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Quantitative mapping of fluorescently tagged cellular proteins using FCS-calibrated four dimensional imaging

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Abstract

The ability to tag a protein at its endogenous locus with a fluorescent protein (FP) enables the quantitative understanding of protein dynamics at the physiological level. Genome editing technology has now made this powerful approach routinely applicable to mammalian cells and many other model systems, opening up the possibility to systematically and quantitatively map the cellular proteome in four dimensions. 3D time-lapse confocal microscopy (4D imaging) is an essential tool to investigate spatial and temporal protein dynamics, however it lacks the required quantitative power to make absolute and comparable measurements required for systems analysis. Fluorescence correlation spectroscopy (FCS) on the other hand provides quantitative proteomic and biophysical parameters such as protein concentration, hydrodynamic radius and oligomerization but lacks the ability for high-throughput application in 4D spatial and temporal imaging. Here, we present an automated experimental and computational workflow that integrates both methods and delivers quantitative 4D imaging data in high throughput. This data is processed to yield a calibration curve relating the fluorescence intensities of image voxels to absolute protein abundance. The calibration curve allows the conversion of the arbitrary fluorescence intensities to protein amounts for all voxels of 4D (3D plus time) imaging stacks. With our workflow the user can acquire and analyze hundreds of FCS-calibrated image series to map their proteins of interest in four dimensions. Compared to other protocols, the current protocol does not require additional calibration standards and provides an automated acquisition pipeline for FCS and imaging data. The protocol can be completed in less than 1 day.

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AUTHOR CONTRIBUTIONS

A.Z.P. performed the experiments, the analysis and designed the software packages. A.Z.P. with the help of Y.C. and M.W. developed the protocol. N.W. tested the protocol and helped in writing the software manuals. B.K. created the homozygous cell line and provided the dextran protocol. M.J.H. provided the code to segment the cells. A.Z.P. with the help of B.K., N.W. and J.E. wrote the protocol.

Competing interests

The authors declare that they have no competing financial interests.

INTRODUCTION

Tagging of proteins with fluorescent markers is essential to study their localization, function, dynamics and interactions in living cells. With the advantage of genome editing technologies such as CRISPR/Cas9¹⁻³, it is now possible to engineer almost any higher eukaryotic cell-type to homozygously express the protein of interest (POI) fused to a fluorescent protein at its physiological levels (see our accompanying Nature Protocol⁴). Genome editing in combination with absolute quantitative fluorescence microscopy extends proteomics methods, which are typically restricted to large cell populations at specific time points, to single cell dynamic proteomics of living cells.

Modern confocal microscope detectors show a linear dependency of fluorophore concentrations and fluorescence intensities within several orders of magnitudes. To perform absolute quantitative imaging and convert relative fluorescence intensities to absolute physical quantities, such as the concentration of the fluorophore bearing protein, one can determine the calibration parameters of this linear dependency. To estimate the fluorophore's concentration, temporal and spatial fluctuations of diffusing species can be used⁵⁻⁷. In this protocol we describe how to use single point confocal fluorescence correlation spectroscopy (FCS), a biophysical single molecule technique that allows to measure concentrations and diffusion coefficients of fluorescently labeled molecules⁷⁻¹⁰. This method is well suited for low to medium abundant proteins (from few pM up to the μ M concentration range), a situation often observed at endogenous expression levels. Furthermore, most of the commercially available confocal microscopes can be equipped with the necessary detectors and hardware to perform FCS, making FCS-calibrated quantitative imaging easily available. In this protocol, FCS is combined with 3D confocal time-lapse imaging to estimate the calibration parameters and convert fluorescence intensities to concentrations in living cells. We previously used this approach to estimate protein numbers on bulk chromatin^{11,12}, on chromosome boundaries¹³, large multi-protein complexes¹⁴, and throughout mitosis for several genome edited cell lines¹⁵. In this protocol, we offer custom software packages and a step-by-step guide based on a Zeiss LSM confocal microscope with FCS capability as a widely used commercial system. The data acquisition software packages (available in Supplementary Software 1-3) simplify the repeated cycles of imaging and FCS measurements required for FCS-calibrated imaging. In combination with the provided online image analysis tools (Box 1), data acquisition can be completely automated¹⁶⁻¹⁸. This allows, without human supervision and in high-throughput, to automatically select cells with the optimal protein expression level and morphology and acquire images and FCS measurements. The data analysis software package (Supplementary Software 4) can be used to automatically extract the fluorescence intensities at FCS measurement points, compute protein concentrations and the FCS calibration parameters, and convert fluorescence image intensities to protein concentrations and absolute protein numbers.

FLUORESCENCE CORRELATION SPECTROSCOPY

For a detailed description on FCS and dual color fluorescence cross-correlation spectroscopy FCCS we refer readers to reviews and protocols (e.g.⁶⁻¹⁰). Here we provide only a short and practically oriented introduction to the topic (Fig. 1). FCS measures the fluorescence

intensity fluctuations of fluorescent molecules in a sub-femtoliter observation volume of a focused laser beam. The intensity changes caused by single molecules moving in and out of the observation volume over time are recorded over a time-frame of 15–30 sec (Fig. 1a-b). The fluorescence intensity fluctuations in the observation volume depend on (i) the concentration of fluorophores, (ii) their mobility, (iii) their photophysics, and (iv), to a lesser extent, the properties of the detector. To analyze the fluctuations, the autocorrelation function (ACF) $G(\tau)$ is computed from the intensity time trace: $I(t) = \langle I \rangle + \delta I(t)$:

$$G(\tau) = \frac{\langle \delta I(t) * \delta I(t + \tau) \rangle}{\langle I \rangle^2}. \quad (1)$$

The brackets $\langle \cdot \rangle$ indicate time-averaged quantities and $\delta I(t)$ the mean of the time-averaged intensity $\langle I \rangle$. The amplitude of $G(\tau)$ is inversely proportional to the number of particles in the observation volume N (Fig. 1c). The decay time τ_D of $G(\tau)$ gives the characteristic time a fluorophore remains in the observation volume, and consequently, is a direct measure of the fluorophore diffusion. These parameters can be extracted by fitting the ACF to a physical diffusion model (see Supplementary Notes 1 and 2). To compute the concentration of the fluorophore at the FCS measurement point, the number of molecules N is divided by an effective confocal volume V_{eff} (Box 2), typically below 1 fl, measured for the used microscopy setup.

OVERVIEW OF THE PROCEDURE

The protocol uses single color FCS to obtain one calibration curve for one type of fluorescent protein (FP) (see Figure 2 for a schematic overview of the procedure). Cells of interest are seeded and prepared for imaging (steps 1|–9|). FCS measurements and fluorescence images are acquired to later determine a calibration curve (steps 10|–21|). Additional 3D confocal time lapse movies can be acquired for further quantification (step 22|). This step does not require additional FCS measurements given that the same imaging settings are maintained. Data are processed to compute a calibration curve (steps 23|–29|). The images acquired in step 22| are converted to concentrations and protein numbers using the estimated calibration parameters (step 30|).

For the sample preparation it is convenient to use a multi-well glass bottom chamber containing separate wells with cells expressing the fluorescently tagged POI (mFP-POI), cells expressing a monomeric form of the fluorescent tag alone (mFP), non-expressing wild type cells (WT), and a droplet of a reference fluorescent dye with a known diffusion coefficient. The fluorescent dye is used to adjust the sample and compute the effective confocal volume V_{eff} (Box 2, steps 17|, 20| and 25|). Cells without a fluorescent tag are used to obtain the background photon counts, for correcting the FCS derived concentrations, and background image fluorescence intensity. Cells expressing the mFP alone are used to calibrate the monomeric fluorophore and estimate whether the POI oligomerizes. Furthermore, they provide a large dynamic range of concentrations and fluorescence intensities at different expression levels for estimation of the calibration parameters.

The objective as well as the bottom of the sample should be carefully cleaned and mounted to avoid any non-planarity between objective and cover-slip (steps 4| and 14|) as this leads to significant aberrations in the point spread function (PSF). Then the correction ring is adjusted and the fluorescent dye is measured (Box 2). The microscope settings, such as pixel-dwell time, laser intensity, and detector gain, are carefully chosen for FCS and imaging the mFP-POI (step 18| and 19|). The imaging settings must be kept the same for all images that need to be quantitatively analyzed and compared. The data for the calibration curve is obtained by placing 2–4 FCS measurement points at specific locations in the cell, typically at the two largest compartments, namely nucleus and cytoplasm (step 21|). We provide macros for the ZEN software to simplify the workflow and link the images with the FCS measurements (Supplementary Software 1–3). Using *FCSRunner* (step 21| Option A, Supplementary Software 1), the image and FCS measurements are automatically acquired with the previously determined imaging and FCS settings. Image pixel coordinates of the FCS measurements are automatically stored for further processing. This procedure is repeated for several cells and the three cell lines WT, mFP, and mFP-POI (Fig. 2c). With *FCSRunner* (Supplementary Software 1) the user manually selects cells and FCS measurement points (step 21| Option A). The protocol also provides an optional fully automated workflow to yield reproducible measurements and a higher throughput (Box 1, step 21| Option B, Supplementary Software 2 and 3). Given that the laser intensity remains stable and that the imaging conditions are not changed, 3D stacks and 4D movies can now be acquired from cells in the multi-well plate without additional FCS calibration measurements (step 22|).

Background counts and effective volume are computed using *FCSFitM* (steps 23|–25|, Supplementary Software 4). For the fluorescent proteins ACFs, bleach and background correction factors are computed using *Fluctuation Analyzer 4G (FA)*¹⁸ (step 26|). At the FCS measurement point the image fluorescence intensity I is calculated in a small region of interest (ROI) (*FCSImageBrowser*, Supplementary Software 4) (step 27|). The ACFs are then fitted to physical models of diffusion to obtain the number of molecules N (see Supplementary Note 2). A bleach and background corrected number of molecules N_c is computed using the previously determined factors (see Supplementary Note 3) to yield a concentration C at the FCS measurement point

$$C = \frac{N_c}{N_A V_{eff}}, \quad 2$$

where N_A is the Avogadro constant. The fitting and concentration estimation can be performed using *FCSFitM* (Supplementary Software 4) (step 28|). Finally, the data is quality controlled to correct for poor fits, measurements at the border of a compartment or on immobile structures (e.g. a nucleoporin on the nuclear envelope) (Fig. 2d and Supplementary Fig. 1) (steps 27|–28|). The concentrations are then plotted against the fluorescence intensity and fitted according to the following linear relationship (Fig. 2e):

$$C = (I - I_b)k. \quad 3$$

Here, I_b is the mean background fluorescent intensity estimated from WT cells and k is the calibration factor. The last steps can be performed using the application *FCSCalibration* (step 29), Supplementary Software 4).

The image pixel fluorescence intensities I_p are converted to concentrations C_p = using Eq. (3) and to protein number per pixel (Fig. 3a) using the physical size of the 3D voxels, characterized by the sampling x and y and the Z-slice interval z :

$$N_p = N_A C_p \Delta x \Delta y \Delta z. \quad 4$$

Fluorescence intensities conversion is automatically performed using the ImageJ plugin *FCSCalibrate* (step 30), Supplementary Software 4). See section “Quantification of FCS-calibrated images” for further details on quantifying protein numbers on cellular structures.

APPLICATIONS OF THE PROTOCOL

The method described in this protocol is generic and can be used for different protein expression systems. Nevertheless, to achieve physiological expression levels, it is preferable to tag proteins at their endogenous locus. The method is not restricted to cell monolayers, however, depending on the system, auto-fluorescence or scattering may impair reliable FCS measurements and imaging. The principles of this protocol do not restrict to the described microscopy setup and can be applied on all systems with photon counting detectors. We extensively tested and applied the method using mEGFP as FP ^{11,13,15,19} and tested mCherry as a red fluorescent protein. Other FP can also be used as long as they are monomeric and have a fast maturation time (see ²⁰ and <http://www.fpvis.org>). As alternative to mEGFP, mNeonGreen has been shown to be brighter and more photostable ²¹. The FP mCherry²² is a good compromise of high photo-stability and fast maturation, but has a rather low quantum yield. Depending on the application other proteins, such as the recent brighter mScarlet ²³, may be better suited. Brighter and more photostable proteins could allow for longer imaging and improved signal to noise ratios.

In combination with image analysis the protocol is a powerful tool to develop quantitative models of cellular processes. For instance, the protocol can be used to estimate the amount of endogenously fluorescently labelled proteins in the different cellular compartments throughout the cell cycle ¹⁵ (see Supplementary Note 4). The protocol can be applied to determine the stoichiometry and number of proteins in multi-protein complexes such as nuclear pore complexes ²⁴, kinetochores ^{25,26}, or centrosomes ²⁷. In this case such structures should be resolved as single point sources in the imaging ¹⁴ (see Supplementary Note 5). The density of proteins on membranes (Fig. 3, see Supplementary Note 6) or at the surface of macro-molecular complexes (e.g. chromosomes) ¹³ can also be estimated using this

protocol. Finally, the number of proteins bound to a large macro-molecular complex can be monitored over time (e.g. condensins bound to chromatin¹²).

LIMITATIONS

The protocol is designed to quantify live-cell images, since FCS measurements cannot be made in fixed cells. The amplitude of the autocorrelation function $G(\tau)$ decreases with the number of particles, thus the sensitivity of FCS decreases for high concentrations of the fluorophore. For a fluorophore such as GFP, concentrations up to 1–2 μM can be measured. This limitation does not apply to the imaging of fluorescence intensity where the time integrated detector signal is used. In this case concentrations above the μM range can be measured as long as the detector is not saturated. At very low fluorophore concentrations, the method is impaired by scattering and auto fluorescence. Nevertheless, by using appropriate corrections for the background, GFP concentrations as low as a few pM can be measured (Supplementary Note 3). A dynamic range of pM to μM concentrations means that over 70% of the proteome of a human cell line is accessible to this method²⁸.

This protocol uses the fluorescence of the FP tagged to the POI as a proxy for the protein distribution. It is important to test that the FP tagged protein is functional (see also accompanying Nature Protocols⁴). Furthermore, all the limitations of FPs such as potential FP induced dimerization, incomplete maturation, and photobleaching also apply to this protocol. Therefore fast maturing, monomeric, photostable, and with a high quantum yield FPs are preferable as tag. The protocol has been tested with mEGFP and mCherry, two FPs that have among the lowest tendency to dimerize, with dissociation constants above 70 nM^{22,29,30}. From the nearly 30 fluorescently tagged proteins we investigated¹⁵, the median whole cell concentration was 80 nM with a 90th percentile interval of [13–415] nM and a maximal value below 20 μM . This indicates that the number of expected dimers is negligible (maximally 1 dimer out of 10.000 proteins).

Photobleaching of the FP leads to underestimation of the number of proteins. In particular, point FCS with its long measuring times can cause strong photobleaching. To milder this issue and derive a reliable calibration curve, we suggest minimizing imaging and FCS time by acquiring one plane and 1 min of FCS measurement per cell. With the settings used for mEGFP bleaching induced by imaging was negligible; whereas the FCS induced photobleaching, estimated by the decrease in photon counts, remained small (1–5%). For mCherry, photobleaching during FCS is stronger (5–15%), however, using the correction method as described in¹⁸ and Supplementary Note 3 the effect of photobleaching can be accounted for. Therefore, with the settings used in this protocol the effect of photobleaching in defining the calibration curve is negligible. For time-lapse 4D movies the bleaching strongly depends on the application and type of imaging. With a time interval of 90 sec, 31 planes per time point (0.25×0.25×0.75 μm pixel resolution), and 40 time points we have not observed a significant photobleaching for POI tagged with mEGFP¹⁵. Higher frame rates, laser power or increased spatial resolution may cause stronger photobleaching that needs to be accounted for. To do this, in a first step, fluorescence intensities are converted to concentrations using the calibration curve. In an additional step the calibrated images are scaled using an estimated bleaching function^{31,32}.

In order to become fluorescent, FPs must undergo a maturation step that can last minutes to hours. The proteins we tested in this protocol, mEGFP and mCherry, have reported maturation times from 10–60 min^{22,23,33–35}. This delay leads to an underestimation of the number of proteins, the degree of which depends on the POI half-life. The fraction of mature fluorescent protein with respect to the total protein can be approximated by $k_M/(k_d + k_M)$ (ref.³⁵), where k_M and k_d is the rate constant for maturation and degradation, respectively. Proteins half-life ($\sim 1/k_d$) vary from over 1 h to several 10 of hours (median 46 h, 90th percentile [8–184] h;³⁶). Therefore, assuming a maturation time of 0.5 h, for 90% of the human proteins the error due to delayed maturation ranges from 0.25% to maximally 6% of the total protein amount.

This protocol is not restricted to the use of mEGFP or mCherry. Depending on the experimental system, issues with photobleaching or maturation time can also be addressed by using more stable and brighter proteins such as mNeonGreen or mScarlet^{21,23}.

COMPARISON TO OTHER METHODS

In the past, several approaches have been developed to obtain quantitative readouts from fluorescent images (reviewed by^{37,38}). With the possibility to express fluorescence proteins from their endogenous locus, these methods can now be used for proteomics studies in almost any eukaryotic cell. Typically, quantitative microscopy relies on the computation of a calibration function to convert fluorescence intensities to physical quantities. The calibration function can be computed using intra or extracellular fluorescent standards^{39,40}, or, as we present in this protocol, direct measurements of the concentrations within cells^{41–43}. Protein numbers can also be directly measured by step-wise photobleaching^{44,45}, photon emission statistics^{45–47}, or, in compartments with freely diffusing components, by analyzing fluorescence fluctuations^{5,6}.

Extracellular fluorescent standards require purified FP, the same FP as used for tagging the POI. Depending on the application, solutions of known concentrations⁴⁸ or diffraction limited complexes with known stoichiometry (e.g. Virus like particles⁴⁹) are measured along with the cells expressing the FP tagged POI. The method is relatively simple but requires additional sample preparation, knowledge of stoichiometry and concentration, and most importantly does not ensure whether the intra- and extracellular excitation and emission properties of the FP are the same³⁸. An alternative is the measurement of total protein amounts with quantitative immunoblotting^{37,39} or Mass-spectrometry^{28,36}. Using additional specialized equipment and reagents, both methods provide an estimate of the total protein amount that can be related to the total cell fluorescence. However, measurements are at the population level and the total amount of protein per cell needs to be extrapolated. Inaccuracies arise in the conversion of immuno reactivities to protein amount, estimate of total number of cells per immunoblot, and when the total protein level strongly differs between cells (e.g. cell cycle variations).

Intracellular calibration standards circumvent the issue of differences in fluorescence in an extracellular environment, but require an a priori knowledge of the stoichiometry or concentration. This method has been extensively used in yeast with proteins on the

kinetochore as calibration standard^{40,48,50}. In mammalian cells an intracellular calibration standard has not been agreed upon making the method not yet applicable.

Direct measurement of protein numbers using discrete photobleaching events is not suited for long time imaging. However, it can be used to define intracellular calibration standards⁴⁵. Counting by photon statistics requires a specialized hardware with at least four detectors and bright fluorescent probes^{46,47}. The possibility to tag proteins with organic dyes in live cells using endogenous expression of SNAP⁵¹, CLIP⁵² or Halo⁵³ tagged proteins could make the method more accessible in the future.

In this protocol the calibration curve is computed from FCS concentration measurements in cells. Compared to methods that use calibration standards the method requires a specialized confocal setup that can then be used for further imaging. Data analysis requires parameter estimation and fitting. Thanks to the availability of software solutions (e.g. QuickFit3 <http://www.dkfz.de/Macromol/quickfit>; SimFCS, <https://www.lfd.uci.edu/globals/>; the Supplementary software in this protocol Supplementary Software 1–4) the user can perform the analysis without a specialized knowledge of FCS.

EXPERIMENTAL DESIGN

Cellular samples.

For FCS-calibrated imaging three different cell line samples are used. To measure the background autofluorescence and background photon counts the cell line of interest is needed as a wild type clone (WT), i.e. not expressing recombinant FPs. Alternatively, measurements in the culture media can be taken. However, this typically underestimates the background by about 10%. To obtain concentration and fluorescence intensity measurements that span a range of up to 2 μM , cells expressing the FP alone are needed. These cells can be generated by transiently transfecting WT cells with a plasmid containing the monomeric form of the FP used for tagging of the POI. To achieve physiological expression of the POI, genome edited cells (ZNF, CRISPR/Cas9) expressing the POI tagged with the FP are preferred (see our accompanying Nature Protocol⁴). The criteria for FPs in FCS are the same as for fluorescent imaging: (i) high molecular brightness, (ii) fast maturation time, (iii) minimal oligomerization, (iv) high photo stability, (v) spectral match with available laser lines and filters. Monomeric EGFP and monomeric EYFP or other bright green/yellow FPs are well suited for FCS-calibrated imaging²¹. The FP mCherry has been tested and is suitable as red FP. Other monomeric and potentially brighter and more photostable red fluorescent proteins can also be used²³.

FCS settings.

The laser intensity should be sufficiently high to obtain good Counts Per Molecule (CPM), as this will lead to a good signal to noise ratio, but also sufficiently low to avoid photobleaching, photodamage and saturation effects during the FCS measurement. First set the fluorescence intensity for the biological sample expressing the FP only to obtain CPMs from 1 to 4 kHz with low photobleaching. In point FCS, photobleaching can be assessed by a continuous decrease in photon counts. Measuring few FCS points (2–4) per cell in

compartments where the POI is freely diffusible minimizes photobleaching. Furthermore, using *FA*, ACFs and protein numbers corrected for photobleaching can be computed (see Supplementary Note 3).

The choice of the pinhole size is a tradeoff between detected signal, sensitivity, and the quality of the fits. Larger pinholes cause a decrease in CPM leading to a decrease in the quality of the fit. This lowers the precision of the diffusion time estimate, a parameter affecting the precision of the effective confocal volume estimate. We found that pinhole sizes from 1 to 1.6 Airy Units (AU) gave reliable confocal volume estimates without impairing the quality of the fits (Supplementary Fig. 2a and ⁵⁴). Since the size of the effective confocal volume depends on the laser power and pinhole size (Supplementary Fig. 2b–c), the same parameters used for the FCS measurements of the POI must also be used for the reference dye.

Imaging settings.

We distinguish between images that are used to compute the calibration curve and images where we apply the image calibration. To define the parameters of the calibration curve, single plane images at single time-points are best suited. Saturation of the image should be avoided, as this will preclude quantification above an upper limit. Furthermore, photobleaching during the acquisition of the calibration images should be minimal. This is typically the case as only one single image per cell is required. To compute an approximation of the protein numbers in a measurement point, the method uses the 3D pixel dimensions of the imaging system. Consequently, images that need to be quantified must be 3D (XYZ) or 4D (XYZT) image stacks. In time-lapse movies, the effect of photobleaching ^{31,32} can affect the quantification. Photobleaching can be reduced by using mild imaging conditions (e.g. shorter pixel dwell-times, less Z-planes) or more photo-stable FPs ^{21,23}. It is recommended, to estimate the photobleaching parameters by fitting a function (e.g. exponential) to the total intensity changes of repeatedly acquired images. These parameters can then be used to correct the whole 4D time lapse after calibration. The spatial sampling must be high enough in order to resolve the structure of interest. For structures that are close to the size of the point spread function (PSF) or non-isotropic, such as membranes, a spatial sampling close to the Nyquist criterion should be used. Similarly to the FCS measurement, the imaging pinhole needs to be adjusted and should be small enough for good confocal sectioning (also 1–1.6 AU). If possible use the same pinhole for imaging and FCS.

Quantification of FCS-calibrated images.

The calibrated images allow an estimate of the protein distribution in specific cellular compartments, organelles or structures. For that purpose additional fluorescent reference markers of the structure of interest imaged in an independent fluorescent channel are required (see also “Markers to aid image segmentation”). For instance, to separate proteins on the nuclear envelope from proteins in the nucleus and cytoplasm a nuclear or DNA marker can be used. Quantification of total protein numbers on structures that are within the size of the PSF, such as large multiprotein complexes or membranes requires particular care ^{13,14} As shown in the supplement the fluorescent signal needs to be integrated throughout the structure in 3D (Supplementary Fig. 3). A diffraction-limited structure requires

integration of the signal in a 3D volume of the size of the PSF (Supplementary Fig. 3c–d). If the size of the PSF is known, one can approximate the total number of proteins from a subset of pixels (Supplementary Fig. 3c–d, squares and diamonds). Furthermore, spatial symmetry properties of the structure of interest can be used. For example, in the equatorial plane of the nucleus, the nuclear membrane can be assumed to be locally isotropic in Z. In this case, to obtain the density of proteins on the nuclear membrane, it is enough to estimate the protein number per unit length on a sufficiently broad nuclear rim at the equatorial plane (Fig. 3 and Supplementary Fig. 3f).

Markers to aid image segmentation.

To quantify the total number of molecules in the whole cell or in specific subcellular compartments or structures, fluorescent markers and image processing are required to define discrete volumes. Cellular compartments useful to be stained for image segmentation and further quantification are the nucleus/chromatin and the cytoplasm. The cytoplasm can be obtained from the difference between a segmented whole cell volume and the segmented nucleus/chromatin compartment. Note that fluorescent markers enabling cellular or subcellular image segmentation must not interfere/cross-talk with the FP tagged to the POI.

DNA counterstains suitable for live cell imaging are SiR-DNA⁵⁵ and Hoechst 33342. SiR-DNA has a low cytotoxicity and can be excited with a 633 nm laser allowing for long-term imaging. Instead of a chemical stain, fluorescently labeled histone 2B (H2B) can be used⁵⁶. An easy solution to segment the cell surface is to use fluorophores that stain the extracellular culture media and cannot penetrate the cellular membrane^{12,15}, such as fluorophores linked to a bulky molecule (e.g. high molecular weight dextran or IgG). This negative stain of the cell boundary has the advantage of being easily segmentable and the fluorescent dye can be adapted to the combination of DNA stain and protein of interest fluorescent probe. For instance, a suitable dye in combination with mEGFP and SiR-DNA is DY481XL. This large stokes-shift dye can be excited together with mEGFP at 488 nm but emits in the red spectral range, this reduces the need for an additional excitation laser and decreases phototoxicity. A list of fluorophores that work well in combination with the commonly used FPs mEGFP and mCherry are provided in Table 1.

Controls needed.

A serial dilution of the fluorescent dye is used to test the linearity of the imaging detector and the FCS concentration measurements. The concentration of the fluorescent dye can be estimated by measuring its absorption using the dye extinction coefficient as provided by the manufacturer. To test the linearity of the imaging detector, measure the different dilutions with the settings used for imaging the POI and plot the average intensity against the dilution factor. Similarly, perform FCS measurements and compute the concentrations (Supplementary Notes 2 and 3). A linear relationship between computed concentrations and dilution factor is expected for low dye concentrations, whereas at high dye concentrations one observes a deviation from linearity. In our setup, and with Alexa488/568, the limit was approximately reached at 2 μM . The linearity range for the imaging detector is larger than for the FCS measurements. This allows estimating the calibration parameters from cells with a low (<1–2 μM) FP concentration. In the calibration curve obtained from *FCSCalibration*

(Supplementary Software 4), deviation from linearity is easily detected by visual inspection. In order to remove cells with a high FP concentration and improve the quality of the calibration curve an upper limit for the concentration can be set in *FCSCalibration*. To test the stability of the microscopy calibration, repeated FCS and imaging data are acquired over a longer period (e.g. 12 h). Such long-term experiments can be performed using the tools for the automated imaging (Supplementary Software 2 and 3). In our setup, we did not observe systematic changes in the calibration function or CPMs over a period of 12 h.

To identify POI oligomerization, a cell line transiently expressing the mFP alone is used. A larger mean CPM in the POI cell line compared to the mean CPM measured in the mFP cell line indicates oligomerization. Inspection of the CPM distribution can be used to identify distinct populations. In *FCSCalibration*, oligomerization is considered by computing a correction factor from the ratio of the two mean CPMs. Cell autofluorescence, can be accounted for by measuring cells that do not express the FP. Finally, to test the precision of the image quantification obtained from this protocol, the user can use a priori knowledge on specific POIs. For instance, in this protocol we used nucleoporins from which we know the NPC density and stoichiometry in HeLa cells. Alternatively, the estimated whole cell protein numbers obtained with this protocol can be compared to results obtained from other quantitative methods (see Comparison to other methods).

MATERIALS

REAGENTS

Cell lines—CAUTION Cell lines used in your research should be regularly checked to ensure they are authentic and not infected with mycoplasma.

- Cell line with a FP labeled POI. Preferably use an endogenously tagged cell line. Here we show data for a HeLa Kyoto cell line with mEGFP-NUP107 (endogenously tagged using a zinc finger nuclease)⁵⁷ The cell line is available upon request.

CRITICAL The FP must be appropriate for the available laser lines and filters. This protocol has been tested with mEGFP and mCherry. Other monomeric fluorescent proteins can also be used.

- Cell line of interest as wild type clone. Here we used a HeLa Kyoto cell line. The HeLa Kyoto cells can be obtained from Dr. S. Narumiya, Department of Pharmacology, Kyoto University.

Mammalian cell culture

- High Glucose Dulbecco's Modified Eagle Medium (DMEM) (Life Technologies, cat. # 41965–039).
- Fetal Bovine Serum (FBS, Life Technologies, cat. # 10270106).
- 10000 units/ml Penicillin–Streptomycin (Life Technologies, cat. # 15140122).
- 200 mM L-Glutamine (Life Technologies, cat. # 25030081).

- NaCl (Sigma Aldrich, cat. # S5886).
- KCl (Sigma Aldrich, cat. # P5405).
- Na₂HPO₄ (Sigma Aldrich, cat. # 255793).
- KH₂PO₄ (Sigma Aldrich, cat. # S5136).
- HCl (Sigma Aldrich, cat. # H1758).

CAUTION HCl is highly corrosive and should be used with the appropriate protective equipment.

- 0.05% (vol/vol) Trypsin (Life Technologies, cat. # 25300054).

Live cell imaging

- Cell culture buffers suitable for imaging. These should be non-fluorescent and have a suitable refractive index. For the protocol presented here, we used CO₂-independent imaging medium without phenol red (Gibco, cat. # ME080051L1).

Confocal volume calibration

- Bright and photostable dyes that match the spectral properties of the FP with known diffusion coefficients are required. For example Alexa488 (Thermo Fisher, NHS ester, cat. # A20000) or Atto488 (Atto-Tec, NHS ester, cat. # AD 488–31) to match mEGFP and Alexa568 (Thermo Fisher, NHS ester, cat. # A20003) to match mCherry.

Transfection

- Plasmid expressing the FP used for tagging the POI under a mammalian promoter. Here we used pmEGFP-C1 kindly provided by J. Lippincott-Schwartz (Addgene plasmid # 54759). mEGFP is the monomeric form of EGFP with the A206K mutation.
- Opti-MEM® I Reduced Serum Medium, GlutaMAX™ Supplement (Life Technologies, cat. #. 51985–026).
- Fugene6 (Promega, cat. # E2691).

Cellular markers (optional)

- SiR-DNA⁵⁵ (also known as SiR-Hoechst) (Spirochrome, cat. #. SC007).
- Hoechst-33342 (Thermo Fisher, cat. # 62249).

CAUTION SiR-DNA and Hoechst-33342 are mutagenic and harmful if swallowed. It causes skin and respiratory irritation. It is suspected of causing genetic defects; handle it while wearing appropriate personal protective equipment. Keep it protected from light.

- Dextran, Amino, 500,000 MW (Life Technologies, cat. # D-7144).
- NaHCO₃ (Sigma Aldrich, cat. # S576).

- Dy481XL-NHS-Ester (Dyomics, cat. # 481XL-01).
- Atto430 LS NHS Ester (Molecular Probes, cat. # AD-430LS-31).
- Molecular Probes® DMSO (Thermo Fisher, cat. # D12345).

Other reagents

- Pure ethanol for cleaning the objective (Sigma Aldrich, cat. # 1009831000).

EQUIPMENT

Hardware requirements

- Scanning confocal microscope with fluorescence correlation setup. The detectors for fluorescence correlation must work in photon counting mode. For imaging the detectors must be in a linear response range (see EQUIPMENT SETUP).
- Temperature control chamber for long time imaging of living cells.
- A high numerical aperture water-immersion objective. On Zeiss systems use C-apochromat Zeiss UV-VIS-IR 40× 1.2 NA, specially selected for FCS (421767–9971-711).
- Observation chamber with coverglass bottom suitable for cell culture with at least 4 separate wells. Imaging plates 96CG, glass bottom (zell-kontakt, cat. # 5241–20), 4 to 8 well IBIDI (IBIDI, cat. # 80427, 80827), 4 to 8 well LabTek (Thermo Fisher, cat. # 155382, 155409, 155383, 155411) both #1 or #1.5 glass thickness can be used.
- For long time imaging an objective immersion micro dispenser system should be used. Leica provides a commercial water immersion micro dispenser.
- 15 cm Nunc dishes for cell culture (Sigma-Aldrich, cat. # D9054).
- PD-10 Sephadex desalting columns (optional, GE Healthcare, cat. #52130800).
- Slide A-Lyzer 10.000 MWCO (optional, Thermo Fisher, cat. # 66810).
- Viva spin 30.000 MWCO (optional, Sigma-Aldrich, cat. # Z614041–25EA).
- 0.22 µm Filters (Sigma-Aldrich, cat. # SLGP033RS).
- Whatman™ lens cleaning tissue, grade 105, 100×150 mm (GE Healthcare Life Sciences; cat. # 2105–841).
- Silicon grease for sealing (KORASILON™ paste, highly viscous Obermeier; cat.# 8000054–99).
- TetraSpeks microspheres, 100 nm (optional, Thermo Fischer, cat. # T7279).
- Workstation with Windows 7 or higher.

Software requirements

CRITICAL All custom software packages are provided as Supplementary Software 1–4. An overview of software requirements is provided in Supplementary Table 1 with links to the most recent source.

- Microscope software licensing for FCS data acquisition.
- For Zeiss microscopes, ZEN black edition (version higher or equal to ZEN 2010, <https://www.zeiss.com/microscopy/us/downloads/zen.html>) to run the VBA macros (Supplementary Software 1 and 2).
- FiJi (<https://fiji.sc>)^{58,59} installation on analysis computer.
- For high-throughput FCS and imaging data acquisition with adaptive feedback (21| Option B), FiJi installation on the computer with the ZEN software controlling the microscope.
- *Fluctuation Analyzer 4G (FA)*,¹⁸ <https://www.ellenberg.embl.de/resources/data-analysis>) on the analysis computer to compute corrected autocorrelation traces from the raw photon counting data.
- The workflow for FCS fitting (*FCSFitM*, Supplementary Software 4) is a MATLAB tool compiled for Windows. This does not require a MATLAB installation. To run the source code on a different operating system, MATLAB with the toolboxes optimization and statistics needs to be installed (R2014a or later, The MathWorks, Inc., Natick, Massachusetts, United States).
- The data analysis tool *FCSCalibration* (Supplementary Software 4) is a R tool (<http://www.R-project.org>) with a *shiny* graphical user interface (<http://cran.r-project.org/package=shiny>). We recommend to use the source code in combination with *RStudio* (<https://www.rstudio.com>). In the packaged executable version for Windows all required dependencies are installed.

REAGENTS SETUP

HeLa growth medium

High Glucose DMEM supplemented with 10 % (vol/vol) FBS, 100 U/ml Penicillin, 100 µg/ml Streptomycin and 2 mM L-Glutamine. Supplemented DMEM medium is abbreviated as complete DMEM. The growth media can be stored up to 1 month at + 4 °C. Pre-warm the solution in a + 37 °C water bath before use.

Imaging medium

CO₂ independent medium without phenol red supplemented with 10% (vol/vol) FBS, 2 mM L-Glutamine, 1 mM sodium pyruvate. Eventually add glucose to achieve a final glucose concentration as in the complete DMEM (4.5 g/liter). Filter through a 0.22 µm membrane to clear the medium of precipitants. The imaging media can be stored up to 1 month at + 4 °C. Pre-warm the solution in a + 37 °C water bath before use.

PBS, 1X

Prepare 137 mM NaCl, 2.7 mM KCl, 8.1 mM Na₂HPO₄ and 1.47 mM KH₂PO₄ in ddH₂O. Adjust the pH to 7.4 with HCl and autoclave. The solution can last several months.

SiR-DNA

Dissolve the SiR-DNA in DMSO and store at - 20 °C according to manufacturer's instructions. The solution can be thawed multiple times and last several months. Aliquoting is not recommended.

Fluorescent dyes for calibration

Dissolve the fluorescent dye in DMSO according to manufacturer's instructions. Dilute a small volume of the stock solution in ddH₂O to the desired concentration (5–50 nM). Store the dye solutions at + 4 °C. The solution lasts for several months.

Dextran labeled with a fluorescent dye

The protocol to generate dextran labeled with a fluorescent dye is given in Box 3. The reagent can be stored for > 1 year at - 20 °C.

EQUIPMENT SETUP**Detectors**

For FCS on Zeiss LSM confocal systems GaAsP or APD detectors (Confocor-3 system or in combination with a PicoQuant upgrade kit) can be used. For Leica microscopes the HyD SMD detectors or APD detectors can be used. The protocol and software is based on Zeiss LSM microscopes (LSM780, LSM880) using ZEN black edition and an inverted Axio Observer. In particular, the VBA macros only work with ZEN black edition. For imaging we tested GaAsP detectors (LSM780 and LSM880) and the AiryScan detector (LSM880).

Microscope objective

Use a water objective with a high numerical aperture and the best chromatic correction as recommended for FCS. Prior to objective purchase we recommend verifying the PSF and confocal volume, using fluorescent dyes (e.g. Alexa488 and Alexa568) and/or fluorescent beads. Use pure water (ddH₂O) as immersion media as this yields the best optical properties. Oil immersion media that match the refractive index of water only do this at a specific temperature (typically 23 °C). Thus due to refractive index mismatch the results with oil immersion media may not have the same quality as the results obtained with water as immersion media.

Supplementary Software Setup

The analysis and acquisition software is listed in Supplementary Table 1. For the installation of the custom packages (Supplementary Software 1–4) follow the instructions included in the software documentation. Most recent versions of the Supplementary Software 1–4 can be found in <https://git.embl.de/grp-ellenberg/fcsranner>, <https://gitembl.de/grp-ellenberg/mypic>, https://git.embl.de/grp-ellenberg/adaptive_feedback_mic_fiji, [*Nat Protoc.* Author manuscript; available in PMC 2019 July 05.](https://git.embl.de/grp-</p></div><div data-bbox=)

[ellenberg/FCSAnalyze](#) respectively. User guides can also be found on the respective Wiki pages, e.g. <https://git.embl.de/grp-ellenberg/fcsranner/wikis>. Install all required software prior to starting the experiment.

Installing Fluctuation Analyzer 4G (FA)

Download the software from <https://www-ellenbergembl.de/resources/data-analysis> ¹⁸. On the data analysis computer with a Windows operating system unpack the software and run the setup.

Installing FiJi

Download the latest version of FiJi from <https://fiji.sc> and install the software on the data analysis computer. For the adaptive feedback pipeline install the software on the computer that runs the software controlling the microscope.

Installing R (optional)

In case *FCSCalibration* packaged for Windows is not used or does not work (Supplementary software 4), you need to install the *R* software. Download *R* (<https://cran.r-project.org>) or preferably *RStudio* (<https://www.rstudio.com>) and install the software on the data analysis computer.

PROCEDURE

Sample preparation TIMING 1 h hands-on, 1 d waiting time, 15 min hands-on

CRITICAL To avoid contamination all steps should be performed in a laminar flow hood.

- 1| For imaging and FCS, a cell density of 50–70% confluence is recommended. Seed the cells on a glass bottom multi-well plate one day before the experiment. Prior to seeding, wash each well twice with PBS. For an 8 well LabTek chamber seed $1.6\text{--}2.0 \times 10^4$ HeLa Kyoto cells into 300 μl complete DMEM per well. Seed cells expressing the fluorescently tagged POI in one well and WT cells in two separate wells. One well should be left empty for the reference FCS measurement with a fluorescent dye.
- 2| Grow cells in complete DMEM at 37 °C, 5% CO₂. Wait for cells to attach (6 h) before transfection in Step 3.
- 3| Transfect the HeLa Kyoto cells with a plasmid containing the mFP of interest 6 h after seeding. Transfection can be carried out using FuGENE reagent. Mix 50 μl Opti-MEM with 1 μl FuGENE6 in an Eppendorf tube. Incubate for 5 min at room temperature (RT, 23 °C). Add 300 ng of plasmid DNA (we use pmEGFP-C1 as an example) and mix well. Incubate the transfection mixture for 15 min at RT. Add 25–50 μl of transfection complex to one well with seeded WT cells. Incubate the cells at 37 °C, 5% CO₂ overnight.

CAUTION Depending on the cell line transfection in the presence of antibiotics may impair cell health and transfection efficiency. If necessary, transfection can be performed in the absence of antibiotics.

CRITICAL STEP Carefully dispense the transfection mix to avoid detaching of cells or contamination of neighboring wells. The amount of cDNA, transfection reagent or the time between transfection and imaging needs to be adapted. The goal is to achieve rather low expression levels of the mFP close to the expression levels of the mFP-POI.

- 4| On the next day, before preparing the sample for imaging (steps 5| to 9|), turn on the microscope, microscope incubator and all required lasers. Allow the confocal microscope system and the incubation chamber to equilibrate and stabilize.

Depending on the setup, this can take over one hour.

- 5| Prior to imaging, change the medium to a phenol red free imaging medium. Use a CO₂ independent medium if your microscope incubator does not allow for CO₂ perfusion or if the CO₂ perfusion impairs the FCS measurements.
- 6| For the automated acquisition of FCS data and imaging data (step 21| Option B), staining of the DNA is required. In order to do this: add SiR-DNA to the cells to achieve a final concentration of 50–100 nM. Incubate for 2 h at 37 °C (without CO₂) until incorporation of SiR-DNA is complete. During the incubation time proceed with steps 7| to 20|.

CRITICAL STEP In combination with a POI tagged with mCherry, the vital dye Hoechst 33342 can be used instead of SiR-DNA. To minimize toxicity, incubate cells for 10 min with 10 nM Hoechst and replace with fresh medium. The staining and cytotoxicity of the DNA dye can differ between cell lines and imaging protocols. This should be carefully tested for example by monitoring the mitotic timing.

- 7| (Optional) If required, add any further fluorescent markers for the cell boundary (see Table 1).

CRITICAL STEP The presence of fluorescently labeled dextran in the cell culture medium reduces the staining of DNA by SiR-DNA. Cell toxicity and optimal SiR-DNA concentration should be tested in combination with the fluorescent marker in the cell culture medium.

- 8| Place a 20–50 µl drop of the diluted fluorescent reference dye (5–50 nM) in the empty chamber well.

CRITICAL STEP This fluorescent dye should have similar spectral properties to the FP of interest and with a known diffusion coefficient. In this example we use Alexa 488.

- 9| To minimize liquid evaporation during the measurements, seal the multi-well plate chamber with silicon grease or use a gas permeable foil suitable for cell culture when using CO₂ dependent imaging media. This step is recommended in case the humidity level is not sufficiently maintained in the microscope incubation chamber during data acquisition.

Microscope setup and effective confocal volume measurement TIMING 1–2 h

- 10] Ensure that the confocal microscope system and the incubation chamber are equilibrated and stabilized (Step 4). Select the beam path and emission filter/range for FCS and sample imaging, achieving the highest match between both. Select the emission filters to get highest emission signal but smallest (in the ideal case no) crosstalk between the individual fluorescence channels. Additional fluorescent channels may be needed for cellular markers (Table 1). The settings for the green (mEGFP) and red (mCherry) FPs as well as the fluorescent dyes Alexa488/568 are shown in Table 2. The emission range can be increased if only one fluorescent marker is present. Note that the protocol uses single color FCS to derive a calibration curve for a FP.
- 11] Clean the water-immersion objective and the glass bottom plate with lens cleaning tissue rinsed with pure ethanol.
CAUTION The objective can be damaged if pressure is applied to the lens.
- 12] Make a rough adjustment of the water objective correction collar to match the glass thickness based on manufacturer's specification (#1 corresponds to 130–160 μm , #1.5 corresponds to 160–190 μm).
- 13] Place a drop of pure water (ddH₂O) on the objective and mount the sample.
- 14] Check correct and planar mounting of the sample. Planarity can be tested using the glass reflection in a XZ scanning mode, with the FCS objective at the smallest zoom or with a lower magnification objective. One should observe a horizontal straight line with negligible deviation in Z (<0.1 degrees tilting) (Fig. 2a). If the reflection of the glass is not planar, adjust your sample or the sample holder accordingly.
- 15] Wait for the system to equilibrate (~30 min).
- 16] Meanwhile, create the folder structure to save the calibration data.
Your_Experiment_Folder\
Calibration\
dye\ contains the FCS measurements of the fluorescent dye
WT\ contains the images and FCS measurements of the WT cells
mFP\ contains the images and FCS measurements of the cells expressing the monomeric FP
POI\ contains the images and FCS measurements of the cells expressing the POI tagged with the FP
- 17] *Pinhole, beam collimator, and objective correction collar adjustments:* Focus the objective on the well with the fluorescent reference dye. Set the pinhole size to 1–1.6 AU. Press the count rate tab. In the ZEN software the count rate window displays average photon counts and average CPM. The latter is calculated from the photon counts divided by an estimated number of molecules. Find the

interface between the glass and the fluorescent solution. This is the Z position where the count rate and the total photon count suddenly increase. Move 30 μm above the glass-surface inside the fluorescent solution. Select the laser power to achieve a CPM above 3 kHz and below approx. 30 kHz. An upper limit of the laser power is reached when the CPM does not increase with increasing laser power. Set the laser power well below (at least 1/2) this upper limit. The final laser power will be set using the FP as reference (step 18). Close the count rate window. When using Zeiss LSM systems, perform both coarse and fine pinhole adjustments in X and Y for the channel of interest. Press the count rate tab and turn the correction collar to achieve the maximal count rate. Verify that this position corresponds to a maximal CPM.

- 18) *Setting FCS laser power using the mFP-POI:* Focus the objective in the well containing the cells expressing the FP-tagged POI. Acquire an image of the FP. Place an FCS measurement point in a region of the cell where the protein is expected to be freely diffusing, homogeneously distributed and free of aggregates (e.g. fluorescent aggregates in juxtannuclear regions) and has a low concentration. When cells exhibit variable expression levels, it is recommended to measure cells with a low expression level of the mFP-POI. Press the count rate tab and change the laser power to achieve a 4 kHz CPM \approx 1 kHz whereby the total counts should not exceed 1000 kHz. Perform a FCS measurement over 30 sec to assess the photobleaching. Photobleaching can be assessed by a decrease in photon counts over time.

CAUTION The APD and GaAsP detectors are very sensitive and can be damaged if too much light is used. Be careful not to exceed the maximal detection range. Stop acquisition if the detector shutter closes.

? TROUBLESHOOTING

- 19) *Setting imaging conditions using the POI:* These imaging settings will be used to compute the FCS calibration parameters (step 29). Set the pinhole size and laser power as close as possible to the values used for the FCS measurements (steps 17–18). Set the pixel size, pixel dwell, and the zoom to be suitable and optimal for your sample. Set the detector gain value so that saturation is avoided. Saturation can be tested by analyzing the intensity histogram or visualized using the range indicator. Set the imaging conditions to acquire a single plane (2D images and no time-lapse). This is the image used to derive the FCS calibration parameters. For 3D imaging (XYZ stacks) use an uneven number of sections in order to always associate the central Z plane with the FCS measurements. Save the settings as a user-defined configuration in ZEN.

CRITICAL STEP The pinhole size should be adjusted to achieve a good confocal sectioning for the structure of interest. The imaging settings determined in this step should be kept for further 3D or 4D imaging (step 22) and calculating concentrations and protein numbers using the FCS calibration curve.

? TROUBLESHOOTING

- 20| *Effective confocal volume measurement:* Move to the well containing the fluorescent reference dye. Focus 30 μm above the glass surface inside the fluorescent solution. Place the FCS measurement point in the center of the field of view. Use FCS settings identical to the settings for the POI (step 18|). Set the measurement time to 30 sec and 6 repetitions. Start the measurement. Save the data with the raw data for further processing to the *Calibration/dye* folder.

CRITICAL STEP It is essential that the laser power and pinhole size are the same as determined in 18|. *FA* computes a bleach-corrected ACF using the raw counting data. Thus it needs to be ensured that this data is saved. For Zeiss LSM select in the Maintain/Confocor options ‘save raw data during measurement’. In the saving directory you should find files of type raw.

? TROUBLESHOOTING

FCS and imaging data acquisition for calibration

- 21| FCS and imaging data can be obtained either by manual acquisition (Option A) or by automated adaptive feedback acquisition (Option B). See the Experimental Design section for more details.

A Manual acquisition TIMING 0.5h setup, 2h measurement

- i. *Background measurements (5–10 cells):* Start the VBA macro *FCSRrunner* (Supplementary Software 1). Specify the output folder as the *Calibration/WT* folder. Define the number of measurement points per cell. By default each point is associated to the cellular compartment nucleus/chromatin or cytoplasm (consult the software documentation for further details). For each compartment at least one point should be measured per cell. Focus in the well with WT cells and search for cells. Set the FCS and imaging settings according to the settings determined in 18| and 19|, 30 sec one repetition. Start the live mode and focus in XYZ to the cell of interest; add points to *FCSRrunner* for the different compartments. Add up to 5 cells. Press ‘Image and FCS’ in the *FCSRrunner* VBA macro to start the acquisition. The VBA macro automatically acquires and saves images, FCS measurements, and FCS measurement coordinates at each cell position. Wait for the acquisition to finish. Delete the positions in *FCSRrunner* and add new positions if more cells are needed.

CRITICAL STEP When adding FCS points to the same object/cell, do not move the XY stage or change the focus.

? TROUBLESHOOTING

- ii. *Measurement of the mFP for the FCS calibration curve (10–20 cells):* Remove previous stage positions from the *FCSRrunner*. Specify the output folder *Calibration/mFP* in *FCSRrunner*. Move the stage to the well containing the cells expressing the monomeric FP construct (mFP). Search for cells exhibiting low mFP expression levels. Switch to the FCS and imaging parameters determined in 18| and 19|, respectively, 30 sec one repetition. Verify that the image does not

saturate and that the total photon counts are low (< 1000 kHz). Start the live mode and focus in XYZ to the cell of interest; add measurement points to *FCSRrunner* for the different compartments. Add up to 5 cells. Press 'Image and FCS' in the *FCSRrunner* VBA macro to start the acquisition. Wait for the acquisition to finish. Delete the positions in *FCSRrunner* and search for more cells to measure. Measure up to 20 cells. Make sure that the FCS raw data are saved.

CRITICAL STEP The APD and GaAsP detectors are sensitive detectors. The protein expression level needs to be low in order to avoid saturation of the imaging intensities and exceeding of the FCS counts beyond 1000 kHz. Make sure that the imaging settings correspond to the imaging settings for the protein of interest. When adding FCS points to a cell do not move the XY stage or the focus.

? TROUBLESHOOTING

- iii. *Measurement of the fluorescently tagged POI (10–20 cells):* Remove previous stage positions in *FCSRrunner*. Specify the output folder *Calibration/POI* in the *FCSRrunner*. Move the stage to the well containing the cells expressing the fluorescently tagged POI. Search for cells that express the fluorescently tagged POI. Switch to the FCS and imaging parameters determined in 18| and 19|, respectively, 30 sec one repetition. Verify that the image does not saturate and that the photon counts are low (< 1000 kHz). For proteins tagged at their endogenous locus the expression levels are typically homogeneous within the population. In this case the previous step can be omitted. Start the live mode and focus in XYZ to the cell of interest; add measurement points to *FCSRrunner* for the different compartments. Add up to 5 cells. Press 'Image and FCS' in the *FCSRrunner* VBA macro to start the acquisition. Wait for the acquisition to finish. Delete the positions in *FCSRrunner* and search for more cells to measure. Measure up to 20 cells. Make sure that the FCS raw data are saved.

CRITICAL STEP Avoid acquiring FCS data in positions where the protein is bound to immobile structures (e.g. nuclear membrane). Indication of a large stable fraction is a strong bleaching during the FCS data acquisition.

? TROUBLESHOOTING

B Automated adaptive feedback acquisition (optional) TIMING 1.5 h setup (first time only), 2 h unattended acquisition

- i. *Loading the MyPiC ZEN Macro.* Start the macro (Supplementary software 2). Click on *Saving* and specify the output directory to *Calibration* directory (see 16|). Click on the *JobSetter* button.
- ii. *Starting the Automated FCS Fiji plugin.* Start *FiJI* and import the automated FCS Fiji plugin (Supplementary software 3). Run the plugin by navigating to *Plugins > EMBL > Automated FCS*.

- iii. *High-resolution imaging settings: HR.* These imaging settings are used for computing the FCS calibration curve and set the FCS measurement points using image analysis. Add an imaging channel for the cellular marker to the settings from 19]. Click the + button on the *JobSetter* to add the imaging job to *MyPiC* and name the job HR (high-resolution).

? TROUBLESHOOTING

- iv. *Low-resolution imaging settings: LR.* These imaging settings are used to automatically detect cells to be imaged and measured with FCS. Change the zoom settings in B.iii to acquire a large field of view. Adjust the pixel size and dwell time to achieve fast acquisition. The quality of the image should be good enough to allow for segmentation of the cellular DNA marker by *Automated FCS*. Use a Z stack if this improves the segmentation quality of the cellular marker. Click the + button in the *JobSetter* to add the imaging job to *MyPiC* and name the job LR (low-resolution).
- v. *Autofocus imaging settings: AF.* To speed up the acquisition and minimize the hardware load it is recommended to keep the same light path settings as in B.iv. Change the settings in B.iv to XZ line-scanning and reflection mode (see also the software manual for Supplementary software 2). Choose a laser line that is not reflected by the main beam splitter (MBS) but detected with the current detector settings. For example use the Argon 514 nm laser line with the MBS 488/561/633 and GFP detection range. Set the laser power and gain to achieve a visible reflection without saturating the detector. Set the number of stacks to cover 20–80 μm with a 100–500 nm Z step size. A small Z step size gives higher precision but requires a longer acquisition time in the absence of a piezo Z stage. Verify that the imaging yields a thin bright line. Click the + button in the *JobSetter* and name the job AF (Autofocus).

CRITICAL STEP With the recommended MBS and laser settings the transmitted light can be high and damage the detector. The user should start with a low laser light and gain and then adjust it accordingly.

- vi. *FCS settings.* Set in ZEN the FCS settings for the mFP-POI (see also 18]). Use one repetition and 30 sec measurement time. In the *JobSetter* click on the FCS tab. Load the FCS settings from ZEN by clicking on the + button and name the job POIFCS. ZEN will prompt the user to save the light path settings.
- vii. *Task 1 of the Default pipeline: Autofocus.* Click on the *Default* button in *MyPiC*. Click the + button and add the AF-job as first task to the default pipeline with a double click. Set *Process Image/Tracking* of the first task to *Center of mass (thr)* and click on *TrackZ*. At each position and repetition *MyPiC* computes the fluorescence center of mass of the upper 20% of the signal and updates the Z position accordingly.
- viii. *Task 2 of the Default pipeline: Low-resolution imaging.* Click the + button of *MyPiC* and add the LR job as second task to the default pipeline with a double click. Test the default pipeline by pressing the play button in the *Pipeline Default*

tasks frame. The image is stored in the directory *Test* with the name DE_2_W0001_P0001_T0001.lsm|czi. Adjust the value of the Z offset for the LR-job to achieve the necessary imaging position from the cover glass. Acquire a final test image. Set *Process Image/Tracking* of the second task to *Online Image Analysis*. At every position and repetition *MyPiC* first acquires the autofocus image followed by a low-resolution image. Then *MyPiC* waits for a command from the *Automated FCS FiJi* plugin (B.xii).

- ix. *Task 1 of the Trigger1 pipeline: Autofocus.* Press the *Trigger1* button in *MyPiC*. Press the + button and add the AF job as first task to the Trigger1 pipeline with a double click. Set *Process Image/Tracking* of the first task to *Center of mass (thr)* and click on *TrackZ*.
- x. *Task 2 of the Trigger1 pipeline: High-resolution imaging.* Press the + button in *MyPiC* and add the HR job as second task to the Trigger1 pipeline with a double click. Test the Trigger1 pipeline by pressing the play button in the *Pipeline Trigger1 tasks* frame. The image is stored in the directory *Test* with the name TR1_2_W0001_P0001_T0001.lsm|czi. Adjust the value of the Z offset for the HR job to achieve the necessary imaging position from the cover glass. Acquire a final test image. Set *Process Image/Tracking* of the second task to *Online Image Analysis*.
- xi. *Task 3 of the Trigger1 pipeline: FCS acquisition.* Press the + button of *MyPiC* and add the POIFCS job as third task to the Trigger1 pipeline.
- xii. *Automatically detect the cell of interest in the Default pipeline.* In the *FiJi* macro *Automated FCS* click on *Parameter Setup*. In the *Job1* column change *Pipeline* to *Default*. Change *Task* to *2* and *Command* to *trigger1*. Specify the channel of the cellular marker in *Channel segmentation*. Set in *Number of particles* the maximum number of cells to process to 4. Specify in *Pick particle* how the cells should be chosen: use *random*. Please refer to the manual of *Automated FCS* (Supplementary Software 3) for a detailed explanation of the software options.
- xiii. *Adjust image analysis settings to detect the cell of interest in the Default pipeline.* Test cell detection by clicking *Run on file*. If cells of interest are detected, an image with the processing results will be generated. To improve segmentation of the cellular marker and separate single cells, the user can change the thresholding method (*Seg. method*) and area size limits. In *Channel intensity filter1* specify the channel of the cellular marker and set a lower and upper value for the mean FI. This avoids cells with an abnormally high FI of the cellular marker (e.g. apoptotic cells). Test the settings with the *Run on file* command. In *Channel intensity filter2* specify the channel of the FP and set a lower and upper value for the mean FI. The lower value should be set so that cells not expressing the FP are avoided. The upper value avoids saturation in the high-resolution image and FCS measurements. Test cell detection by clicking *Run on file* and select the image generated in B.viii. Acquire additional images of cells expressing the mFP and endogenously mFP-POI to further test the image analysis settings.

? TROUBLESHOOTING

- xiv.** *Automatically place FCS measurements in the Trigger1 Pipeline.* In the *Job2* column of the *Parameter Setup* window change *Pipeline* to *Trigger1*. Change *Task* to *2* and *Command* to *setFcsPos*. Specify the cellular marker channel in *Channel segmentation*. Specify the number of FCS measurement points and the pixel distance from the boundary of the segmented object (*FCS pts. region/2* and *# oper. (erode <0, dilate >0)*, respectively). Use at least one point inside (*# oper < 0, nucleus*) and one point outside (*# oper > 0, cytoplasm*) of the segmented cell marker.
- xv.** *Adjust image analysis settings to place FCS points in the Trigger1 pipeline.* Verify detection of cells and the placement of FCS points by clicking *Run on file* in *Parameter Setup* and selecting the image generated in B.x. If objects of interest are detected, an image of the processing results will be generated. As for step B.xiii modify the *Seg. method*, the area size limits, the *Channel intensity filterland Channel intensity filter2* if required. If necessary, adjust the pixel distance to the object border to ensure correct placement of the FCS measurement points (*# oper. (erode <0, dilate >0)*). Acquire additional images of cells expressing the mFP and mFP-POI to further test the image analysis settings.

? TROUBLESHOOTING

- xvi.** *Automated measurement of the FP (10–20 cells).* In *MyPiC* specify the output directory to *Calibration/mFP*. Press *Default Positions*, click on *Multiple* and mark 3–10 positions in the mFP well. To improve the throughput manually select positions where several cells in the field of view express the mFP construct. In *Automated FCS* set the *Directory to monitor* to *Calibration/mFP*. Press start in *Automated FCS* and *MyPiC*. When the acquisition is finished press *Stop* in *Automated FCS*.

CRITICAL STEP When this step is performed for the first time the user should supervise the acquisition to ensure that the cells obtained from the analysis express the FP at the appropriate level. Furthermore, images should not be saturated and the FCS counts should be below 1000 kHz. See also troubleshooting on how to adjust the settings in *Automated FCS* (step B.xv).

? TROUBLESHOOTING

- xvii.** *Automated measurement of the fluorescently tagged POI (10–20 cells).* In *MyPiC* specify the output directory to *Calibration/POI*. Press *Default Positions*, delete previous positions and mark 3–10 positions in the POI well. To improve the throughput, manually select positions where several cells in the field of view express the mFP-POI. In *Automated FCS* specify the current directory to *Calibration/POI*. Press start in *Automated FCS* and *MyPiC*. When the acquisition is finished press *Stop* in *Automated FCS*.

CRITICAL STEP Same considerations as for step B.xvi apply for this step.

? TROUBLESHOOTING

- xviii. *Automated background measurements* (5–10 cells). In *MyPiC* specify the output directory to *Calibration/WT*. Press *Default Positions*, delete previous positions and mark 2–3 positions in the WT well. In *Automated FCS* set the current directory to *Calibration/WT*. Change *Channel intensity filter2* to *None*. Press start in *Automated FCS* and start in *MyPiC*. When the acquisition is finished press *Stop* in *Automated FCS*.
- 22 | *(Optional) Additional imaging*: Perform additional imaging using the same detector and laser settings as specified in 19|. The fluorescence intensity in these images can be converted to concentrations and protein number using the calibration parameters (see step 30|).

CRITICAL STEP To convert fluorescence intensities to protein numbers the image needs to be a 3D (XYZ) or 4D (XYZT) stack.

Data Processing TIMING 10 min setup, 0.5–1h of unattended computation (depending on the number of measurements), 30 min quality control, 0.5–1h image processing

- 23 | Start the tool FCSFitM (Supplementary Software 4).

CRITICAL STEP See the manual of the software for detailed explanations.

- 24 | *Computing the background photon counts with FCSFitM*. Change to tab *FCS background in FCSFitM*. Specify the directory *Calibration/WT*. Press *SelectAll* and *Compute* to calculate the average photon counts for all measurements. For the background value no distinction is made between measurements in the nucleus and the cytoplasm. Write down the counts of Ch1 (or Ch2). This will be needed in step 26|.

CRITICAL STEP In this and all subsequent analysis steps Ch1 indicates the channel with the lowest wave length (e.g. GFP, Alexa488) and Ch2 the channel with the higher wave length (e.g. mCherry, Alexa568). If APDs are used the ZEN notation of Ch2 and Ch1 corresponds in the analysis to Ch1 and Ch2, respectively.

? TROUBLESHOOTING

- 25 | *Effective confocal volume estimation from the fluorescent dye using FCSFitM*. Change to tab *Fit Dye*. Specify the diffusion coefficient of the fluorescent dye used for Ch1 or Ch2. The default values are for Alexa488 and Alexa568 at 37 °C. See also Supplementary Note 2. Specify the directory that contains the fluorescent dye data (*Calibration/dye*). Press *SelectAll* and *Compute*. Wait until the computation is finished. The program computes a best estimate for w_0 and κ by performing repeated rounds of parameter optimization. Finally a mean effective confocal volume V_{eff} is computed (Supplementary Note 1 and Box 2). Check that results have been saved in *Calibration/dye/focalVolume.txt* and *Calibration/dye/optimisedFit*.

? TROUBLESHOOTING

- 26 | *Computing correlations and corrections using FA.* Start *FA*. Specify the raw data format *Zeiss/ZEN (*.zen)*. Specify the import settings according to the used detectors. See also the manual for the Supplementary Software 4. Select the path to *Calibration* and click *Include subdirectories*. Add all measurements and name the session *2c* for later reuse. Press *Check files* and wait for all files being processed. In *Modify and correlate* set the base frequency to 100.000 Hz (i.e. 10 μ s time interval). Press *Calculate all* and wait for the end of the computation. In the *Intensity corrections* tab enter the background values obtained from 24| in the *Offset Ch1* text element. Press *Apply to* and *Calculate All*. Wait for the computation to finish. Perform a visual quality check (QC) of the data according to the requirements in Supplementary Fig. 1 and annotate traces that do not pass the QC with a 'x'. Parameter-based QC are applied at step 29|. In the *Save, export and report* tab press *Save all* button and the *FA format* button. This saves the data to the tab delimited result table *Calibration/2c.res* (see also documentation to Supplementary Software 4).

CRITICAL STEP On a Zeiss Confocor, to maintain the channel convention (see 24|) when loading data acquired using APDs you may need to swap the order of the channels (see also documentation to Supplementary Software 4).

- 27 | *Extracting fluorescence intensities at the FCS measurement points.* Start *FiJi*. Start the macro *Plugins > EMBL > FCSImageBrowser*. Press *Load .res file* and load the result table created in 26|. Click on *Report* to generate the table *2c.fint*. The table contains the fluorescence intensities in ROIs of different sizes at the FCS measurement points. If images have been acquired using the adaptive feedback pipeline, perform an additional visual QC of the images. Omit points located at the cell boundary. Annotate the FCS points that do not pass the visual QC of the images with a 'x' in *FA*. In the *Save, export and report* tab press *Save all*. Press *FA format* button to save the updated results table.
- 28 | *Fitting the ACFs functions using FCSFitM.* Press *Add res file* and load the result table from 26| into *FCSFitM*. Press *Run fit* to perform fits to the autocorrelation data. The program generates two separate tables, *lcopt.res* and *2c_opt.res* for one- and two-component anomalous diffusion models, respectively. *FCSFitM* uses the data from 25| and 27|. The generated tables summarize the fit parameters, the concentrations, and fluorescence intensities at the FCS measurement points (see documentation of Supplementary Software 4). The ACF fit results can be viewed in *FA* using the session name *lc_opt* or *2c_opt* (*lc_opt_w* and *2c_opt_w* if a weighted fit is used).

? TROUBLESHOOTING

- 29 | *Computing FCS calibration curve using FCSCalibration.* All data to compute the calibration curve are stored in the tables generated in step 28|. The *R* application *FCSCalibration* provides a graphical interface to further QC the data and automatically computes the calibration curve. Start the *FCSCalibration* and specify the directory containing the *2c_opt.res* (28|). If required, change the name of *Dye*, *POI*, and *mFP* entries to match the name of the directories of the

current experiment. Choose the ROI size and the compartment (nucleus, cytoplasm or all) to use. Choose the filtering parameters by investigating their distribution. Typically the listed default parameters work for most applications. Press *Report* to save the calibration data to the folder *Results/Calibration*.

? TROUBLESHOOTING

- 30 |** *Converting fluorescence intensities to concentrations and protein numbers.* For the linear transformation of the fluorescence intensities to concentration the user only needs the parameters of the FCS calibration curve step 29|. To convert concentrations into number of proteins the voxel volume is required, which is stored in the image metadata. Conversion in protein number per pixel is only performed for 3D and 4D data. The FiJi plugin *FCSCalibrateImage* (Supplementary Software 4) reads the image, the metadata, and the calibration data to generate a calibrated image for the channel of interest.

TIMING

Steps 1|–9|: Sample preparation: 1 h hands-on, 1 d waiting time, 15 min hands-on.

Steps 10|–20|: Microscope setup and effective confocal volume measurement: 1 h to 2 h depending on the experience.

Step 21| Option A: Manual acquisition: 0.5 h setup, 2 h measurement.

Step 21| Option B: Automated adaptive feedback acquisition: 1.5 h setup (first time only) and 2 h unattended acquisitions. When settings are reused, the setup time reduces to less than 10 min.

Step 22|: The time required for acquiring additional imaging depends on the application (2D, 3D or 4D imaging).

Steps 23|–30|: Data Processing: 10 min setup, 0.5–1 h of unattended computation (depending on the number of measurements), 30 min quality control, 0.5–1 h image processing.

Box 2: Estimate the effective confocal volume using a fluorescent dye: Acquisition and data processing 30 min.

Box 3: Generating dextran labeled with a fluorescent dye: Preparation 3h and overnight dialysis

ANTICIPATED RESULTS

The workflow described in this Protocol provides a calibration coefficient for converting fluorescence intensity data to protein concentrations and numbers. The coefficient needs to be calculated for every experiment. Depending on the stability of the system, its value may remain similar over time. In combination with further image analysis steps using image analysis software such as MATLAB or ImageJ the stoichiometry of the protein of interest on

cellular structures can be computed (Fig. 3). The adaptive feedback imaging workflow can be used to acquire additional FCS measurements and 4D image movies in high-throughput. The FCS measurements can be further used to study the biophysical properties of the POI (diffusion coefficient, aggregation). In our laboratory we used this method to generate FCS-calibrated 4D image movies of cells during mitosis¹⁵.

Figure 3 shows an example for the quantification of the protein density on the nuclear envelope. A cell line expressing NUP107, a nucleoporin which is a constitutive part of the nuclear pore complex (NPC), endogenously tagged with mEGFP has been used. Based on a DNA marker the nucleus was segmented in 3D to extract the localization of the nuclear envelope. For further processing the equatorial plane of the nucleus has been used (Fig. 3b) and the mean fluorescence intensity in the nuclear rim has been extracted. Depending on imaging settings and microscope, the value of the arbitrary fluorescent intensity units on the nuclear envelope can be significantly different (Fig. 3c). However, after conversion to absolute protein numbers using a separate calibration curve for each microscope, comparable results between the different imaging systems are obtained (Fig. 3d). The quantification of NUP107 using this method yielded a protein density at the nuclear envelope that is in good agreement with the density predicted from the NUP107 stoichiometry (32 proteins/NPC,²⁴) and the NPC density in interphase (10.66 ± 2.69 NPCs/ μm^2 ,⁵⁷) (Fig. 3d, gray shaded area).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Automated FCS calibrated imaging

This box gives an outline of the requirements for acquiring FCS and imaging data automatically. The two supplementary software *MyPiC* (Supplementary Software 2) and *Automated FCS* (Supplementary Software 3) are used. With an adaptive feedback workflow^{17,18} cells are automatically selected, according to their morphology and protein expression level, imaged and measured using FCS. This yields reproducible measurements and a higher throughput. The procedure requires a fluorescent marker, such as a DNA marker, or any other segmentable fluorescent marker, to identify cells. If the signal of the POI is sufficiently good, no additional markers are required. To perform automated acquisition three different imaging settings are used:

- A high-resolution imaging of the fluorescent marker and FP to place FCS measurements and acquire the reference image (Step 21| B.iii).
- A low-resolution imaging of the fluorescent marker and FP to identify cell of interest (Step 21| B.iv).
- An autofocus XZ scan to detect the cover glass reflection and correct for drift in Z (Step 21| B.v).

Refer to the corresponding manuals for Supplementary Software 2–3 for a detailed description. After the first setup settings can be reused. This reduces the setup time to less than 10 min.

Estimate the effective confocal volume using a fluorescent dye.

Timing 30 min This box describes how to compute the parameters for the effective confocal volume. For FCS calibrated imaging this step is particularly important since it is used to scale all measurements. Use a fluorescent dye in aqueous solution (5–50 nM, ⁶⁰) with high molecular brightness, spectral match with available laser lines and filters, spectral match with the FP of interest, and a known diffusion coefficient. To investigate proteins tagged with green/yellow FPs (e.g. GFP, YFP, mNeonGreen), Alexa488 is a suitable dye. For proteins tagged with a red FP (e.g. mCherry, mScarlet), Alexa568 is a suitable dye.

Procedure:

1. Adjust the high NA water immersion objective to the thickness of the cover slip glass by turning the correction ring (Fig. 2a).
2. Assess the optimal adjustment by measuring the counts per molecule (CPM), the photon counts, and the reflection at the immersion water glass interface in a XZ scan. At the optimal adjustment, CPM and photon counts are maximal and the reflection line is the thinnest (Fig. 2a).
3. Acquire repeated FCS measurements (up to 6, 30 sec) of the reference dye.
4. Fit the ACFs (Fig. 2b) to a one component model of diffusion (see also Supplementary Notes 1 and 2)

$$G(\tau) \sim \left(1 + \left(\frac{\tau}{\tau_D}\right)\right)^{-1} \left(1 + \kappa^{-2} \left(\frac{\tau}{\tau_D}\right)\right)^{-1/2}.$$

5. Obtain the diffusion time τ_D from the fit of the ACFs, that is the average time the fluorophore remains in the confocal volume, and the ratio of axial to lateral radius of the confocal volume κ (typically between 4–8). Use the diffusion time to compute a lateral radius

$$w_0 = 2\sqrt{D_{dye}\tau_D}.$$

The diffusion coefficients D_{dye} for Alexa488 and Alexa568 at 27 °C are $D_{A488} = 365 \pm 20 \mu\text{m}^2/\text{sec}$ and $D_{A568} = 410 \pm 40 \mu\text{m}^2/\text{sec}$ (M. Wachsmuth personal communication and ⁶¹) yielding at 37 °C a mean of $D_{A488} = 464.23 \mu\text{m}^2/\text{sec}$ and $D_{A568} = 521.46 \mu\text{m}^2/\text{sec}$ (Supplementary Note 2, Eq. S7). Diffusion coefficients for additional fluorescent dyes can be found in ⁶². Typical values for w_0 are 190–250 nm.

6. Determine the effective confocal volume by

$$V_{eff} = \pi^{\frac{3}{2}} w_0^3 \kappa.$$

The processing and fitting steps 4–6 can be performed using *FCSFitM* (Supplementary Software 4).

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Generating dextran labeled with a fluorescent dye.

Timing 3 h and overnight dialysis This box describes the protocol to label dextran with a fluorescent dye. The reagent can be used to stain the extracellular medium and provide a marker for segmenting the cellular boundary.

Procedure:

1. Dissolve 125 mg dextran, Amino, 500 000 MW, in 6.25 ml of 0.2 molar NaHCO_3 (pH 8.0–8.3, freshly prepared).
2. Mix well by stirring.
3. Dissolve 10 mg Dy481XL-NHS-Ester (Dyomics 481XL-01) in 0.7 ml of DMSO.
4. Add the Dy481XL solution directly into the dextran solution (from step 1) as quickly as possible since it is not stable.
5. Mix at RT, stirring with ~500 rpm for 1 h.
6. Use 3x PD-10 Sephadex desalting columns to separate unreacted Dy481XL following the suppliers manual and elute with 3.5 ml water per column.
7. Dialyze against water (sterile MonoQ) using Slide A-Lyzer 10000 MWCO at 4°C. Exchange the water after ~2 h. Further dialyze overnight.
8. Concentrate the labeled dextran using Viva Spin 30.000 PES as described in the supplier's manual.
9. Measure the amount at the wavelength of 515 nm and calculate the concentration of the fluorescently labeled dextran using the following equation:

$$\text{Concentration} = \frac{\text{Absorbance}}{\text{extinction} * \text{pathlength}} .$$

10. Aliquot and store at - 20 °C. The reagent can be stored for > 1 year.

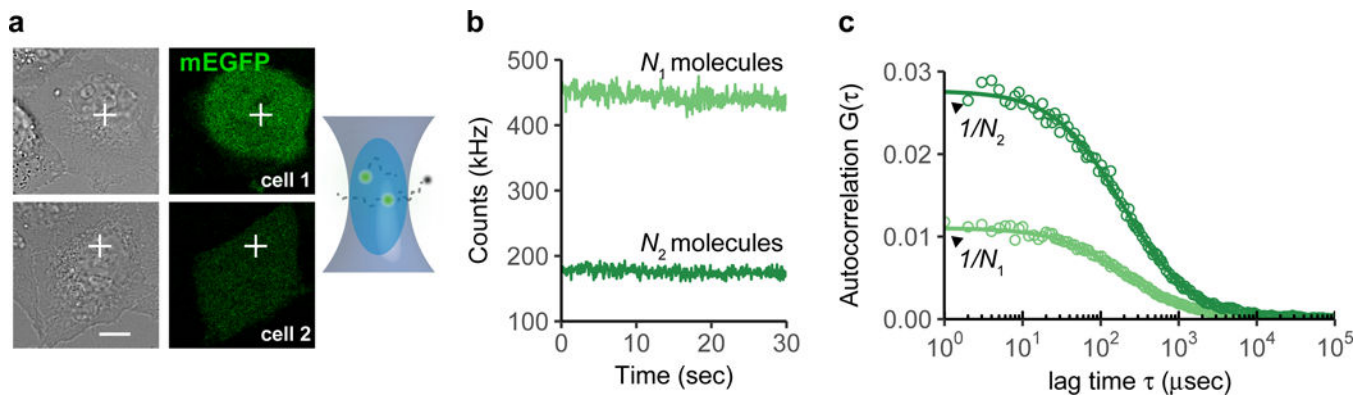


Figure 1: Principle of fluorescence correlation spectroscopy (FCS) for point confocal microscopes.

(a) The excitation laser beam is positioned to a specific location of a cell. Fluorophores entering the confocal volume are excited and the number of photons emitted is recorded. Shown are two cells expressing different levels of the fluorescent protein mEGFP. Scale bar 10 μm .

(b) The fluctuations of the fluorophores diffusing in and out of the confocal volume cause fluctuations in the number of photons.

(c) Computing the self-similarity of the fluctuations in (b) yields auto-correlation functions (ACFs). The amplitude of the ACF is directly proportional to the inverse of the number of molecules observed on average in the confocal volume.

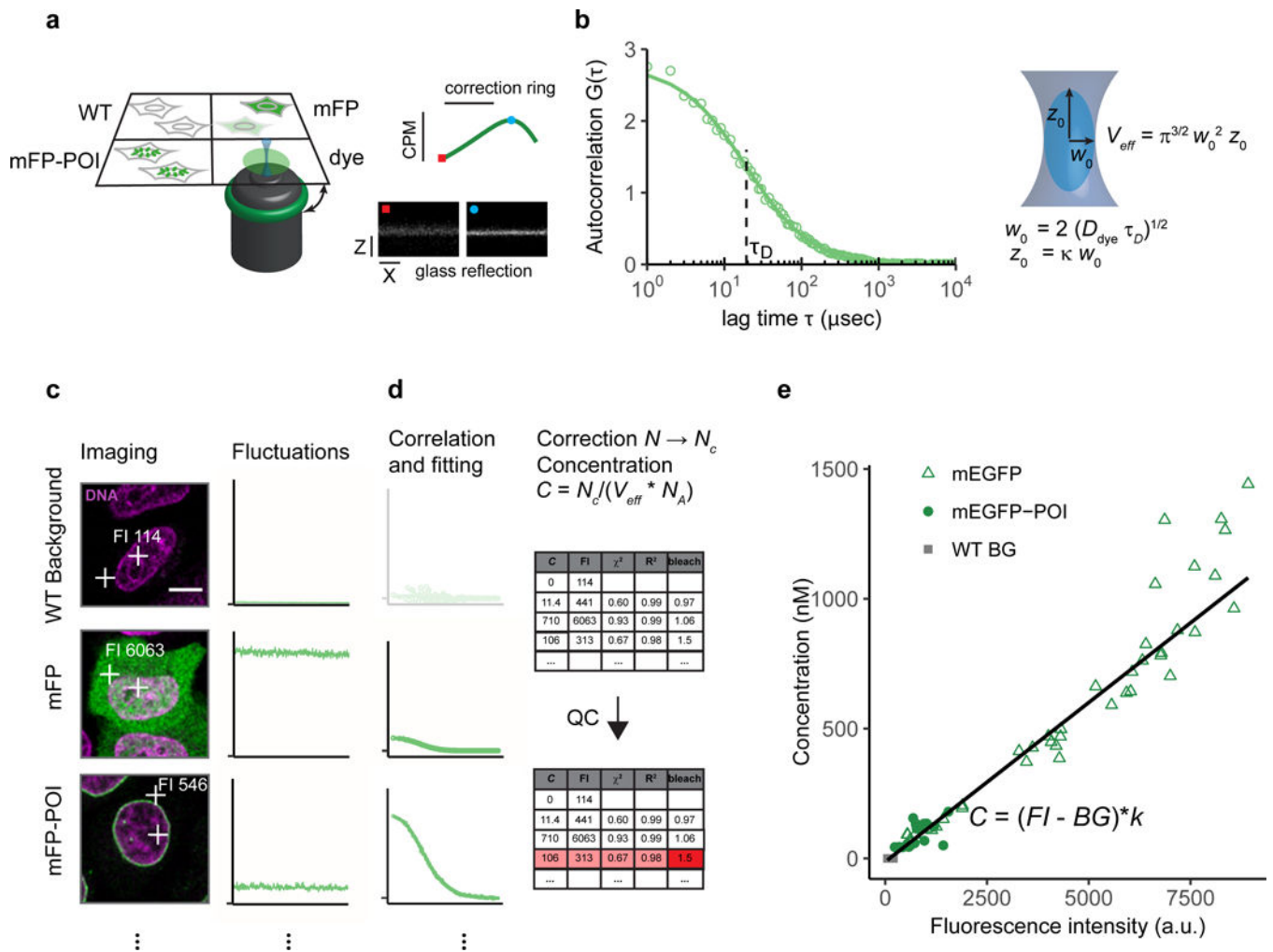


Figure 2: Workflow for FCS-calibrated imaging

(a) Setup of the multi-well chamber with four different samples and optimization of the water objective correction for glass-thickness and sample mounting using a fluorescent dye and the reflection of the cover glass. dye: fluorescent dye to estimate the effective confocal volume, WT: WT cells, mFP: WT cells expressing the monomeric form of the FP, mFP-POI: cell expressing the fluorescently labeled POI.

(b) Computation of parameters of the effective confocal volume using a fluorescent dye with known diffusion coefficient D_{dye} and similar spectral properties as the fluorescent protein. Fitting of the ACF to a physical model of diffusion yields the diffusion time τ_D and structural parameter κ . The two parameters are used to compute the focal radius w_0 and the effective volume V_{eff} (Box 2).

(c) Images and photon counts fluctuations are acquired for cells that do not express a fluorescent protein (WT), cells expressing the mFP alone (mEGFP), and cells expressing the tagged version of the POI, mFP-POI (mEGFP-NUP107). The image fluorescence intensity (FI) at the point of the FCS measurement is recorded. Scale bar 10 μm .

(d) The ACF is fitted to a physical model to yield the number of molecules in the effective volume. WT concentrations are set to 0. After background and bleach correction

concentrations are computed using the V_{eff} estimated in (a-b) and the Avogadro constant N_a . Data is quality controlled with respect to the quality of the fit and the amount of photobleaching using objective parameters.

(e) Concentrations estimated from FCS and image fluorescence intensities are plotted against each other to obtain a calibration curve (black line).

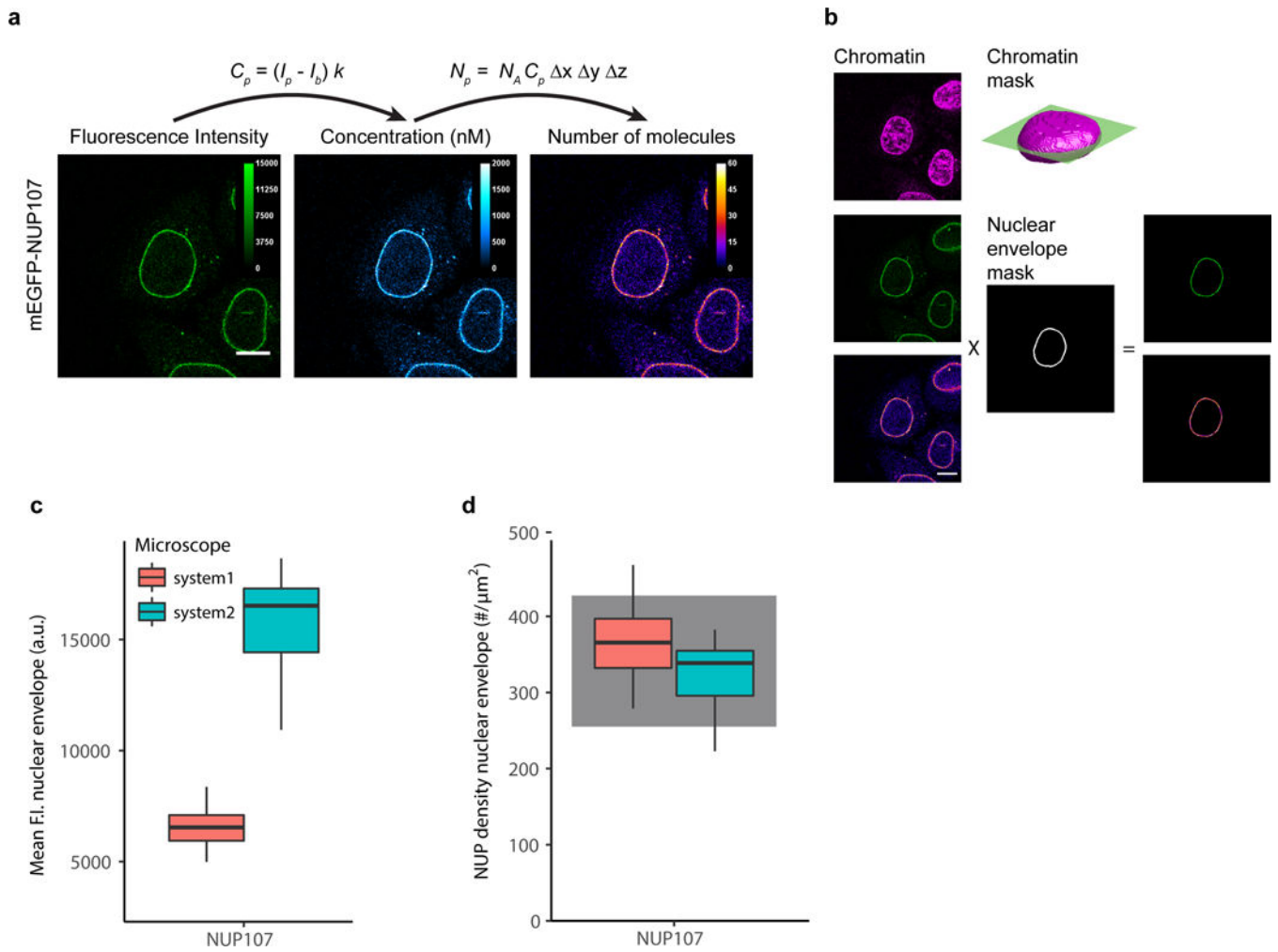


Figure 3: Example of image quantification using FCS calibration.

(a) The fluorescence intensity at each pixel, I_p , is converted to concentrations and protein numbers (for 3D and 4D images). The images are acquired with the same imaging parameters as the images used to compute the FCS calibration curve. Shown is data for a HeLa Kyoto cell line with endogenous NUP107 tagged N-terminally with mEGFP. Scale bar 10 μm .

(b) The quantitative distribution of a POI can be derived by using markers for cellular structures. DNA stained with SiR-DNA is used to compute a 3D chromatin mask of the nucleus. In the equatorial plane, a three pixel wide rim defines the nuclear envelope. Fluorescence intensities and protein numbers on the nuclear envelope can then be calculated. Scale bar 10 μm .

(c) mEGFP-NUP107 average fluorescence intensity on the nuclear envelope. Data shows results obtained on two different microscopes. The boxplots show median, interquartile range (IQR), and 1.5*IQR (whiskers), for $n = 16\text{--}22$ cells and 2 independent experiments on each microscope. System 1: LSM880, FCS and imaging using the 32 channel GaAsP detector. System2: LSM780, FCS using the APD detector (Confocor-3), imaging using the 32 channel GaAsP detector.

(d) Conversion of fluorescence intensity to protein numbers using the corresponding calibration curve obtained for each experiment and microscope system (not shown). The protein density on the nuclear envelope has been computed according to Eq. S24 (Supplementary Note 6). The gray shadowed boxes show the expected protein numbers for NUP107^{18,19}.

Table 1:

Combination of fluorescent proteins and dyes that can be used for tagging the POI and staining cellular landmarks.

Fluorescent protein for POI	Dye/fluorescently tagged protein or nuclear landmark	Dye for cell boundary landmark	Remarks
<i>mEGFP</i> Ex: 488 Em: 492–544	<i>SiR-DNA</i> Ex: 633 Em: 640–695	<i>Dy481XL-01</i> Ex: 488 Em: 640–695	
<i>mCherry</i> Ex: 561 Em: 588–690	<i>H2B-mCerulean</i> Ex: 458 Em: 458–492	<i>Atto430LS-31</i> Ex: 458 Em: 588–690	Small bleed-through of mCerulean in the Atto430LS-31 channel
<i>mEGFP</i> Ex: 488 Em: 492–544	<i>H2B-mCherry</i> Ex: 488 or 561 Em: 588–624	<i>Dy481XL-01</i> Ex: 488 Em: 625–695	Cross-talk between mCherry and Dy481XL-01 channel
<i>mCherry</i> Ex: 561 Em: 588–690	<i>Hoechst 33342</i> Ex: 405 Em: 410 – 480	<i>Alexa488</i> Ex: 488 Em: 492–544	Phototoxicity of 405 nm laser upon long-term imaging
<i>mCherry</i> Ex: 561 Em: 588–610 (small emission range to minimize bleed-through)	<i>SiR-DNA</i> Ex: 633 Em: 640–695	<i>Atto430LS-31</i> Ex: 458 Em: 588–690 or <i>Alexa488</i> Ex: 488 Em: 492–544	Bleed-through of SiR-DNA in the mCherry channel when excited with 561 nm. This impairs the FCS measurements at low protein numbers

The first 3 entries represent optimal combinations. For low expressing proteins the chromatin stain should not cross-talk into the FP channel. The emission range is given as a guideline for minimizing crosstalk. Ex: Excitation, Em: Emission.

Table 2:

Imaging and FCS settings to be used for green and red fluorescent proteins and dyes. The settings are given for Zeiss LSM microscopes.

Alexa-488/GFP settings	
FCS APD detectors	
Main beam splitter (MBS)	488, 488/561, 488/561/633
Second beam splitter (NFT)	NFT 565
Green channel emission filter (channel 2)	505–540, 505–550
FCS GaAsP detectors	
Main dichroic	488, 488/561, 488/561/633
ChS1	501–544
Zeiss LSM, Imaging detectors	
Main dichroic	488, 488/561, 488/561/633
Ch1/ChS1	492–544
Alexa-568/mCherry settings	
FCS APD detectors	
Main beam splitter (MBS)	488/561, 488/561/633
Second beam splitter (NFT)	NFT 565
Red channel emission filter (channel 1)	600–650
FCS GaAsP detectors	
Main beam splitter (MBS)	488/561, 488/561/633
ChS2	595–655
Zeiss LSM, Imaging detectors	
Main dichroic	488, 488/561, 488/561/633
Ch2/ChS2	595–655

Table 3:

Troubleshooting table.

Step	Problem	Possible reason(s)	Solution
18	High count rates (exceeding 500 kHz in FCS).	High laser power.High protein expression.	Decrease the laser power. Decrease the pinhole size.Find cells with a lower protein expression.
19	Saturation of the image.	High protein expression. High laser power.High gain.	Find a cell with lower protein expression. Reduce the laser power and/or detector gain.
20 , 21 A.ii, A.iii	No FCS raw data are saved.	<i>Maintain</i> options are not set properly.	Make sure <i>save raw data during measurement</i> is checked in the <i>Maintain</i> tab.
21 Ai, Aii, Aiii	Trend or large fluctuations in the photon counts leading to poor quality FCS measurements (e.g. Supplementary Fig. 1).	Focus drift when acquiring multiple cells.Focus drift during acquisition.FCS measurement point is close to cell boundary.	Acquire only 5 cells at a time that are close enough in space.Wait for the system to equilibrate before starting the experiment.Optimize the Z position for the FCS measurement. Start a live FCS measurement and move the Z position to reach a maximal count rate. This indicates a point far from the cell boundary.
21 B.iii	Weak staining with SiR-DNA.	Short incubation time or too low SiR-DNA concentration. The uptake may depend on the cell line.	Increase the incubation time and/or the SiR-DNA concentration. Follow manufacturer instructions to increase the staining.
21 B.xiii, B.xv	<i>Automated FCS</i> does not detect cells from the low-resolution imaging and/or high-resolution imaging.	Segmentation parameters are not properly set.Cell(s) are close to the image boundary.Poor imaging or staining conditions.	Change the segmentation parameters. Verify that the fluorescence limits match your sample and image bit depth. The default maximal fluorescence value is set for 8bit images.Increase image size.Change the imaging settings to get better imaging. Optimize the staining conditions for the cellular marker.
21 B.xvi	Image saturation of the high-resolution image. Photon counts > 1000 kHz. Imaged cells do not express the fluorescent protein. Drift in the photon counts.	Parameters in <i>Automated FCS</i> are not set correctly. Parameters in <i>Automated FCS</i> are not set correctly. Imaging Z position may be too high or too low. This often occurs for points in the cytoplasm.	Decrease the maximal value for the FI (<i>Max intensity</i>) of the fluorescent protein. Increase the minimal value for the FI (<i>Min intensity</i>) of the fluorescent protein. In <i>MyPiC</i> change the Z offset of the high-resolution imaging (Trigger1, task 2). Use the cells expressing the mFP and manually vary the Z position and verify that the photon counts are maximized.
21 B.xvii	No cells expressing the POI are automatically selected.	Low FI of the mFP-POI in the segmented area.	Decrease the minimal value for the fluorescence intensity (<i>Min intensity</i>).
24 25	List remains empty.	GUI responsiveness.	After clicking <i>Select All</i> the files name should appear.
28	Warning: "No focal volume found!...". High concentrations for low fluorescence values of WT measurements.	<i>Fit Dye</i> has not been performed. WT measurements have not been properly included.	Perform the step <i>Fit Dye</i> or click on <i>Use local volume file</i> if the confocal volume has already been calculated. The file is stored as <i>focalVolume.txt</i> in the dye directory. For WT FCS measurements make sure that the number of molecules has been set to 0. This should occur automatically if the name of the WT folder is correctly specified in <i>FCSFitM</i> .
29	No data is shown. Data strongly deviates from linear relationship. POI data does not show the same linear relationship as mFP data.	The filtering parameters are too strict. Low protein concentration. Strong bleaching.High concentration. Protein multimerization	Change or remove filtering parameters. Try to use only nucleus or cytoplasmic compartment.Set limits to the concentrations. Normalize the POI data using the mFP data. For this, activate the option in <i>FCSCalibration</i> .