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Reliability and accuracy of single-molecule FRET studies for characterization of structural dynamics and distances in proteins

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Single-molecule Förster-resonance energy transfer (smFRET) experiments allow the study of biomolecular structure and dynamics in vitro and in vivo. We performed an international blind study involving 19 laboratories to assess the uncertainty of FRET experiments for proteins with respect to the measured FRET efficiency histograms, determination of distances, and the detection and quantification of structural dynamics. Using two protein systems with distinct conformational changes and dynamics, we obtained an uncertainty of the FRET efficiency \leq 0.06, corresponding to an interdye distance precision of \leq 2 Å and accuracy of \leq 5 Å. We further discuss the limits for detecting fluctuations in this distance range and how to identify dye perturbations. Our work demonstrates the ability of smFRET experiments to simultaneously measure distances and avoid the averaging of conformational dynamics for realistic protein systems, highlighting its importance in the expanding toolbox of integrative structural biology.

Förster-resonance energy transfer (FRET) studies have become a widely used approach to complement classical structural biology techniques ¹⁻⁴. They provide information on the structure and conformational heterogeneity of biomolecules over a distance range of 30 to 120 Å and, when performed on single molecules, contribute additional information regarding conformational dynamics on the timescales of nanoseconds to seconds ^{1,2,5-10}. They also allow for quantitative assessment of structural dynamics and heterogeneity of conformational ensembles. This information is not easily accessible by X-ray crystallography, cryogenic-electron microscopy or cross-linking mass-spectrometry, which provide structural information of solution structures but lack temporal information. FRET can also be used to resolve (parts of) structures in an integrative manner (refs. 11–17) and has the unique ability to provide correlated information on structure and dynamics ^{1,2}.

Hellenkamp et al. presented a quantitative multilaboratory smFRET blind study assessing the validity of using smFRET for structural measurements. This study used static double-stranded DNA

(dsDNA) that demonstrated a high reproducibility between the different laboratories with an uncertainty of ≤ 6 Å for the FRET-derived distances 18 . These results strongly supported the idea that standardized smFRET measurements in combination with standardized data analysis routines are a useful addition to the integrative modeling of static biomolecular structures 12,19,20 .

Here, we assessed whether the established procedures translate to more flexible biomacromolecules such as proteins that undergo conformational changes. Compared to dsDNA, proteins are more challenging systems, because the local environments and flexibility of the tethered dyes can vary considerably. Site-specific dye labeling of proteins usually requires the introduction of point mutations (for example, cysteines or nonnatural amino acids), which can affect its structure and function¹. Moreover, proteins require careful handling and storage due to sample instability and aggregation, and are sensitive to experimental conditions, buffer composition, pH, temperature, surface interactions and so on. In a blind comparison study involving

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19 laboratories using diffusion-based confocal smFRET, we investigated the maltose-binding protein (MalE) and the U2 Auxiliary Factor 2 (U2AF2), which display conformational dynamics on different time and length scales. We addressed two key questions: (1) how consistently can smFRET efficiency histograms (and the derived distances) be determined by different laboratories for protein samples prepared with stochastic fluorophore labeling? (2) How reliably can smFRET measurements detect structural dynamics in these proteins and what are the minimal structural fluctuations detectible?

Our study confirmed the reproducibility of accurate FRET efficiency histograms and the ability of smFRET to detect and quantify conformational dynamics on the submillisecond timescale. We demonstrate reproducible FRET efficiency values with uncertainties ≤0.06 corresponding to a distance precision of ≤ 2 Å and an accuracy ≤ 5 Å in MalE. Moreover, we compare the variability of setup-dependent parameters and identified the main sources of calibration uncertainty. To push the detection limits for structural dynamics, we refined established experimental and data analysis procedures and studied distinct dye pairs to identify and eliminate dye-specific effects. With this, we could detect distance fluctuations on the order of 5 Å in the FRET-sensitive range. Our work demonstrates that smFRET is able to characterize challenging and realistic protein systems with conformational dynamics on timescales from nanoseconds to seconds, highlighting its importance in the expanding toolbox of integrative structural biology¹⁹⁻²¹.

Results

We chose two protein systems with conformational dynamics on different timescales. Our first target was the MalE protein of *Escherichia coli*, the periplasmic component of the ATP binding cassette transporter MalFGK₂-E (refs. 22–24). MalE exhibits a typical periplasmic-binding protein fold^{25,26} composed of two rigid domains connected by a flexible two-segment ß-stranded hinge (Fig. 1a). This structure enables an allosterically driven motion from an open to closed state upon maltose binding on the subsecond timescale (Supplementary Fig. 1). As a second system, we chose the large subunit of U2AF2 from the pre-messenger RNA (mRNA) splicing machinery²⁷. Its two RNA recognition motif domains (RRM1,2) are connected by a long flexible linker and bind single-stranded Py-tract RNA²⁸. For U2AF2, the two domains fluctuate between an ensemble of detached conformations and a compact conformation in the apo state²⁹, whereas ligand binding stabilizes an open conformation (Fig. 2a)³⁰.

SmFRET experiments were blindly performed by 19 laboratories for MalE and by seven laboratories for U2AF2 using different implementations of diffusion-based confocal spectroscopy with alternating excitation, that is, microsecond-ALEX (alternating laser excitation mode)³¹ for intensity-based analysis and nsALEX³² or pulsed-interleaved excitation (PIE)³³ for intensity- and lifetime-based analyses (Supplementary Fig. 2). To avoid additional complexity and to restrict any preknowledge regarding the samples, the proteins were labeled and checked for functionality before being delivered to the participants. Information regarding the identity of the proteins and ligands, labeling positions, labeling efficiency, and expected FRET efficiencies and changes were not provided. The laboratories were informed about which fluorophores were coupled. We adapted a data analysis routine similar to ref. 18 to determine setup-independent accurate FRET efficiency E values from the photon counts detected in the donor (D) and acceptor (A) detection channels during a single-molecule event. The procedure is described in the Methods and includes subtraction of background signals from all channels and the determination of four correction factors: (α) for spectral crosstalk of D fluorescence into the A channel, (β) for normalization of direct D and A excitation fluxes, (y) for differences in D and A quantum yields and detection efficiencies and (δ) for the ratio of indirect and direct A excitation (Supplementary Fig. 3 and Supplementary Tables 1 and 2)³⁴. The use of ALEX or PIE (Supplementary Note 1) was crucial for corrections of the photon counts to reflect the actual D and A signal and exclusion of single-molecule events from incompletely labeled molecules or ones showing photo-blinking and bleaching 8,9,18,34,35.

MalE

We prepared three double-cysteine variants of MalE with interresidue distances that cover a large part of the dynamic range of FRET (Fig. 1b, Methods and Supplementary Fig. 4). The variants were designed to show a decrease (MalE-1, K29C-S352C), an increase (MalE-2, D87C-A186C) or an unaltered interdye distance (MalE-3, A134C-A186C) upon maltose binding. All variants of MalE were stochastically labeled in one of the laboratories at the given positions with the donor Alexa Fluor 546 (Alexa546) and acceptor Alexa Fluor 647 (Alexa647). Before shipment, we confirmed the functionality of the labeled protein by ligand titrations using smFRET and microscale thermophoresis, and verified that maltose did not affect the dye properties (Supplementary Figs. 5 and 6). To allow a comparison, participants were asked to provide mean FRET efficiencies using Gaussian fits for apo and holo FRET efficiency histograms (Fig. 1c) and to determine a global y value for all measurement conditions (Supplementary Note 2 and Supplementary Fig. 3). For this workflow, participants used custom or publicly available software packages.

FRET efficiency histograms for representative experiments on MalE in the apo (no ligand) and the holo state (1 mM maltose) are shown in Fig. 1c with mean values reported by 16 laboratories. They show very good agreement and reproducibility. It was not possible to extract accurate FRET efficiency values from three laboratories due to, for example, missing or suboptimal laser lines (Supplementary Table 1 and Supplementary Note 2). All laboratories observed the expected changes for MalE-1, MalE-2 and no shift for MalE-3. This indicates that the samples did not degrade during shipment on dry ice and storage in the laboratories at 4 °C. MalE-1 showed an average FRET efficiency of 0.49 ± 0.06 in the apo state that increased to 0.67 ± 0.05 in the holo state. MalE-2 showed the expected decrease in FRET efficiency from 0.83 ± 0.03 to 0.71 ± 0.05 in the apo and holo states, respectively (Fig. 1c). MalE-3, with both labels on one lobe, showed no significant change in FRET efficiency ($E_{apo} = 0.91 \pm 0.02$, $E_{holo} = 0.92 \pm 0.02$).

The standard deviation of the determined mean FRET efficiency over all laboratories was less than ± 0.06 , similar to the precision found for dsDNA previously 18 (Extended Data Table 1 and Supplementary Table 3). We observe the highest standard deviation for MalE-1 and the lowest values of ± 0.02 for MalE-3, which also has the highest FRET efficiency. We observed systematic deviations of the reported FRET efficiency values for the apo and holo states from the mean value. Hence, we analyze the individual FRET efficiency differences, $\langle E_{\rm holo} \rangle - \langle E_{\rm apo} \rangle$, between the apo and holo states for the different laboratories (Fig. 1d). The distributions indeed narrow for all samples by approximately twofold because systematic deviations cancel out $(\sigma_{\langle E_{\rm holo} \rangle - \langle E_{\rm apo} \rangle}$ for MalE-1 ± 0.02 , MalE-2 ± 0.02 and MalE-3 ± 0.01 : Fig. 1d, Extended Data Table 1 and Supplementary Table 3).

U2AF2

For the second protein, U2AF2, we chose the published double-cysteine variant L187C-G326C of the minimal RRM1,2 construct, where we previously verified that protein function is not affected by labeling (Fig. 2a)^{36,37}. The construct was labeled stochastically on the two RRM domains with the dye pair Atto532–Atto643. A subset of seven groups measured the sample. To investigate the consistency of the obtained FRET efficiency histograms, we plotted the smFRET histograms from individual laboratories (Fig. 2b,c, row 1) as well as the average distribution illustrated by the mean and standard deviation (row 2). All groups found a single broad distribution (Fig. 2b, row 1, apo) with an average $E = 0.74 \pm 0.03$ (row 2). In the presence of 5 μ M ligand (U9 RNA, K_d of roughly 1.3 μ M), a second narrower peak at lower E appears

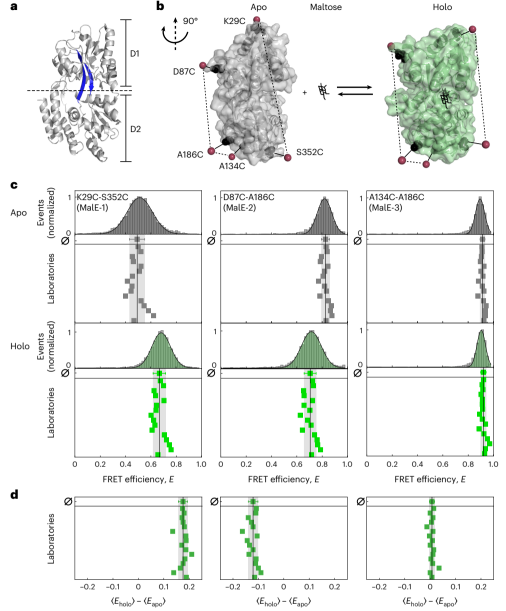


Fig. 1 | **Experimental design of MalE as a protein model system for smFRET studies. a**, Crystal structure of MalE in its ligand-free apo state (PDB ID 10MP) with domains D1 and D2 linked by flexible beta sheets (highlighted in blue). **b**, The crystal structure of MalE (rotated by 90° as compared to **a** in the apo (gray, PDB ID 10MP) and holo (green, PDB ID 1ANF) states with mutations at K29C-S352C (MalE-1), D87C-A186C (MalE-2) and A134C-A186C (MalE-3) indicated in black. Note, each mutant only contains one cysteine pair and was measured using the Alexa546-Alexa647 FRET pair. The estimated mean position of the fluorophores from AV calculations are shown as red spheres. **c**, FRET efficiency *E* histograms for three MalE mutants, MalE-1 (left), MalE-2 (middle) and MalE-3 (right), in the

absence and presence of 1 mM maltose (bottom, green) for one exemplary dataset measured in laboratory 1. The distribution is fitted to a Gaussian distribution. The reported mean FRET efficiencies for 16 laboratories are shown below (due to experimental difficulties, the results of three laboratories were excluded; Supplementary Table 1). The mean FRET efficiency and the standard deviation of all 16 laboratories are given by the black line and gray area. **d**, Individual FRET efficiency differences for each laboratory, between the apo and holo states, $\langle E_{\rm holo} \rangle - \langle E_{\rm apo} \rangle$, for MalE-1 (left), MalE-2 (middle) and MalE-3 (right). The mean FRET efficiency difference and the standard deviation of all 16 laboratories are given by the black line and gray area.

(Fig. 2c, row 1) with an average $E = 0.46 \pm 0.04$ (row 2) as expected for the open conformation of the holo state^{30,36}. Notably, a fraction of around 15% of ligand-free protein remains in the sample at the RNA concentration used (Supplementary Fig. 7).

For the apo state, we obtained a similar standard deviation of ± 0.03 as found for MalE, however, a clear outlier was apparent (Supplementary Table 4). To test whether user bias affected the reported results, a single person reanalyzed the datasets. This person developed an optimal procedure for determining the correction factors for this challenging sample (Supplementary Note 3) and improved the

agreement to a standard deviation of ± 0.008 with no change in mean E (Fig. 2d,e and Supplementary Table 4). The reanalysis revealed the detection correction factor γ to be the main cause of the deviations between the measurements. As a single population of the apo state did not allow for a robust determination of the γ factor $\gamma^{34,35}$, it was best to estimate the γ factor from a global analysis of the apo and holo measurements. This was possible since the quantum yield of the fluorophores remained unchanged upon RNA binding (Supplementary Table 5). We also reanalyzed data from the same seven laboratories for MalE-1 apo and obtained nearly identical mean FRET efficiencies

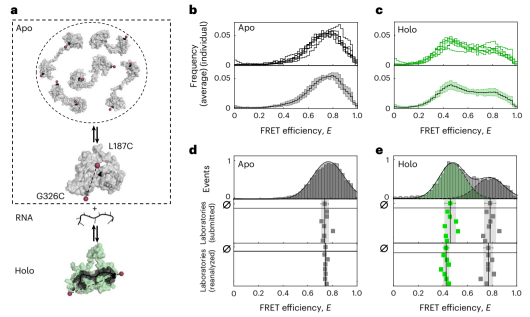


Fig. 2 | The experimental system of U2AF2 (RRM1, 2) and a comparison of FRET efficiency histograms from seven different laboratories. a, Schematic of the dynamics of U2AF2. The apo state (in gray, top) undergoes fast exchange between an ensemble of detached structures of which five representative structures are displayed. A slower exchange occurs between the dynamic detached ensemble and a compact conformation (PDB ID 2YHO) shown below. The holo state (in green, PDB ID 2YH1) bound to a U9 RNA ligand (in dark gray) assumes a well-defined, open conformation. Positions of cysteine mutations introduced for labeling (L187 in RRM1 and G326 in RRM2) are depicted as black spheres with the mean dye position determined by AV calculations indicated by red spheres. b,c, SmFRET efficiency histograms reported by the seven participating laboratories for apo (b) and holo (c) measurements of U2AF2. The top shows the individual FRET efficiency histograms and the bottom shows the average FRET efficiency histogram (solid line) with standard deviation (light area). d, SmFRET efficiency E histograms of U2AF2 in the apo state. The top shows a representative 1D FRET efficiency histogram with a Gaussian fit (laboratory 1).

The middle shows the reported mean FRET efficiencies reported by seven laboratories. The mean value from all datasets is 0.739 ± 0.029 , shown above with the corresponding standard deviation in gray. The bottom shows the extracted mean FRET values after reanalysis of the collected data. After reanalysis, the agreement improved to 0.742 ± 0.008 . **e**, SmFRET efficiency histogram comparisons of U2AF2 in the holo state. $5 \mu M$ of U9 RNA was used to obtain the holo state. The top shows a representative 1D FRET efficiency histogram of laboratory 1 fitted to two Gaussian distributions to determine the FRET efficiencies of the different subpopulations, yielding mean FRET efficiencies of 0.44 for RNA-bound and 0.76 for the RNA-free conformation. The middle shows the mean FRET efficiencies reported by the seven laboratories. The mean values from all seven of the datasets were 0.45 ± 0.04 for the RNA-bound conformation (in green) and 0.78 ± 0.04 for the RNA-free conformation (in gray). The bottom shows the reanalysis of the holo measurements yielding values of 0.42 ± 0.02 and 0.77 ± 0.03 for RNA-bound and RNA-free fractions, respectively.

and standard deviations (0.49 \pm 0.05 versus 0.47 \pm 0.06, Supplementary Fig. 8). This indicates that user bias was less pronounced when a global, well-defined analysis procedure for determining γ was provided over several samples covering a substantial fraction of the FRET range (Supplementary Note 2).

For the holo state of U2AF2, good agreement between laboratories was obtained for the peak positions with a standard deviation of ± 0.03 and ± 0.02 for the high- and low-FRET peaks, respectively. A minimal improvement resulted from the reanalysis (Supplementary Table 4). In contrast to the agreement in FRET efficiency, we observed variations in the relative amplitudes of the two populations: 0.58 ± 0.08 for the holo state and 0.42 ± 0.08 for the apo population (Fig. 2c and Supplementary Table 4). We attribute this to potentially reduced protein activity, degradation of the RNA ligand and sensitivity of conformational dynamics to the experimental conditions, for example, temperature, ligand concentration, buffer composition, salt concentration or the presence of stabilizers such as bovine serum albumin (BSA) (Supplementary Fig. 7).

Setup-dependent parameters and correction factors

The quality of smFRET experiments is determined by the statistics of the measurement and the performance of the setup to maximize photon collection and thereby minimize shot noise. To this end, we quantified the number of bursts, average photon count rate, burst duration and the number of photons in the D and A channels for the MalE measurements from eight laboratories (Fig. 3a and Supplementary

Fig. 9). On average, participants collected 6.000 bursts (minimum 500, maximum 21,000) of molecules carrying both fluorophores. The required number of bursts for a smFRET analysis depends on the goal of the experiment. To determine the average FRET efficiency from a single population, as performed for MalE, roughly 1,000 bursts of double-labeled molecules may be sufficient. For advanced analysis methods such as time-correlated single photon counting (TCSPC) for lifetime analysis, burst-wise fluorescence correlation spectroscopy (FCS) or a photon distribution analysis (PDA) that are applied to subensembles, higher burst numbers of >5,000 are desired. Typical count rates per single-molecule event were found to be 60 ± 20 kHz, with an average burst of 90 \pm 40 photons and 1.7 \pm 0.9 ms duration (Fig. 3a and Supplementary Fig. 9). The average count rate and burst duration depend on the size of the confocal volume, where smaller sizes result in higher count rates but shorter burst durations. We observe a negative correlation between burst duration and average count rate (Fig. 3b, Pearson's r = -0.58 and Supplementary Fig. 10). The large spread of the burst duration arises from the fact that some participants applied a diffraction-limited observation volume, while others underfilled the objective lens to create a larger confocal volume with a diameter of roughly 1 μm (assuming that the detection pinhole corresponds to the excitation volume). We also observed a small positive correlation between detected photon numbers and burst duration (Fig. 3c, Pearson's r = 0.54 and Supplementary Fig. 10). This suggests that larger volumes, in combination with high irradiances,

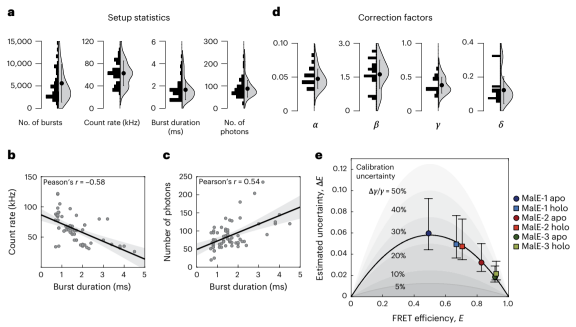


Fig. 3 | **Setup-dependent parameters and calibration uncertainty. a**, The distribution of the parameters quantifying the statistics of the measurements and the performance of the setups used for both MalE and U2AF2 measurements are shown as histograms and violin plots for the measurements from eight laboratories. The circle and whiskers in the violin plot indicate the mean and standard deviation (n = 64, averaged over eight samples measured in the eight different laboratories). Sample-dependent distributions of the shown parameters are given in Supplementary Fig. 9. b, c, Pairwise plots of the average count rate (**b**) and the number of photons (**c**) against the burst duration. The same datasets are plotted as used for **a**. While the count rate decreases slightly for longer burst durations, a positive correlation is observed for the acquired number of photons per burst and the burst duration, indicating that larger observation volumes result in a higher accumulated signal per molecule. Correlations between all parameters are shown in Supplementary Fig. 10.

Error bands indicate the 95% confidence intervals of the regression. **d**, The distributions of the four correction factors for the calculation of accurate FRET efficiencies for all the MalE measurements are shown as histograms and violin plots for the measurements from all laboratories. The circle and whiskers in the violin plot indicate the mean and standard deviation (n = 64, averaged over eight samples measured in the eight different laboratories). **e**, A plot of the standard deviation of the reported FRET efficiencies from 16 laboratories (as a measure of the experimental uncertainty) against the average FRET efficiency for the MalE mutants 1–3 reveals that lower uncertainties are observed for higher FRET efficiencies. The black line represents a fit of the estimated uncertainties under the assumption that the variations arise solely due to uncertainty in the γ factor (equation (1)). The inferred relative uncertainty of the γ factor is around 23%. Shaded areas indicate relative uncertainties of 5–50%. Error bars indicate 95% confidence intervals around the average value.

yield the highest number of photons per burst³⁸. Smaller volumes generally allow for higher burst collection rates with higher count rates and thus shorter interphoton times, enabling fast transitions on the sub-µs timescale to be resolved^{39,40}. Longer burst durations offer the benefit that slower dynamics can be studied.

For an accurate analysis, the correction factors for donor spectral crosstalk (α), excitation flux (β), detection efficiency and quantum yields (y) and direct acceptor excitation (δ) must be determined (see ref. 18, Supplementary Table 5). We plot the distribution of the correction factors used to determine accurate FRET efficiencies for MalE in Fig. 3d from 16 laboratories (Supplementary Table 1). Besides fluorophore properties, these also depend on setup-specific parameters including dichroic mirrors, emission filters, detectors, excitation wavelengths and laser power. Nonetheless, we observed a well-defined distribution for α of 0.05 \pm 0.01, which is determined by the emission filters and detectors in both detection channels. A larger spread was observed for β values of 1.6 \pm 0.6 and δ of 0.12 \pm 0.08. These depend on the ratio of the excitation powers, where most participants used about half the laser power for direct acceptor excitation (45 \pm 27 μ W) in comparison to the donor excitation (78 \pm 58 μ W), resulting in similar count rates after donor and acceptor excitation. The agreement between the reported FRET efficiency values clearly shows that the diverse experimental settings are compensated by the correction procedure applied here.

For γ , which is the most difficult factor to determine, we observed an average of 0.4 \pm 0.1 (Supplementary Fig. 3b). It depends on the

acceptor-to-donor ratio of the detection efficiencies, g, and the effective fluorescence quantum yields, ϕ_F , as $\gamma = g_A \phi_{F,A}/g_D \phi_{F,D}$ (ref. 18). Similar to crosstalk, y strongly depends on the emission filters and the type of detectors used. Due to $\phi_{\rm F,A}$ of roughly 0.32 (acceptor) and $\phi_{\rm ED}$ of roughly 0.72 (donor), all laboratories reported y factors below 1. Despite the large spread in the reported values, we observed very good agreement for the reported FRET efficiencies in our blind study. Our analysis identified y as the key factor limiting the consistency, which is supported by the following arguments: (1) in Fig. 1d, the spread of $\langle E_{\text{holo}} \rangle - \langle E_{\text{apo}} \rangle$ is smaller (for example, 0.06 to 0.02 for MalE-1) than for absolute E values in Fig. 1c, suggesting that errors in E are systematic rather than random. (2) The observed spread in reported FRET efficiencies depends on the absolute FRET efficiency measured for MalE (Fig. 1c,d). (3) We also calculated the uncertainty in the FRET efficiency calculation using error propagation for crosstalk, direct excitation and background correction in the donor and acceptor channels. The reported uncertainty can be attributed mainly to the y factor (Fig. 3e and Supplementary Note 4) with the error of the y factor, $\Delta \gamma$, that propagates into an uncertainty in the reported FRET efficiencies, ΔE :

$$\Delta E = E(1 - E) \frac{\Delta \gamma}{\gamma} \tag{1}$$

Notably, the observed experimental ΔE is well described by equation (1) (black line in Fig. 3e), yielding a relative uncertainty of

 $\Delta\gamma/\gamma=23\%$ corresponding to $\Delta\gamma\cong0.07$. The improved agreement between measurements on reanalysis for U2AF2 (Fig. 2d) suggests that the accuracy of the analysis could be improved by standardized procedures for the determination of all correction factors, which differ depending on the number of populations in the measurement and whether the FRET efficiency peak is dynamically averaged (Supplementary Note 2).

Detection and quantification of conformational dynamics

Fluorescence trajectories of immobilized molecules provide access to kinetics on the millisecond to second timescales via a dwell-time analysis (Supplementary Fig. 1)41-43. For freely diffusing molecules, millisecond dynamics can be studied in the same fashion when molecules diffuse slowly^{44,45}. The detection and quantification of submillisecond conformational dynamics in quickly diffusing molecules (with the maximum timescale limited by the burst duration) is possible via FRET-FCS^{44,46,47}, filtered-FCS^{48,49}, burst-variance analysis (BVA)⁵⁰, FRETtwo-channel kernel-based density distribution estimator⁵¹, dynamic PDA⁵², FRET efficiency *E* versus fluorescence-weighted average donor lifetime $\langle \tau_{D(A)} \rangle_{E}$ analysis $(E-\tau \text{ plots})^{52,53}$, nanosecond-FCS⁵⁴, recurrence analysis of single particles⁵⁵, photon-by-photon maximum likelihood approaches^{40,56-59} and Monte Carlo diffusion-enhanced photon inference (MC-DEPI)⁶⁰. To assess how consistently dynamics can be detected in smFRET measurements, we asked the participants to evaluate whether the proteins were static or dynamic on the (sub-) millisecond timescale and which method they used to come to this conclusion (Supplementary Table 6).

BVA and E-τ plots are frequently used techniques to visualize FRET dynamics by comparing the measured data to theoretical expectations. BVA detects dynamics by estimating the standard deviation of the FRET efficiency over individual bursts, using a predefined photon window (typically ≥100 µs depending on the molecular brightness). Due to FRET dynamics, the standard deviation of the FRET signal within a burst (red line in Fig. 4a) can be higher than expected from shot noise (black semicircle in Fig. 4a), which becomes visible as a deviation or apparent dynamic shift, ds⁵⁰. In the E- τ plots, the observed FRET efficiency determined via intensity (Fig. 4b) is a species-weighted average and, in the presence of dynamics, the position along the yaxis depends on the fraction of time spent in the respective states. The fluorescence lifetime of the donor (Fig. 4b, $\langle \tau_{D(A)} \rangle_F$, x axis) is a photon-weighted average, because only a single lifetime is determined. Hence, it is weighted toward the lifetime of low-FRET states as they emit more donor photons^{52,53}, shifting the data to the right of the 'static' FRET line. E-τ plots can detect dynamics on the nanosecond to millisecond timescale. Here, we have included an additional correction that considers distance fluctuations of the flexible dye linkers (6 Å) resulting in a slightly curved 'static' FRET line 52,61. To quantify dynamics between two distinct states, a theoretical 'dynamic' FRET line (red, Fig. 4b) is overlaid. Again, ds is defined as the deviation of the observed data from the theoretical static line (Fig. 4b and Supplementary Note 6). It is important to mention that FRET dynamics, and the related ds, are not always of conformational origin.

MalE exhibits slow ligand-driven dynamics on the subsecond timescale between high- and low-FRET states (Supplementary Fig. 1) 62 . Here, we investigated whether the apo and/or holo states undergo dynamics faster than the timescale of diffusion. Both techniques reveal that MalE exhibits no large FRET-fluctuations on the ms timescale (Fig. 4c,d and Supplementary Fig. 11). Almost all groups confirmed this assessment and only three groups concluded that MalE is dynamic without further justification (Supplementary Table 6). To investigate the presence of potential dynamics in more detail, we determined the ds for a subset of the data (eight laboratories for BVA, Fig. 4e and five for E– τ , Fig. 4f, Supplementary Note 5, and Supplementary Table 7). As a static control, we determined the ds of the dsDNA rulers used in ref. 18 (mean \pm 1 s.d. as determined from

laboratories 1 and 2) shown in gray in Fig. 4e, f (Supplementary Table 8). The ds did not exceed that of the static reference when using BVA for all MalE mutants. From the E- τ plots, however, ds was higher than for dsDNA, especially for MalE-1. This sample clearly exceeds what is predicted for a static system or even what is predicted for dynamics between the apo and holo states (Fig. 4f, red lines and Supplementary Note 6). Hence, some laboratories categorized MalE as dynamic. The cause of this ds, which must originate from FRET dynamics that are faster than around 100 μ s, will be discussed in detail below.

In contrast to MalE, all groups found U2AF2 to be dynamic as was expected for two domains connected by a flexible linker (Fig. 4c–f and Supplementary Table 6). The ligand-free apo state shows pronounced deviations from the behavior for static molecules both in the BVA and $E-\tau$ plots, while the RNA-bound holo state shows a notable ds for BVA but not for the $E-\tau$ analysis (Fig. 4c–f). It was challenging to assess whether the holo state is truly static or dynamic since it contained a measurable fraction of apo protein, which overlaps with the holo population. Hence, U2AF2 is a challenging test case, yet, dynamics were unambiguously detected in all laboratories demonstrating the reliability of smFRET for investigating dynamic systems.

Accuracy of FRET-derived distances and structural modeling

Accurate FRET efficiencies need to be converted into distances for comparison with structures or to use them as distance constraints in $integrative \ FRET-assisted \ structural \ modeling \ ^{1,5,7,15,63,64}. \ SmFRET \ experimental \ constant \ experimental \ constant \ experimental \ constant \ experimental \ experim$ ments yield FRET efficiencies as a result of dynamically, nonlinearly averaged distances due to the flexible fluorophore linkers. To assess the accuracy of our measurements, we applied the accessible volumes (AV) approach^{5,6,6,4,65}, which uses a coarse-grained dye model to estimate the FRET efficiency averaged model distance $R_{(F)}^{\text{model}}$ between the two dyes. For this, all possible positions of the fluorophores are averaged, taking into account linker conformations and steric hindrances (Fig. 5a-c, Methods and ref. 6). For AV calculations, we assume fast rotational and slow positional averaging with respect to the fluorescence lifetime. Prediction of measured distances via FRET values based on the flexibility and attachment points of a fluorophore is an area of active research and alternative methods are being developed, for example, rotamer libraries⁶⁶ or molecular dynamics simulations^{67,68}.

The average experimental FRET efficiencies from the individual smFRET histograms $\langle E \rangle$ for MalE (Fig. 2) were used to determine $R_{\langle E \rangle}$ for each laboratory (Extended Data Table 1 and Supplementary Table 3) using the Förster equation (equation (2)):

$$R_{\langle E \rangle} = R_0 \left(\frac{1}{\langle E \rangle} - 1 \right)^{\frac{1}{6}} \tag{2}$$

The Förster radius of Alexa546–Alexa647 on MalE was determined to be $R_0 = 65 \pm 3$ Å (Supplementary Note 7). Figure 5d displays the correlation between the experimental observable $R_{(E)}$ and predicted $R_{(E)}^{\rm model}$ using apo and holo structures exhibiting an uncertainty of 3–5 Å over all variants. In agreement with the predictions by Peulen et al.⁶⁹, this accuracy is achieved despite stochastic protein labeling, which could result in different charge environments and AVs of the fluorophores depending on the labeling positions. This is evident by the varying dye behavior at different locations (Fig. 5b). During the study, three laboratories studied additional MalE variants (MalE-4, K34C-N205C and MalE-5, T36C-N205C) with a larger FRET efficiency contrast between the apo and holo states, complementing the results of the other variants (Extended Data Table 1).

Figure 5d reveals the largest deviation between experimental and predicted distances for MalE-1, which also had the highest ds values (Fig. 4f and Supplementary Fig. 11). Therefore, we investigated the role of dye-protein interactions using single-cysteine variants of MalE by measuring the fluorescence lifetimes, and time-resolved and steady-state anisotropies (Supplementary Note 8, Supplementary

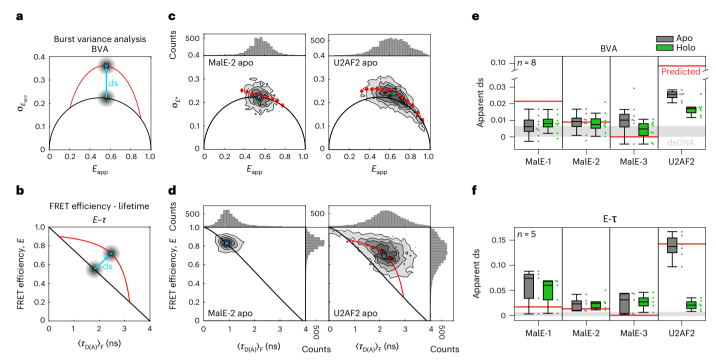


Fig. 4 | **Detection and characterization of conformational dynamics on the submillisecond timescale in MalE and U2AF2.** a,b, Schematic representations of BVA (a) and $E^-\tau$ (b) plots. The ds is defined as the excess standard deviation compared to the static line (shown in black). Dynamic FRET lines are indicated in red. c, BVA of MalE-2 labeled with Alexa546–Alexa647 without maltose (apo, left) and U2AF2 labeled with Atto532–Atto643 without RNA (apo, right). Here, the BVA is based on a photon binning of five photons. Red diamonds indicate the average standard deviation of all bursts within a FRET efficiency range of 0.05. The mean positions of the populations (cyan crosses) were determined by fitting a two-dimensional Gaussian distribution to the data (Supplementary Note 5). **d**, The plots of the FRET efficiency E versus intensity-weighted average donor lifetime $\langle \tau_{D(A)} \rangle_F$ of the same measurement as in **c**. The donor-only population was excluded from the plot. For MalE-2, the population falls on the static FRET line, while a clear ds is observed for U2AF2. The endpoints of the dynamic FRET line for

U2AF2 were determined from a subensemble analysis of the fluorescence decay. **e**, **f**, The apparent ds of the peak of the population was determined graphically from BVA (eight laboratories for MalE and seven laboratories for U2AF2, respectively) (**e**) and E- τ (five laboratories) (**f**) plots (Methods). For U2AF2 in the holo state, the ds was assessed only for the low-FRET RNA-bound population. Boxes indicate the median and 25/75% quartiles of the data. Whiskers extend to the lowest or highest data point within 1.5 times the interquartile range. The gray area indicates the ds obtained for the dsDNA used in a previous study ¹⁸ based on measurements performed in laboratory 1 for BVA (ds_{DNA} = 0.0033 ± 0.0033) and laboratory 2 for the E- τ plot (ds_{DNA} = 0.0026 ± 0.0044). The horizontal red lines indicate the expected ds for a potential conformational exchange between the apo and holo states. We computed the expected change in FRET efficiency using their structural models in the PDB (Supplementary Note 6 and Supplementary Table 9).

Tables 5, 11 and 12 and Fig. 5b). Labeling at residue 352 promotes dye sticking to the protein surface indicated by multiexponential fluorescence lifetimes and a high residual anisotropy, r_{∞} , for both fluorophores ($r_{\infty} > 0.25$). Labeling at residue 29 only shows sticking for the donor ($r_{\infty,D} > 0.30$, $r_{\infty,A}$ roughly 0.12). At other positions (for example, residue 186), free rotation is possible for both dyes (Supplementary Tables 5 and 11). These position-specific interactions can cause the observed deviations between the experiment and structural model (Fig. 5d and Extended Data Table 1) and high ds values for MalE-1 (Fig. 4f). By using the accessible contact volume (ACV) approach⁶³, which accounts for dye–protein interactions, the root-mean-average deviation between the structural model and experimental values decreased from 3 Å for AV to 2 Å (Fig. 5c). For protein labeling on opposite sides, dye–protein interactions in the ACV model result in reduced model distances and improved accuracy for all outliers (Fig. 5d and Extended Data Table 1).

It was suggested to use the combined residual anisotropy of D and A $(r_{c,\infty} = \sqrt{r_{\infty,D}} \ r_{\infty,A})$ for filtering out dye-related artifacts in FRET-assisted structural modeling with an empirical threshold of $r_{c,\infty} < 0.2$ (refs. 13,70). To further investigate dye-specific sticking, three laboratories studied MalE mutants with the additional dye pairs Alexa546–AbbSTAR635P, Atto532–Atto643 and Alexa Fluor 488 (Alexa488)–Alexa647 and determined the residual anisotropies and distance uncertainties based on the orientation factor κ^2 (Fig. 5e, top, Supplementary Tables 13 and 14 and Supplementary Notes 8 and 9).

The dve pair Alexa546-Alexa647 showed the highest combined anisotropies (Supplementary Fig. 12a and Supplementary Table 13), which is attributed to the donor Alexa546 as the combined anisotropy also remains high for Alexa546-AbbSTAR635P but is reduced for Alexa488-Alexa647. To derive a robust and well-defined threshold for recognizing measurements with dye artifacts, we determined the uncertainty in the FRET-derived distances, $\Delta R_{\rm ann}(\kappa^2)$, that originates from the uncertainty of the orientation factor κ^2 . Previous approaches estimated the uncertainty in κ^2 from the residual anisotropy in terms of rotational restrictions (wobbling-in-a-cone model)⁷⁰⁻⁷³. Here, we used a 'diffusion with traps' model, which assumes two dye populations (free and trapped) and relates the residual anisotropies to the fraction of dyes interacting with the surface of the biomolecule (Supplementary Note 9). Based on the estimated distance uncertainty, we propose a threshold of $\Delta R_{app}(\kappa^2)$ < 10% to identify measurements with dye-related artifacts (Fig. 5e, bottom). This threshold corresponds to a combined residual anisotropy of 0.25, similar to the previously suggested empirical threshold value of around 0.2 (refs. 13,70).

Next, we investigated whether dye sticking could cause the ds in the E- τ plot for MalE-1 with Alexa546–Alexa647 (Fig. 4f). To be observable in the E- τ plot, the exchange between the free and trapped dye species must occur faster than the diffusion time of roughly 1 ms, otherwise the two species would be observable as individual peaks. We observed a correlation between the laboratory-averaged $\langle ds \rangle$ and $\langle r_{c,\omega} \rangle$

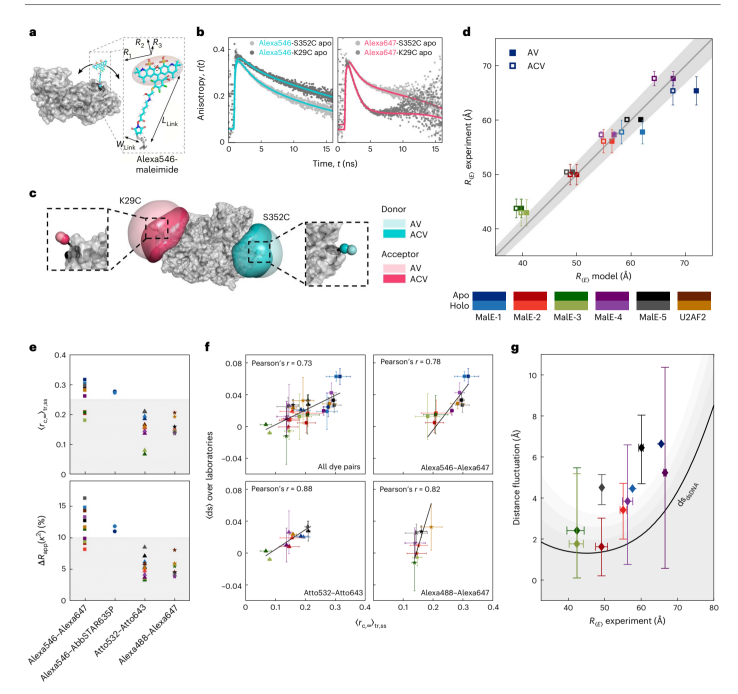


Fig. 5 | Assessing the accuracy of smFRET-derived distances in MalE. a-d, AV calculations and model-based interdye distances. a, Schematic of Alexa546 attached to MalE (PDB 10MP) showing the parameters needed for the AV calculations using the AV3 model⁶ (Supplementary Table 10). **b**, Fluorescence anisotropy decays of single-cysteine mutants for the donor (Alexa546, left) and acceptor (Alexa647, right) at the labeling positions K29C and S352C. Solid lines represent fits to a model with two or three rotational components (Supplementary Tables 11 and 12 and Supplementary Note 8). c, AV (light color) and ACV (dark color) calculations for Alexa546 (cyan) and Alexa647 (pink) at labeling positions 352 and 29. The zoom-ins show the mean positions of the dyes based on the AV (light shade) and ACV (darker shade) models. d, Comparison of the experimentally obtained FRET-averaged distance $R_{\langle E \rangle}$ with the theoretical model distances using the AV (filled squares) and ACV (empty squares) calculations. Errors represent the standard deviation in experimental distances (n = 16 laboratories for MalE mutants 1 - 3, n = 2 laboratories for MalE mutants 4 - 5,n = 7 laboratories for U2AF). The solid line represents a 1:1 relation and the gray area indicates an uncertainty of ± 3 Å for a Förster radius of $R_0 = 65$ Å. MalE-4 and -5

were measured by two laboratories. e, Detection of dye-specific protein interactions. Top shows the five MalE mutants and U2AF2 labeled with different dye combinations to determine the donor-acceptor-combined residual anisotropy, $\langle r_{c,\infty} \rangle_{\rm tr,ss}$ (n=3 laboratories). Bottom shows the distance uncertainty relating to κ^2 , $\Delta R_{\rm app}$ (κ^2), estimated (Supplementary Note 8). A maximum allowed distance uncertainty of $\leq 10\%$ (shaded gray region) in $\Delta R_{\rm app}$ (κ^2) leads to a dye-independent threshold of 0.25 for $\langle r_{c,\infty} \rangle$. **f**, The apparent dynamic shift $\langle ds \rangle$ versus the combined residual anisotropy $\langle r_{\rm c, \circ} \rangle$ is shown for all measured dye pairs (top left) and individually. Error bars of the apparent ds represent the standard deviation over n = 3 laboratories. For the combined residual anisotropy, the propagated 1σ uncertainty (Supplementary Note 8). **g**, The structural flexibility of MalE estimated after filtering using the distance uncertainty threshold shown in **e** (Supplementary Note 12). Error bars represent the 1σ percentiles averaged over all dye pairs (n = 1, MalE-1; n = 7, MalE-2 and MalE-3; n = 4, MalE-4 and n = 5, $MalE-5). \ The \ residual \ distance \ fluctuations \ obtained \ from \ control \ measurements$ on dsDNA in one laboratory (ds $_{\text{dsDNA}}$ = 0.0026 \pm 0.0044) are shown as a black line (gray areas represent confidence intervals of 1σ , 2σ and 3σ).

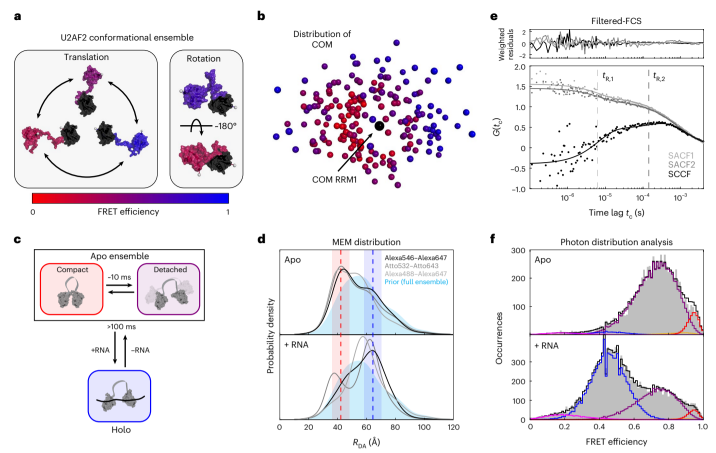


Fig. 6 | **Structural characterization of U2AF2.** a, Structural flexibility of U2AF2 is given by translational (left) and rotational (right) movement of the two domains. Representative structures are taken from the ensemble determined using NMR and SAXS measurements²⁹. **b**, Degeneracy of structural states in FRET measurements. The position of the two domains of U2AF2 is illustrated by the COM of the C_{α} atoms in RRM2 (residues 260–329, colored) with respect to RRM1 (residues 150–227, black) for the 200 structures of the conformational ensemble²⁹. The COM of RRM2 is color-coded according to the FRET efficiency determined using AV3 calculations. **c**, A schematic of the kinetic model used for the global dynamic PDA of U2AF2 (Supplementary Note 17). **d**, Distance distributions obtained from a donor fluorescence decay analysis by a model-free MEM approach (Supplementary Note 15). The distance distribution from the NMR–SAXS ensemble²⁹ (light blue) was used as the prior distribution. The expected interdye distances for the compact apo and open holo states are shown as red and blue dashed lines (PDB 2YH0 and 2YH1). Shaded areas indicate the

distance broadening due to the flexible dye linkers of 6 Å. The distribution in the donor–acceptor distance $R_{\rm DA}$ for different dye pairs is shown. **e**, Filtered-FCS reveals conformational dynamics in the U2AF2 apo ensemble on two timescales, $t_{R,1} = 9 \pm 3$ and $t_{R,2} = 300 \pm 90$ µs, average and standard deviation (n = 3, results from laboratory 1 are shown). The two species were defined at the lower and upper edge of the FRET efficiency histogram shown in Fig. 2b, top panel (see Methods and Supplementary Note 16 for details). The species autocorrelation functions (SACFs) and one of the two species cross-correlation functions (SCCFs) are shown. The weighted residuals are shown above. **f**, The PDA analysis was conducted globally over both apo (top) and holo (bottom) measurements using time windows of 0.5, 1.0, 1.5 and 2.0 ms (the 1.0 ms time window histograms are shown). A relaxation time of roughly 10 ms for the dynamics between the detached ensemble and compact apo state with a small amplitude was determined (orange curve) (Supplementary Fig. 16 and Supplementary Note 17).

over all dye pairs (Pearson's r = 0.73), with a stronger correlation when each dye pair is investigated individually (Fig. 5f). As conformational dynamics should be label independent, dye sticking is likely responsible for the observed ds values. The x intercept of the linear fit is between 0.1 and 0.2, suggesting a dye-dependent anisotropy threshold needs to be considered. When applying the criteria $\langle r_{\rm c, \infty} \rangle <$ 0.25 to MalE-1 (Supplementary Fig. 12b), only the dye pair Atto532–Atto643 should be used for distance determination, which also showed a markedly reduced ds (Supplementary Fig. 12c). Lifetime analysis of MalE-1 donor-only molecules showed donor quenching only at position 352, which confirms that labeling at this position is problematic (Supplementary Fig. 12c, Supplementary Note 10 and Supplementary Table 5).

Using the above criteria of $\langle r_{\rm c,\infty} \rangle < 0.25$ to minimize the influence of dye artifacts on the ds, we hypothesized that the remaining ds could be indicative of low-amplitude, fast conformational fluctuations. A P test analysis between the ds for dsDNA and protein samples (P < 0.05) indicated that the ds is still significant for various protein

variants after filtering out dye artifacts (Supplementary Note 11, Supplementary Table 8). To estimate the conformational fluctuations necessary to generate the observed ds (Fig. 4f and Supplementary Table 8), we assume that dynamics occur between two nearby states with interdye distances of $R_{(E)} \pm \delta R$ where δR is the amplitude of the fluctuation⁶¹ (Fig. 5g, Supplementary Note 12 and Supplementary Table 8). This inferred fluctuation provides an upper bound for the conformational flexibility because factors such as calibration errors. dye blinking or photoisomerization could contribute to the observed ds. We consider the ds obtained from dsDNA as the lower limit (black line in Fig. 5g, ds_{DNA} = 0.0026 \pm 0.0044: Supplementary Note 12), which defines the current detection limit for dynamics in smFRET experiments. The MalE variants 1, 4 and 5 exceed the ds for dsDNA by 2-3 Å (Fig. 5g, Supplementary Fig. 13 and Supplementary Table 8). Consistent with the smFRET results, all-atom molecular dynamics simulations of MalE using the ff14SB force field⁷⁴ (Supplementary Note 13) suggest thermally induced conformational fluctuations with a standard

deviation up to roughly 3 Å at the labeled residues in MalE-1, MalE-4 and MalE-5. This is larger than the typical fluctuations of about 1 Å (ref. 75) and leads to a broadening of the interresidue distance distributions for these FRET pairs. We conclude that the observed ds in the experiments can be explained by a combination of measurement uncertainty and small-scale structural fluctuations. Note that such small-scale fluctuations can be amplified in FRET experiments when the dye linker acts as a lever arm for appropriate labeling positions. A detailed discussion of the theoretical limits for detecting dynamics in smFRET experiments using BVA or the $E-\tau$ is given in Supplementary Note 14.

Quantitative analysis of U2AF2

The structural characterization of U2AF2 is more complex than for MalE and a simple distance comparison is not possible. Nonetheless, we asked what information smFRET measurements could provide for such a dynamic system. We first surveyed the structural information available on apo U2AF2 from nuclear magnetic resonance (NMR) and small-angle X-ray scattering (SAXS) data²⁹. The highly flexible linker allows for a heterogeneous ensemble of U2AF2 conformations (Fig. 6a). To assess how this translates into a smFRET distribution, we quantified the FRET efficiency using AV calculations for all 200 conformers from the NMR–SAXS-derived ensemble of apo U2AF2 (ref. 29). Notably, conformations with similar center-of-mass (COM) distances between the domains showed different FRET efficiencies (Fig. 6a,b), because domain rotations result in distinct interdye distances for identical COM (Fig. 6a, right). Due to this degeneracy, a single-distance probe is insufficient to capture the full structural complexity.

The observed ds in the apo state suggests the presence of conformational dynamics (Fig. 4d–f). To decipher the underlying kinetics and their temporal hierarchy, we applied three analyses. First, we investigated the interdye distance distribution of the apo and holo states from the donor lifetime using a model-free maximum entropy method (MEM) (Fig. 6c,d and Supplementary Note 15) 76 . As a prior, we used the NMR–SAXS structural ensemble. This analysis yielded consistent results for all three dye pairs studied for U2AF2. The MEM analysis revealed peaks in the probability density at the expected distances for the compact apo conformation and RNA-bound holo structure (Fig. 6d, dashed lines). We note that the fluorescence lifetime analysis resolves states on the nanosecond timescale and is therefore less sensitive to dynamic averaging.

Second, to assess the dynamics on the microsecond timescale, three groups performed filtered-FCS and found at least two relaxation times (9 \pm 3 and 300 \pm 90 μs ; Fig. 6e, Supplementary Table 15 and Supplementary Note 16), which were independent of the dyes used (Supplementary Fig. 15). We assign the fast process to dynamics within the detached domains and the slower process to interconversion between compact conformations within the conformational ensemble.

Last, we investigated dynamics on the millisecond timescale using a dynamic PDA. A global analysis of the apo and holo measurements was performed using the kinetic model shown in Fig. 6c (Supplementary Note 17 and Supplementary Table 16). The apo state was treated as a two-state system with slow dynamics between a detached ensemble and a well-defined, compact apo conformation. The rapid dynamics within the detached ensemble is empirically described using a broad, static distribution. For the holo measurement, we account for the residual population of apo molecules. Exchange between the holo and apo states is irrelevant as the binding and dissociation of RNA occurs on timescales of more than 100 ms (ref. 36). This model incorporates all information and is sufficient to describe the smFRET efficiency histograms. The dynamic PDA analysis returned a relaxation time of roughly 10 ms for the dynamics between the detached ensemble and compact apo state (Fig. 6f, orange curve, Supplementary Fig. 16 and Supplementary Table 16). We also determined an interdye distance of $R_{(F)} = 61 \,\text{Å}$ in the RNA-bound holo state, which is in good agreement with 63 Å from the RNA-bound conformation (Protein Data Bank (PDB) 2YH1).

Discussion

We show that smFRET can provide accurate distances of conformational states and reliable information on conformational dynamics in proteins. Since all experiments were performed using established techniques and analyzed with freely available software $^{5,6,34,35,77-79}$, such information is accessible to any group with similar expertise. Despite the challenges of protein samples, we achieved a similar precision in FRET efficiencies as reported for dsDNA 18 (between ± 0.02 and ± 0.06) (Extended Data Table 1). The reproducibility in excluding large-scale conformational dynamics for MalE on a timescale <10 ms while detecting large-scale submillisecond dynamics in U2AF2 shows that the community can deal with dynamic protein systems. In addition, we could consistently establish the timescales and hierarchy of the exchange dynamics in such a complex protein system as U2AF2. The study of complex dynamics is improvable by probing additional distances $^{5,13,17,80-83}$.

The high level of agreement is notable given the diversity of the setups (Fig. 3 and Supplementary Fig. 2) and the number of possible pitfalls. A large contribution to the spread in the reported mean FRET efficiencies was caused by systematic errors in the data analysis. This is supported by a comparison of the FRET efficiency changes $(\langle E_{\text{holo}} \rangle - \langle E_{\text{apo}} \rangle)$ instead of absolute FRET efficiency values (Fig. 1d), which reduced the spread of roughly threefold. Having a single person reanalyze the data led to a similar decrease in the uncertainty of the FRET efficiency for the apo state of U2AF2 (Fig. 2d). Determination of y was most crucial and the optimal approach depends on the details of the studied system (Supplementary Note 2). In the intensity-based approach of Lee et al.34, multiple samples with uniform fluorophore properties are required or individual corrections need to be made. When using the approach of Kudryavtsev et al. 35 via $E-\tau$ calibration, the system needs to be static and a single population suffices. A protocol with unambiguous instructions for the calibration steps and minimized number of user-dependent steps would enhance the accuracy of FRET measurements.

From accurate FRET efficiencies, we obtained reproducible interdye distances with a precision of 3 Å and an accuracy of 5 Å against structural models of MalE (Extended Data Table 1). This is similar to what was determined for dsDNA samples. This is a very positive outcome, given that dsDNA features a consistent, homogenous chemical environment for each labeling position, in contrast to the variable dye environment experienced in proteins. The distance determination could be improved by including the interaction of the fluorophores with the protein surface using ACV calculations (Fig. 5d and Methods) 63 . Furthermore, we give experimental support (Fig. 5f) for only using dyes with a combined residual anisotropy of $r_{c,\infty} < 0.25$, as suggested previously 13,70 . Proteins often exist within a family of conformations as we observed for U2AF2 (Fig. 6d). Determining how to best deal with distance distributions for conformational ensembles is one of the challenges for structural biology.

Investigating different dye pairs allowed us to reduce dye artifacts, leading to more accurate FRET efficiencies and reliable detection of the dynamics. Hence, we investigated the detection limits for ds and studied its relation to conformational dynamics with a subset of laboratories. Besides conformational motions, dynamic FRET shifts can occur in different directions and have several origins including structural instabilities³⁷ or photophysics (as shown in Fig. 5f)⁴⁴. Thus, it is advisable to verify the key findings in smFRET measurements with at least two dye pairs and/or with different residue combinations in the protein. Once the non-FRET-dynamic contributions are minimized, we still observed significant residual ds for MalE. Consistent with molecular dynamics simulations (Supplementary Note 13), we interpret these shifts as small-scale conformational dynamics and established a current lower limit for the detection of structural changes via smFRET on the order of ≤5 Å. In summary, the consensus of smFRET experiments on two protein systems exhibiting dynamic behavior on different spatiotemporal scales obtained blindly from 19 laboratories offers strong

support for its use as a robust, versatile and quantitative tool for the coming age of dynamic structural biology. In this context, it will be crucial to integrate the correlated structural and dynamic information provided by smFRET¹ with structural information provided by other experimental techniques as well as artificial intelligence-based protein structural prediction8³. Considering that protein structure prediction has reached the single-structure frontier8⁴, the information from smFRET experiments could leverage the power of artificial intelligence to resolve more complex multi-state and ensemble structural models8³. Vice versa, the power of artificial intelligence and deep learning can be used to increase the throughput for the design and analysis of smFRET experiments8⁵-87.

Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41592-023-01807-0.

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Methods

Sample preparation of proteins

Double-cysteine mutants of MalE were prepared and labeled using established protocols⁶². Human RRM1,2 L187C-G326C mutant (U2AF2-148-342) was obtained and purified as described in Mackereth et al.³⁰.

Fluorescence labeling of proteins

All fluorophores were purchased as maleimide derivatives from commercial suppliers as listed in Supplementary Table 19. MalE was stochastically labeled as described previously⁸⁸ with fluorophores as indicated in the text with a combined labeling efficiency higher than 70% resulting in a donor–acceptor pairing of at least 20%. Protein stability and functionality (ligand binding) was verified by affinity measurements using microscale thermophoresis⁸⁹. All preparations, that is, MalE-wildtype, unlabeled cysteine mutants and fluorophore-labeled variants, showed an affinity for maltose between roughly 1 and 2 μ M (Supplementary Fig. 5) consistent with previously published $K_{\rm d}$ values for wildtype MalE^{90,91}. The stability and labeling of the sample were verified by FCS (Supplementary Fig. 18), which excluded the presence of larger aggregates in the samples and confirms that MalE is functional.

U2AF2 was stochastically labeled as described previously in Voith von Voithenberg et al. 36 . The combined labeling efficiencies for the labeling reactions were 20 and 14% for the Alexa546–Alexa647 and Atto532–Atto643 pairs, respectively. For Alexa488–Alexa647, the combined labeling efficiency was found to be 10%. The functionality of the labeled U2AF protein was checked with affinity measurements for U9 RNA, which was found to be $1.2~\mu\text{M}$ (ref. 30), consistent with the previous reports 36 (Supplementary Fig. 7d).

Sample handling

Both protein systems required special handling due to sample instability or aggregate formation, which are both problematic for long-term storage and shipping. The labeled MalE proteins were stored in 50 mM Tris-HCl pH 7.4, 50 mM KCl with 1 mg ml⁻¹ BSA at 4 °C for less than 7 d. U2AF2 was stored in 20 mM potassium phosphate buffer pH 6.5, 50 mM NaCl and kept in the fridge until used. Both samples were loaded in low-binding Eppendorf tubes (Eppendorf Germany, catalog no. 0030108094) and shipped on ice in a cooling box with overnight shipping to avoid unnecessary freezing and thawing. MalE stock solutions were on the order of 10 to 100 nM concentration and the sent stock solution of U2AF2 was 5-10 uM concentration. Dilution buffers for apo and holo measurements were provided. SmFRET experiments were carried out by diluting the labeled proteins to concentrations of roughly 50 pM in 50 mM Tris-HCl pH 7.4, 50 mM KCl supplemented with the ligand maltose at 1 mM concentration. Labeled U2AF2 protein was measured at roughly 40–100 pM in 20 mM potassium phosphate buffer pH 6.5, 50 mM NaCl. Purchased U9 RNA (Biomers.net GmbH and IBA Solutions for Life Sciences) was dissolved in RNA-free water and added directly to the solution at a final concentration of 5 µM for the holo measurements. Both proteins were studied on coverslips typically passivated with 1 mg ml⁻¹BSA in buffer before adding the sample. The measurements were performed without any photostabilizer to keep the measurements as simple as possible to avoid any further source for discrepancies between the groups, for example, degradation of photostabilizer or use of different photostabilizer concentrations.

SmFRET data acquisition and analysis

Data acquisition and correction procedures were performed for confocal measurements as described by Hellenkamp et al.¹⁸. The samples were measured using ALEX or PIE on a confocal microscope as sketched in Supplementary Fig. 2. A description of the experimental procedures of all laboratories is given in Supplementary Note 18.

Briefly, the three recorded intensity time traces for each single-molecule event are:

donor emission after donor excitation: ⁱI_{Dem|Dex},

acceptor emission after donor excitation (FRET signal): ⁱI_{AemiDex},

and acceptor emission after acceptor excitation : ${}^{i}I_{Aem|Dex}$.

The apparent (raw) FRET efficiency is computed as:

$$E_{\rm app} = \frac{{}^{\rm i}I_{\rm Aem|Dex}}{{}^{\rm i}I_{\rm Dem|Dex} + {}^{\rm i}I_{\rm Aem|Dex}},\tag{3}$$

Recorded intensities were corrected for background contributions as:

$${}^{ii}I_{\text{Dem}|\text{Dex}} = {}^{i}I_{\text{Dem}|\text{Dex}} - {}^{i}I_{\text{Dem}|\text{Dex}}^{(\text{BG})}, \tag{4}$$

$$^{ii}I_{\text{Aem}|\text{Dex}} = {}^{i}I_{\text{Aem}|\text{Dex}} - I_{\text{Aem}|\text{Dex}}^{(\text{BG})}, \tag{5}$$

$$^{ii}I_{\text{Aem}|\text{Aex}} = {}^{i}I_{\text{Aem}|\text{Aex}} - I_{\text{Aem}|\text{Aex}}^{(\text{BG})}, \tag{6}$$

where $I_{\text{Dem}|\text{Dex}}^{(\text{BG})}$ $I_{\text{Aem}|\text{Dex}}^{(\text{BG})}$ and $I_{\text{Aem}|\text{Aex}}^{(\text{BG})}$ are the respective background signals. Correction factors for spectral crosstalk, α and direct excitation, δ , were determined from the donor- and acceptor-only populations³⁴. The corrected acceptor fluorescence after donor excitation, $F_{\text{A}|\text{D}}$, is computed as:

$$F_{\text{A|D}} = {}^{\text{ii}}I_{\text{Aem|Dex}} - \alpha {}^{\text{ii}}I_{\text{Dem|Dex}} - \delta {}^{\text{ii}}I_{\text{Aem|Aex}}$$
 (7)

The γ and β factors, correcting for differences in the detection yield and excitation fluxes of the donor and acceptor dyes, were estimated using a global correction procedure following the approach of Lee et al. (Supplementary Fig. 3)³⁴. Alternatively, when pulsed excitation was used and the sample is known to be static, the γ factor can be determined by fitting the measured population to the static FRET line^{35,92}. This allows a robust determination of the γ factor when only a single species is present but requires a static sample and the appropriate static FRET line (Supplementary Note 2).

The accurate FRET efficiency \boldsymbol{E} and stoichiometry \boldsymbol{S} values were then calculated as:

$$E = \frac{F_{A|D}}{\gamma^{ii}I_{Dem|Dex} + F_{A|D}},$$
(8)

$$S = \frac{\gamma^{ii}I_{\text{Dem}|\text{Dex}} + F_{\text{A}|\text{D}}}{\gamma^{ii}I_{\text{Dem}|\text{Dex}} + F_{\text{A}|\text{D}} + ii}I_{\text{Aem}|\text{Aex}}/\beta}.$$
 (9)

Conversion of accurate FRET efficiencies into distances were done using equation (2) with Förster radii determined as described in Supplementary Note 7.

Detection of protein dynamics

In this work, we used the following two approaches to detect conformational dynamics:

BVA. In BVA, the presence of dynamics is determined by looking for excess variance in the FRET efficiency data beyond the shot-noise limit. The standard deviation ($\sigma_{E_{app}}$) of the apparent FRET efficiency (E_{app}) is calculated using a fixed photon window of n=5 over the time period of the individual bursts given by:

$$\sigma_{E_{app}} = \sqrt{\frac{E_{app}(1 - E_{app})}{n}},\tag{10}$$

The shot-noise limited standard deviation of the apparent FRET efficiency is generally described by a semicircle 50 (Fig. 4a and Supplementary Fig. 11a–d). In the presence of dynamics, the standard deviation for the FRET efficiency within a burst becomes higher than that expected from shot noise. Photophysical effects such as photobleaching and blinking also give rise to the higher standard deviation beyond the shot-noise limit. Typically, BVA is sensitive to fluctuations in the FRET signal of ${\gtrsim}100~\mu\text{s}$, but this depends on the brightness of the burst and the photon window used.

FRET efficiency versus fluorescence-weighted average donor lifetime analysis (E– τ plots). Two-dimensional histograms of the FRET efficiency E and donor fluorescence lifetime $\langle \tau_{D(A)} \rangle_F$ (Fig. 4b and Supplementary Fig. 11e–h) were created for single-molecule measurements using multiparameter fluorescence detection (MFD) in combination with PIE³⁵, described below. Static FRET lines were calculated using the following equation:

$$E = 1 - \frac{\tau_{\rm D(A)}}{\tau_{\rm D(0)}} \tag{11}$$

and further modified for linker dynamics⁶¹. Deviations of FRET populations from the static FRET line can indicate FRET dynamics, which can be due to conformational fluctuations or photophysical dynamics. In addition, a time-resolved FRET analysis of TCSPC data can accurately resolve the distance heterogeneities by revealing multiple components in the decay curve and recovering their specific species fractions and FRET rate constants⁶⁹. Dynamics are thus detected from the presence of multiple components in the subensemble decay of a single FRET population. In addition, dynamics that are slower than the fluorescence lifetime (roughly 5 ns) are not averaged in the FRET lifetime analysis leading to the detection of the full conformational distribution.

MFD with PIE

MFD, introduced by Eggeling et al. 93 , combines spectral and polarized detection with picosecond pulsed lasers and TCSPC, allowing the simultaneous detection of intensity, lifetime, anisotropy and spectral range of the fluorescence signal of single molecules. nsALEX or PIE additionally provides the acceptor lifetime information 35 . Due to the availability of the lifetime information when using pulsed excitation, this approach is well suited for using E- τ -based analyses.

Dye simulations (AV and ACV)

The AV approach uses a simple coarse-grained dye model⁶⁴ defined by five parameters: the width and length of the linker, and three radii that define the fluorophore volume (Fig. 5a and Supplementary Table 10). Using these parameters, AV simulations for both fluorophores were calculated by considering the linker flexibility and steric hindrances of the labeled molecule (Fig. 5a). In the ACV model⁶³, the position of the dyes is biased toward the protein surface, resulting in a reduction of the interdye distance for the given labeling positions. To do this, the residual anisotropy was used to estimate the fraction of sticking dyes. In the computation of the FRET-averaged model distances, the occupancy of a thin surface layer (roughly 3 Å) was then increased such that its fraction matches the amount of interacting dye detected in the experiment (Fig. 5b and Supplementary Table 10).

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

The data for all figures, all supplementary figures, the raw data for MalE measurements from all laboratories (with the exception of one mutant from one laboratory) and the raw data for all U2AF2 measurements have been uploaded to Zenodo (https://doi.org/10.5281/zenodo.7472900). PDB IDs used are 1OMP, 1ANF, 2YHO and 2YH1. Source data are provided with this paper.

Code availability

The software used for data analysis are available from the respective laboratories: laboratory no. 1, PAM (PIE Analysis with MATLAB) software package (ref. 79 in the main text); laboratory no. 2, Home-written LabView-based software (ref. 15 in Supplementary Information); Laboratory no. 3. FRETBursts toolkit (ref. 14 in Supplementary Information): laboratory no. 4, PAM (PIE Analysis with MATLAB) software package; laboratory no. 5, PAM (PIE Analysis with MATLAB) software package; laboratory no. 6, PAM (PIE Analysis with MATLAB) software package; laboratory no. 7, PAM (PIE Analysis with MATLAB) software package; laboratory no. 8, PAM (PIE Analysis with MATLAB) software package v.2.0; laboratory no. 9, PAM (PIE Analysis with MATLAB) software package; laboratory no. 11, data were analyzed with the burst analysis toolbox (BAT, V2018 and V2019) and filtered and visualized with T3ee (V2018, V2019) (ref. 84 in Supplementary Information); laboratory no. 12: PAM (PIE Analysis with MATLAB) software package; laboratory no. 13, IgorPro 8 (Wavemetrics); laboratory no. 14, PAM (PIE Analysis with MATLAB) software package; laboratory no. 15, Software Package for Multiparameter Fluorescence Spectroscopy, Full Correlation and Multiparameter Fluorescence Imaging developed in C.A.M. Seidel's laboratory (http://www.mpc.uni-duesseldorf.de/seidel/); laboratory no. 16, ALEX-suite software package (ref. 88 of Supplementary Information); laboratory no. 17, FRETBursts analysis software (ref. 14 of Supplementary Information) and laboratory no. 18, FRETBursts toolkit (ref. 14 of Supplementary Information).

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Author contributions

T.C. initiated the study and D.C.L. coordinated the study. G.A., C.G., A.B., C.A.M.S., D.C.L. and T.C. designed research. A.B., C.A.M.S., D.C.L. and T.C. supervised the project. R.M. cloned and purified MalE variants. H.-S.K. and M.S. provided U2AF2. C.G. and G.A. performed labeling of MalE and U2AF2 variants, respectively, for shipment to participating laboratories. M.P., J.F. and C.A.M.S. designed MalE mutants 4 and

5 in silico. M.P. and J.F. performed initial measurements on MalE mutants 4 and 5. G.A. reperformed the analysis on the provided raw data for U2AF2 and MalE-1 from eight laboratories. G.A., A.B. and M.P. performed ds estimation. C.G. performed FCS experiments on MalE variants, time-resolved anisotropy experiments and R_0 -determination. M.P. performed time-resolved anisotropy analysis of single labeled MalE cysteine mutants from ensemble measurements as well of all MalE and U2AF2 dye combinations from smFRET measurements. C.G., G.A. and M.P. performed measurements of MalE mutants with additional dye combinations and M.P. and A.B. performed statistical analysis of ds and anisotropies. G.A. and A.B. performed estimation of setup-dependent parameters and PDA of U2AF2. G.A. performed the filtered-FCS and A.B. performed TCSPC analysis of U2AF2. C.G., G.A. and M.P. performed smFRET measurements on dsDNA rulers. M.P. performed AV and ACV modeling of dye distributions for MalE and U2AF2. G.G.M.M. performed microscale thermophoresis experiments. M.d.B. performed confocal scanning experiments for surface-immobilized MalE. All authors were involved in performing comparison experiments and analyzing smFRET data. G.A. and C.G. consolidated data collection of participating laboratories. G.A. and C.G. designed Fig. 1. G.A., C.G. and M.P. designed Fig. 2. A.B. designed Figs. 3 and 4. A.B. and M.P. designed Figs. 5 and 6. G.A., C.G., M.P., A.B., C.A.M.S., D.C.L. and T.C. interpreted data and wrote the manuscript in consultation with all authors.

Competing interests

T.D.C. and A.N.K. are founders of different companies selling single-molecule fluorescence microscopes (Exciting Instruments, Oxford Nanoimager). The other authors declare no competing interests.

Additional information

Extended data is available for this paper at https://doi.org/10.1038/s41592-023-01807-0.

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41592-023-01807-0.

Correspondence and requests for materials should be addressed to Anders Barth. Claus A. M. Seidel. Don C. Lamb or Thorben Cordes.

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Extended Data Table. 1 | Average of mean FRET efficiency and standard deviation for MalE and U2AF2 samples reported by the participating laboratories

		F	Experimental Valu	Modeled Distances		
Sample Condition		$\mu_{\langle E angle} \pm \sigma_{\langle E angle}$	$egin{aligned} oldsymbol{\mu}_{\langle E_{ m holo} angle - \langle E_{ m apo} angle} \ &\pm oldsymbol{\sigma}_{\langle E_{ m holo} angle - \langle E_{ m apo} angle} \end{aligned}$	$egin{aligned} oldsymbol{\mu_{R_{\langle E angle}}} & \pm \ oldsymbol{\sigma_{R_{\langle E angle}}} \left[ext{Å} ight] \end{aligned}$	$R_{\langle E \rangle}^{ m AV}$ [Å]	$R_{\langle E \rangle}^{ m ACV} [m \AA]$
MalE-1	apo	0.49 ± 0.06	0.177±0.019	65.4±2.6	72.0	67.7
Maie-i	holo	0.67 ± 0.05	0.17/±0.019	57.8 ± 2.1	62.1	58.3
MalE-2	apo	0.83 ± 0.03	-0.121±0.019	50.0±1.9	50.1	48.8
Maie-2	holo	0.71±0.05	-0.121±0.019	56.1±2.1	56.5	55.0
MalE-3	apo	0.913±0.019	0.007±0.010	43.8±1.7	39.9	38.9
Maie-3	holo	0.920 ± 0.021	0.007±0.010	43.0±2.4	40.8	39.8
MalE-4*	apo	0.442±0.025		67.6 ± 1.2	67.8	64.3
IVIaIE-4	holo	0.678 ± 0.017		57.4±0.7	56.9	54.6
MalE-5*	apo	0.613 ± 0.003		60.22 ± 0.15	61.8	59.3
	holo	0.821±0.001		50.43 ± 0.08	49.3	48.2
U2AF2**	apo	0.74 ± 0.03		49.6±1.3		
	holo***	0.46 ± 0.04		60.8±1.7		

The calculated average $\mu_{(E)}$ and standard deviation $\sigma_{(E)}$ of the mean FRET efficiency values provided by the participating labs are given for all three studied mutants of MalE labeled with Alexa546 and Alexa647 under both apo and holo conditions (see Supplementary Table 3). The calculated mean and standard deviation of the difference in the reported mean FRET efficiency between the apo and holo $(\langle E_{\text{holo}}\rangle - \langle E_{\text{apo}}\rangle)$ for the three MalE mutants are given by $\mu_{(E_{\text{holo}})-(E_{\text{apo}})}$ and $\sigma_{(E_{\text{holo}})-(E_{\text{apo}})}$ respectively (see Supplementary Table 3). The calculated average $\mu_{R_{(E)}}$ and standard deviation $\sigma_{R_{(E)}}$ of the mean distances were derived according to Eq. 2. The modeled distances $R_{(E)}^{\text{ACV}}$ are derived using accessible volume (ACV) and accessible contact volume (ACV) calculations respectively, as described in the Methods. We also give the average and standard deviation for the FRET values determined for U2AF2 labeled with Atto532-Atto643 under both apo and holo conditions (Supplementary Table 4). *Only studied by two laboratories. **Due to the fast-structural dynamics in the sample, only 7 labs studied this mutant and distances were not determined. *** Only the holo state under holo condition was considered.

nature research

Corresponding author(s):	Barth, Seidel, Lamb, Cordes
Last updated by author(s):	Dec 21, 2022

Reporting Summary

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For	all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.
n/a	Confirmed
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	A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
	The statistical test(s) used AND whether they are one- or two-sided Only common tests should be described solely by name; describe more complex techniques in the Methods section.
	A description of all covariates tested
\boxtimes	A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
	A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

Lab#1: SPCM software (Becker & Hickl GmbH) and HydrHarp400 (PicoQuant)

Lab#2: HydrHarp400 (PicoQuant)

Lab#3: National Instruments-Card PCI-6602 (National Instruments, USA)

Lab#4: SymPhoTime64, (PicoQuant) Lab#5: HydraHarp 400 (PicoQuant)

Lab#6: HydraHarp 400 and Symphotime 32 software (PicoQuant).

Lab#7: National Instruments card PCle-6353 with acquisition controlled using custom software (see ref. 64 of Supplementary Information)

Lab#8: HydraHarp 400 (PicoQuant)

Lab#9: Igor-Program (Wavemetrics) (see ref. 64 of Supplementary Information)

Lab#10: LabVIEW software (see ref. 63-71 of Supplementary Information)

Lab#11: HydraHarp 400 (PicoQuant)

Lab#12: SymPhoTime64 software package (PicoQuant)

Lab#13: TimeHarp200 (PicoQuant)

Lab#14: SPCM software (Becker & Hickl GmbH)

Lab#16: National Instruments-Card PCI-6602 (National Instruments, USA).

Lab#17: VistaVision software (version 4.2.095, 64-bit, ISSTM, USA) (see ref. 82 of Supplementary Information)

Lab#18: TimeHarp 200 (PicoQuant)

Data analysis

Lab#1: PAM (PIE Analysis with Matlab) software package (see ref. 79 in the main text)

Lab#2: Home-written LabView-based software (see ref. 15 of Supplementary Information)

Lab#3: FRETBursts toolkit (see ref. 14 of Supplementary Information)

Lab#4: PAM (PIE Analysis with Matlab) software package Lab#5: PAM (PIE Analysis with Matlab) software package

Lab#6: PAM (PIE Analysis with Matlab) software package

Lab#7: PAM (PIE Analysis with Matlab) software package Lab#8: PAM (PIE Analysis with Matlab) software package v2.0 Lab#9: PAM (PIE Analysis with Matlab) software package

Lab#11: Burst Analysis Toolbox (BAT) V2018 and V2019, and T3ee V2018 and V2019 (see ref. 84 of Supplementary Information)

Lab#12: PAM (PIE Analysis with Matlab) software package Lab#13: IgorPro 8 (Wavemetrics, Portland OR, USA). Lab#14: PAM (PIE Analysis with Matlab) software package

 $Lab \#15: Software\ Package\ for\ Multiparameter\ Fluorescence\ Spectroscopy,\ Full\ Correlation\ and\ Multiparameter\ Fluorescence\ Imaging\ Spectroscopy,\ Full\ Correlation\ and\ Multiparameter\ Fluorescence\ Imaging\ Spectroscopy\ Full\ Correlation\ and\ Multiparameter\ Fluorescence\ Imaging\ Spectroscopy\ Full\ Correlation\ Application\ Fluorescence\ Imaging\ Spectroscopy\ Full\ Correlation\ Fluorescence\ Imaging\ Spectroscopy\ Full\ Fluorescence\ Imaging\ Spectroscopy\ Full\ Fluorescence\ Fluorescence\ Imaging\ Fluorescence\ Fluorescenc$

developed in C.A.M. Seidel's lab (http://www.mpc.uni-duesseldorf.de/seidel/). Lab#16: ALEX-suite software package (see ref. 88 of Supplementary Information) Lab#17: FRET Bursts analysis software (see ref. 14 of Supplementary Information)

Lab#18: FRETBursts toolkit (see ref. 14 of Supplementary Information)

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- A list of figures that have associated raw data
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The data for all figures, all supplementary figures, the raw data for all MalE measurements (with the exception of one mutant from one laboratory) and the raw data for all U2AF2 measurements have been uploaded to Zenodo (https://doi.org/10.5281/zenodo.7472900). PDB-ID of protein structures: 10MP, 1ANF, 2YH0, 2YH1

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Life sciences study design

Study description

Research sample

Sampling strategy

the selections study design				
All studies must dis	sclose on these points even when the disclosure is negative.			
Sample size	19 Laboratories. These are the groups that responded to the invitation to participate in the study.			
Data exclusions	Data from three labs were excluded due to excessive photobleaching during the measurement, inappropriate experimental setups for the provided fluorophores, or difficulties with data collection (see notes on Table S1 for more details)			
Replication	Replication was performed by measuring samples in the different participating laboratories. Replicates were examined for consistency.			
Randomization	Randomization is not applicable because we tested the consistency of the results between all participating labs for the same sample .			
Blinding	This was a blind study where the individual groups (with the exception of those providing the proteins) were not informed regarding what protein sample they were measuring. Groups were informed regarding the appropriate conditions for the different samples.			

Behavioural & social sciences study design

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i studies must disclose on these points even when the disclosure is negative.

quantitative experimental, mixed-methods case study).

State the research sample (e.g. Harvard university undergraduates, villagers in rural India) and provide relevant demographic information (e.g. age, sex) and indicate whether the sample is representative. Provide a rationale for the study sample chosen. For studies involving existing datasets, placed describe the dataset and source.

Briefly describe the study type including whether data are quantitative, qualitative, or mixed-methods (e.g., qualitative cross-sectional,

studies involving existing datasets, please describe the dataset and source.

| Constitution | C

Describe the sampling procedure (e.g. random, snowball, stratified, convenience). Describe the statistical methods that were used to predetermine sample size OR if no sample-size calculation was performed, describe how sample sizes were chosen and provide a rationale for why these sample sizes are sufficient. For qualitative data, please indicate whether data saturation was considered, and what criteria were used to decide that no further sampling was needed.

Data collection Provide details about the data collection procedure, including the instruments or devices used to record the data (e.g. pen and paper,

2

Data collection	computer, eye tracker, video or audio equipment) whether anyone was present besides the participant(s) and the researcher, and whether the researcher was blind to experimental condition and/or the study hypothesis during data collection.
Timing	Indicate the start and stop dates of data collection. If there is a gap between collection periods, state the dates for each sample cohort.
Data exclusions	If no data were excluded from the analyses, state so OR if data were excluded, provide the exact number of exclusions and the rationale behind them, indicating whether exclusion criteria were pre-established.
Non-participation	State how many participants dropped out/declined participation and the reason(s) given OR provide response rate OR state that no participants dropped out/declined participation.

If participants were not allocated into experimental groups, state so OR describe how participants were allocated to groups, and if

Ecological, evolutionary & environmental sciences study design

allocation was not random, describe how covariates were controlled.

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Randomization

Reproducibility

Randomization

Blinding

Study description

Briefly describe the study. For quantitative data include treatment factors and interactions, design structure (e.g. factorial, nested, hierarchical), nature and number of experimental units and replicates.

Research sample

Describe the research sample (e.g. a group of tagged Passer domesticus, all Stenocereus thurberi within Organ Pipe Cactus National Monument), and provide a rationale for the sample choice. When relevant, describe the organism taxa, source, sex, age range and any manipulations. State what population the sample is meant to represent when applicable. For studies involving existing datasets, describe the data and its source.

Sampling strategy

Note the sampling procedure. Describe the statistical methods that were used to predetermine sample size OR if no sample-size calculation was performed, describe how sample sizes were chosen and provide a rationale for why these sample sizes are sufficient.

Data collection Describe the data collection procedure, including who recorded the data and how.

Timing and spatial scale Indicate the start and stop dates of data collection, noting the frequency and periodicity of sampling and providing a rationale for these choices. If there is a gap between collection periods, state the dates for each sample cohort. Specify the spatial scale from which the data are taken

Data exclusions If no data were excluded from the analyses, state so OR if data were excluded, describe the exclusions and the rationale behind them, indicating whether exclusion criteria were pre-established.

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Describe the extent of blinding used during data acquisition and analysis. If blinding was not possible, describe why OR explain why blinding was not relevant to your study.

Did the study involve field work? Yes No

Field work, collection and transport

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Materials & experime	ntal s	ystems Methods		
n/a Involved in the study		n/a Involved in the study		
Antibodies		ChIP-seq		
Eukaryotic cell lines		Flow cytometry		
Palaeontology and a				
Animals and other o				
Human research par	rticipant	S		
Clinical data Dual use research or	f concer	n		
Dual use research o	redricer			
Antibodies				
Antibodies used	Describ	be all antibodies used in the study; as applicable, provide supplier name, catalog number, clone name, and lot number.		
Validation		be the validation of each primary antibody for the species and application, noting any validation statements on the acturer's website, relevant citations, antibody profiles in online databases, or data provided in the manuscript.		
Eukaryotic cell lin	۵۲			
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Mycoplasma contaminati	on	firm that all cell lines tested negative for mycoplasma contamination OR describe the results of the testing for oplasma contamination OR declare that the cell lines were not tested for mycoplasma contamination.		
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Palaeontology an	d Arc	chaeology		
Specimen provenance		Provide provenance information for specimens and describe permits that were obtained for the work (including the name of the issuing authority, the date of issue, and any identifying information).		
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Ethics oversight Identify the organization(s) to was required and explain wh		withe organization(s) that approved or provided guidance on the study protocol, OR state that no ethical approval or guidance quired and explain why not.		
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Animals and othe	r org	anisms		
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Laboratory animals For laboratory animals, repo		oratory animals, report species, strain, sex and age OR state that the study did not involve laboratory animals.		
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Ethics oversight

Human research participants

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Population characteristics

Describe the covariate-relevant population characteristics of the human research participants (e.g. age, gender, genotypic information, past and current diagnosis and treatment categories). If you filled out the behavioural & social sciences study design questions and have nothing to add here, write "See above."

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ChIP-seq

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Confirm that you have deposited or provided access to graph files (e.g. BED files) for the called peaks.

Data access links For "Initial submission" or "Revised version" documents, provide reviewer access links. For your "Final submission" document, provide a link to the deposited data. May remain private before publication.

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Describe the software used to collect and analyze the ChIP-seq data. For custom code that has been deposited into a community

Methodology

Replicates Describe the experimental replicates, specifying number, type and replicate agreement. Describe the sequencing depth for each experiment, providing the total number of reads, uniquely mapped reads, length of reads and Sequencing depth whether they were paired- or single-end. Antibodies Describe the antibodies used for the ChIP-seq experiments; as applicable, provide supplier name, catalog number, clone name, and lot Peak calling parameters Specify the command line program and parameters used for read mapping and peak calling, including the ChIP, control and index files Data quality Describe the methods used to ensure data quality in full detail, including how many peaks are at FDR 5% and above 5-fold enrichment.

Flow Cytometry

Plots

Software

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Describe the sample preparation, detailing the biological source of the cells and any tissue processing steps used. Sample preparation Identify the instrument used for data collection, specifying make and model number. Instrument Describe the software used to collect and analyze the flow cytometry data. For custom code that has been deposited into a Software community repository, provide accession details. Describe the abundance of the relevant cell populations within post-sort fractions, providing details on the purity of the Cell population abundance samples and how it was determined Describe the gating strategy used for all relevant experiments, specifying the preliminary FSC/SSC gates of the starting cell Gating strategy population, indicating where boundaries between "positive" and "negative" staining cell populations are defined.

Tick this box to confirm that a figure exemplifying the gating strategy is provided in the Supplementary Information.

Magnetic resonance imaging

Experimental design

Indicate task or resting state; event-related or block design. Design type

repository, provide accession details.

Design specifications Specify the number of blocks, trials or experimental units per session and/or subject, and specify the length of each trial

or block (if trials are blocked) and interval between trials

Behavioral performance measures State number and/or type of variables recorded (e.g. correct button press, response time) and what statistics were used

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Statistical modeling & infere	nce				
Model type and settings	Specify type (mass univariate, multivariate, RSA, predictive, etc.) and describe essential details of the model at the first and second levels (e.g. fixed, random or mixed effects; drift or auto-correlation).				
Effect(s) tested	Define precise effect in terms of the task or stimulus conditions instead of psychological concepts and indicate whether ANOVA or factorial designs were used.				
Specify type of analysis: W	hole brain ROI-based Both				
Statistic type for inference (See Eklund et al. 2016)	Specify voxel-wise or cluster-wise and report all relevant parameters for cluster-wise methods.				
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Functional and/or effective conn	Report the measures of dependence used and the model details (e.g. Pearson correlation, partial correlation, mutual information).				
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