Förster Resonance Energy Transfer beyond 10 nm - Exploiting the Triplet State

Kinetics of Organic Fluorophores

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ABSTRACT

Inter- or intramolecular distances of biomolecules can be studied by Förster resonance energy transfer (FRET). For most FRET methods, the observable range of distances is limited to 1-10 nm and the labeling efficiency has to be controlled carefully to obtain accurate distance determinations, especially for intensity-based methods. In this study, we exploit the triplet state of the acceptor fluorophore as a FRET readout using fluorescence correlation spectroscopy and transient state monitoring. The influence of donor fluorescence leaking into the acceptor channel is minimized by a novel suppression algorithm for spectral bleed-through, thereby tolerating a high excess (up to 100 fold) of donor-only labeled samples. The suppression algorithm and the high sensitivity of the triplet state to small changes in the fluorophore excitation rate make it possible to extend the observable range of FRET efficiencies by up to 50 % in the presence of large donor-only populations. Given this increased range of FRET efficiencies, its compatibility with organic fluorophores and the low requirements on the labeling efficiency and instrumentation, we foresee that this approach will be attractive for in-vitro and in-vivo FRET-based spectroscopy and imaging.

KEYWORDS

FRET, fluorescence correlation spectroscopy, transient state monitoring, triplet state, iodide quenching, spectral bleed-through/cross-talk
Introduction

During Förster (or fluorescence) resonance energy transfer (FRET), energy is transferred via a non-radiative dipole-dipole interaction from a donor (D) to an acceptor (A) molecule in a strongly distance-dependent manner. This has made FRET a widely used method to determine inter- or intra-molecular distances in the range of typically 1-10 nm.¹

Techniques for FRET estimation can be divided into those applicable to conventional microscopes (intensity-based) and techniques that require dedicated instrumentation (e.g. time-resolved; fluorescence lifetime imaging microscopy (FLIM), single molecule measurements).

The most popular intensity-based approach is steady-state monitoring of the D and A-emission upon D-excitation.²⁻⁵ Due to the spectral overlap of D and A, the detected A-signal is influenced by D-emission leaking into the A-channel and direct excitation of A. As a result of the spectral bleed-through, additional control measurements have to be acquired of D- and A-emission for D-only and A-only labeled samples. For most intensity-based approaches, the apparent FRET efficiency can be affected by incompletely labeled FRET pairs, which exist to a substantial amount in most FRET experiments. Therefore, careful calibration measurements are necessary to control the labeling efficiencies and the relative concentrations of D and A.

FLIM is a robust but technically challenging method for FRET estimation.⁶ By monitoring changes in the D-fluorescence lifetime, it is immune to spectral bleed-through and A-only labeled samples. FLIM can also differentiate between D-only and D-A labeled samples at high and intermediate FRET efficiencies. At low FRET efficiencies, however, the donor lifetimes of D-only and D-A labeled samples are similar and hence difficult to distinguish. The resulting FRET efficiencies can then be underestimated to a significant degree.

Similar to FLIM, single-molecule multiparameter fluorescence detection (smMFD) has difficulties to estimate low FRET efficiencies. To differentiate between fluorescent
trajectories of low-FRET species and D-only labeled species, a threshold has to be applied on the detected acceptor signal that often cuts off signals originating from FRET efficiencies < 10 %. On the other hand, by analyzing multiple parameters (e.g. intensity, lifetime, anisotropy) from individual molecules simultaneously, smMFD can reach a high precision and accuracy in the FRET measurements, identify unspecific quenching effects and distinguish between D- and/or A-labeled specimen.7-9

An alternative method to detect low FRET efficiencies in cell-free systems is luminescence resonance energy transfer (LRET), which uses lanthanides as donors.10,11 Due to the long lifetime (ms) of lanthanides, the large Stokes shift and Förster radii up to 9 nm, LRET efficiencies can be determined over a wide range of distances (1-14 nm). However, a wider applicability to cell measurements has to date been limited due to the lack of suitable bright and photo-stable lanthanide complexes and difficulties to specifically attach these probes to proteins or subcellular structures.12 In addition, LRET measurements require a modulated, high intensity UV excitation source and a time-gated detection rarely offered by standard microscopes.

In this study, we present an approach for FRET efficiency estimation that exploits the triplet state of the A-fluorophore. The triplet state population of any fluorophore is sensitive to its excitation rate, independent of whether the fluorophore is excited directly or via FRET. For low FRET efficiencies, also small absolute changes in E, and thus also of the excitation rate of A, can strongly influence the population of the A-triplet state due to its relatively long lifetime (μs). This makes it possible to distinguish between low FRET efficiencies generated from D-A distances of more than 10 nm, also when standard organic fluorophores are used. The approach is demonstrated by determining FRET efficiencies of DNA samples labeled with the fluorophores ATTO 488 and Alexa Fluor 610. First, we present the basic principle of the method by using fluorescence correlation spectroscopy (FCS) to monitor the A-triplet state. In a second step, we look into the possibility of applying FRET-triplet state
measurements to a broader range of microscopic techniques. This is done by adapting the recently developed transient state (TRAST) monitoring technique to the FRET-triplet state measurements.\textsuperscript{13-15} In TRAST, the A-triplet state is deduced by modulating the excitation of D in a well-defined on-off fashion and detecting the A-fluorescence for different pulse settings. The same kind of on-off modulation can be obtained by scanning a laser beam over the image region with different scanning speeds.\textsuperscript{16} This indicates that our approach can determine the A-triplet state, and hence FRET efficiencies, in a wide range of microscopy settings, including confocal laser scanning microscopy and fluorescence-based plate-readers. The determined FRET efficiencies are compared with theoretical values obtained from a standard DNA model\textsuperscript{17} and with fluorescence lifetime measurements of the donor.

Due to a novel suppression algorithm for D-bleed-through, substantial fractions of incompletely labeled FRET molecules can be tolerated and D-A pairs with large Förster radii can be used with only minor concerns regarding the spectral overlap of D and A. Taken together, A-triplet state measurements are able to distinguish FRET efficiencies even below 5% and D-A distances well beyond the 10 nm limit of conventional FRET methods can be detected.

**Theory**

**Electronic state model.** To compute the FRET efficiencies, we use an electronic state model consisting of a D and A-fluorophore coupled by energy transfer with a rate $k_{\text{FRET}}$ (Fig. 1). Each individual molecule comprises a ground and an excited singlet state ($S_0$ and $S_1$), the lowest triplet state ($T_1$) and a photo-oxidized ($\hat{R}^+$) or -reduced ($\hat{R}^-$) radical state. Transitions between $S_0$ and $S_1$ are quantified by the fluorescence decay rate $k_{10} = 1/\tau_F$ ($\tau_F$ is the fluorescence lifetime) and by the excitation rate $k_{01}(\vec{r}, t) = \sigma_{\text{exc}} I_{\text{exc}}(\vec{r}, t)$. Here, $\sigma_{\text{exc}}$ is the excitation cross section for transitions from $S_0$ to $S_1$ and $I_{\text{exc}}(\vec{r}, t)$ is the excitation intensity at
a location \( \vec{r} \) within the laser beam. \( k_{\text{ISC}}, k_T, k_{\text{ox}} \) and \( k_{\text{red}} \) denote the intersystem crossing rate, the triplet relaxation rate, the oxidation rate and the reduction rate, respectively. Oxidation is usually assumed to originate from \( S_1 \) and \( T_1 \). However, since the populations of \( S_1 \) and \( T_1 \) only differ by a scaling factor and \( k_{\text{ox}} \) and \( k_{\text{red}} \) are well separated in time from the triplet transitions, oxidation is for simplicity assumed to occur only from \( T_1 \). For an individual fluorophore, the probability of being in a certain state (\( S_0, S_1, T_1 \) or \( \hat{R}^+ \)) at a location \( \vec{r} \) and time \( t \) can be calculated by a system of four first-order differential equations.

\[
\frac{d}{dt} \begin{pmatrix}
S_0(\vec{r},t) \\
S_1(\vec{r},t) \\
T_1(\vec{r},t) \\
\hat{R}^+(\vec{r},t)
\end{pmatrix} = \begin{pmatrix}
-k_{01}(\vec{r},t) & k_{10} & k_T & k_{\text{red}} \\
0 & -(k_{10}+k_{\text{ISC}}) & 0 & 0 \\
0 & k_{\text{ISC}} & -(k_T+k_{\text{ox}}) & 0 \\
0 & 0 & k_{\text{ox}} & -k_{\text{red}}
\end{pmatrix} \begin{pmatrix}
S_0(\vec{r},t) \\
S_1(\vec{r},t) \\
T_1(\vec{r},t) \\
\hat{R}^+(\vec{r},t)
\end{pmatrix}
\]

For a fluorophore with a photo-reduced radical state \( \hat{R}^- \), Eq. 1 can be modified by exchanging \( \hat{R}^+ \) with \( \hat{R}^- \) and interchanging \( k_{\text{ox}} \) and \( k_{\text{red}} \). Equation 1 can also be applied to the D or A of a FRET-pair with energy transfer occurring from \( S_1^D \) to \( S_0^A \) as well as to \( T_1^A \), but not to the radical state of A. For the donor, the fluorescence decay rate is substituted with

\[
k_{10}^{\text{D, tot}} \equiv k_{10}^D + k_{\text{FRET}} S_0^A(\vec{r},t) + T_1^A(\vec{r},t) \approx k_{10}^D + k_{\text{FRET}}
\]

while the excitation rate of A is replaced by

\[
k_{01}^{A, \text{tot}}(\vec{r},t) \equiv k_{01}^A(\vec{r},t) + k_{\text{FRET}} S_1^D(\vec{r},t), \text{ with} \\
S_1^D(\vec{r},t) = \frac{k_{01}^D(\vec{r},t) k_T^D}{k_{01}^D(\vec{r},t) (k_{\text{ISC}} + k_T^D) + k_{10}^{\text{D, tot}} k_T^D}.
\]

(Fig. 1). In Eq. 2-3 and from now on, upper case letters \( D,A \) indicate, whether a rate or a state belongs to the donor or acceptor, respectively. Under our experimental conditions, \( \hat{R}^{+A} \) can be neglected in the A-triplet state analysis in FCS (The typical transit time of the studied
molecules through the laser excitation beam was 300 μs and the time range of the $\hat{R}^{+A}$ state build-up was in the range of ms. Eq. 1 can then be approximated as a three-state model of $S_0$, $S_1$ and $T_1$) and TRAST (Following onset of excitation, the relaxation time for the A-triplet state build-up was 3 μs. The corresponding relaxation time for the $\hat{R}^{+A}$ state build-up was in the millisecond time scale i.e. well after the equilibration of the triplet state.). Given that $S_1^A$ is typically < 0.05, the boundary condition of Eq. 1 ($S_0^A + S_1^A + T_1^A + \hat{R}^{+A} = 1$) can be simplified to $S_0^A + T_1^A \approx 1$. Excitation of $S_1^A$ and $T_1^A$ into higher excited singlet and triplet states, $S_n^A$ and $T_n^A$, is excluded from the electronic state model of Eq. 1, because $S_n^A$ and $T_n^A$ are only sparsely populated due to the moderate excitation irradiances used (< 120 kW/cm²)²⁰,²¹ and the short lifetimes of those states (fs).²⁰

FRET efficiencies $E$ can be calculated from the FRET rate by

$$E = \left( \frac{k_{10}^D}{k_{FRET}} + 1 \right)^{-1}$$

4

**Triplet state kinetics from FCS.** In FCS, fluorescence fluctuations from individual molecules are studied as they diffuse in and out of an observation volume defined by the focus of a laser beam. The detected fluorescence $\bar{F}(t)$ depends on the detection quantum yield of the instrument ($\Phi_{det}$), the fluorescence quantum yield of the fluorophore ($\Phi_F$), the collection efficiency function $CEF(\bar{r})$ and the fluorophore concentration $c(\bar{r}, t)$.

$$\bar{F}(t) = \int F(\bar{r}, t) dV = \int \Phi_{det} \Phi_F CEF(\bar{r}) \sigma_{exc} I_{exc}(\bar{r}, t) c(\bar{r}, t) dV$$

5

The intrinsic fluorescence fluctuations, $\delta F$, of $\bar{F}(t)$ under a constant excitation intensity $(I_{exc}(\bar{r}, t) := I_{exc}(\bar{r}))$ can be described by the autocorrelation function $G(\tau)$. 


\[ G(\tau) = \frac{\langle F(t) F(t + \tau) \rangle}{\langle F(t) \rangle^2} - 1 = \frac{\langle \delta F(t) \delta F(t + \tau) \rangle}{\langle F(t) \rangle^2} \]
\[
= \frac{1}{N} \left( 1 - \frac{1}{1 + \tau/\tau_D} \left( \frac{1}{1 + (\omega_0 / \omega_z)^2 \tau/\tau_D} \right)^2 \right) \cdot \prod_{x=1}^{n} G^x_T(\tau)
\]
\[
G^x_T(\tau) := 1 - \bar{T}^x + \bar{T}^x \exp \left( -\tau / \bar{\tau}^x \right)
\]

Here, \( G(\tau) \) is shown for one diffusing species and \( n \) different independent dark states \( G^i_T(\tau) \).

The angle brackets indicate time-averages, \( \tau_D = \omega_0^2 / 4D \) the diffusion time of the studied molecules with a diffusion coefficient \( D \), \( N \) the average number of molecules in the detection volume and \( \omega_0 \) and \( \omega_z \) denote the distance from the center of the detection volume in the radial and axial direction, respectively, at which \( F(t) \) is decreased by a factor of \( 1/e^2 \). \( G^i_T(\tau) \) contains information about the average population of molecules in the detection volume that are in the dark state, \( \bar{T}^x \), and their average relaxation time \( \bar{\tau}^x \).

In the FCS analysis, A and D are described by a three state model of \( S_0, S_1 \) and \( T_1 \) and the resulting triplet state rates \( k_{\text{ISC}} \) and \( k_T \) can be derived from the average triplet state population \( \bar{T} \) and the corresponding relaxation time \( \bar{\tau}_T \) as described in \(^{22}\).

The A-triplet state is dependent on the A-excitation rate, which in turn is proportional to the FRET efficiency. Therefore, the A-triplet state parameters, \( \bar{T}^A \) and \( \bar{\tau}_T^A \), determined from A-correlation curves, are not only related to the triplet state rates \( k_{\text{ISC}}^A \) and \( k_T^A \), but also to the FRET rate \( k_{\text{FRET}} \) in analogy to the FRET-modified electronic state model in Eq. 1.
Here, $T^A(\vec{r})$ and $\tau^A_T(\vec{r})$ are defined at any location $\vec{r}$ in the excitation volume. Subsequent integration of $T^A(\vec{r})$ and $\tau^A_T(\vec{r})$ over the whole detection volume, weighted with $<F(t)>^2$, yields $\bar{T}^A$ and $\bar{\tau}^A_T$, respectively.\textsuperscript{22} By knowing $k_{ISC}^A, k_T^A, k_{10}^A$ and $\sigma_{exc}$ of A and D beforehand, one can solve Eq. 3, 4 and 7 to yield $k_{FRET}$ and the corresponding FRET efficiency.

**Triplet state and redox kinetics from TRAST.** Transient state monitoring (TRAST) has been applied previously to determine triplet and radical state properties of fluorophores.\textsuperscript{14-16} In TRAST, the excitation intensity is modulated in an on-off fashion while recording the time-averaged fluorescence response. During each excitation pulse, the triplet and radical state populations build up over time, leading to a decrease in fluorescence. By investigating the change in average fluorescence for different pulse widths and/or pulse periods, the populations of non-fluorescent states can be monitored.

TRAST is applied as described by Spielmann \textit{et al.}\textsuperscript{15} but in a confocal instead of a total-internal-reflection microscope setting. The Gaussian-Lorentzian excitation profile is approximated by a rectangular homogenous excitation profile yielding the same overall excitation power: $k_{01}(\vec{r}, t) := \bar{k}_{01}(t)$. Thus, $\bar{k}_{01}$ is the averaged excitation rate contributing to the fluorescence $\bar{F}(t)$, where $\bar{F}(t)$ is integrated over the focal spot as described in Eq. 5. This approximation has been used previously for FCS measurements\textsuperscript{20,23} and allows extracting reasonable estimates of triplet parameters from TRAST measurements while simplifying the analysis substantially.
The time-resolved fluorescence response, $\bar{F}(t)$, during a given excitation pulse can then be calculated by solving the rate equations of Eq. 1 for the excited singlet state population, $S_1(t)$. The duty cycle, which is the ratio of the pulse width to the pulse period, was kept low to limit bleaching and depletion effects. A low duty cycle also ensures that the system has completely relaxed at the onset of every new pulse in the pulse train (i.e. all molecules then reside in $S_0$). As the singlet, triplet and redox kinetics are well separated in time (ns, µs and ms), they can be treated independently. Moreover, the singlet state kinetics are generally too fast to be resolved by the present instrumentation. Therefore, for any individual fluorophore, the average fluorescence emitted during a pulse of width $w$ can be approximated by:

$$
\bar{F}(w) = \bar{F}_0 \left[ \frac{\bar{k}_{ISC}'}{-\left(\bar{k}_{ISC} + k_T\right)^2 w} \left( e^{-(\bar{k}_{ISC} + k_T)w} - 1 \right) + \frac{k_T}{\bar{k}_{ISC}'} + \frac{\bar{k}_{ox}'}{-\left(\bar{k}_{ox} + k_{red}\right)^2 w} \left( e^{-(\bar{k}_{ox} + k_{red})w} - 1 \right) + \frac{k_{red}}{\bar{k}_{ox}'} \right]
$$

In Eq. 8, $\bar{k}_{ISC}'$ denotes the equivalent intersystem crossing from the two singlet states ($S_0$ and $S_1$) to the triplet state $T_1$ and $\bar{k}_{ox}'$ the equivalent oxidation rate from $T_1$ to the radical state $R^+$:

$$
\bar{k}_{ISC}' = k_{ISC} \frac{\bar{k}_{01}}{\bar{k}_{01} + k_{10}}
$$

$$
\bar{k}_{ox}' = k_{ox} \frac{\bar{k}_{ISC}'}{\bar{k}_{ISC} + k_T}
$$

Moreover, in Eq. 8, $\bar{F}_0$ represents the average fluorescence count rate in the absence of transient populations of $T_1$ and $R^+$ and is proportional to the detection efficiency $\Phi_D$, the fluorophore concentration $\bar{c}$, the excitation volume $V_{exc}$ and the fluorescence quantum yield $\Phi_F$. 

10
\[
\bar{F}_0 = \Phi_D \bar{c} V_{\text{exc}} \Phi_F \frac{k_{10} \bar{k}_{01}}{k_{10} + \bar{k}_{01}}.
\]

10

Here, we are only interested in relative changes in \( \bar{F}(w) \) that originate from the build-up of transient states (e.g. \( T_1 \) and \( \hat{R}^+ \)). Therefore, \( \bar{F}(w) \) is normalized to unity for \( w \) shorter than the triplet relaxation time.

To determine FRET efficiencies, \( \bar{k}_{01} \) in Eq. 9 and 10 is substituted by \( \bar{k}_{01}^{A_{\text{tot}}} (\hat{r}, t) \) in analogy to Eq. 3. Furthermore, the expression for \( \bar{S}_1^D \) can be simplified in the analysis, because the build-up of the D-triplet state can be neglected in the presence of millimolar concentrations of potassium iodide\(^{24} \) and for the moderate excitation rates used in the TRAST measurements (\( k_{\text{ISC}}^D < k_T^D \)).

**Materials and Methods**

**FCS.** Measurements were performed on an epi-illuminated confocal microscope (Leitz).\(^{25} \) The light of an argon-ion laser (488 nm; Lasos, Jena, Germany) or an argon-krypton ion laser (488 nm, 568 nm; Melles Griot, Carlsbad, CA, USA) was filtered (488/10 nm, 568/10 nm, Chroma Technology Corp., Rockingham, VT, USA), focused in front of the microscope, reflected by a dichroic mirror (Z488RDC, Chroma Technology Corp.; FF576/661, Semrock, Rochester, NY, USA) and focused on the image plane by a water-immersion objective (63x/1.2 Plan-Neofluar, 160 mm tube length; Zeiss, Oberkochen, Germany). The intensity distribution is assumed to be Gaussian distributed in the radial direction and Lorentzian distributed in the axial direction, with a focal diameter of \( \sim 0.7 \mu\text{m} \).

The resulting fluorescence was collected by the same objective, focused onto a 50 \( \mu\text{m} \) pinhole, band-pass filtered (532/70 nm, filter 1: 675/135 nm, blue-shifted filter 2: 640/115 nm; Chroma Technology Corp.), split by a 50/50 beam splitter (Thorlabs, Newton, NJ, USA), and focused onto two single-photon-counting avalanche photodiodes (SPCM-AQR-14,
Perkin-Elmer Optoelectronics, Waltham, LA, USA). The signals of the two detectors were processed by a correlator (ALV-5000, ALV GmbH, Langen, Germany) and then further analyzed using custom software. The oxygen concentration in the buffer solution was strongly reduced by applying a flow of argon over the sample droplet.

**TRAST measurements.** The instrumentation was based on a previously described set-up for total internal reflection fluorescence microscopy, here used in a confocal setting. Briefly, a diode laser (Calypso, Cobolt AB, Solna, Sweden; 491 nm) was focused onto an acousto-optic modulator (AOM, MQ 180 –AO, 25-VIS, AA Opto-Electronic, Orsay, France) to yield the on-off modulation. After the AOM, the laser beam was collimated and reflected by a dichroic mirror (F500-Di01, Semrock Inc.) onto the objective (40x/1.2 C-Apochromat, water immersion, Zeiss). The resulting radial focus spot diameter was estimated to ~ 2 µm. The emitted fluorescence was collected with the same objective, band-pass filtered (675/135 nm, 530/50 nm, Chroma Technology Corp.) and focused onto an electron multiplying CCD (Luca, Andor Technology, Belfast, Northern Ireland, United Kingdom; used in quarter screen mode, 330 by 248 pixels). The modulation scheme of the AOM was set by a National Instruments (Austin, TX, USA) PCI 6602 card connected to the modulator controller (AA Opto Electronic). The camera, diode laser and National Instruments card were controlled via custom Matlab software.

Reference TRAST measurements on Rhodamine 110 (Rh110) were performed in order to relate the used laser intensities to the $k_{01}^{D}$ rate in Eq. 9 and 10. For the excitation intensities used in this study, the Rh110 excitation rate $k_{01}^{Rh110}$ was calculated by fitting measurements to Eq. 8 and 9 with fixed photo-physical parameters ($k_{ISC}^{Rh110} = 10^6$ s$^{-1}$, $k_{T}^{Rh110} = 0.45 \cdot 10^6$ s$^{-1}$, $k_{10}^{Rh110} = 244 \cdot 10^6$ s$^{-1}$). The D-excitation rate was then calculated according to: $k_{01}^{D} = \frac{k_{D}^{exc,491} \kappa_{Rh110}^{Rh110}}{\sigma_{exc,491}^{Rh110}}$, with $\sigma_{exc,491}^{Rh110} = 2.9 \cdot 10^{-16}$ cm$^2$ and $\sigma_{exc,491}^{D} = 2.5 \cdot 10^{-16}$ cm$^2$. 
The direct excitation \( \tilde{k}_{01}^A \) of A by laser light was estimated using \( \sigma_{\text{exc},491}^A = 0.08 \cdot 10^{-16} \text{cm}^2 \). 

\( \sigma_{\text{exc},491}^{D,A} \) and \( \sigma_{\text{exc},491}^{\text{Rh110}} \) were obtained by using literature values of the maximum extinction coefficient (as specified by Invitrogen Inc. and ATTO-TEC GmbH) and then calculating the extinction coefficient of a certain wavelength in relation to the measured fluorescence spectra. Extinction coefficients for FCS were determined in a similar fashion to

\[
\sigma_{\text{exc},488}^D = 2.243 \cdot 10^{-16} \text{cm}^2 \quad \text{and} \quad \sigma_{\text{exc},488}^A = 0.079 \cdot 10^{-16} \text{cm}^2.
\]

**Fluorescence lifetime and spectrum measurements.** Measurements were performed on a spectrofluorometer capable of time correlated single photon counting (FluoroMax-3, Horiba, Longjumeau, France; excitation source for lifetime measurements: NanoLED 495 nm). Lifetime decay data were analyzed by least squares deconvolution of a one-component exponential decay for D-only samples (\( \tau_{\text{F.D}} \)) and deconvolution of a 2-3 exponential decay model for D-A samples (DAS6, Horiba) with lifetimes \( \tau_i \) and amplitude \( A_i \). For the donor fluorophore in the presence of an acceptor, an average lifetime

\[
\tilde{\tau}_{\text{F,DA}} = \tau_1 A_1 + \tau_2 A_2 + \tau_3 A_3
\]

was calculated and used for further FRET efficiency calculations

\[
E = 1 - \frac{\tau_{\text{F,D}}}{\tilde{\tau}_{\text{F,DA}}}
\]

11

The fluorescence decay rate in the absence of FRET was calculated from the fluorescence lifetime of A-only or D-only labeled samples by \( k_{10}^{D,A} = \tau_f^{-1} \).

**Sample preparation.** As a model system, we used highly-purified double-stranded DNA (Purimex, Grebenstein, Germany) with ATTO 488 (ATTO-TEC GmbH, Siegen, Germany) and Alexa Fluor 610 (Invitrogen Inc., Eugene, OR, USA) labeled onto different positions of the DNA strand. The DNA lead strand CCT TTC AGG AGG AAG GAG AGA
GAG GAA AAT CAA ATC CTC CTA AGG G was either unmodified (control measurements) or modified at position three with ATTO 488 attached via a C₆-linker. The complementary strand CCC TTA GGA GGA TTT GAT TTT CCT CTC TCT CTT CCC TCC TGA AAG G was unmodified or modified with Alexa Fluor 610 attached via a C₆-linker at position 13, 18, 23, 27, 29, 32 or 44. Hybridization of double stranded DNA (dsDNA) was performed by adding lead and complementary strands in a concentration ratio of 2:1 (FCS) or 1:3 (lifetime measurements) to a tris-buffer (20 mM tris, 50 mM NaCl, 10 mM MgCl₂, pH 7.5). The DNA strands were incubated at 95 ºC for 5 min and gradually cooled to room temperature over three hours. dsDNA was diluted to nanomolar concentration in tris-buffer with or without potassium iodide (KI; Sigma-Aldrich, St. Louis, MO, USA). If not stated otherwise, samples were measured in air-saturated tris-buffer with 5 mM KI.

**DNA model.** FRET efficiencies were estimated from a standard DNA model with D and A fluorophore attached to a double-stranded helix of B-DNA.¹⁷ The helix had a width of 2 nm, a helical turn, \( \Delta \alpha \), of 36 degrees per base pair and a distance \( d_{bp} = 0.338 \) nm between two consecutive base pairs. The dyes were linked to the DNA by a carbon linker consisting of six carbon atoms (estimated length: \( C₆ = 0.6 \) nm) protruding perpendicular from the DNA body. The average distance \( d_{DA} \) between the donor and acceptor with \( \Delta bp \) base pairs between them was calculated according to

\[
d_{DA} = \left[ 2(1 + C₆)^2 (1 - \cos(\Delta bp \Delta \alpha)) + \Delta bp^2 d_{bp}^2 \right]^{1/2}.
\]

Subsequently, the corresponding FRET efficiency

\[
E = \left[ 1 + \left( \frac{d_{DA}}{R₀} \right)^6 \right]^{-1}
\]
was computed. Here, $R_0$ denotes the Förster radius of the FRET pair.

$$R_0 = \left[8.79 \times 10^{-4} J(\lambda) \Phi_F^D n^{-4} \kappa^2 \right]^{1/6}$$

15

$R_0$ depends on the spectral overlap integral $J(\lambda)$ between the fluorescence spectrum of D and the absorption spectrum of A, $\Phi_F$ of D in the absence of A, the refractive index $n$ of the medium and the orientation factor of D- and A-dipole moments, $\kappa^2$. For ATTO 488 and Alexa 610, the Förster radius was computed to 5.6 nm ($\Phi_F^D = 0.7$, assumed random orientation of D-A dipole moments: $\kappa^2 = 2/3$, $n = 1.337$, $J(\lambda) = 2.344 \times 10^{-13}$ cm$^3$ M$^{-1}$).

**Results**

1. **Effects of potassium iodide on the FRET pair.**

The triplet state population of an individual fluorophore strongly depends on the excitation rate, no matter how the fluorophore is excited (Eq. 7). Hence, when an acceptor is excited by FRET, its triplet state population can be used as a measure of the FRET efficiency and the corresponding D-A distance. To ensure a substantial triplet state build-up of A, relatively high $I_{\text{exc}}$ are required. This may lead to a considerable triplet build-up of D. When D is in its triplet state, no FRET will occur between D and A. The singlet-triplet state transitions of D thereby superimpose a modulation on the A-excitation and on its fluorescence. The resulting A-fluorescence fluctuations are then difficult to separate from those resulting from A-triplet state transitions. These complications can be circumvented by addition of potassium iodide (KI) to the solvent.

In millimolar concentrations, KI is known to promote intersystem crossing to the triplet state due to a heavy atom effect in all dyes. For many dyes with an excitation maximum in the blue range, it additionally enhances the triplet state decay rate by a charge-transfer mediated interaction between the fluorophore and iodide. Taken together, addition of
KI in millimolar concentrations can result in a reduced triplet state build-up for dyes in the blue excitation range and an increased triplet state build-up for dyes with excitation maxima in the green range or longer. In this study, we exploited the effect of KI to markedly reduce the triplet state build-up of D and minimize fluctuations in the A-excitation. The KI-mediated increase in the triplet state build-up of A makes it more sensitive to changes in excitation intensity and hence to differences in FRET efficiency.

The triplet state kinetics of the two dyes responded to potassium iodide (KI) as reported in 24, including also a KI-induced build-up of a photo-reduced radical state \( \dot{R}^- \) of ATTO 488 in proportion to the applied KI concentration. A concentration of 5 mM KI was identified as the best compromise between maintaining a low amount of \( \dot{R}^- \) formation while decreasing the ATTO 488 triplet state population and increasing the Alexa 610 triplet state population (Fig. 2). When KI was added, the measured triplet fraction of ATTO 488 bound to DNA was reduced from maximally 40 % in the absence to maximally 10 % in the presence of KI. The corresponding triplet relaxation time shifted from several \( \mu \)s to 0.2 \( \mu \)s for all \( I_{exc} \) applied. The largest \( \dot{R}^- \) population was 10 % with relaxation times of 10 \( \mu \)s. In contrast, the triplet fraction of Alexa 610 bound to dsDNA increased from 50 % to 60 % after addition of KI, while the triplet relaxation time remained constant (2-3 \( \mu \)s). The relaxation times of the triplet states of ATTO 488 and Alexa 610 as well as of the ATTO 488 radical state were each separated by an order of magnitude and could therefore be clearly distinguished and separated in the analysis.

\[
k_{isc}^{DA}, k_T^{DA}, k_{red}^D \text{ and } k_{ox}^D \text{ were obtained by direct excitation of ATTO 488-only or ATTO488-Alexa 610 labeled dsDNA (excitation: 488 nm or 568 nm). The correlation curves for ten } I_{exc} (6-500 kW/cm^2) \text{ were each analyzed by Eq. 6 using one diffusion component and one (acceptor; } T_1 \text{) or two (donor; } T_1, \dot{R}^- \text{) dark states. The resulting } \bar{T} \text{ and } \bar{T}_T \text{ for all } I_{exc} \text{ were simultaneously fitted to Eq. 7 according to the electronic state model of a single}
\]
fluorophore (Eq. 1). In these global fits, $k_{ISC}^{DA}$ and $k_{T}^{DA}$ were free to vary, while

$$\sigma_{exc,488}^D = 2.243 \cdot 10^{-16} \text{cm}^2, \sigma_{exc,568}^A = 1.667 \cdot 10^{-16} \text{cm}^2, k_{10}^D = 265 \cdot 10^6 \text{s}^{-1} \text{ and } k_{10}^A = 255 \cdot 10^6 \text{s}^{-1}$$

were kept constant. At 5 mM KI, the triplet rates were found to be $k_{ISC}^D = 2.0 \cdot 10^6 \text{s}^{-1}$,

$$k_T^D = 5.1 \cdot 10^6 \text{s}^{-1}, k_{ISC}^A = 1.4 \cdot 10^6 \text{s}^{-1} \text{ and } k_T^A = 0.13 \cdot 10^6 \text{s}^{-1}. k_{red}^D \text{ and } k_{ox}^D \text{ were determined from the redox state amplitude and relaxation time in analogy to}^{18,24} \text{ to}$$

$$k_{red}^D = 0.025 \pm 0.003 \cdot 10^6 \text{s}^{-1} \text{ and } k_{ox}^D = 0.1 \pm 0.01 \cdot 10^6 \text{s}^{-1}.$$  

2. FRET efficiencies determined by FCS

**Acceptor FCS curves show a FRET dependent triplet state population.** By addition of KI, the D-triplet state was sufficiently suppressed and hence, an almost continuous FRET-mediated excitation of A could be ensured. In the measurements, D was directly excited and A-autocorrelation curves were recorded for different $I_{exc}$ (6-120 kW/cm²) and D-A distances (10-41 Δbp). In addition to the diffusion of the molecules, three dark state relaxation processes were identified in the autocorrelation curves: the D-triplet ($\tau_T^D \approx 0.2 \mu$s), the A-triplet ($\tau_T^A \approx 3 \mu$s ) and the D-radical state ($\bar{\tau}_R^D \approx 15 \mu$s ). Examples of the resulting A-autocorrelation curves are shown in Fig. 3. As expected, the triplet state population increased and the triplet state relaxation time decreased with shorter D-A distances. However, for certain D-A distances, differences observed in the autocorrelation curves were smaller than expected. For instance, the correlation curves from samples 41 Δbp and 26 Δbp as well as 10 Δbp and 15 Δbp are relatively closely spaced in Fig. 3. A possible reason can be large subfractions of D-only or A-only labeled species that will bias the detected average triplet state population to lower values. The influence of incompletely labeled FRET pairs is a problem for most FRET measurements and usually requires extended control measurements to compensate for. In the following chapter, we will propose a procedure that will correct for...
D-only labeled samples without the need for complicated control measurements or additional samples.

Apart from addition of KI, oxygen can be removed from the buffer solution to increase the FRET-sensitivity of A-triplet state measurements (Fig. 3, inset). Oxygen is known to promote the triplet state transitions, in particular the relaxation from $T_1$ to $S_0$. Hence, removal of oxygen from the buffer typically leads to an increase in the triplet state population of the dye. When applying 5 mM KI and diminishing the oxygen concentration in the buffer, the effect of KI on the dyes predominated: $k_{\text{ISC}}^{D\rightarrow A}, k_T^{D}$ did not drastically change while $k_T^{A}$ was reduced to $k_T^{A} = 0.06 \cdot 10^6 \text{s}^{-1}$. Hence, the D-triplet state population was found to remain low while the A-triplet state population was found to clearly increase upon low oxygen concentrations. From Eq. 7 it follows that differences in low excitation rates then lead to larger absolute differences in $\bar{T}^A$. In line with this, we noted that upon reduction of oxygen and addition of KI, samples with low FRET efficiencies could be far better distinguished from each other ($\sim$ 150 % increase in the difference of the A-triplet fraction between the 41 Δbp and 26 Δbp sample). A complete removal of oxygen, however, may also lead to a stronger build-up of $\hat{R}^{-D}$, since oxygen can quench this state$^{18}$. Therefore, oxygen was not fully removed in these measurements.

**Donor bleed-through correction.** To enhance the resolution and minimize the influence of D-only labeled specimen on the FRET efficiency, a procedure was introduced to correct for the spectral bleed-through of D-fluorescence.

The detected emission signal consists of A-fluorescence and D-fluorescence leaking into the detection channel (Fig. 4A). Both, D- and A-fluorescence, contribute to the diffusion part of the autocorrelation curve. While the D-triplet state population has a negligible amplitude with $\tau_T^{D} \approx 0.2\,\mu\text{s}$, A has a prominent triplet state population with $\tau_T^{A} \approx 2\,\mu\text{s}$. Because the autocorrelation analysis does not differentiate between D and A-fluorescence, the
detected triplet state in the \(\mu s\) range will be an average of the D and A-triplet state, each
weighted with the molecular brightness of the corresponding fluorophore in the detection
channel. Hence, due to D-bleed-through, the measured A-triplet amplitude \(\bar{T}_m^A\) will be
reduced compared to the actual average triplet state population \(\bar{T}^A\) of the acceptor. \(^{27,28}\)

\[
\bar{T}_m^A = \frac{\bar{T}^A \left(1 - Q\right)^2}{1 + Q^2 \bar{T}^A \left(1 - \bar{T}^A\right)}.
\]

Here, \(Q\) is defined as the ratio of the detected D-fluorescence to the A-fluorescence.

\[
Q = \frac{d \cdot F_D}{d \cdot F_D + a \cdot F_A}
\]

Here, \(d\) and \(a\) denote the fractions of the D and A-fluorescence intensities, \(F_D\) and \(F_A\), that are
transmitted through the filter. \(d\) and \(a\) were determined from the emission spectra of D and A
and the transmission spectra of the corresponding emission filter and dichroic mirror used in
the experiment. The real \(\bar{T}^A\) can be calculated by first measuring the A-triplet amplitude \(\bar{T}_m^A\)
of the A-fluorophore with an emission filter 1 matched to the spectral region of the A-
fluorescence \((\rightarrow \bar{T}_m^A)\), followed by a second measurement with a slightly blue-shifted A-
emission filter 2 \((\rightarrow \bar{T}_m^A)\). \(\bar{T}_m^A\) will be lower than \(\bar{T}_m^A\) because of the stronger D-bleed-
through for filter 2. Using \(\bar{T}_m^A\), \(\bar{T}_m^A\) and the values determined for \(a_{1,2}\) and \(d_{1,2}\), \(\bar{T}^A\) can be
calculated by solving Eq. 16 and 17 for the two filters simultaneously.

**D-bleed-through correction compensates for different A-labeling efficiencies.** The
devised correction of \(\bar{T}^A\) is not only valid for D-bleed-through of D-A labeled samples, but
for incompletely labeled D-only samples as well. The correction procedure was tested on
samples of D-only and D-A labeled dsDNA strands added in different concentration ratios
(D-only/D-A = 2-144) and \(\bar{T}_m^A\), \(\bar{T}_m^A\) and \(\bar{T}^A\) were determined for each sample (Fig. 4B).
With an increasing amount of D-only labeled dsDNA, the measured triplet fraction of A decreased rapidly. Applying the slightly blue-shifted emission filter decreased $T_m^A$ even further. However, by applying the bleed-through correction procedure, $T^A$ was recovered over the full range of tested D-A ratios. Consequently, by this procedure, A-labeling efficiencies down to 1% could be tolerated while still performing robust FRET measurements.

**Acceptor triplet state measurements and the FRET electronic state model.** Being able to correct for donor bleed-through and varying A-labeling efficiencies, we investigated D-A samples for different FRET efficiencies. The donor was excited for seven $I_{exc}$ (7 - 120 kW/cm²) and the autocorrelation curve $G(\tau)$ of the A-channel was measured consecutively for the two A-emission filters 1 and 2. $G(\tau)$ was then fitted according to Eq. 6 including three dark states corresponding to the D-triplet, the A-triplet and the D-photo-reduced state. For each $I_{exc}$, the measured triplet fractions $T_m^A$ of filter 1 and 2 were used to calculate the real $T^A$ according to Eq. 16 and 17. Subsequently, $T^A$ and $\tau^A$ from all $I_{exc}$ were fitted to Eq. 7 according to the FRET electronic state model depicted in Fig. 1. In this fitting procedure, E was the only free fitting parameter. $k_{ISC}^D$, $k_T^D$, $k_{ISC}^A$ and $k_T^A$ were obtained by FCS measurements by direct excitation of each fluorophore (on dsDNA; see section 1 above) and were kept constant during the fitting procedure. As an alternative, or to circumvent the need for a red laser line, $k_{ISC}^A$ and $k_T^A$ can be determined by exciting the D of a FRET sample with known E and detecting the A-autocorrelation curve. In this case, the A-triplet state properties $T^A$ and $\tau^A$ can be determined from the A-autocorrelation curve and then fitted to Eq. 1 and 7 (now with the known $E$ as a constant value) to obtain $k_{ISC}^A$ and $k_T^A$.

The experimentally determined $T^A$ and $\tau^A$ are in good agreement with the fitted data (Fig. 5). For a single FRET sample, $T^A$ increases with increasing $I_{exc}$ while $\tau^A$
The efficiency, $\tilde{F}$, simultaneously decreases. At constant $I_{\text{exc}}$, $\tilde{T}$ increases with $E$ while $\tilde{\tau}$ is reduced with increasing $E$. The different FRET samples can be clearly resolved.

It should be noted that in the presented data analysis $E$ is the only free fitting parameter. This makes the analysis robust and, theoretically, a single data point (sample excited with one $I_{\text{exc}}$) is sufficient to calculate the FRET efficiency. However, to improve the accuracy of $E$, multiple data points were recorded for each sample at different $I_{\text{exc}}$.

**FRET efficiencies**. The FRET efficiencies obtained from the fits of $\tilde{T}$ and $\tilde{\tau}$ (Fig. 5) are shown in Fig. 6. The corresponding standard deviations, determined from 4 measurements, are about 10%.

For comparison, theoretical FRET efficiencies were determined based on the DNA model of a double-helix B-DNA, where the distances between the fluorophores ATTO 488 and Alexa 610 and the corresponding $E$ were calculated based on a Förster radius of 5.6 nm (see Materials and Methods) and according to Eq. 13-15. The FRET efficiencies determined with FCS were found to be in adequate agreement with these theoretically calculated FRET efficiencies.

**3. FRET efficiencies determined by TRAST.**

Sequentially, the donor was excited with ~50 different pulse trains whose pulse widths, $w$, were varied between 100 ns and 10 ms. The pulse height (e.g. the laser intensity) and the pulse train duration were kept constant ($I_{\text{exc}}$ between 30 and 80 kW/cm$^2$ and camera exposure times between 4 and 10 s) at a duty cycle of 1 %. For each $w$, the corrected A-fluorescence images were integrated over a circular region ($\varnothing$ 1.5 μm) in the center of the laser focus. After every fourth pulse train, a calibration measurement was taken at the shortest pulse width to account for concentration changes in the sample.
**Donor bleed-through correction.** As in the FCS measurements, the detected intensity $\bar{F}$ consisted of D and A-fluorescence, $\bar{F} = a \cdot \bar{F}_A + d \cdot \bar{F}_D$. To remove the D-bleed-through, the measurement procedure was performed with two emission filters successively: filter 1 matched the A-emission spectrum ($\bar{F}_1$) and filter 2 collected only D-fluorescence ($\bar{F}_2$; filter 1: 675/135 nm, $d_1 = 3.2\%$, $a_1 = 82.7\%$; filter 2: 530/50 nm, $d_2 = 57.6\%$, $a_2 = 0\%$). By knowing the spectral properties of the dyes, the dichroic mirror and the two filters, $\bar{F}_A$ was computed for each pulse width according to

$$\bar{F}_A(w) = \frac{1}{a_1} \left( \bar{F}_1(w) - \frac{d_1}{d_2} \bar{F}_2(w) \right)$$

**TRAST confirms FRET dependent triplet state build-up.** The integrated A-fluorescence ($\bar{F}_A$) is plotted against the pulse width ($w$) for various D-A distances measured at the same $I_{\text{exc}}$ (Fig. 7). A fast and a slow decrease of $\bar{F}_A$ can be seen for increasing $w$. These two decreases correspond to two photo-induced processes: the fast process ($w \sim \mu s$) originates from A-triplet state formation, while the slow process ($w \sim ms$) arises from transitions into the photo-oxidized radical state $\hat{R}^+$ of A. At similar $I_{\text{exc}}$, these transient state populations increase significantly with decreasing D-A distances.

For comparison, the corresponding D-fluorescence ($\bar{F}_D$) for each D-A sample is plotted in red dots in Fig. 7. $\bar{F}_D$ is approximately constant for $w \sim \mu s$ and increases only slightly for the high FRET sample (10 Δbp). The slight increase in $\bar{F}_D$ suggests that the A-triplet state is less FRET-active than the A-singlet state, which would contradict the assumption made in Eq. 2 and 3. One also has to consider that the increase in $\bar{F}_D$, resulting from D-A pairs, is diminished compared to the theoretically expected value due to a D-only labeled sub-fraction of DNA. Still, the increase in $\bar{F}_D$ is smaller than theoretically expected from the decrease in $\bar{F}_A$. To clarify how FRET-active the A-triplet state is, additional
fluorescence cross-correlation experiments were performed. Given that the A-triplet state
would be FRET-inactive, a transition of A into the triplet state would de-quench and increase
the D-fluorescence while A becomes non-fluorescent. The resulting cross-correlation between
the D- and A-fluorescence would show an anti-correlation in the time range of the A-triplet
state. No such anti-correlation could be observed in the μs time range (data not shown).
Hence, the A-triplet state is FRET-active, although supposedly with a slightly reduced
efficiency a compared to the \( S^D_1 \rightarrow S^A_0 \) efficiency. This finding thereby supports the
approximation made in Eq. 2.

In addition to the electronic states depicted in Fig. 1, a photo-oxidized radical state of
D, \( \hat{R}^{+D} \), is observed in the TRAST measurements (Fig. 7). \( \hat{R}^{+D} \) builds up for \( w \sim \text{ms} \) and
reduces the average \( \bar{F}_D \). On the other hand, there is also a build-up of the A-radical state \( \hat{R}^{+A} \),
which is FRET-inactive. This means that D cannot transfer energy to A any longer and \( F_D \) is
increased. As can be seen in Fig. 7, the influences of both phenomena on \( \bar{F}_D \) largely cancel
each other out. However, at low \( E \), the first process slightly dominates, whereas at higher \( E \),
the effect of the \( \hat{R}^{+A} \) build-up on \( \bar{F}_D \) is more prominent.

The effect of A-transient states is much more prominent on the \( \bar{F}_A \) curve than on the
\( \bar{F}_D \) curve (Fig. 7). Thus, especially for low FRET efficiencies, \( \bar{F}_A \) is more sensitive to FRET
and hence more suitable as a read-out parameter for FRET determination by TRAST.

**Determination of FRET efficiencies.** FRET efficiencies were calculated by fitting
Eq. 8 and 9 to the bleed-through corrected A-fluorescence \( \bar{F}_A(w) \). The thereby obtained \( k_{ISC}' \)
is then used to calculate the FRET efficiency from Eq. 3 and 4. In Eq. 8 and 9, \( k_{ISC}' \), \( k_{ox}' \) and
\( k_{ed} \) are free fitting parameters, while the parameters \( k_{ISC}^A \), \( k_T^A \), \( k_{10}^{DA} \) and \( \sigma^{DA}_{exc,491} \) were kept
constant for all samples. Here, \( k_{ISC}^A \) and \( k_T^A \) were determined from TRAST measurements of a
D-A labeled reference sample (15 Δbp, E = 0.4) by fitting the measured fluorescence to Eq. 8 and 9.

The obtained fits were in good agreement with the experimental data (Fig. 7). The obtained redox rates ($k_{ox}^A \approx 1.4 \cdot 10^3 s^{-1}$, $k_{red}^A \approx 0.5 \cdot 10^3 s^{-1}$) were similar for all FRET samples. The FRET efficiencies computed from TRAST measurements were in good agreement with FRET efficiencies obtained from FCS and from the DNA model (Fig. 6).

For the TRAST measurements, the error-margins in Fig. 6 were calculated by propagating the 95 % confidence interval obtained by curve fitting Eq. 8 and 9 to the computed $F_A$ and by assuming 5 % uncertainty in the laser power measurement.

4. FRET efficiencies determined by D-lifetime measurements.

For comparison, D-lifetime measurements were also performed using time correlated single photon counting. In the absence of KI and for optimal labeling conditions (almost each D has an A), the D-only lifetime could be fitted with a single-exponential decay having a lifetime of 4.05 ± 0.01 ns. D-lifetimes in the presence of A showed a multi-exponential decay behavior that could be fitted satisfactorily only with a two to three component lifetime model. One component with a decay time of $\tau_{F1} \sim 2.5$ ns was observed for all samples, while another component with a decay time of $\tau_{F2} \sim 4$ ns, corresponding to D-only labeled dsDNA, could only be distinguished from the FRET-reduced lifetimes ($\tau_{F3}$) measured in D-A samples with Δbp < 24. Calculating $E$ from $\tau_{F3}$ did not result in reasonable $E$ values. Therefore an average fluorescence lifetime (Eq. 11) was computed from $\tau_{F1}$ and $\tau_{F3}$ and used for further FRET evaluations (Eq. 12). The resulting FRET efficiencies are plotted in Fig. 6.

Discussion

Our results suggest that, even in the presence of a large D-only labeled sample fraction, A-triplet state measurements can extend the useful range of intra- or intermolecular distances beyond the 10 nm limit often stated for FRET-based methods. We have
shown that FCS and TRAST measurements could unambiguously differentiate between samples of 0.4 %, 3 % and 5 % FRET efficiency, which corresponded to D-A distances of 14 nm, 10 nm and 9 nm in this study. The presented approach monitors the triplet state build-up of the acceptor, which in turn depends on the FRET-mediated excitation of the dye. The general strategy of exploiting photo-induced, long-lived transient states of A was previously demonstrated by FRET-FCS measurements using the trans-cis isomerization kinetics of the cyanine dye Cy5 as a readout.\textsuperscript{30} One advantage of exploiting the triplet state kinetics instead, as done in this study, is that this state is inherent to practically all organic fluorophores, which thus broadens the choice of available A-dyes. In addition, because of an additional spectral correction used to compensate for D-bleed-through, also FRET pairs with higher spectral overlap and possibly larger Förster radii can be applied. This should make it possible to observe D-A distances even beyond 15 nm. In this study, KI was used to suppress the D-triplet state and increase the sensitivity of the A-triplet state. The same effect can be obtained by using an A with a pronounced triplet state build-up.\textsuperscript{31} In this case, the excitation intensities can be reduced in order to avoid a D-triplet state build-up. The resulting decreased molecular brightness of A is not an issue for the TRAST method as it can be compensated for by using higher concentrations or longer acquisition times.\textsuperscript{14}

The results also predict that the technique is applicable to imaging using a standard laser scanning microscope or a wide-field microscope equipped with a time-modulated excitation. As a proof of principle, we have performed TRAST measurements in a one point modus. The presented TRAST method showed that accurate FRET efficiencies could be calculated by modulating the excitation light in an on-off fashion using an AOM. However, the same kind of on-off modulation can also be obtained by scanning a laser beam over the sample and taking images with different scanning speeds, which result in different pixel dwell times.\textsuperscript{16} Therefore, the same theory and data analysis that has been developed here can be applied for both TRAST concepts, independent of the experimental realization used.
Applying FCS for solution measurements or TRAST for FRET-imaging has several advantages. We could show that, in the A-triplet state measurements, influences of D-bleed-through and therefore also of D-only labeled sub-fractions can be fully corrected for. By the proposed correction procedure, D-only sub-populations as high as 99% could be tolerated when calculating the FRET efficiencies of the D-A samples. The influence of direct excitation of A in a D-A pair is included in the analysis. However, the influence of directly excited A-only labeled sub-fractions can only be disregarded as long as the direct excitation of A is negligible. In our measurements, a small fraction of A-only labeled samples did not change the FRET efficiency notably. However, for A-only fractions larger than 50%, we observed a moderate decrease in the apparent FRET efficiency.

In comparison, the corrections for spectral bleed-through in intensity-based FRET measurements are less robust. Here, the FRET efficiencies are determined from the ratio of the fluorescence intensities detected in the A- and in the D-channel. The A-D intensity ratio has to be corrected for D-bleed-through into the A-channel and direct excitation of A. The D bleed-through can be determined from the ratio of the D-fluorescence of a D-only labeled sample in the D- and A-channel. The extent of direct excitation of A is estimated by exciting A of an A-only labeled sample with the laser line used for FRET measurements and one suitable for direct excitation of A. However, the latter ratio will only be an estimate, because the A-intensity obtained by direct excitation might not be proportional to the A-intensity obtained under different FRET conditions. For intensity-based FRET measurements, $E$ will be an average of $E$ from the D-A labeled sub-population and the D-only labeled sub-population with $E = 0$. Only an apparent, reduced $E$ can be determined that will be dependent on the amount of D-only labeled samples. In this case, $E$ can only be compared between different probes given that the sample concentrations and labeling efficiencies are comparable from experiment to experiment.
In contrast, by A-triplet state measurements, one can suppress the influence of D-only labeled sub-populations by a simple reference measurement on the same sample. A D-only labeled reference sample is still required to determine the D-triplet state properties. However, the A-only labeled reference sample can be replaced with a sample of known FRET efficiency.

FRET lifetime measurements, on the other hand, have no constraints concerning spectral bleed-through. $E$ is determined from the D-fluorescence only, which, for appropriate filter conditions, will not be contaminated by the red-shifted A-fluorescence. For the same reason, A-fluorescence of directly excited A will not influence the D-lifetime measurements. D-Lifetime measurements are able to distinguish D-A and D-only labeled sample fractions for high and intermediate $E$ and reasonable D-only/D-A ratios. However, for lower $E$, the FRET-quenched D-lifetime becomes comparable to the unquenched D-lifetime and the two components can not be properly separated. The resulting $E$ will then be reduced compared to the actual D-A FRET efficiency. For high D-only/D-A ratios, the contribution of the quenched in relation to the unquenched D-fluorescence will be minimal and difficult to distinguish in the data analysis.

The FRET D-lifetime measurements, presented in this study, were performed for optimal labeling conditions (ideally each D has an A). Given this, we were able to distinguish also low FRET efficiencies from each other (Fig. 6). This would not have been possible for samples with a varying amount of D-only labeled sub-populations.

Interestingly, FCS, TRAST, as well as lifetime measurements underestimated the FRET efficiency of the high FRET sample (10 Δbp, $E_{\text{model}} = 0.95$). Strongly reduced FRET efficiencies for similar D-A distances have been reported before.\textsuperscript{19} Possible reasons are (i) dimerization of D and A into a non-fluorescent complex, (ii) charge transfer between D-A through space interactions or (iii) higher anisotropy values of D and A compared to the theoretical values used for random orientation of the fluorophores. Contrary to Dietrich et
we did not observe dimerization (no shift in the absorption and emission spectra of the fluorophores bound to dsDNA). However, in the FCS measurements, we did observe a strongly reduced molecular brightness of A for the 10 ∆bp sample in comparison to the 15 ∆bp sample. This reduced molecular A-brightness can, at least partially, be attributed to a strong bleaching of Alexa 610 due to the high FRET rate and simultaneous direct laser excitation. This is in analogy to a previous study\textsuperscript{29}, where the photobleaching of red fluorophores was increased in the presence of a blue-shifted excitation outside of the excitation spectrum of the fluorophores. Hence, the population of higher excited states and consecutive bleaching from those states may be an underlying mechanism.

The multi-exponential decay in our D-lifetime measurements suggests additional energy transfer pathways from D to A, e.g. complex formation of the fluorophore with the base guanosine\textsuperscript{32} or electron hopping \textit{via} the guanosine bases of the DNA.\textsuperscript{33,34} These processes would increase the apparent FRET efficiency, which is consistent with the FRET efficiencies determined from the D-lifetime measurements in Fig. 6. On the other hand, these additional processes can be included in the A-triplet state measurements, if $k^A_{\text{ISC}}$ and $k^A_T$ are determined on a D-A labeled reference sample with known FRET efficiency. At the same time, the influence of excitation saturation and bleaching artifacts on the FRET efficiency analysis is minimized.

Saturation and bleaching influence the determined $k^A_{\text{ISC}}$ and $k^A_T$ differently for FCS ($k^A_{\text{ISC}} = 2.1 \pm 0.1 \cdot 10^6 \text{s}^{-1}$, $k^A_T = 0.2 \pm 0.01 \cdot 10^6 \text{s}^{-1}$) and TRAST ($k^A_{\text{ISC}} = 3.6 \pm 0.13 \cdot 10^6 \text{s}^{-1}$, $k^A_T = 0.14 \pm 0.01 \cdot 10^6 \text{s}^{-1}$) measurements. For the applied excitation intensities in the upper range, FCS will tend to underestimate $k^A_{\text{ISC}}$.\textsuperscript{22} This is reflected in the apparently decreased diffusion times observed in the correlation curves for higher excitation rates and indicates bleaching and thus depletion of singlet state fluorophores in the center of the focus. In TRAST, on the other hand, lower excitation intensities were used than for FCS. In addition,
over the course of the whole pulse train a small fraction of A will bleach away that will be larger for longer pulse widths than for shorter pulse widths. As the diffusion of the DNA strands is quite slow, the bleached fluorophores will not totally renew within the off periods of the light modulation. This will lead to an apparently larger A-triplet fraction and therefore to an overestimation of $k_{ISC}^A$. In order to obtain absolute FRET efficiencies using FCS or TRAST, it is thus recommended to calibrate the A-triplet rates for a given FRET pair by using a reference sample with known FRET efficiency. Very often however, only relative distances are to be measured, and these are very well reflected in the A-triplet state population without the need of an additional calibration step.

In conclusion, A-triplet state measurements provide a powerful tool to investigate long range FRET efficiencies. Combined with a novel suppression algorithm for spectral bleed-through, the technique estimates accurately even low FRET efficiencies $\leq 5\%$, a FRET regime where donor lifetime or intensity based measurements are usually biased by D-only labeled specimen or by additional energy transfer processes. We predict that the approach will prove useful also on conventional laser scanning microscopes and wide-field microscopy settings.

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References


**Figure 1:** Electronic state model of D and A coupled via FRET with rate constant $k_{\text{FRET}}$. Each molecule consists of a ground and an excited singlet state, $S_0$ and $S_1$, the first excited triplet state, $T_1$, and a radical state $\cdot R$. $k_{01}$, $k_{10}$, $k_{\text{ISC}}$, $k_T$, $k_{\text{ox}}$ and $k_{\text{red}}$ denote the rate constants for excitation and de-excitation of the singlet state, intersystem crossing and de-excitation of the triplet state and oxidation and reduction rate of the radical state, respectively. Upper indices D, A relate the corresponding rates or states to the donor or acceptor molecule.

**Figure 2:** Influence of potassium iodide (KI) on D and A bound to dsDNA. Experimental (□) and fitted (□) correlation curves are shown for D and A in the absence and presence of KI. A: In the presence of 5 mM KI, the triplet state population of ATTO 488 is decreased (indicated by the arrow) and shifted to faster times (2.5 µs → 0.2 µs). $I_{\text{exc, 488 nm}}$: 189 kW/cm$^2$, emission filter: 532/70 nm. B: The triplet fraction of Alexa 610 increases upon addition of KI (indicated by the arrow) while the triplet relaxation time remains constant at 3 µs. $I_{\text{exc, 568 nm}}$: 28 kW/cm$^2$, emission filter: 640/115 nm.

**Figure 3:** A-correlation curves for FRET mediated excitation of oligonucleotides with different D-A base pair separations Δbp. The experimental (□) and fitted (Δbp = 10 □, 15 □, 20 □, 24 □, 26 □, 41 □) correlation curves (Eq. 4) consist of one diffusion component (Diff., $\tau_D \sim 350$ µs) and three dark states corresponding to the D-triplet ($T_1^D$, $\tau_T^D \sim 0.2$ µs), the A-triplet ($T_1^A$, $\tau_T^A \sim 3$ µs) and the D-radical state ($\cdot R^D$, $\tau_{\cdot R}^D \sim 10$ µs). Corresponding residuals are shown below. The A-triplet state population increases in proportion to the FRET.
efficiency $E_{exc,488\text{ nm}}$: 119 kW/cm$^2$, emission filter: 675/135 nm. **Inset:** A-correlation curves for different FRET samples in a low oxygen solution under argon flow. Applying an argon flow markedly increases the triplet state population and the difference in the triplet state populations especially for low-FRET samples.

**Figure 4:** Influence of D-only labeled samples on the A-triplet amplitude. **A:** Fluorescence signal detected in the A-channel. The signal consists of A-fluorescence, $F_A$, and D-fluorescence, $F_D$, leaking into the channel. The modulation in the signal originates mainly from singlet-triplet transitions ($S^A_1 - T^A_1$) of A. The singlet-triplet transitions of D are suppressed by KI, thereby ensuring an almost constant excitation of A via FRET. **B:** Correction of $\bar{T}^A$, which was determined from A-autocorrelation curves for FRET-mediated excitation. Because of spectral bleed-through, the average triplet state population of Alexa 610 (▼, filter 1) decreased upon addition of D-only labeled dsDNA. Applying a slightly blue-shifted emission filter 2 decreased the triplet state population (▼) even further. The influence of the spectral bleed-through could be eliminated by applying the correction procedure described in the text. The resulting real $\bar{T}^A$ (○) could be recovered over the full range of D/A concentrations. The dashed line represents the mean value of all recovered $\bar{T}^A$. Data points represent the mean value of 6 measurements and the corresponding standard deviation in those. $I_{exc,488\text{ nm}}$: 55 kW/cm$^2$, emission filter 1: 675/135 nm, $d_1 = 3.45 \%$, $a_1 = 79.7 \%$; emission filter 2: 640/115 nm, $d_2 = 9.01 \%$, $a_2 = 80.4 \%$; sample: 15 Δbp.

**Figure 5:** Experimental (filled dots) and fitted (empty dots connected by line) triplet state populations (A) and triplet state relaxation times (B) of A for different FRET oligonucleotides (Δbp = 10 ●, 15 ●, 20 ●, 24 ●, 26 ●, 29 ●, 41 ●). Autocorrelation curves of A were recorded for various $I_{exc,488\text{ nm}}$ (7 – 120 kW/cm$^2$) for the emission filters 675/135 nm and 640/115 nm.
and analysed (Eq. 5). The real $\bar{T}^A$ were calculated by applying the correction procedure described in the text. $\bar{T}^A$ and $\bar{\tau}_T^A$ were fitted with the electronic state model depicted in Figure 1. Here, $k_{\text{ISC}}^A$ and $k_T^A$ were determined from a reference sample with known FRET efficiency (15 Δbp, $E = 0.4$; $k_{\text{ISC}}^A = 2.1 \cdot 10^6 \text{s}^{-1}$, $k_T^A = 0.2 \cdot 10^6 \text{s}^{-1}$). For a single FRET sample, $\bar{T}^A$ increases in proportion to $I_{\text{exc}}$ whereas $\bar{\tau}_T^A$ decreases with $I_{\text{exc}}$. At a constant $I_{\text{exc}}$, $\bar{T}^A$ increases with increasing $E$ whereas $\bar{\tau}_T^A$ is reduced at the same time, as indicated by the arrows.

**Figure 6:** FRET efficiencies obtained for optimal D-A labeling conditions from a theoretical DNA-model (□), A-FCS (●), A-TRAST (■) and D-lifetime (▼) measurements. D-A distances (upper x-axis) were calculated with the dsDNA model according to the D-A base pair difference and a Förster radius of $R_0 = 5.6$ nm. The corresponding standard errors of the mean of $E$ were determined from 4 (FCS) or 10 (lifetime) measurements. The A-FCS FRET efficiencies agree well with the $E$ calculated from the DNA-model whereas lifetime measurements tend to overestimate $E$, especially for lower $E$. The high FRET sample (10 Δbp) is underestimated by both FCS and lifetime measurements due to bleaching (FCS) and a secondary charge transfer process between the D and A through space interactions (FCS, lifetime).

**Figure 7:** TRAST measurements of different FRET samples with Δbp = 10, 20, 26 and 41 at excitation intensities of $I_{\text{exc}} = 50$ kV/cm². The normalized measured (□) and fitted (□) A-fluorescence decrease with increasing excitation pulse width according to the triplet (3 µs) and redox state (2 ms) build-up. The D-fluorescence (•) increases with excitation pulse width due to a reduced energy transfer from D to A because of the triplet and redox state build-up of A. Samples at increasing FRET efficiencies ($E$) are indicated by the arrows. At similar donor
excitation intensities, the decreases in A-fluorescence and the increases in D-fluorescence become more pronounced for higher $E$. Fitting rates used here: $k_{ISC}^A = 3.6 \times 10^6 \text{s}^{-1}$, $k_t^A = 0.14 \times 10^6 \text{s}^{-1}$, $k_{10}^D = 261.78 \times 10^6 \text{s}^{-1}$, $k_{10}^A = 255.1 \times 10^6 \text{s}^{-1}$.

Figure 1:

![Figure 1 diagram]

Figure 2:

![Figure 2 graph]

Figure 3:

![Figure 3 graph]

Figure 4:
Figure 5:

A

B
Figure 6:

![Graph showing FRET efficiency vs. distance D-A (nm) with data points for DNA model, FCS, TRAST, and Lifetime.](image)

Figure 7:

![Graph showing normalized fluorescence vs. pulse width (ms) with different Δbp values.](image)