binding and molecular cluster formation might offer one technique for a more complete characterization of lipid and protein dynamics in the near future. For example, reaction processes such as transient binding events of transmembrane proteins to associated structures such as the actin cytoskeleton or the extracellular matrix could also be quantified with the analysis pipeline. In light of the broad applicability, straightforward implementation, usage of standard fluorescent dyes at turn-key microscopes, we envisage statistical sFCS analysis to become an important tool for characterizing nanoscale heterogeneity of diffusion dynamics in the plasma membrane of living cells.

**METHODS**

**Preparation of Supported Lipid Bilayers.** SLBs were prepared by spin-coating as described elsewhere.

A solution of 1 mg/mL total lipid concentration (either 100% DOPC (Avanti Polar Lipids, USA) or 50% DOPC/50% cholesterol (Avanti Polar Lipids)) both doped with DPPE-Abberior STAR Red (Abberior, Germany) at a weight on weight ratio of 1:2000 in 1:2 methanol/chloroform was spin-coated at 3200 rpm onto a clean 25 mm #1.5 glass coverslip (VWR, UK). The resulting lipid film was hydrated with SLB buffer (150 mM NaCl, 10 mM 4-(2-hydroxyethyl)-1-piperazinethanesulfonic acid (HEPES) pH 7.4) and washed multiple times.

Glass coverslips were cleaned using piranha edging (3:2 sulfuric acid/hydrogen peroxide) for 2 h and not stored for longer than a week. For microscopy experiments, the cells were seeded onto 25 mm glass coverslips and allowed to grow to a confluence of about 75%. Cells were washed once with hypotonic (30%) then with 100% GPMV buffer (containing 150 mM NaCl, 10 mM HEPES, 2 mM CaCl$_2$, pH 7.4) and incubated for 3 h in 2 mM diethiothreitol and 25 mM parafomaldehyde at 37 °C. The

**Preparations of Giant Plasma Membrane Vesicles.** GPMVs from Ptk2 cells were prepared as described previously. Cells were allowed to grow to a confluence of about 75%. Cells were washed once with hypotonic (30%) then with 100% GPMV buffer (containing 150 mM NaCl, 10 mM HEPES, 2 mM CaCl$_2$, pH 7.4) and incubated for 3 h in 2 mM diethiothreitol and 25 mM parafomaldehyde at 37 °C. The
Microscopy. All imaging and sFCS data were acquired on a Zeiss 780 confocal laser scanning microscope (Carl Zeiss, Germany) equipped with a 40x C-Apochromat NA 1.2 W Corr FCS objective (Carl Zeiss). All sFCS experiments were performed using the GaAsP detector (Channel S) in photon counting mode, and the pinhole was set to 1 airy unit. For the GFP experiments, fluorescence was excited using a 488 nm argon laser, and a single 488 dichroic was chosen; for the Atto647N-labeled lipids, the fluorescence was excited using a 633 nm He–Ne laser, and the 488/594/633 MBS was used. sFCS experiments were acquired in line scan mode at a zoom of 40 with maximum time-lapse repetitions at highest scanning speed. These settings resulted in a line of 52 pixels (with a pixel size of 100 nm), a scanning frequency of 2081 Hz, a pixel dwell time of 3.94 μs, and an overall measurement time of about 47 s. The measurements were saved as .lsm5 files.

Simulations. Computer simulations were performed as Monte Carlo simulations in the programming language Python using the nanosimpy repository on GitHub.20,38 For free Brownian diffusion, we generated random tracks in a box of 2 μm × 8 μm. One hundred particles were randomly distributed, and their movement was simulated with diffusion coefficients D varying from 0.01 to 1.00 μm²/s. The tracks were simulated for t = 45 s with time steps of 3.94 μs (same as the pixel dwell time) using periodic boundary conditions. The scanned line of the sFCS experiments (hereafter referred to as sFCS line) was placed in the center of the simulation box (microns away from the boundaries). The sFCS line consisted of 52 pixels (length of 5.2 μm), where at every time step the molecules were passed through a Gaussian-shaped observation spot (resembling the microscopes’ point spread function) to generate the intensity fluctuations. The full width at half-maximum (FWHM) of the Gaussian-shaped observation spot was set to 200 nm to mimic confocal recordings. sFCS simulations were sampled at 500, 2081, and 3500 Hz. The obtained transit times, τ, for free Brownian diffusion, were allowed to sink down and equilibrate for 15 min at 37 °C. The coverslips were washed three times with GPMV buffer. PLL coating was performed by incubating the coverslip with 0.01% PLL solution (Sigma-Aldrich) for 1 h at room temperature. GPMVs were dropped onto poly- L-lysine (PLL)-coated 25 mm diameter coverslips. PLL coating was performed by incubating the coverslip with 0.01% PLL solution (Sigma-Aldrich) for 1 h at room temperature. GPMVs were allowed to sink down and equilibrate for 15 min at 37 °C. The diffusion was measured at the top membrane of the GPMVs. For the simulations of hindered hop diffusion, we adapted the simulations from point STED-FCS simulations for sFCS.31,32 To this end, we calculated a random meshwork with distinct barriers using Voronoi transformation of randomly seeded points. This spatial hopping map was superimposed with the simulation box and the sFCS measurement (compare Supporting Information Figure S5a). Particles at the boundary had a given probability (p_{trap}) to transition to the neighboring compartment or to reside in the same mesh compartment. For a diffusion coefficient of D = 0.2 μm²/s, we explored the parameter space employing shorter (30 s) scanning STED-FCS measurements with a fwhm of 40 nm in STED (yielding D_{on}) with 100 particles in the simulation. Ultimately, we simulated hop hindered diffusion with p_{trap} = 0.01, with a characteristic length scale of the mesh size of L = 100 nm (yielding a Voronoi region size given as ψ×area of 186 nm) of 150 particles for 45 s at a scanning frequency of 2081 Hz and a pixel dwell time of 3.94 μs. The respective intensity carpets from the simulations were outputted as .tif files.

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