# Cross Talk Free Fluorescence Cross Correlation Spectroscopy in Live Cells

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ABSTRACT Fluorescence correlation spectroscopy (FCS) is now a widely used technique to measure small ensembles of labeled biomolecules with single molecule detection sensitivity (e.g., low endogenous concentrations). Fluorescence cross correlation spectroscopy (FCCS) is a derivative of this technique that detects the synchronous movement of two biomolecules with different fluorescence labels. Both methods can be applied to live cells and, therefore, can be used to address a variety of unsolved questions in cell biology. Applications of FCCS with autofluorescent proteins (AFPs) have been hampered so far by cross talk between the detector channels due to the large spectral overlap of the fluorophores. Here we present a new method that combines advantages of these techniques to analyze binding behavior of proteins in live cells. To achieve this, we have used dual color excitation of a common pair of AFPs, ECFP and EYFP, being discriminated in excitation rather than in emission. This is made possible by pulsed excitation and detection on a shorter timescale compared to the average residence time of particles in the FCS volume element. By this technique we were able to eliminate cross talk in the detector channels and obtain an undisturbed cross correlation signal. The setup was tested with ECFP/EYFP lysates as well as chimeras as negative and positive controls and demonstrated to work in live HeLa cells coexpressing the two fusion proteins ECFP-connexin and EYFP-connexin.

# INTRODUCTION

The method of fluorescence correlation spectroscopy (FCS) analyzes the fluctuations of the fluorescence intensity of molecules diffusing in and out of the subfemtoliter volume of a strongly focused excitation beam in a confocal microscope. From the correlation function of the fluctuations it is possible to derive the concentration, diffusion coefficient, and intra-molecular properties of the fluorophore influencing its fluorescence properties. After the introduction of FCS in the early 1970s (1,2) it took two decades before the method became popular especially due to the improvement of the signal/noise ratio and detector sensitivity down to the single molecule level (3).

Life science applications are investigated to an increasing degree: binding studies of protein subunits (4), molecular identification (5), observation of conformation changes (6,7), or microsecond protein dynamics (8).

Due to its noninvasive character, FCS studies were also rapidly expanded to live cell investigations (9–11). Applications include protein oligomerization (12), diffusion processes in the nucleus (13), or determination of the hybridization state of oligonucleotides (14). Other studies have focused on technical aspects of working in live cells such as cellular autofluorescence (15) or the confinement of the excitation volume (16).

The extension of FCS, fluorescence cross correlation spectroscopy (FCCS), deals with the problem of measuring the colocalization of two diffusing species on a molecular level. In FCCS, two species must be labeled with spectrally distinct fluorophores and the fluctuations of the fluorescence must be

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recorded in two separate channels. In the absence of cross talk, the cross correlation of the two channels reveals whether the two species are linked to one another or not. Hence, FCCS deals with a colocalization on a molecular level sometimes referred to as codiffusion.

FCCS has been applied to test for irreversible association kinetics of DNA renaturation (17), aggregation of prion proteins (18), vesicle fusion (19), determination of the gene expression by quantification of mRNA (20), simultaneous binding of two DNA duplexes to the NtrC-enhancer complex (21), and enzyme kinetics (22) where the method is expanded to a high throughput screening tool (23).

Though FCCS is suited for a number of unsolved questions in cell biology, there are only a few reports of in vivo studies to date. Bacia et al. have shown that subunit A and B of cholera toxin do not separate along the endocyclic pathway until reaching the Golgi apparatus (24). Kim et al. studied the binding kinetics of calmodulin and calmodulin-kinase II in HEK293 cells (25).

Particularly, the latter report suggests the possibilities of FCCS in the field of cell signaling. Most signaling cascades comprise various receptors, first and second messengers building a highly complex network where diffusion of molecular components and binding-unbinding reactions between these components are the crucial parameters for understanding the cellular response. Due to its single molecule sensitivity, FCCS studies can be performed with physiological concentrations of the molecular compounds (normally proteins) that participate in a signal cascade. It is, therefore, possible to avoid artifacts from overexpression, which might perturb the delicate balance within the cascade.

The main reason for the lack of cross correlation studies in live cells is related to the different photophysical

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requirements on the fluorophores used. Today, the dominant fluorophores for cell biology applications are autofluorescent proteins (AFPs; (26)). AFPs can be used for specific labeling of defined gene products; they have low cytotoxicity and do not require microinjection or other invasive techniques for entering a cell. Unfortunately, AFPs are not well suited for FCCS studies, and to our knowledge there are only two FCCS works in live cells relying exclusively on AFPs (27,28). Apart from the lower photostability of AFPs compared to that of artificial dyes, the most popular AFPs, namely, ECFP, EGFP, and EYFP, show a large spectral overlap in the fluorescence spectra (see Fig. 1 for ECFP and EYFP). This leads to cross talk in the correlation signal that is hard to correct. In live cell measurements, where the concentration of the fluorophores is not known a priori and where background fluorescence is high, cross talk usually does not allow for performing FCCS.

Alternative methods based on, e.g., fluorescence resonance energy transfer (FRET) proved hard to implement in cell environments due to cross talk, unknown concentrations of the fluorophores, and unknown maximal FRET efficiencies (29,30).

### THEORY

In FCS the concentration fluctuations of fluorescing particles in an effective volume element are used to determine the diffusion constants and concentrations of the dissolved species. In practice this is achieved by focusing an excitation beam into a solution or a live cell specimen containing the particles of interest. The focus constitutes an open volume through which particles can diffuse in and out, leading to concentration fluctuations and in turn to fluctuations of the fluorescence intensity signal F. These fluctuations are correlated in time yielding a correlation function

$$g_{ij}^{(2)}(\tau) = \frac{\langle F_i(t)F_j(t+\tau)\rangle}{\langle F_i(t)\rangle\langle F_i(t)\rangle},\tag{1}$$

where  $F_i(t)$  is the fluorescence intensity of the species *i* (i = 1, 2 for ECFP and EYFP) at time *t* and the brackets denote time averaging. In the case of i = j this is the autocorrelation function.



FIGURE 1 Excitation and emission spectra of ECFP and EYFP (from Cubitt et al. (41)). In the shaded area are the excitation wavelengths chosen to prevent cross talk and the undetected spectral range due to the trichroic beamsplitter. The circle stresses the spectral region in which, normally, cross talk arises.

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By modeling the focal volume with a three-dimensional (3D) Gaussian profile, it is possible to give an analytical model for the correlation function of free translational 3D diffusion (31):

$$g_{ii}^{(2)}(\tau) = \underbrace{\frac{1}{\langle N_{ii} \rangle}}_{1} \underbrace{\left(1 - \frac{F_{\text{bg},i}}{F_{\text{tot},i}}\right)^{2}}_{2} \underbrace{\left(1 + \frac{P_{i}\exp(-\tau/\tau_{\text{p},i})}{1 - P_{i}}\right)}_{3} \\ \times \underbrace{\left(1 + \frac{\tau}{\tau_{\text{diff},i}}\right)^{-1}}_{4} \underbrace{\left(1 + \frac{\tau}{S_{i}^{2}\tau_{\text{diff},i}}\right)^{-\frac{1}{2}}}_{5}.$$
 (2)

In this equation, terms 4 and 5 describe the diffusion, term 3 describes fluctuations due to protonation/deprotonation processes observed in some AFP fluorophores (32), and term 2 is a correction for uncorrelated background light. In detail,  $\langle N_{ii} \rangle$  is the average number of particles within the focus,  $F_{tot,i}$  is the total fluorescence intensity,  $F_{bg,i}$  is the uncorrelated background intensity,  $P_i$  the fraction of molecules being in the protonated state,  $1/\tau_{p,i}$  the protonation rate,  $\tau_{diff,i}$  is the diffusion time, and  $S_i$  is the structural factor of the focal volume, which is defined as  $S_i = (r_{z,i}/r_{xy,i})$ . The diffusion time is inversely proportional to the diffusion constant  $\tau_{diff,i} = r_{xy,i}^2/4D_i$ , with  $r_{xy,i}$  the width of the Gaussian profile and  $D_i$  the diffusion constant.

If the diffusion of the fluorescing particles is restricted to a planar sheet (e.g., a cell membrane), Eq. 2 can be simplified to a two-dimensional (2D) model:

$$g_{ii}^{(2)}(\tau) = \frac{1}{\langle N_{ii} \rangle} \left( 1 - \frac{F_{bg,i}}{F_{tot,i}} \right)^2 \\ \times \left( 1 + \frac{P_i \exp(-\tau/\tau_{p,i})}{1 - P_i} \right) \left( 1 + \frac{\tau}{\tau_{diff,i}} \right)^{-1}.$$
 (3)

In the case of different fluorophores (i.e.,  $i \neq j$ ), Eq. 1 holds for cross correlation analysis. If the excitation volumes of the two exciting laser beams overlap perfectly and the  $1/\tau_{p,i}$  rates of the two dyes do not correlate, the cross correlation function in 3D is written as

$$g_{ij}^{(2)}(\tau) = \frac{1}{\langle N_{ij} \rangle} \left( 1 - \frac{F_{bg,i}}{F_{tot,i}} \right) \left( 1 - \frac{F_{bg,j}}{F_{tot,j}} \right) \\ \times \left( 1 + \frac{\tau}{\tau_{diff,ij}} \right)^{-1} \left( 1 + \frac{\tau}{S_i^2 \tau_{diff,ij}} \right)^{-\frac{1}{2}}, \qquad (4)$$

and for 2D diffusion the last term can be omitted.

One of the protein complexes used for our FCCS experiments was a gap-junction hemichannel or connexon. A single connexon comprises six protein subunits called connexins. In our experiments, we have used cells that coexpress connexins labeled with ECFP and EYFP. In this case the hexamers consist of a mixture of ECFP and EYFP labeled monomers. The distribution of labels is assumed to be binomial:

$$P(k) = \binom{6}{k} p^{k} (1-p)^{6-k},$$
(5)

with P(k) the probability of having a connexon consisting of k ECFP-connexins and 6 - k EYFP-connexins.

For the sake of clarity, we will first consider the simpler case of a dimer. Here, the auto- and cross correlation amplitude measure two different concentrations of the dimer because the binomial distribution gives two homolabeled dimers which do not contribute to the cross correlation amplitude and only one homolabeled dimer that does not contribute to the autocorrelation amplitude. Therefore, we introduce factors  $C_{ij}$  correcting this systematic error. In the dimer case, e.g., the amplitude of the cross correlation function is reduced by a factor of 2 and hence the correction factor has to be  $C_{12,dimer} = \text{all}/\text{detected} = 4/2 = 2$  as it is also calculated by Eq. 6 (see below).

To derive the correction factors, Eqs. 2 and 4 are examined at  $\tau = 0$  and the additional effect of different brightnesses (distribution of heterolabeled connexons) of the hexamers has to be taken into account (24). Then the cross correlation amplitude correction factor is given by

$$C_{\rm ij,hexamer} = \frac{36p(1-p)}{\sum_{k=0}^{6} k(6-k) \binom{6}{k} p^{k} (1-p)^{6-k}},$$
(6)

for the probability of having an ECFP or a EYFP molecule assumed to be the same ( $p_{\text{ECFP}} = p_{\text{EYFP}} = 0.5$ ),  $C_{ij,\text{hexamer}} = 1.2$ . A similar calculation gives amplitude correction factors for the autocorrelation amplitudes  $C_{ii,\text{hexamer}} = 0.76$  (24).

### MATERIALS AND METHODS

#### **ECFP-EYFP** lysates

For the preparation of ECFP and EYFP from cell lysates we have used HEK 293 cells that were transiently transfected with the plasmids pECFP or pEYFP. In brief, cells (800  $\mu$ l;  $1.5 \times 10^7$ /ml) were harvested and 15  $\mu$ g of plasmid was added to the suspension. The cells were then electroporated at 250 V and 1800  $\mu$ F in a 0.4 cm cuvette (Peqbio Easyject Plus, Peqlab, Erlangen, Germany). After electroporation, the cells were immediately transferred into 15 cm cell culture dishes and grown overnight. For lysis, the cells were harvested in 500  $\mu$ l buffer (10 mM HEPES pH 7.9, 10 mM KCl, 0.1 mM EDTA, 0.1 mM EGTA, 1 mM DTT, 1 mM phenylmethylsulfonyl fluoride), followed by incubation on ice for 20 min. Then 30  $\mu$ l 10% Nonidet-40 was added and incubated for 2 min at 4°C while shaking. Nuclei were pelleted at 16,000 × g for 3 min at 4°C, and supernatant was used for FCCS experiments.

### Cell culture and transient transfection

For transient expression of an AFP labeled membrane protein, we used C-terminal fusion constructs of ECFP and EYFP to the gap-junction protein connexin46. HeLa cells (ECACC No. 96112022) were cultivated in Dulbecco's modified Eagle's medium (DMEM) containing 10% newborn calf serum (NCS) in a humidified CO<sub>2</sub>-incubator at 37°C. For transient transfection, we followed the protocol provided for the lipofectin reagent (Life Technologies,

Karlsruhe, Germany). Briefly, the cells were grown in 35 mm glass bottom petri dishes (MatTek, Ashwell, MA) to 50–75% confluency. We used 35 mm glass bottom petri dishes because they allow a better observation of the cell monolayers in an inverted microscope. The cells were cotransfected in Opti-MEM1 medium (Life Technologies) containing 7–10  $\mu$ l lipofectin and 1–2  $\mu$ g of plasmid DNA (pECFP-N1-rCx46 and pEYFP-N1-rCx46 either alone or in 1:1 mixtures) for 6 h at 37°C. After this time the DNA/lipofectin suspension was removed and replaced with normal DMEM.

The cells were then transferred into an incubator set to a temperature of 27°C for 24–48 h for a better insertion of the protein in the plasma membrane.

Transfection efficiencies >30% were routinely obtained with this method for HeLa cells.

## Experimental setup for cross talk free FCCS

FCCS experiments were performed using a modified Olympus IX-70 inverted microscope with a high numerical aperture objective (UPLAPO  $60 \times$ 1.2 W, Olympus Europe, Hamburg, Germany). For confocal excitation a trichroic beamsplitter (Z425/514R at 10°, Chroma Technology, Rockingham, VT) reflects the excitation beams into the side port of the microscope. The collected photons pass the beamsplitter and a pinhole (diameter 50  $\mu$ m) and are detected by an actively quenched avalanche photo diode (APD, SPCM-AQR 14, PerkinElmer Optoelectronics, Fremont, CA). The excitation wavelengths were supplied by an argon-ion laser (at 515 nm, Innova 90C, Coherent, Palo Alto, CA) and a frequency doubled (at 425 nm, BBO crystal, Fujian Castech Crystals, Fujian, China) passively mode-locked titan sapphire oscillator (at 850 nm, Mira 900 F, Coherent) with a pulse width of 100 fs and a repetition rate of 76 MHz, which was pumped by a frequency doubled Nd:YAG laser (Verdi V5, Coherent). The pulse repetition rate was reduced to 7.6 MHz by an acousto-optic modulator (AOM)-based pulse picker system (PulseSelect, APE, Berlin, Germany). It also provided the delayed pulse signal for the second AOM (350 MHz - 5 ns rise time, Crystal Technology, Palo Alto, CA), generating a 10 ns green laser pulse 50 ns after the blue one. The excitation power was adjusted to 50  $\mu$ W for the blue and 5  $\mu$ W for the green laser (measured continuous wave). The detector signal was recorded with a time correlated single photon counter card (TCSPC; SPC630 Fifo Mode, Becker & Hickl, Berlin, Germany) and analyzed by custom software including a software correlator. Confocal images were recorded by scanning the sample through the focus using a three-axis piezo actor (PiMars P-527.3CL, Physik Instrumente, Karlsruhe, Germany) mounted on the microscope stage. Fitting was performed with nonlinear least-square Levenberg-Marquardt routines.

The adjustment procedure followed this scheme: first, the blue and green excitation beams were brought into congruence at the beam combiner and right before the microscope. Then the detection path of the setup was adjusted with blue excitation light only and a thick solution of rhodamine 6G solvated in ethylenglycol as a sample. Alternating the fine adjustment of the incoming excitation beam and the detection pathway maximized the countrate at the detector. Only the green excitation beam was fine adjusted for maximum intensity at the same pinhole position as for the blue beam alone. A last adjusting step was to gain maximum parfocality of both excitation colors. This was achieved by changing the divergence of the blue beam with the last curved mirror of the pulse picker. Inspection of the two laser spots at the ceiling of the room (no objective in the revolver) showed a perfect overlap.

### RESULTS

### Main principle of cross talk free measurements using FCCS

The aim of this study is to measure the colocalization of proteins in live cells using the fluorophore pair ECFP and

EYFP. This is not possible by standard FCCS setups due to cross talk caused by the overlap of the emission spectra of these fluorophores (Fig. 1). Indeed, using our setup with a standard filter set for ECFP and EYFP, we always see fluorescence of ECFP in the EYFP detection channel leading to cross correlation without EYFP fluorescence at all. This problem is addressed by using a pulsed excitation pattern of alternating blue and green laser pulses, which exclusively excite the cyan fluorescent protein (ECFP) with a wavelength of 425 nm and the yellow fluorescent protein (EYFP) with 515 nm (shaded bars in Fig. 1). We use a single detector to count the emitted photons of both fluorophores, which are afterwards rearranged by the software to form time traces of the two labels. For this purpose a TCSPC stores each photon with an additional time tag giving the arrival time of this photon relative to a synchronization pulse (see Fig. 2 A). The pulse repetition rate is optimized to allow for fluorescence decay after excitation to prevent cross talk in the time domain. In fact, the delay between succeeding pulses (50 ns) is much larger than the fluorescence lifetime of the AFPs  $(\sim 3 \text{ ns})$ . The length of the blue pulse of 100 fs is given by the TiSa laser and would in principle allow additional analysis of the fluorescence lifetime of ECFP although this is not required for the colocalization studies. The cross correlation setup would also work with a continuous wave (cw) laser around 425 nm together with a fast AOM and a pulse generator to synchronize the AOMs for blue and green light.

For testing the setup we have used samples of ECFP and EYFP and an ECFP/EYFP chimera (both proteins connected by a calcium sensitive calmodulin linker) in solution. In addition we have used live HeLa cells cotransfected with ECFP-connexin and EYFP-connexin fusion proteins. These connexin-AFP fusion proteins form hexameric membrane

pulse А pulse 100 fs 10 ns blue laser O @425 nm off green laseron @515 nm off fluorescence 0 pulse delay 50 ns (pulse rep rate)<sup>1</sup> 130 ns sync pulse в trichroic beamsplitter AOM blue laser Q beam AOM microscope combiner green lase objective pinhole photodiode delay APD timer: time-resolved pulse picker. data recording

FIGURE 2 (A) Pulsed excitation scheme for cross talk free FCCS. Fluorescence time intervals for data analysis can be selected by software. (B) Schematic overview of the FCCS setup.

channels that are inserted into the membrane in the endoplasmatic reticulum. The channels are then transported into the plasma membrane via the Golgi apparatus. These channels form a pool of free diffusing particles in the plasma membrane. The particles of this pool show a distribution of different numbers of ECFP and EYFP labels that is assumed to be binomial. Therefore the majority of the channels should contain both labels (see Theory).

# Signal enhancement by filtering in the time domain

To separate fluorescence of ECFP and EYFP, we determine the arrival time of all fluorescence photons with respect to the blue laser pulses. The blue pulses were chosen for synchronization due to the much better time resolution of these pulses (100 fs) compared to the green pulses generated by switching an AOM (10 ns).

Fig. 3 A shows the histogram of the arrival times. It exhibits two main peaks, a sharp one at t = 0 ns and a broader peak around t = 50 ns, corresponding to the fluorescence excited by the blue laser and the green laser, respectively. The smaller peak at t = 13 ns corresponds to an after pulse, which could not be totally suppressed by the pulse picker. The delay between main pulse and after pulse reflects the inverse repetition rate of the Titan Sapphire oscillator. Due to the time resolution of 100 fs of the blue laser, it is possible to determine the fluorescence lifetime of the fluorescence component excited by the blue laser. Here, we were able to determine the lifetime of ECFP to 3.1 ns, which is in good agreement with published values for this fluorophore (33). The fluorescence peak of the EYFP fluorescence is asymmetric and can be described as a convolution of a Gaussian and an exponential decay curve. Considering only the right wing, it is possible to observe a decay in the range of 3 ns corresponding to the lifetime of EYFP (34).

The histogram was measured from a solution containing exactly the same amount of ECFP and EYFP. Taking into account the absorption cross sections and quantum yields of ECFP and EYFP, the excitation power, and detection efficiencies of ECFP and EYFP, one would expect two times more photons in the EYFP channel than in the ECFP channel. Instead we observed 1.6 times more counts in the ECFP channel. These additional background photons in the ECFP channel could arise from autofluorescence or stray light from within the cell. Stray processes are fast and should appear in the histogram at t = 0 ns. For fluorescence processes from components other than ECFP, we would expect longer or shorter fluorescence lifetimes than for ECFP. In any case, it is possible to enhance the signal/noise ratio by disregarding time channels with high background fluorescence. Unfortunately, the main part of autofluorescence detected in the spectral range of ECFP also has a lifetime comparable to ECFP, resulting in a poor enhancement of the signal/noise ratio for the ECFP channel.



FIGURE 3 (*A*) Histogram of the photon arrival times. Two main peaks corresponding to photons originating from ECFP (*left*) and EYFP (*right*) are observable. (*B*) Background corrected FCS curves of ECFP, EYFP, and the FCCS curve for a mixture of both fluorophores in solution (*cyan*, CFP autocorrelation; *yellow*, YFP autocorrelation; *black*, cross correlation; *red*, fit functions). (*C* and *D*) FCS curves of ECFP, EYFP, and the FCCS curve for the ECFP/EYFP chimera in solution. (*C*) Measured curves. (*D*) Background corrected curves. (*E* and *F*) FCS curves of ECFP, EYFP, and the FCCS curve for coexpressed ECFP/Cx46 and EYFP/Cx46 in live HeLa cells. (*E*) Measured curves. (*F*) Background and permutation corrected curves. (*Inset*) Schematic top view of six connexins fused to three ECFPs' respective EYFPs.

Except for this autofluorescence, which obviously can be excited by 425 nm, other sources of background have been systematically eliminated by time resolved measurements of the buffer, several different coverslips (BK7, SUPRASIL I and II), and the setup without any specimens (data not shown).

### Signal/noise ratio

In this section we want to discuss the signal/noise ratio of the correlation curve using cw and pulsed excitation, respectively. First assume time intervals of the inverse repetition rate ( $\Delta t = 130$  ns) and cw excitation, i.e., pulse lengths of 130 ns, too. Using a typical count rate of 10 kcps and a detection efficiency of 10% of the microscope, the probability for emitting a photon in the focus per inverse repetition rate  $\Delta t$  is 1.3%. Provided that the number of photons follows a Poisson distribution, the probability that two photons are emitted is <0.01%, i.e., per 150 intervals containing one photon there is one interval containing two photons. Because of the missing photon pairs with short time distance, marginally lower signal/noise values for short correlation

times  $\tau$  would be expected in pulsed excitation. To check this, we recorded 50 correlation curves with cw and pulsed (pulse width 10 ns) excitation, respectively, using the same count rate of 4 kcps. The signal/noise ratio as a function of time is  $\sqrt{\operatorname{var}(g(\tau) - 1)}/g(\tau) - 1$ . For short times, the signal/noise ratio is slightly better for cw excitation (factor of 1.4), for times around the diffusion time there is no difference, and for times longer then the diffusion time the signal/noise ratio is slightly better for pulsed excitation (factor 1.4). The differences become smaller for longer pulse widths.

### Deviation of the diffusion time of ECFP and EYFP

All ECFP autocorrelation curves showed a systematic error toward apparently longer diffusion times (see Fig. 3 *B*, *D*, and *F*). Two effects can account for this enlargement of the confocal volume: i), the short excitation pulses ( $\sim$ 100 fs) with a high peak power caused saturation of the fluorescence emission in the Gaussian excitation profile, effectively broadening the focus (35); and ii), due to chromatic aberrations of

the objective for the emission wavelength of ECFP, the detected focal volume for ECFP is shifted in the *z* direction with respect to the effective detection volume for EYFP (spanned by  $r_{xy,2}, r_{z,2}$ ). This results in an effective detection volume for ECFP that is bigger than the focal volume for EYFP.

To estimate the impact of the first effect, we measured the diffusion times of a diluted ECFP lysate excited with 100 fs pulses as a function of the mean excitation power. The diffusion time started to increase for excitation powers >10  $\mu$ W and rose until a maximum of a factor of 1.9 was reached at 70  $\mu$ W, comparable to data published before (35). A further increase of the excitation power yielded a decrease of the diffusion time due to bleaching of the fluorophore in the excitation volume. The correlation time measured at lowest power is close to the value obtained with cw excitation. We want to compare the ECFP diffusion with the EYFP diffusion excited with 10 ns pulses. For such long pulses the saturation effect is relatively small (35), i.e., the correlation times measured for EYFP are close to values of cw measurements, too.

The correlation curves measured in live cells (see Fig. 3 *F*) were excited with 80  $\mu$ W and show diffusion times of  $\tau_{\rm YFP} = 7.5$  ms and  $\tau_{\rm CFP} = 16.4$  ms, i.e., the diffusion time observed with short pulses is a factor of 2.2 longer than with long pulses or cw. Compared to this result, a factor of 1.8 was found for the diffusion of ECFP excited with 5  $\mu$ W (comparable to cw) and 70  $\mu$ W in the lysate. For the correlation curves of the chimera in lysate, which was excited with 50 $\mu$ W/blue, the difference is smaller. Here, an increase of the diffusion time of a factor of 1.2 ( $\tau_{\rm YFP} = 0.18$  ms and  $\tau_{\rm CFP} = 0.22$  ms) was observed compared to 1.3 for ECFP excited with 5  $\mu$ W and 50  $\mu$ W. Hence, most of the increase of the diffusion time can be explained by saturation effects due to short pulsed excitation.

### **ECFP-EYFP** in lysate

The first step to test our experimental approach is to check whether there is any cross correlation between unbound ECFP and EYFP in solution. For this we have used either single or mixed solutions of free ECFP and EYFP from cell lysates. The lysates were diluted with phosphate-buffered saline to concentrations of  $3 \times 10^{-9}$  mol/l, i.e., to a concentration where on average a single molecule is found in the focal volume.

Experiments that test for channel cross talk with solutions containing only ECFP or EYFP showed no emission of ECFP with excitation by 515 nm laser light (EYFP excitation) and vice versa (data not shown).

Fig. 3 *B* shows the corresponding background corrected autocorrelation and cross correlation curves for freely diffusing ECFP and EYFP in mixed solution. The FCS curves are obtained by taking all photons of the arrival time histogram into account. The curves were least square fitted with

the diffusion term of Eq. 2 (all errors given originate from the fitting procedure). The diffusion constants have similar values of  $D_1 = 7.7 \pm 0.2 \times 10^{-7}$  cm<sup>2</sup>/s and  $D_2 = 1.2 \pm 0.1 \times 10^{-6}$  cm<sup>2</sup>/s, which are reasonable values for small proteins in solution (13,36). A deviation of the fit function compared to the measured curve for the EYFP at longer times  $\tau$  most probably arises from small aggregates with other proteins in the lysate. No cross correlation is observable (*noisy black line* in Fig. 3 *B*) as expected for this solution of unbound ECFP and EYFP labels.

# **ECFP/EYFP** chimera in lysate

Fig. 3 C shows the correlation curves for the ECFP/EYFP chimera in lysate, which is diluted with phosphate-buffered saline (no Ca<sup>2+</sup> ions included to prevent FRET in the chimera). Here, the diffusion constants are slightly smaller  $(D_1 = 8.1 \pm 0.3 \times 10^{-7} \text{ cm}^2/\text{s} \text{ and } D_2 = 7.3 \pm 0.2 \times 10^{-7}$ cm<sup>2</sup>/s) than in the mixture sample of ECFP-EYFP due to the bigger hydrodynamic radius of the fusion protein. As expected this system shows a cross correlation amplitude. The value for the diffusion constant of the cross correlation curve is slightly lower ( $D_{12} = 6.1 \pm 0.2 \times 10^{-7} \text{ cm}^2/\text{s}$ ) than for the autocorrelation. Applying the fluorescence background correction term of Eq. 2 (Eq. 4 for the cross correlation) results in a cross correlation amplitude of half the theoretical maximum value, assuming the green channel is free of background fluorescence (see Fig. 3 D). Possible reasons are a nonideal overlap of the excitation volumes and nonfunctional AFPs in the construct.

### ECFP-connexin and EYFP-connexin in live cells

Fig. 3 *E* shows the autocorrelation and cross correlation curves for ECFP- and EYFP-connexin fusion proteins in the plasma membrane of live HeLa cells. Connexins form a hexameric complex called connexon, schematically shown in Fig. 3 *F* (*inset*). For the correlation curve of EYFP, the protonation term has to be taken into account (Eqs. 3 and 4 for cross correlation without the last term) to achieve satisfactory fitting. The values derived for  $P_2$  and  $\tau_{p,2}$  are in good agreement with previously reported values (32). In the measurement results shown in Fig. 3, *B–D*, this blinking was obscured because it is in the same time region as the diffusion of the fast chromophores in solution. The ECFP curve, on the other hand, does not show this protonation behavior.

The diffusion constants of  $D_1 = 1.2 \pm 0.1 \times 10^{-8} \text{ cm}^2/\text{s}$  for ECFP and  $D_2 = 1.2 \pm 0.1 \times 10^{-8} \text{ cm}^2/\text{s}$  for EYFP correspond well with membrane protein diffusion in live cells (37,38). The cross correlation diffusion constant is again close to the autocorrelation values ( $D_{12} = 1.0 \pm 0.1 \times 10^{-8} \text{ cm}^2/\text{s}$ ), as expected. After fluorescence background correction of the data and taking into account the various compositions of a connexon (homo- and several heterolabeled hexamers, see Theory), an amplitude of the cross correlation

curve (see Fig. 3 *F*) of 1.86, compared to a maximum of 2.18, shows the spectral overlap of the excitation volumes to be  $\sim$ 86%.

# CONCLUSIONS

We were able to design and test a new setup with pulsed excitation lasers to prevent cross talk in the detection pathway to measure the colocalization of proteins on a single molecule level. With this setup we have successfully demonstrated colocalization of membrane proteins labeled with the living colors ECFP and EYFP at molecular level in live cells. We were also able to prove the concept with positive and negative control experiments using unbound ECFP and EYFP from cell lysates in buffer. The major problem in all measurements was autofluorescence in the ECFP channel. Our method provides the advantage of filtering in the time regime, i.e., suppressing all fluorescence photons arriving from faster (e.g., Raman scattering) and slower decay processes (e.g., fluorescence of color centers in glass). Unfortunately, in the biological specimens investigated in this study, most of the autofluorescence shows the same decay time as the ECFP molecules. However, working with other cell lines may result in other fluorescing components and, hence, in less autofluorescence from fluorophores exactly matching the photophysical properties of ECFP.

For measurements in the presence of high autofluorescence, multiple ECFP labels (2–3) per protein proved to be useful to detect this protein in the ECFP channel above background.

Future developments could improve the investigations of colocalizations in live cells. Obviously, the development of AFPs with properly separated emission spectra would allow the application of the standard FCCS method using two detectors. Nowadays available monomeric red fluorescent proteins show weak fluorescence (39) that is difficult to detect effectively above the background in cells. Nevertheless, Saito et al. (27) and Baudendistel et al. (28) succeeded recently in this task by using mRFP1 fused to the sample protein and measured FCCS curves in live cells. A very promising technique for all live cell studies would be the application of fusion proteins in combination with artificial dyes that have much better photophysical properties than AFPs. First results on this technique were shown recently (40).

Today, AFPs (and among them ECFP and EYFP) are the dominant fluorophores in cell biology applications and are widely used in many laboratories. Our new technique of FCCS might support the investigations of protein-protein interactions on a single molecule level within a live cell. For this an easier to use and cheaper setup can be used utilizing two pulsed laser diodes or two AOMs and suitable laser sources for excitation. The relatively short data acquisition time of 30 s allows time resolved observation of protein interactions, e.g., in a signaling cascade. This will shed new light on cell functions, which now can be monitored under physiological conditions in live cells with resolution below diffraction limit. This should work in cellular membranes and in the cytosol as well as in any other larger compartment of cells.

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