Automated suppression of sample-related artifacts in Fluorescence Correlation Spectroscopy

Jonas Ries1,4,*, Mathias Bayer2, Gábor Csúcs2, Ronald Dirx3, Michele Solimena3, Helge Ewers1, and Petra Schwille4

1 ETH Zürich, Laboratory of Physical Chemistry, Wolfgang-Pauli-Str. 10, 8093 Zürich, Switzerland
2 ETH Zürich, Institute for Biochemistry, Schafmattstr. 18, 8093 Zürich, Switzerland
3 Molecular Diabetology, Paul Langerhans Institute Dresden, School of Medicine and University Clinic ‘Carl Gustav Carus’, Dresden University of Technology, 01307 Dresden, Germany
4 Biotechnological Center of the TU Dresden, Tatzberg 47-51, 01307 Dresden, Germany
* jries@ethz.ch

Abstract: Fluorescence Correlation Spectroscopy (FCS) in cells often suffers from artifacts caused by bright aggregates or vesicles, depletion of fluorophores or bleaching of a fluorescent background. The common practice of manually discarding distorted curves is time consuming and subjective. Here we demonstrate the feasibility of automated FCS data analysis with efficient rejection of corrupted parts of the signal. As test systems we use a solution of fluorescent molecules, contaminated with bright fluorescent beads, as well as cells expressing a fluorescent protein (ICA512-EGFP), which partitions into bright secretory granules. This approach improves the accuracy of FCS measurements in biological samples, extends its applicability to especially challenging systems and greatly simplifies and accelerates the data analysis.

© 2010 Optical Society of America

OCIS codes: (180.0180) Microscopy; (170.0170) Medical optics and biotechnology; (170.1420) Biology; (170.1530) Cell analysis; (170.6280) Spectroscopy, fluorescence and luminescence; (170.1790) Confocal microscopy; (170.2520) Fluorescence microscopy.

References and links
1. E. L. Elson and D. Magde, “Fluorescence correlation spectroscopy. I. Conceptual basis and theory,” Biopolymers 13 (1), 1–27 (1974).
2. R. Rigler and E. Elson, Fluorescence Correlation Spectroscopy: Theory and Applications, (Springer, 2001).
3. E. P. Petrov and P. Schwille, “State of the art and novel trends in fluorescence correlation spectroscopy, in: Standardization in Fluorometry: State of the Art and Future Challenges, (Springer, Berlin Heidelberg New York, 2007).
4. K. Bacia and P. Schwille, “A dynamic view of cellular processes by in vivo fluorescence auto-and cross-correlation spectroscopy,” Methods 29(1), 74–85 (2003).
5. T. Dertinger, V. Pacheco, I. von der Hocht, R. Hartmann, I. Gregor, and J. Enderlein, “Two-Focus Fluorescence Correlation Spectroscopy: A New Tool for Accurate and Absolute Diffusion Measurements,” ChemPhysChem 8(3), 433–443 (2007).
6. S. Kim, K. Heinze, and P. Schwille, “Fluorescence correlation spectroscopy in living cells,” Nat. Methods 4(11), 963–974 (2007).
7. K. Bacia, S. Kim, and P. Schwille, “Fluorescence cross-correlation spectroscopy in living cells,” Nat. Methods 3(2), 83–89 (2006).
1. Introduction

In Fluorescence Correlation Spectroscopy (FCS) the fluorescence fluctuations, caused by fluorophores diffusing through a small ($\approx fL$) detection volume, are analyzed in terms of their auto-correlation function. It is a powerful technique to measure local concentrations, translational and rotational diffusion coefficients, binding and reaction kinetics and photodynamics in vitro as well as in vivo [1–4]. The use of FCS has been promoted by the introduction of commercial FCS systems, so that it can now be considered as a well established technique. This holds true especially for measurements in bulk solution where FCS can reach remarkable accuracy [5]. On the other hand, measurements in cells [6, 7], biological membranes [8] or whole organisms [9] often suffer from imperfections of the system. Aggregates of fluorescent molecules or their association to vesicles result in spikes in the fluorescence intensity. Photobleaching of the auto-fluorescent background or of immobilized molecules, depletion due to photobleaching or a change in the local environment due to sample movements may lead to slow changes of the fluorescence signal in time which superimpose the fluctuations from single molecule dynamics resulting in distorted correlation curves [Figs. 1(a)–1(c)]. This complicates the determination of concentrations, diffusion coefficients and interactions of single biomolecules, the usual application of FCS.

The most common way to reduce the impact of the above mentioned imperfections is [4] (1) to take several short measurements instead of one long measurement, (2) to manually discard distorted correlation curves and (3) to fit the average of the remaining curves with a model function describing one additional mobile species. This additional component in the correlation function, sometimes combined with an overall offset, approximates the distorted part of the experimental correlation curve at larger lag-times.

This approach is not optimal. Hand-selection of curves is often the most time-consuming step in FCS data analysis and it is often ambiguous [Fig. 1(d)] with the danger of introducing a subjective bias. In addition, the distorted parts of the correlation curve can often not be described well by only one additional component. As a result, during the fit the component describing the single molecule dynamics may still contain features of the distortions. This can lead to a strong error and bias in the parameters of interest such as diffusion coefficients and concentrations of single molecules [Figs. 1(b) and 1(c)]. In addition, the introduced additional free fitting parameters can render the fit results ambiguous. Note that even in optimal in vivo systems, often a two-component diffusion or anomalous diffusion has to be assumed. Including a third component for distortions usually results in too many free parameters and indefinite fit results. Finally, large parts of the data are discarded leading to additional noise on the correlation curves.
Here we describe a simple approach for automated FCS data analysis. The idea is to divide one long measurement into many short measurements, to automatically discard parts of the data leading to distorted curves and to calculate undistorted correlation curves from the remaining part. This automation has several advantages. It avoids the subjective hand-selection and leads to more objective, simpler and faster data analysis. The use of shorter time intervals allows the extraction of a larger portion of the measured data and extends the use of FCS to very difficult systems, e.g. with many bright aggregates. Finally, it enables completely automated FCS measurements in non-ideal systems — a prerequisite for \textit{in vivo} high-throughput FCS screens.

2. Theory

2.1. Model functions

In FCS, the fluorescence intensity $I(t)$ is recorded with a high temporal resolution. From this signal, the auto-correlation curve $G(\tau)$, which measures its self similarity, can be calculated:

$$G(\tau) = \frac{\langle \delta I(t) \delta I(t+\tau) \rangle}{\langle I(t) \rangle^2}$$  \hspace{1cm} (1)

Here $\langle \rangle$ denotes the time average, $\delta I(t) = I(t) - \langle I(t) \rangle$. $\tau$ is called the lag time. The calculation of the auto-correlation curve from the fluorescence intensity [Eq. (1)] can be performed efficiently on a quasi-logarithmic time scale with a ‘multiple tau’ correlation algorithm [10].

To obtain the parameters of interest, the auto-correlation curve is fitted with a mathematical function which takes into account the sources for the intensity fluctuations. A commonly used model for pure Brownian diffusion through a three-dimensional Gaussian detection volume is [3]:

$$G(\tau) = \frac{1}{N} \left( 1 + \frac{\tau}{\tau_D} \right)^{-1/2} \left( 1 + \frac{\tau}{S^2 \tau_D} \right)^{-1/2} = \frac{1}{N} GD(\tau)$$  \hspace{1cm} (2)

Here $N = V_{\text{eff}}C$ is the number of particles in the detection volume $V_{\text{eff}} = \pi^{3/2} Sw_0^3$, $w_0$ is the $1/e^2$-radius of the laser focus and structure parameter $S = w_z/w_0$ measures the aspect ratio of the Gaussian detection volume. $\tau_D = w_0^2/4D$ is the diffusion time and a measure for the diffusion coefficient $D$.

For two diffusing species with different diffusion times $\tau_{D1}$ and $\tau_{D2}$, taking into account triplet/blinking kinetics and assuming the same brightness of the molecules the correlation function is:

$$G(\tau) = \frac{1}{N} \left( 1 + \frac{T}{1-T} e^{-\tau/\tau_\tau} \right) (FG_{D1}(\tau) + (1-F)G_{D2}(\tau))$$ \hspace{1cm} (3)

$N$ is the total number of particles in the detection volume, $F$ denotes the fraction of molecules with the diffusion coefficient $D_1$, $T$ is the fraction of the molecules in the dark state and $\tau_\tau$ is connected to the lifetime of the dark state [3]. Often, this formula is also used if the molecular brightnesses of the two components are not equal (e.g. free fluorophores and bright vesicles). In this case $N$ is not directly related to the concentrations any longer.

2.2. Automated analysis of FCS data

The typical steps involved in hand-selection of distorted curves are: acquisition of several curves (e.g. $10 \times 10$ s); comparison of the curves; rejection of distorted curves, identified by a deviation from the majority of the other curves; and calculation of the average of the undistorted curves. Our approach attempts to formalize and automate these steps. Accordingly, the
Fig. 1. **FCS measurements in complex systems.**

**a:** Image of ICA512-EGFP in Ins-1 cells. Bright secretory granules are clearly visible.

**b:** Fluorescence intensity measured on ICA512-EGFP. Bright granules cause spikes in the fluorescence intensity and bleaching of an auto-fluorescent background or depletion of fluorophores lead to a slow change in the intensity over time. As a result, the auto-correlation curve is severely distorted (○). Even a two-component fit (—) cannot accurately resolve the fast diffusion of the single proteins ($\tau_D = 1.0 \text{ ms, } N = 4.2$). A common way to reduce the distortions is to acquire several short correlation curves and discard distorted curves by hand-selection. The average of the best two curves calculated on 10 s long intervals of the intensity (○) resembles the control measurement (cell without bright vesicles, amplitude normalized for visualization) much better, albeit with larger noise on the curve.

**c:** 10 curves of 10 s. Obviously distorted curves (···) are discarded and the average of good curves (—) is used for further analysis ($N = 10.5, \tau_D = 0.73 \text{ ms}$). The inclusion of ambiguous curves (−·−) changes the result of the fit ($N = 13.0, \tau_D = 0.99 \text{ ms}$).
algorithm we propose consists of the following parts: calculation of correlation curves corresponding to short intervals of the fluorescence intensity $I(t)$, ordering of the curves based on their deviation from the other curves and averaging of the curves with the smallest deviation:

1. Division of the fluorescence intensity trace $I(t)$ of length $T_M$ into $n = T_M/\Delta T$ short intervals (time windows) $I_k(t)$ of length $\Delta T$. $\Delta T$ can be much smaller and $n$ much larger than practical for hand-selection.

2. Calculation of $n$ correlation curves $G_k(\tau)$ from the short intensity traces $I_k(t)$ with a reduced time resolution using a multiple tau correlation algorithm. The choice of a minimal lag-time $\tau_{\text{min}} \approx 0.1 \tau_{D1}$ has the advantage that calculation times are reduced and that the parts at smaller $\tau$, where the shot noise (random noise on the curve) dominates the correlation curve, is not considered for the following ordering step.

3. Ordering of the curves according to their deviation from the average:
   (a) Make a list of all curves.
   (b) For all $k$ compare $G_k(\tau)$ with the average of all other $G_{j\neq k}(\tau)$ in the list. As a measure for the difference we use:
   $$dG_k := \langle (G_k(\tau) - \langle G_j(\tau) \rangle_{j \neq k})^2 \rangle_i$$
   (4)
   $\langle \rangle_{j \neq k}$ denotes the average over all curves $j \neq k$, $\langle \rangle_i$ is the average over all lag times $\tau_i$.
   (c) Determine maximum difference $dG_m = \max(dG_k)$.
   (d) Store $dG_m$ and the index $m$.
   (e) Remove $m$ from the list.
   (f) Continue with step 3b until the list is empty.

At the end of step 3 all curves are sorted according to their quantitative deviation from the average.

4. Chose maximum allowed $dG^{\text{max}}$. How to chose $dG^{\text{max}}$ will be discussed in more detail below. After this step we have eliminated the irregular curves.

5. For all $dG_m < dG^{\text{max}}$ calculate the corresponding correlation curves $\tilde{G}_m(\tau)$ with the full time resolution.

6. Average all $\tilde{G}_m(\tau_i)$.

We implemented the algorithm based on the raw data of photon arrival times, but it can be implemented equally well if many short correlation curves are acquired, e.g. by using a hardware correlator.

A larger usable part of the data can be obtained by choosing overlapping time intervals, e.g. by using 1 s time intervals at 0.5 s spacing. Stronger oversampling (smaller spacing) further increases the usable part of the data. However, this comes at the expense of increased calculation times.
2.3. **Length of time interval \( \Delta T \)**

How long should the time interval \( \Delta T \) be? A small \( \Delta T \) increases the number of correlation curves and therefore the usable portion of the data. In addition, a residual change of the average intensity during \( \Delta T \) will be reduced. However, when \( \Delta T \) is too small the shot noise becomes the dominating noise on the correlation curve and conceals the distortions. As a result the sorting algorithm fails. Also, traces that are too short (\( \Delta T \ll 10^5 \tau_D \), where \( \tau_D \) is the timescale of interest, e.g. the diffusion time of the single molecules) result in a systematic bias \([11]\). Therefore, a \( \Delta T \) should be chosen which is large enough that this bias is avoided and for which the shot noise is not larger than the deviations due to distortions on the time scale evaluated during the sorting algorithm. Visual inspection of correlation curves from a typical measurement, calculated on different time intervals \( \Delta T \), will help finding an optimal value for this parameter.

2.4. **Maximum difference \( dG_{\text{max}} \)**

The parameter \( dG_{\text{max}} \) defines what distortions are still allowed. A large \( dG_{\text{max}} \) results in better statistics and lower noise on the correlation curves at the expense of a larger influence of distortions. If \( dG_{\text{max}} \) is so small that only a few curves are left, the average curve will be very noisy and a possible bias can occur [Figs. 2(e)–2(g)].

To determine the optimal \( dG_{\text{max}} \) we suggest plotting the fitted parameters of interest (e.g. \( \tau_D \), \( N \)) in dependence of \( dG_{\text{max}} \) for a few curves of a dataset and to determine the range in which these parameters are constant [Figs. 2(e)–2(g)]. Another option is to measure homogeneous control samples to determine the range of naturally occurring \( dG_m \). We found that the optimal \( dG_{\text{max}} \) was usually about one order of magnitude above the minimum of \( dG_m \) [Fig. 2(h)]. It is a merit of this approach that the dependence of the parameters of interest (e.g. \( \tau_D \), \( N \)) on the exact choice of \( dG_{\text{max}} \) is very small.

3. **Results and Discussion**

3.1. **FCS on Streptavidin-Atto565 with fluorescent beads**

As a well defined test system to investigate the performance of the automated analysis algorithm we chose a 2 nM solution of Streptavidin-Atto565 contaminated with 100 nm fluorescent beads which mimic bright vesicles or aggregates. The transit of the bright beads leads to spikes in the fluorescence intensity [Fig. 2(a)]. These spikes occurred so frequently that all curves calculated on 10 s parts of the intensity trace were affected. Therefore, even hand-selection could not recover undistorted correlation curves in this case. Auto-correlation curves, calculated on longer (10 s) or on shorter (1 s) intervals, showed severe distortions [Fig. 2(b)]. The two-component fits, although visually acceptable, resulted in diffusion times (\( \tau_{D1} = 0.09 \) ms, 10 s intervals and \( \tau_{D1} = 0.10 \) ms, 1 s intervals) very different from the control consisting of only Streptavidin-Atto565 without beads (\( \tau_D = 0.185 \) ms). The curve extracted with the automated selection algorithm described above resulted in a correlation curve hardly distinguishable from the control (\( \tau_D = 0.185 \) ms).

Measurements on samples with varying amounts of fluorescent beads exhibited a large spread in the diffusion times of the uncorrected curves (\( \tau_D = 0.76 \pm 0.95 \) ms, 10 s interval and \( \tau_D = 0.16 \pm 0.09 \) ms, 1 s interval), whereas the automatically processed curves lead to very reproducible parameter estimates (\( \tau_D = 0.181 \pm 0.007 \) ms, \( F = 0.953 \pm 0.016 \)). Even for the samples with a large amount of beads, where only 10% of the data could be used to construct the final correlation curve, the fit parameters were consistent (\( \tau_D = 0.180 \) ms, \( F = 0.951 \)).
3.2. FCS on ICA512-EGFP in Ins-1 cells

To demonstrate the automated selection algorithm on biological samples we chose a Ins-1 cell line [12] stably expressing ICA512-EGFP [13], a protein that partitions into secretory granules. These granules are bright, slowly moving entities that cause spikes in the fluorescence intensity [Fig. 2(c)]. In addition, the average intensity decreases slightly over time due to depletion of ICA512-EGFP and the bleaching of immobile fluorescent proteins and the auto-fluorescent background. Also here the resulting correlation curve was severely distorted [Fig. 2(d)]. The automated selection algorithm resulted in a curve with greatly reduced distortions, although the comparison with the control (ICA512-EGFP in a cell showing only very few secretory granules) revealed a residual slow component. The reason could be dim granules diffusing through the periphery of the detection volume which were not bright enough to cause distortions comparable to the residual noise on the curves. However, compared to the uncorrected curves the slow component was reduced from 75% (10 s intervals) and 57% (2 s intervals) to 15% and had a smaller impact on the parameter estimates.

3.3. Dependence of fit parameters on \( dG_{\text{max}} \)

To investigate the influence of the exact choice of \( dG_{\text{max}} \) on the parameter estimates we used the selection algorithm to order the curves calculated on short intervals of the fluorescence intensity, determined the average of a fraction of the curves with the smallest \( dG_m \) and fitted this average with a two-component fit including triplet (Streptavidin-Atto565 with fluorescent beads and ICA512-EGFP with bright granules) or a one-component fit with fixed triplet (controls).

For the control samples, the parameter estimates hardly depended on the fraction of used curves [Figs. 2(e)–2(f)]. Only the use of a fraction which was too small, as when it was comprised of just a handful of remaining curves, lead to high noise on the average curve and less accurate parameter estimates. The parameter \( dG_m \), a measure for the difference between the last curve used and the average of the other curves [Eq. (4)], spanned about one order of magnitude [Fig. 2(h)]. This can be considered the variation of \( dG_m \) in samples not influenced by distortions, suggesting a choice of \( dG_{\text{max}} \approx 10 \times \min(dG_m) \).

For measurements on Streptavidin-Atto565 with fluorescent beads and on ICA512-EGFP with bright granules \( dG_m \) spanned several orders of magnitude [Fig. 2(h)]. Due to the higher number of free fitting parameters (in addition to \( N \) and \( \tau_D \) in the control case now also the triplet fraction \( T \), the fast fraction \( F \) and the diffusion time of the slow fraction \( \tau_{D2} \) were varied during the fit), their variation was in general increased. Apart from this noise, \( N \) and \( \tau_D \) remained constant over a surprisingly large range of \( dG_{\text{max}} \), even for values of \( dG_{\text{max}} \) significantly larger than the suggested \( \approx 10 \times \min(dG_m) \) where the slow fraction became significant. Only inclusion of more than 70% of the curves lead to a significant deviation of the parameters.

Compared to the control, a systematic variation of \( \tau_D \) was visible already much earlier when more than 70% of the curves were discarded. The reason was mainly due to the triplet/blinking part: if fewer curves are considered, the higher noise on the curves results in a poorly defined triplet/blinking fraction \( T \). Accordingly, in this range \( T \) showed strong deviations (data not shown). Fixing \( T \) during the fitting reduced these deviations. However, this also decreased the range over which \( \tau_D \) remained unaltered — due to the large number of fluorophores in vesicles or fluorescent beads these entities do not exhibit a significant triplet/blinking. If their contribution to the correlation curve becomes larger, the overall triplet/blinking fraction \( T \) is reduced.

3.4. Limitations

The suggested method will ease the data analysis in most cases where hand-selection is otherwise required. The density of bright events limits the length of the time intervals \( \Delta T \), since
only unaffected parts of the data are used to construct the final curve. If the density of bright aggregates or vesicles is so high that every time interval is affected, this approach fails. On the other hand, a minimum length of the time windows is required to reduce the shot noise on the correlation curves sufficiently, such that distortions become detectable. If bright events are sparse, the time windows can be chosen longer and samples with lower molecular brightness and hence stronger noise on the correlation curves can be analyzed.

A strong change in the mean intensity (depletion due to photobleaching, bleaching of a background) leads to distortions even if curves are calculated using short time windows. In addition, the individual curves will exhibit different amplitudes and might be discarded by the algorithm since they deviate from the average correlation curve. In this case, a correction of the raw intensity trace [8] might be helpful. However, care must be taken that bright events are excluded from the intensity trace when calculating its smoothed approximation.

Due to rejection of the parts of the data affected by bright aggregates, information about this slowly mobile species is lost. In addition, the short time intervals lead to a cut-off of slow dynamics. Although of potential interest, this component is usually not accessible by FCS since rare events require immensely long measurement times for statistical accuracy, otherwise repeated measurements will exhibit a large spread. To recover some information about the bright events, additional methods like counting the spikes and comparison of the average fluorescence intensity of the selected parts with the overall intensity can be attempted.

3.5. Comparison to other approaches

Although hand-selection of undistorted curves is not optimal, there are only a few reports on automated FCS data analysis. This might be partially due to the complexity of previous approaches with a set of empirical parameters which had to be adapted to the experimental conditions. These methods rely on judging the quality of the fit and rejection of those curves with irregular residuals [14, 15]. They require individual curves of reasonably low noise and therefore comparably long time windows $\Delta T$. The influence of the exact choice of the parameters and the model on the outcome of the selection is not clear. A direct selection of curves prior to fitting might therefore be more suited to distinguish between different possible models.

A straight-forward approach to reject distortions due to single bright events is the identification of peaks in the intensity. We tested an algorithm which rejects short intervals if the maximum of the intensity, sampled at a time resolution $t_b$, is larger than a cut-off $I_{\text{max}}$. The performance of this algorithm in rejecting distortions was only slightly below that of the sorting algorithm presented in this work, but required a careful choice of $t_b$ and $I_{\text{max}}$ for every set of measurements. With parameters $t_b$ and $I_{\text{max}}$ optimized for every curve, the peak-finding algorithm resulted in a fast fraction $F$ a few percent lower than that obtained with the sorting algorithm. $\tau_D$ was similar within the fitting error. Apart from its dependence on empirical parameters a drawback of the peak-finding algorithm is that it might fail to detect peripheral transits or low brightness vesicles, mistake shot noise in the intensity for an irregular event and does not remove distortions due to instabilities or photobleaching.

The new sorting approach presented here has a good performance in rejecting distorted curves, it requires only short time windows $\Delta T$ and can therefore sample large parts of the data and, most importantly, it does not depend strongly on empirical parameters which renders it easy to use.

4. Materials and Methods

4.1. Fluorescence Correlation Spectroscopy

All FCS measurements were performed on a Zeiss LSM 710 Confocor 3 System using a 40 x NA 1.2 UV-VIS-IR C Apochromat water-immersion objective. For measurements on
Fig. 2. a,b: FCS on Streptavidin-Atto565 with 100 nm fluorescent beads. The beads cause spikes in the fluorescence intensity (a), leading to a distorted correlation curve (b, □, fit: $\tau_{D1} = 0.09$ ms). The average of many correlation curves calculated with short 1 s time intervals is less distorted (○), but the fit values ($\tau_{D1} = 0.10$ ms) still differ significantly from the control measurement consisting only of Streptavidin-Atto565 (+, $\tau_{D1} = 0.185$ ms). The automated selection algorithm discards the parts of the intensity trace with spikes (blue in a), the corresponding correlation curve (○, $\tau_{D1} = 0.185$ ms) is hardly distinguishable from the control (amplitude normalized for illustration). c,d: FCS on ICA512-EGFP in Ins-1 cells. The ICA512-EGFP partitions partially into secretory granules which cause spikes in the fluorescence intensity (c). In addition, bleaching of the fluorescent background and depletion of ICA512-EGFP due to photobleaching results in a decay of the average intensity. d: The correlation curves, calculated on 10 s long intensity traces (□) and with short 2 s parts of the intensity trace (○), are severely affected. The automated selection algorithm greatly reduces distortions (○), although some residual slow component is visible when compared to the control (+), measured on ICA512-EGFP in cells showing only very few secretory granules and amplitude normalized to the corrected curve. e-h: Dependence of fit parameters on $dG_{max}$. Curves were ordered according to their deviation from the other curves $dG_m$ and a fraction of the curves with the smallest $dG_m$ was averaged. Results from the fit of these average curves (e: number of particles $N$ normalized for illustration, f: diffusion time of the fast component $\tau_{D1}$ and g: fast fraction $F$) and the maximum $dG_m$ (h) were plotted in dependence on the fraction of curves used for the average.
Streptavidin-Atto565 (Atto-Tec, Siegen, Germany) mixed with 100 nm fluorescent beads (FluoroSpheres 505/515, Molecular Probes, Eugene, OR) 6.4 μW of the 561 laser line and 3.2 μW of the 488 laser line were used. For measurements on ICA512-EGFP 8.1 μW of the 488 nm laser line were used. The raw data of photon arrival times were stored and further processed.

4.2. Data analysis

Data analysis routines, as described in the section 'Automated analysis of FCS data' were implemented with MATLAB (Mathworks, Natick, MA) with some time critical routines (binning of the photon arrival times, multiple tau correlation algorithm) written in C. FCS curves were fitted with a nonlinear least squares fitting algorithm. The time required to automatically analyze a typical 100 s measurement is a few seconds. The majority of the time is required to calculate the final correlation curve with a high temporal resolution.

4.3. Ins-1 cells

Ins-1 cells stably expressing ICA512 fused to the n-terminus of EGFP were described previously [13]. Cells were cultured in complete media (RPMI supplemented with 10% heat inactivated fetal calf serum, 10 mM HEPES pH 7.4, 2 mM L-glutamine, 100 U/ml penicillin, 100 mg/ml streptomycin, 1 mM sodium pyruvate, 50 mM 2-ME and 600 mg/ml G-418 Sulphate) at 37 °C, 5% CO2 in a humidified incubator. For FCS measurements, cells were plated in complete media on 24 mm glass cover slides. Measurements were performed at room temperature (25 °C) in extracellular solution (ECS, 145 mM NaCl, 5 mM KCl, 2 mM CaCl2, 1 mM MgCl2, 10 mM HEPES, 10 mM Glucose) supplemented with 0.2 % bovine serum albumin at pH 7.4.

5. Conclusion

We presented a method to automatically analyze FCS data sets and to discard corrupted parts which otherwise lead to distortions of the correlation curves. This method replaces the time-consuming and subjective hand-selection of distorted curves, opening up a way to high-throughput FCS screens on biological samples. In addition, by evaluating short parts of the measurement, it extends the applicability of FCS to very noisy systems where even hand-selection does not result in undistorted curves.

The rejection of distorted curves is determined by essentially one parameter. Since the results depend only weakly on the choice of this parameter, it does not have to be adapted to the individual measurement. This renders the application of this method very easy with a low risk of introducing additional errors. Especially when implemented into the analysis software of commercial FCS instruments, this method will simplify FCS measurements in biological samples and further promote the application of this technique.

Acknowledgements

This work has been partially supported from the DFG (grant SO 818/1-1) to MS and PS.