

Dissecting Mechanisms of Ligand Binding and Conformational Changes in the Glutamine-Binding Protein

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Abstract

Ligand binding and conformational changes of biomacromolecules play a central role in the regulation of cellular processes. It is important to understand how both are coupled and what their role is in biological function. The biochemical properties, conformational states, and structural dynamics of periplasmic substrate-binding proteins (abbreviated SBPs or PBPs), which are associated with a wide range of membrane proteins, have been extensively studied over the past decades. Their ligand-binding mechanism, i.e., the temporal order of ligand-protein interactions and conformational changes, however, remains a subject of controversial discussion. We here present a biochemical and biophysical analysis of the *E. coli* glutamine-binding protein GlnBP concerning ligand binding and its coupling to conformational changes. For this, we used a combination of experimental techniques including isothermal titration calorimetry, single-molecule Förster resonance energy transfer, and surface-plasmon resonance spectroscopy. We found that both apo- and holo-GlnBP show no detectable exchange between open and (semi-)closed conformations on timescales between 100 ns and 10 ms. Furthermore, we also demonstrate that ligand binding and conformational changes in GlnBP are highly correlated. A global analysis of our results is consistent with a dominant induced-fit mechanism, where the ligand binds GlnBP prior to conformational rearrangements. Importantly, we suggest that the rigorous experimental and theoretical framework used here can be applied to other protein systems where the coupling mechanism of conformational changes and ligand binding is yet unclear or where doubts prevail.

eLife assessment

This **important** study combines a range of biophysical techniques to carry out a series of **compelling** experiments to explore whether glutamine binding protein binds glutamine via an induced fit or a conformational selection process. The evidence supporting the major conclusion of the work is **convincing**, although it may not be generalized to other protein-ligand or protein-protein systems. The work will be of broad interest to biochemists and biophysicists.

Introduction

Periplasmic substrate-binding proteins (SBPs)^{1,2,3,4,5,6} are small, soluble proteins (molecular weight <100 kDa) that are often associated with membrane complexes, including the superfamily of ATP-binding cassette (ABC) transporters⁷. SBPs recognize and bind numerous classes of substrates including (but not limited to) ions, vitamins, co-factors, sugars, peptides, amino acids, system effectors, and virulence factors^{8,9}. Major biological functions of SBPs are to facilitate membrane transport by delivery of substrate molecules to a transmembrane component or to signal the presence of a ligand^{8,9}. They are ubiquitous in archaea, prokaryotes, and eukaryotes and possess a highly-conserved three-dimensional architecture with two rigid domains, D1/D2, that are linked by a flexible hinge composed of β -sheets (**Figure 1A**), α -helices, or smaller sub-domains^{8,9}. The available crystal structures of SBPs reveal that many exist in two conformations, a ligand-free open (apo) and a ligand-bound closed state (holo; **Figure 1B**).

Several recent studies focused on the characterization of structural dynamics and conformational heterogeneity as well as the ligand-binding mechanisms that underlie SBP function^{13,14,21}. Based on crystal structures, it was proposed that SBPs use an induced fit, IF ligand binding mechanism (**Figure 1C**)^{8,9}, where the event of ligand binding triggers a functionally-relevant conformational change. This intuitive model was challenged by nuclear magnetic resonance (NMR) based paramagnetic relaxation enhancement (PRE) experiments²², molecular dynamics (MD) simulations^{23,24} and X-ray crystallography^{25,26,27} revealing the existence of unliganded closed or semi-closed states and their dynamic exchange with the respective open (apo) conformation. Similar findings on ligand-independent conformational changes were presented for the maltose-binding protein, MalE^{23,24,28}, histidine-binding protein (HisJ)²⁹, D-glucose/D-galactose-binding protein (GGBP)^{25,27,30,31}, ferric-binding protein (FBP)³², choline/acetylcholine substrate-binding protein (ChoX)³³, the Lysine-, Arginine-, Ornithine-binding (LAO) protein³⁴ and glutamine-binding protein (GlnBP)^{18,35,36}. For MalE, NMR techniques revealed a low abundance (<10%) of a semi-closed state that is in rapid dynamic equilibrium with the open apo conformation on the <50 μ s timescale. The existence of closed, unliganded state(s) allow for an alternative mechanism via conformational selection (CS)³⁷, where conformational changes occur intrinsically prior to ligand binding, and, upon ligand binding, the closed holo state is stabilized (**Figure 1C**).

Ligand-binding mechanisms have also been the focus of several studies of GlnBP^{18,19,38}. GlnBP is part of an ABC transporter system in *E. coli* and binds *L*-glutamine with sub-micromolar affinity^{39,40} and arginine with millimolar affinity⁴¹. It is monomeric and comprised of two globular domains: the large domain (residues 5 – 84, 186 – 224) and the small domain (residues 90 – 180), linked via a flexible hinge (residues 85 – 89, 181 and 189). GlnBP was crystallized in two distinct conformational states: open (apo, ligand-free)¹¹ and closed (holo, ligand-bound)^{12,39} (**Figure 1B**). Recent studies of GlnBP^{18,19,38} used a combination of single-molecule Förster resonance energy transfer (smFRET) measurements, NMR residual dipolar coupling

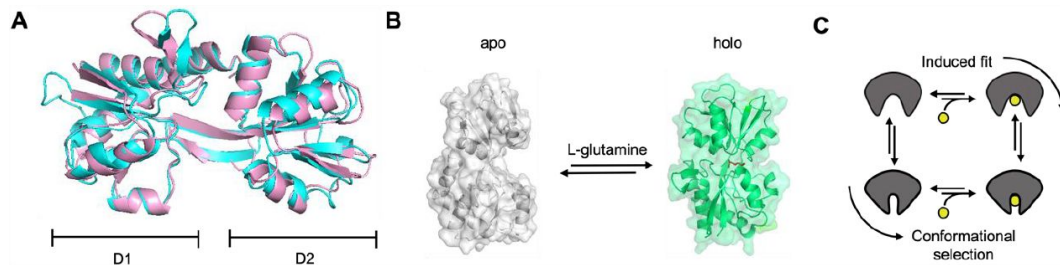


Figure 1.

Conformational states and possible ligand binding mechanisms of typical SBPs.

(A) Structural comparison of SBD2 from *Lactococcus Lactis* (PDB file:4KR5¹⁰; cyan) and glutamine-binding protein GlnBP from *E. coli* (pink). SBD2 and GlnBP share 34% sequence identity with a TM-score of 0.90, indicating high structural similarity. (B) Crystal structures of the ligand-free (PDB file:1GGG¹¹; grey) and ligand-bound (PDB file:1WDN¹²; green) state of GlnBP from *E. coli*. (C) Sketch of ligand binding via induced-fit (IF) and conformational selection (CS).

(RDC)¹⁸ experiments, MD simulations^{35, 36} and Markov state models (MSMs)³⁸. Based on the results, it was proposed that GlnBP undergoes pronounced conformational changes both in the absence¹⁸ and in the presence¹⁹ of substrate, involving a total of four to six conformational states. These findings and the interpretation that ligand binding in GlnBP might occur by means of a combination of CS and IF³⁸ seemed puzzling to us in light of the following arguments: (i) Kooshapur and co-workers demonstrated that NMR experiments on GlnBP do not support the idea of intrinsic conformational dynamics in apo-GlnBP⁴². (ii) The existence of multiple semi-closed conformers of apo- and holo-GlnBP contradict the findings from MD simulations⁴³ and smFRET work studying the ligand binding domains SBD1 and SBD2 (from the amino acid transporter GlnPQ^{20, 44}), which structurally resembles GlnBP (**Figure 1A**; SBD2 shows ~34% sequence identity with GlnBP, TM-score of 0.90). (iii) Ligand binding to the closed state of the GlnBP conformation seems unlikely, considering the limited accessibility of the binding site, which is also seen for related proteins such as MalE⁴⁵.

These controversial findings and arguments reveal a central problem in the study of ligand-binding mechanisms, which is the availability of sufficient experimental evidence to distinguish one mechanism from the other. Importantly, both IF and CS imply temporal order of ligand-protein interactions and conformational changes and thus require kinetic data for univocal identification^{37, 46-50}. The existence of a ligand-free protein conformation, which structurally resembles a ligand-bound form, is necessary^{27, 51, 52} but by itself not sufficient evidence for a CS mechanism, as ligand binding may not proceed via this conformation at all. Vice versa, the inability to experimentally detect ligand-free closed conformations cannot be taken as an indicator for IF as a dominant pathway^{37, 46-50} since only very few techniques are able to detect low abundance (high free energy) conformers and their exchange kinetics with the stable ones. Whether, e.g., a ligand-free closed (or near-closed) conformation⁵³ can be observed depends on the magnitude of its equilibrium probability²⁰ as well as the sensitivity of the techniques used to probe it. Single-molecule fluorescence approaches can provide such information¹⁴⁻²¹, yet often suffer from photon-limited time-resolution and potential labeling artefacts. Also, the analysis of ensemble-averaged relaxation rates from nonequilibrium stopped-flow kinetics or equilibrium NMR experiments can be inconclusive under certain experimental conditions^{50, 54}, e.g., under the pseudo-first-order condition of high ligand concentrations in stopped-flow experiments^{37, 46-50}. Consequently, to validate or rule-out the presence of a certain ligand-binding mechanism such as IF and CS, a set of complementary and consistent structural, thermodynamic, and kinetic data of the protein system is required⁵⁰.

Here, we revisit the question of IF versus CS mechanisms for GlnBP by biochemical and biophysical analyses of ligand binding and its coupling to conformational changes. For this, we used a combination of isothermal titration calorimetry (ITC), smFRET and surface-plasmon resonance (SPR) spectroscopy to derive sufficient evidence that can support the (in)compatibility of the data with of such mechanisms. Using smFRET, we observed that apo- and holo-GlnBP show no detectable exchange of open and (partially-) closed states on timescales from as slow as 10 ms to as rapid as 100 ns, and any observed FRET dynamics could be traced back to photophysical origins rather than to conformational changes. Importantly, in all our smFRET assays, ligand binding and conformational dynamics were highly correlated. The global analysis of all (kinetic) parameters from smFRET and SPR finally allowed us to form an argument that the CS model is incompatible with GlnBP, but all data is fully compatible with the IF mechanism. The rigorous experimental and theoretical framework used here can further be applied to other biomacromolecular systems, where the coupling of conformational changes and ligand-binding is unclear.

Results

Biochemical characterization of GlnBP and ligand binding

For our study of the thermodynamic and kinetic aspects of ligand binding in GlnBP, we produced wild-type protein (GlnBP WT) and two double-cysteine variants for analysis of conformational states via smFRET: GlnBP(111C-192C) with point mutations at V111C and G192C (**Figure S1A** [↗](#)), and GlnBP(59C-130C) with point mutations at T59C and T130C (the latter was adapted from refs.¹⁸ [↗](#), ¹⁹ [↗](#); **Figure S1C** [↗](#)). All protein variants were expressed in *E. coli* and purified using affinity chromatography (see Methods for details). Protein purity was assessed by Coomassie-stained SDS-PAGE analysis (**Figure 2A** [↗](#)). As reported previously, GlnBP co-purifies with bound glutamine⁵⁵ [↗](#), which was removed by unfolding and refolding of the purified protein. We verified the monomeric state and proper folding of the resulting protein using size-exclusion chromatography (SEC, **Figure 2B** [↗](#)) by comparing the elution volume and shape of the monodisperse peak of GlnBP before and after the procedure (**Figure S2** [↗](#)).

To assess the binding affinity of GlnBP WT and the two GlnBP cysteine variants for *L*-glutamine, we performed isothermal titration calorimetry (ITC)⁵⁶ [↗](#). Refolded GlnBP WT showed a K_d for *L*-glutamine of 22 ± 7 nM (**Figure S3A** [↗](#)) and K_d values of 31 ± 3 nM and 35 ± 5 nM for the two cysteine variants (**Figure 2C** [↗](#), **Figure S3B** [↗](#)). These values are in agreement with previously published data⁵⁷ [↗](#). This verifies that the unfolding and refolding process as well as cysteine substitutions did not impact the biochemical properties of unlabeled GlnBP.

Analysis of conformational states of freely-diffusing GlnBP via smFRET

After assessing the thermodynamic properties of GlnBP, we characterized the conformational states and changes associated to ligand binding via smFRET. With smFRET, it is possible to study biomacromolecules in aqueous solution at ambient temperature, and identify conformational changes, heterogeneity, small sub-populations and determine microscopic rates of conformational changes.⁵⁸ [↗](#)–⁶⁰ [↗](#) We performed smFRET experiments on freely-diffusing (**Figure 3** [↗](#)) and surface-immobilized GlnBP using the refolded variants GlnBP(111C-192C) and GlnBP(59C-130C) labeled with two different dye pair combinations, AF555/AF647 and ATTO 532/ATTO 643, to minimize any position- and fluorophore-dependent effects. The smFRET assays were designed such that the inter-dye-distance of the apo state results in a lower FRET efficiency as compared to the holo state of the protein (**Figure S1A/C** [↗](#)).

Solution-based μ sALEX⁵⁹ [↗](#) data of GlnBP(111C-192C) labeled with AF555/AF647 are shown in **Figure 3B** [↗](#) after an all-photon burst search.⁶¹ [↗](#) Both apo and holo states, in the absence and presence of saturation levels of glutamine, respectively, show a clear predominant population of donor-acceptor-labeled protein at S^* -values of ~ 0.5 , with two distinct mean apparent E^* values for the apo (mid FRET, 0.51) and holo (high FRET, 0.68) states (**Figure 3B/C** [↗](#), **Figure S4A** [↗](#)). This can be interpreted as a transition from open (apo) to closed (holo) GlnBP conformations upon the addition of the ligand. Similar results were obtained for the second double-cysteine variant (GlnBP(59C-130C), **Figure 3D** [↗](#), **Figure S5** [↗](#)) and from measurements with a different pair of fluorescent dyes for GlnBP(111C-192C) (ATTO 532/ATTO 643; **Figure S4B** [↗](#)). Notably, further analysis and a comparison of mean accurate FRET efficiencies and the inter-dye distances, show good agreement with simulated inter-dye distances of ± 0.2 nm using the open- and closed-GlnBP crystal structures except for the holo-state of GlnBP(59C-130C), which deviated by ~ 0.8 nm (**Figure S1B** [↗](#)/D).

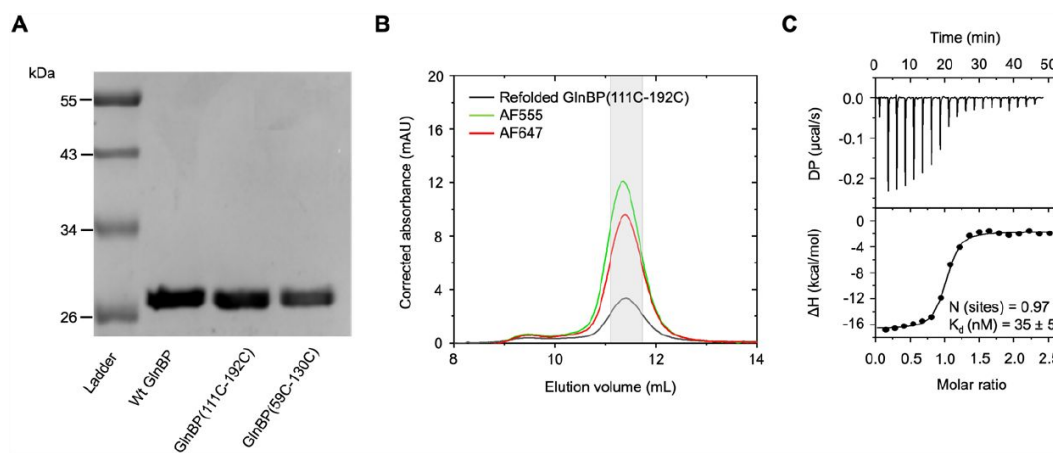


Figure 2.

Biochemical characterization, fluorescence labeling and thermodynamic characterization of ligand binding of GlnBP.

(A) SDS-PAGE analysis of GlnBP purity with coomassie-staining. Lane 1, molecular mass ladder with sizes of proteins indicated in kDa; lane 2, purified GlnBP WT; lane 3, purified double-cysteine variant GlnBP(111C-192C); lane 4, purified double-cysteine variant GlnBP(59C-130C). (B) SEC was used to further purify the fluorescently-labeled proteins. The protein absorption was monitored at 280 nm (black curve), the donor dye absorption (AF555) at 555 nm, and the acceptor dye absorption (AF647) at 647 nm. The labeling efficiency of AF555 and AF647 were estimated to be about 71% and 59%, respectively. For the solution-based smFRET measurements, the used protein fractions are indicated in grey. (C) Ligand-binding affinities of refolded, unlabeled GlnBP(111C-192C) was determined by ITC with a $K_d = 35 \pm 5$ nM for L-glutamine (mean value from $N = 3$ with standard deviation), which is in agreement with previous reports^{57,60,61}. The free energy of binding was $\Delta G = -42.6$ kcal/mol with the enthalpy $\Delta H = -62.3$ kcal/mol and entropy contributions $-T\Delta S = 19.9$ kcal/mol.

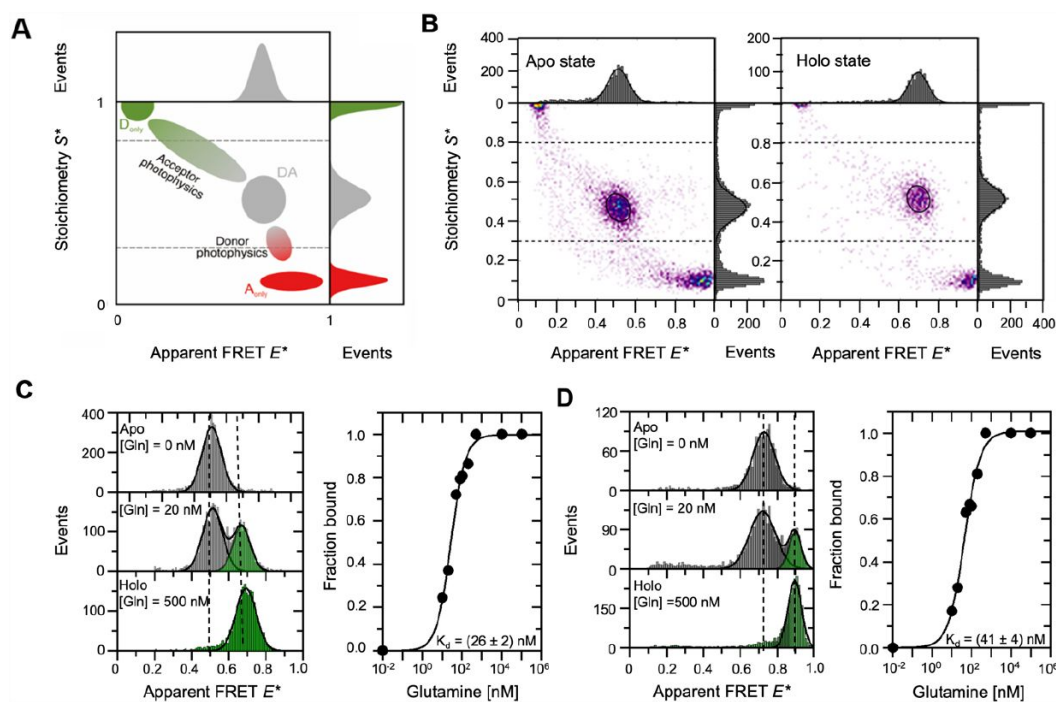


Figure 3.

smFRET analysis of GlnBP using diffusion-based μ sALEX.

(A) Graphical depiction of an E^*-S^* histogram obtained by μ sALEX; panel adapted from ref. 110. Using μ sALEX, the stoichiometry S^* can be used to separate donor-only ($S > 0.8$, D_{only}), acceptor-only ($S < 0.3$, A_{only}), and the FRET molecular species with both donor and acceptor fluorescently-active fluorophore (S^* between 0.3-0.8, DA). Bridge artifacts or smearing caused by donor or acceptor photophysics (photoblinking and/or photobleaching) can cause artificial broadening of the FRET population or a shift of the extracted mean apparent FRET efficiency. (B) μ sALEX-based E^*-S^* histograms of the refolded GlnBP(111C-192C) double-cysteine variant labeled with AF555 and AF647. (C, D) Diffusion-based single-molecule analysis and ligand-binding affinity measurements with μ sALEX of doubly-labeled GlnBP(111C-192C) (C) and GlnBP(59C-130C) variants (D) at different ligand concentrations. Values provided are mean $\pm$ SD ($N = 3$). For plotting purposes the concentration of glutamine in the apo state was set artificially to a value of 0.01 nM in the right parts of panels C/D. Data fitting of the fraction of the high-FRET subpopulation as a function of ligand concentration was performed with the Hill equation, which is a valid approximation for describing the bound fraction of GlnBP as a function of glutamine in the case where $[GlnBP] \ll K_d$.

Importantly, a quantitative analysis of the fraction of the closed state (high-FRET) subpopulation as a function of ligand concentration (**Figure 3C, D**) with the Hill equation ($n = 1$) provides K_d values in the 20-50 nM range for all labeled GlnBP variants, which is fully consistent with ITC results (**Figure 2C**, **Figure S3/S6**). Interestingly, we found that arginine, the non-cognate ligand of GlnBP, induces hardly any FRET shifts at mM concentration of ligand (**Figure S7**) despite its binding to GlnBP at these concentrations (**Figure S8**).

We were also unable to identify a clear high-FRET subpopulation in the absence of a ligand, which would indicate slow intrinsic exchange of apo/open GlnBP with a (partially) closed conformation on timescales slower than the burst duration, >10 ms (**Figure 3B**, apo). To estimate the upper bound of the fraction of a potential low abundance state that was present but poorly sampled due to statistics, we compared the number of bursts within $\langle E^* \rangle \pm \sigma$ between the characteristic regions of the identified high and low FRET subpopulations from representative data sets of GlnBP(111C-192C) in the absence of a ligand. For this, the FRET populations were fitted with Gaussian functions (with mean values and σ), which serves as a good approximation for mean E^* values that are not close to 0 or 1. For representative data sets of AF dyes, we found a ratio of ~12% ($E^*_{\text{holo}} = 0.64$, $\sigma = 0.061$, $N = 626$; $E^*_{\text{apo}} = 0.47$, $\sigma = 0.070$, $N = 5,013$) and for ATTO dyes, a ratio of ~4% ($E^*_{\text{holo}} = 0.56$, $\sigma = 0.056$, $N = 124$; $E^*_{\text{apo}} = 0.37$, $\sigma = 0.047$, $N = 2,908$). This suggests an upper bound of 5-10% for the subpopulation of (partially) closed conformations exchanging with apo GlnBP. Thus, our results agree with the idea that GlnBP mainly exists in a predominant state – the open conformation – in the absence of the cognate ligand, glutamine. The agreement of the ligand concentration dependence in ITC and smFRET strongly suggests that ligand binding and conformational change (into the closed state) are correlated.

Screening for rapid conformational dynamics via analysis of “within-burst” FRET dynamics

Next, we analyzed our smFRET data for “within-burst” dynamics using burst-variance analysis (BVA)⁶², multi-parameter photon-by-photon hidden Markov modeling (mpH²MM)⁶³, intensity-based FRET efficiency versus donor lifetime (E - τ ; E stands for FRET efficiency, τ is lifetime) plots⁶⁴ and burst-wise fluorescence correlation spectroscopy (FCS). These analyses provide access to FRET-dynamics that occur on timescales from a few milliseconds down to the sub- μ s regime. This allows us to assess whether the observed FRET populations represent stable conformational states or time averages of (rapidly) interconverting states.

We first performed BVA of GlnBP(119-192) data with ATTO 532/ATTO 643 as a dye pair using a dual-channel burst search (DCBS)⁶¹. In BVA, within-burst E^* -dynamics are identified as an elevated standard deviation of the apparent FRET efficiencies, $\sigma(E^*)$, beyond what is expected from photon statistics, i.e., $\sigma(E^*)$ values larger than the theoretical semicircle (**Figure S9-S11**, panels A). Our analysis indicates that, for each of the different ligand concentrations, at least some of the recorded single molecules undergo dynamic changes in E^* while diffusing through the confocal spot (**Figure S9**). The within-burst dynamics are more prominent for AF555 and AF647 as fluorescent labels (**Figure S10**) and become most abundant in GlnBP(59C-130C), which was used in previous studies^{18, 19, 38}; **Figure S11**. It is important to note that dynamic changes in apparent FRET efficiency can have photophysical origins and do not necessarily confirm the presence of conformation dynamics. For example, the apparent dynamic changes in E^* might represent within-burst dynamics between FRET-active sub-populations (i.e., $S^* \sim 0.5$) and FRET-inactive subpopulations (e.g., donor-only, acceptor-only). Therefore, it is essential to quantify the BVA observed dynamics and identify the corresponding E^* - S^* subpopulations between which the dynamic transitions occur.

For this purpose, we used multi-parameter photon-by-photon hidden Markov modelling (mpH²MM)^{63, 65} to identify the most-likely state model that describes the experimental results based on how E^* and S^* values change within single-molecule bursts. Such analysis can

provide rates of exchange between distinct states of E^*/S^* and its interpretation is described in detail in Supplementary Note 1. The $mpH^{2\text{mm}}$ analyses can differentiate whether apparent dynamic changes in E^* arise from two conformational sub-populations or from photophysical transitions that do not represent conformational dynamics of GlnBP. Our analysis in **Figure 4B** shows clear signatures for donor- and acceptor-blinking between bright and dark states of the fluorophores (**Figure 4**, **Figure S9**–**11**), i.e., the FRET species with intermediate S^* exchange with species of very high and low S^* values, respectively. $mpH^{2\text{mm}}$ identifies single and static apo FRET-active mid- E^* state in the absence of a ligand and a single and static FRET-active high- E^* state in the presence of saturating levels of ligand, which describe the open (mid- E^*) and closed (high- E^*) conformations of GlnBP. It is only in the presence of low concentrations of glutamine (around its K_d) where two FRET-active sub-populations, representing two distinct conformational states, are identified that could interconvert on timescales slower than 10 ms (i.e., slower than typical burst durations). In conclusion, if intrinsic conformational dynamics existed in apo or holo GlnBP, it could only be between the highly-populated FRET conformation we identify and another conformation that is populated significantly below the sensitivity of our measurement and analysis (i.e., a minor population with a fraction <5-10%) or these transitions would have to occur much faster than the time resolution of our experiments (< 100 μ s), dictated by the alternation periods in the μ sALEX experiment.

To check for the presence of faster dynamics, we used multiparameter fluorescence detection with pulsed interleaved excitation (MFD-PIE)⁶⁶, GlnBP(119-192) labeled with ATTO 532/ATTO 643 were used since this combination of cysteines and dyes showed the least photophysical artifacts. In **Figure 5**, we first show two dimensional plots of FRET efficiency (E) versus donor fluorescence lifetime values in the presence of acceptor ($\tau_{D(A)}$) for apo and holo GlnBP. The theoretical linear relationship between E and $\tau_{D(A)}$ defines the static FRET line (**Figure 5A**, black lines). When the labeled molecules exhibit dynamics faster than the diffusion time, the fluorescence-weighted-average of the donor lifetime becomes biased towards longer donor lifetimes due to the higher brightness values of low-FRET species.⁶⁴ Therefore, fast conformational switching is seen as bursts with distinct FRET efficiency values exhibiting a population shift towards the right of the static FRET line. As can be observed from the E - τ plots (**Figure 5A**), the center-of-mass of the FRET populations for both apo and holo GlnBP are coinciding with the static FRET line, suggesting the absence of conformational changes on timescales faster than ms in line with data in **Figure 4**.

We also looked for dynamics using burst-wise FCS analysis (**Figure 5B**). For this, bursts containing signal from both fluorophores were selected, padded with 50 ms before and after burst identification and the fluorescence autocorrelation functions of donor (**Figure 5B**, green curves) and acceptor signals (**Figure 5B**, red curves) as well as for the fluorescence cross-correlation functions between donor and acceptor signals (**Figure 5B**, black curves) were calculated. Conformational dynamics are expected to manifest themselves as an anticorrelation contribution in the cross-correlation function between donor and acceptor channels due to fluctuations in FRET efficiencies that occur faster than the translational diffusion component of the correlation functions (~ 1 ms on our setup).⁶⁷ The burst-wise FCS analysis at times <100 μ s resulted in plateaued cross-correlation functions (**Figure 5B**, black lines) for apo and holo states indicating the lack of dynamics down to the time-resolution of the experiments, i.e., the typical clock time of the photon time tagging on the order of 100 ns.

Studies of surface-immobilized GlnBP via TIRF microscopy

Next, we characterize GlnBP and its conformational dynamics on timescales beyond the residence time of molecules in the confocal excitation volume (i.e., >1-10 ms) with the hope to obtain information on rare conformational events. We consequently conducted smFRET with NTA-based surface-immobilization of the GlnBP His-tag using TIRF microscopy (see Supplementary Note 2 and accompanying Figures **Figure S12**–**16** for details). We reasoned that this would also allow the direct comparison of our results to those of Wang, Yan and co-workers^{18, 19, 38},

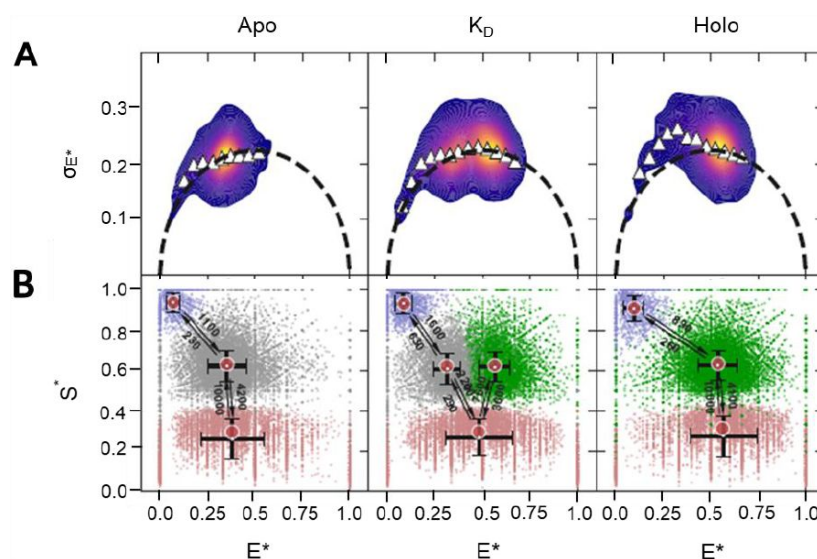


Figure 4.

Screening GlnBP for rapid within-burst FRET dynamics.

(A) Burst Variance Analysis (BVA) showing a weak signature of within-burst FRET dynamics in the low E^* regime. (B) Two-dimensional E^* versus S^* scatter plots of dwells in mpH²MM-detected states within bursts detected by the Viterbi algorithm. Arrows and adjacent numbers indicate transition rates in s^{-1} . Transitions with rates $<100 s^{-1}$ are omitted since such long dwells in a state before transitions are improbable to occur within single-molecule bursts with durations <10 ms and are most probably a mathematical outcome of the mpH²MM optimization framework. The dispersion of the E^* and S^* values of dwells in mpH²MM-detected states are due to the short dwell times in these states, where the shorter the dwell time in a state is, the lower the number of photons it will include, and hence the larger the uncertainty will be in the calculation of E^* and S^* values of dwells. E^* and S^* are E^* and S^* values uncorrected for background, since in mpH²MM all burst photons are considered, including ones that might be due to background. Full analysis shown in [Figure S9](#).

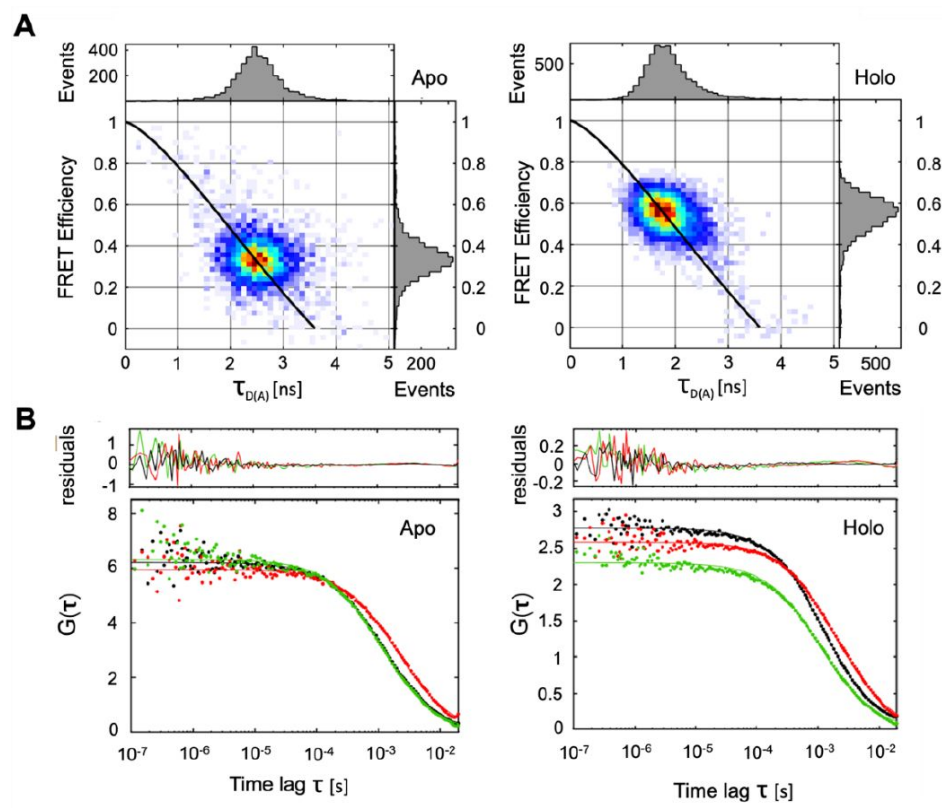


Figure 5.

Screening GlnBP for rapid dynamics within single molecule bursts using E- τ and burst-wise FCS analyses.

(A) Two-dimensional histogram of FRET efficiency (E) versus donor lifetime in the presence of acceptor ($\tau_{D(A)}$) for apo (left) and holo (right) GlnBP. The FRET populations coincide well with the theoretical static FRET line (black) indicating the absence of conformational dynamics taking place at timescales faster than ms. (B) Analysis of FRET conformational dynamics using burst-wise FCS for apo and holo states on bursts exhibiting photoactive donor and acceptor fluorophores. The fluorescence autocorrelation functions of the detected donor (DDxDD) and acceptor signal (AAxAA) are displayed in green and red, respectively. The fluorescence cross-correlation function between donor and acceptor signals (DDxDA) is shown in black.

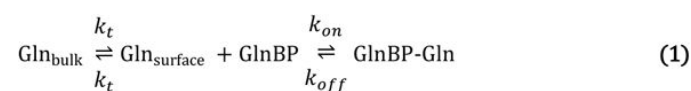
Importantly, in our analysis, we found that various buffer additives used for oxygen depletion have the same effect on GlnBP as the addition of glutamine (i.e., apo-GlnBP becomes artificially “closed” in the presence of the additives) as we demonstrated in solution-based μ sALEX experiments (Figure S12). Consequently, these additives were omitted since their effects mimic that of substrate binding. Strikingly, the conformational states of GlnBP were also partially altered upon surface immobilization (Figure S13), i.e., the E^* values of GlnBP in apo/holo-state were significantly higher than in solution (Figure S13, S15). Furthermore, GlnBP did not retain its biochemical activity on the glass coverslips (Figure S13), i.e., only ~50 % of all GlnBP molecules showed the expected shift towards higher FRET values upon addition of the ligand (Figure S13F). To validate our setup and immobilization approach, we additionally tested dsDNA (Figure S13A, C) and the two previously studied proteins SBD1 and SBD2 (Figure S16). Here, we did not observe discrepancies in FRET efficiency or biochemical activity, and the data of freely-diffusing and surface-immobilized species were consistent.

Our combined smFRET analysis of GlnBP under different biochemical conditions suggests that conformational changes are tightly coupled to the ligand glutamine (Figure 3). We can also rule-out fast conformational dynamics on timescales between 100 ns and 10 ms of apo and holo GlnBP via mpH²-MM, MFD-PIE, and burst-wise FCS (Figure 4, 5). Furthermore, our analysis suggests that apo GlnBP does not adopt (partially) closed conformations on the timescale >10 ms that are of high abundance, i.e., >5-10 % (Figure 3-5). While these results provide valuable information on ligand binding affinity, conformational heterogeneity and timescales of conformational dynamics in GlnBP, they are insufficient to exclude one or the other ligand-binding mechanisms (IF vs. CS). We thus decided to integrate the information into a general theoretical framework for analysis of ligand-binding mechanisms^{37, 46, 47}, for which knowledge of the association and dissociation rates of ligand binding are required.

Insights on ligand binding kinetics from bulk spectroscopy

Such kinetic information is available from surface plasmon resonance spectroscopy (SPR). We immobilized GlnBP via its His-tag on a sensor chip and monitored its interaction with glutamine as a function of time. Even though GlnBP became partially inactive during immobilization for smFRET in TIRF microscopy (Figure S12-S16), we reasoned that non-functional GlnBP will not be observed in SPR since only functional protein can contribute to the signal changes. The assumption that GlnBP remains functional on SPR-chips was validated by the match of ligand-binding characteristics obtained from ITC (Figure 2C, Figure S3), smFRET (Figure 3C, D) and SPR (Figure 6A).

In SPR, GlnBP showed specific and stable interaction with glutamine based on the magnitude of the equilibrium RU response as a function of glutamine concentration (Figure 6A). Analysis of the concentration-dependent maximal RU units yields a K_d of 10 nM (Figure 6A). The overall maximal response of around 3-4 RU indicates a 1:1 stoichiometry of glutamine assuming a monomeric state of GlnBP (Figure 2C, Figure 6). Kinetic association and dissociation experiments were conducted under pseudo-first order conditions, i.e., the assumption of constant glutamine concentrations during an SPR run, due to the applied flow of buffer. The data were analyzed with the standard two-step reaction scheme^{68, 69}.



This includes a mass-transport step between the bulk solution of the applied flow and the sensor surface with transport rate k_t in both directions, and a binding step with effective on- and off-rate constants, k_{on} and k_{off} . Because of the dominance of mass transport, fits of this reaction scheme to the SPR sensorgrams using fit parameters k_t and k_{on} (after substituting k_{off} with $K_d k_{\text{on}}$ in the scheme) do not allow determination of k_{on} within reasonable error bounds. However, fits with

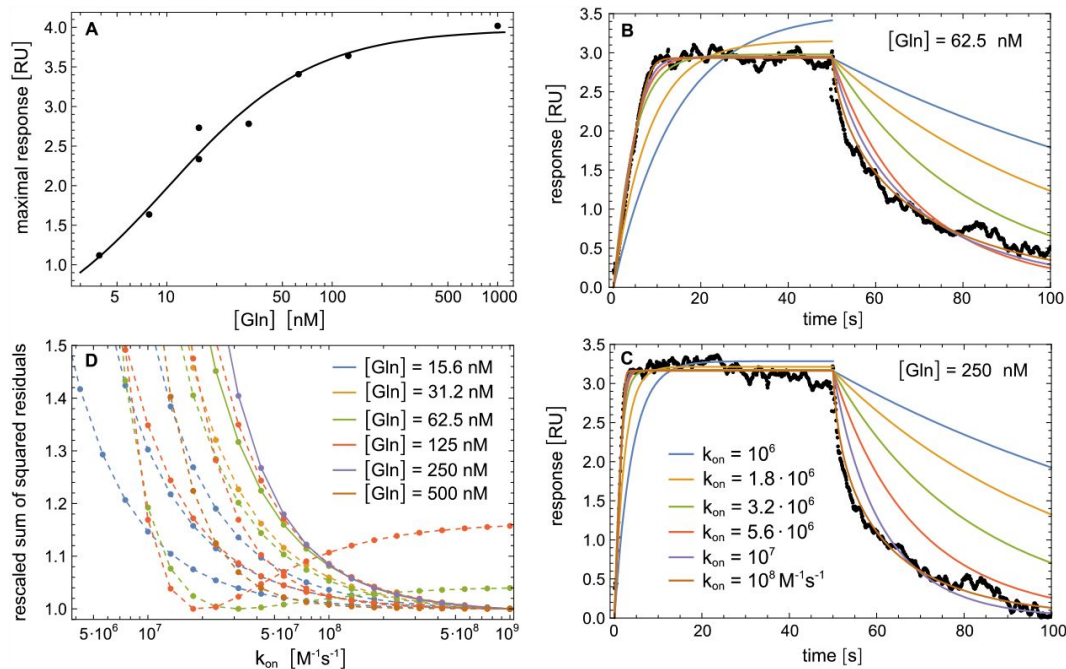


Figure 6.

Kinetic analysis of ligand binding and dissociation in GlnBP using SPR.

(A) Fitting of maximal responses in sensorgrams from a measurement set with [Gln] concentrations from 7.8 to 1000 nM (data points) with $f = c/(1 + K_d/[Gln])$ leads to $K_d = 10 \pm 1$ nM. (B, C) SPR sensorgrams with an association phase of 50 s at the indicated glutamine concentrations [Gln] followed by a dissociation phase of 50 s with [Gln] = 0 in the bulk flow (data points), and fits of the sensorgrams with the reaction scheme (1) for different values of the effective on-rate constant k_{on} (see Methods for details). (D) Rescaled sum of squared residuals versus k_{on} for fits of sensorgrams with different values of [Gln] in the association phase. Note that multiple repeats for the ligand concentrations [Gln] = 15.6 nM, 62.5 nM, and 125 nM are plotted. The two curves with full lines correspond to fits in panels B and C. The 11 curves with dashed lines correspond to the fits in [Figure S17](#).

fixed values of k_{on} indicate that effective on-rate constants smaller than $10^{-7} \text{ M}^{-1} \text{ s}^{-1}$ are incompatible with the sensorgrams (**Figures 6B, C** and **Figure S17**). More precisely, plots of the rescaled sum of squared residuals for these fits versus k_{on} (**Figure 6D**) indicate a lower bound of at least $3 \cdot 10^{-7} \text{ M}^{-1} \text{ s}^{-1}$ for k_{on} ; this implies $k_{\text{off}} = K_d k_{\text{on}} > 0.3 \text{ s}^{-1}$ (with $K_d = 10 \text{ nM}$). Among the 13 plots in **Figure 6D**, and among the 4 plots for $[\text{Gln}] = 125 \text{ nM}$, only one plot exhibits a minimum of the sum of squared residuals below this bound and is therefore likely an outlier.

Discussion & conclusion

Conformational states of macromolecular complexes and changes thereof govern numerous cellular processes including replication⁷⁰, transcription^{71, 72}, translation⁷³, signal transduction⁷⁴⁻⁷⁶, membrane transport^{77, 78}, regulation of enzymatic activity⁷⁹⁻⁸², and the mode of action of molecular motors^{83, 84}. While many conformational changes that are triggered by ligand binding have been characterized extensively, it has also become evident that proteins exhibit prominent intrinsic structural dynamics without the involvement of ligands or other biomacromolecules^{1-6, 85-90}. In a four-state system (**Figure 1C**), ligand-binding can occur via two ‘extreme’ mechanistic pathways, i.e., ligand binding occurs before conformational change (induced fit, IF) or conformational change occurs before ligand binding (conformational selection, CS). The clear temporal ordering of ligand binding and conformational change along these pathways implies that the binding transition time, i.e., the time the ligand needs to enter and exit the protein binding pocket, are small compared to the dwell times of the protein in the two conformations, which is plausible for small ligands⁴⁶. Here, we dissected the ligand-binding processes and conformational dynamics in GlnBP using complementary techniques. We used smFRET experiments to monitor dynamics of conformational changes, SPR to monitor ligand binding and dissociation kinetics, and we obtained ligand affinity values from ITC, SPR and smFRET. We finally ask the question, which binding mechanism is compatible with the combined kinetic and thermodynamic data.

Elucidation of the ligand binding mechanism

To elucidate the binding mechanism, we considered all available information from smFRET in combination with kinetic analysis of SPR data. We hereby follow a published theoretical framework that aims at an unambiguous assignment of the reaction schemes via kinetic rate analysis^{37, 46, 47}. In essence, we ask whether the experimental parameters are compatible both with the IF pathway and the CS pathway of **Figure 7A, B** or only one of them. Both pathways are shown with conformational excitation and relaxation rates, k_e and k_r , and with association and dissociation rate constants, k_+ and k_- , for the binding-competent conformation.

Our smFRET analysis indicates that ligand binding is correlated to a conformational change from an open to a closed state of GlnBP and gives detailed information on the conformational dynamics. It excludes structural dynamics of both apo- and holo-GlnBP on timescales between 100 ns and 10 ms. We were also able to estimate an upper bound of <5-10% for the population of the potential ligand-free (partially-)closed conformations that might exchange with apo-GlnBP. The analysis of SPR sensorgrams leads to the bounds $k_{\text{on}} > 3 \cdot 10^{-7} \text{ M}^{-1} \text{ s}^{-1}$ and $k_{\text{off}} = K_d k_{\text{on}} > 0.3 \text{ s}^{-1}$ (with $K_d = 10 \text{ nM}$) for the effective on- and off-rate constants k_{on} and k_{off} at all considered ligand concentrations of glutamine up to 500 nM.

We first discuss the scenario of a dominant CS pathway in GlnBP. To relate it to the effective on- and off-rates of the SRP analysis, we note that the relaxation rate $k_{\text{obs}}^{\text{CS}}$ of the CS reaction scheme in **Figure 7B** can be well-approximated by

$$k_{\text{obs}}^{\text{CS}} \approx k_{\text{on}}[\text{L}] + k_{\text{off}} \quad (2)$$

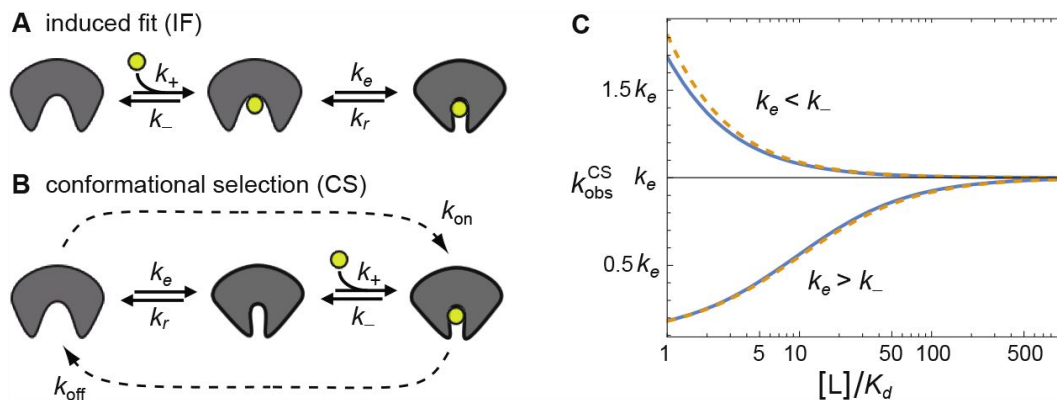


Figure 7.

Dominant relaxation rate k_{obs} of binding pathways.

(A, B) Induced-fit and conformational-selection pathways with conformational excitation and relaxation rates, k_e and k_r , and with association and dissociation rate constants, k_+ and k_- , for the binding-competent conformation of the pathway. (C) Dominant relaxation rate, k_{obs}^{CS} , of the conformational-selection pathways versus ligand concentration $[L]$. Blue lines represent the exact pseudo-first-order result $k_{obs}^{CS} = \frac{1}{2}(S - \sqrt{S^2 - 4(k_e(k_+[L] + k_-) + k_r k_-)})$ with $S = k_e + k_r + k_+[L] + k_-$ and $K_d = k_-(k_e + k_r)/k_+ k_e$ for $k_- = 10 k_e$ and $k_r = 9 k_e$ (upper curve) and $k_- = 0.1 k_e$ and $k_r = 9 k_e$ (lower curve). The dashed yellow lines represent the approximate result from equation 2. For the induced-fit pathway, the dominant relaxation rate $k_{obs}^{IF} = \frac{1}{2}(S - \sqrt{S^2 - 4(k_+[L](k_e + k_r) + k_e k_-)})$ with S as above is monotonously increasing (similar to k_{obs}^{CS} for $k_e > k_-$) and has the limiting value $k_e + k_r$ at large ligand concentration^{37, 47}.

with effective on- and off-rate constants $k_{on}=k_e k_+/(k_e+k_+[L])$ and $k_{off}=k_r k_-/(k_e+k_+[L])$ that depend on the conformational transition rates, k_e and k_r , between the open and closed conformation in an unbound GlnPB and on the rates, k_+ and k_- , for the binding step in the closed conformation along this pathway. This approximation holds for small populations of the closed conformation in ligand-free GlnPB with upper bound of 5-10% from the smFRET analysis and for ligand concentrations $[L] > K_d$ and, thus, for all the concentrations shown in **Figure 6**. At the largest ligand concentration of 500 nM of the SPR sensorgrams, we obtain $k_{obs}^{CS} > 15 \text{ s}^{-1}$ from this equation. Eqn. 2 can be further simplified to

$$k_{obs}^{CS} \approx \frac{k_e k_- (1 + [L]/K_d)}{k_e + k_- [L]/K_d} \quad (3)$$

with $K_d = k_r k_- / k_+ k_e$. The limiting value of k_{obs}^{CS} at large ligand concentration $[L]$ obtained from this equation is k_e . To conclude the argument, we now consider two cases: (1) for $k_e > k_-$, the relaxation rate k_{obs}^{CS} increases with $[L]$ as seen in **Figure 7c** (lower curve). The limiting value k_e of $k_{obs}^{CS}([L])$ is therefore larger than 15 s^{-1} , because $k_{obs}^{CS} > 15 \text{ s}^{-1}$ at $[L] = 500 \text{ nM}$ (see above). (2) for $k_e < k_-$, the relaxation rate k_{obs}^{CS} decreases with $[L]$ as seen in **Figure 6c** (upper curve). In this case, $k_{obs}^{CS}([L])$ is already very close to its limiting value k_e at $[L] = 500 \text{ nM}$ for $K_d = 10 \text{ nM}$. In both cases, we thus obtain $k_e > 15 \text{ s}^{-1}$, and from this, $k_r > 9 k_e > 135 \text{ s}^{-1}$ for an upper bound of 10% of the population $k_e/k_e + k_r$ of the closed conformation in ligand-free GlnBP. However, rates $k_r > 135 \text{ s}^{-1}$ correspond to transition timescales $< 7.4 \text{ ms}$, which are timescales for conformational dynamics excluded by the smFRET results. Alternatively, timescales smaller than 100 ns are “allowed” for conformational exchange between the open and closed state. However, we consider this timescale unrealistically fast for the overall inter-domain conformational changes of all substrate binding domains and for the herein case of GlnBP. We thus conclude that the CS pathway is incompatible with our results. In contrast, IF is fully compatible with all experimental data presented here (see Supplementary Note 3). We thus propose that IF is the dominant pathway for GlnBP, further supported by the notion that the open conformation is much more likely to bind substrate than the closed one based on steric arguments (see Supplementary Note 4).

What implications do our results and the proposed integrative strategy for determining (or excluding) ligand binding mechanisms have for other protein systems? Generally, we encourage the use of similar strategies for other biomacromolecular systems. A potential improvement would be to obtain relaxation kinetics without the mass transport limitations in SPR, which is particularly relevant for small ligand molecules. Thus, stopped-flow (FRET) experiments, which have already been used in the 1970s for binding-rate determination in GlnBP, would be a more direct approach that could complement smFRET data nicely and lead to the same conclusions as presented above.

An interesting next target would be the comparison of binding mechanisms in type I and type II SBPs, which differ in their overall core topology and in the composition of their hinge domain. The hinge domain allows for conformational change between open and closed conformational states and is composed of two β -strands for type II, but three strands for the type I family. Additionally, we recently showed that C-terminal embellishments in members of the type II family have an impact on the conformational states and dynamics, all of which can directly impact and alter the binding mechanism, e.g., from lock-and-key towards an induced-fit.

We finally think that various SBP systems (and their binding mechanisms) should be revisited. This is relevant since there are many findings and controversial interpretations whenever intrinsic conformational motion or closed-unliganded conformations were identified for the maltose binding protein MalE, histidine binding protein (HisJ), D-glucose/D-galactose-binding protein (GGBP), ferric-binding protein (FBP), choline/acetylcholine substrate binding protein (ChoX) and the Lysine-, Arginine-, Ornithine-

binding (LAO) protein³⁴. Also the advent of single-molecule approaches, such as nanopore-recordings¹³ and single-molecule Förster-resonance energy transfer (smFRET)^{14–21} provided a large pool of data for various ABC transporter-related SBPs^{20, 44} with a wide range of distinct ligands such as metal ions^{20, 93}, osmolytes^{20, 94}, amino acids^{16–21}, peptides²⁰, sugars^{7, 20, 63, 95, 96}, siderophores⁹⁷, and other small molecules⁹⁸ – for most of which additional data is required to univocally assign a ligand-binding mechanism.

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Author contributions

N.Z. and T.C. conceived and designed the study and supervised the project. Z.H., S.P., M.R., A.H. and D.G. performed molecular biology and produced GlnBP. Z.H., S.P., A.H. and A.N. performed biochemical characterization. Z.H. established labeling protocols. S.B. and K.J. provided SPR data. Z.H., S.P. and A.H. performed μ ALEX experiments, E.B. performed PIE experiments. E.B. and D.C.L. analyzed PIE data. Z.H. and M.I. performed TIRF experiments, O.B. supported TIRF data analysis. O.B., P.D.H. and E.L. performed smFRET data analysis using mpH²MM. E.L. performed Gln docking calculations onto GlnBP structures. M.I., T.W. and T.C. analyzed SPR data. T.W. and T.C. analyzed and interpreted kinetic rate schemes. Z.H. prepared figures. Z.H., T.W. and T.C. wrote the manuscript in consultation with all authors. All authors contributed to discussion and interpretation of the results and approved the final version of the manuscript.

Material and methods

All commercially obtained reagents were used as received, unless stated otherwise. The following grades were used: Guanidine hydrochloride (99%, Sigma Aldrich), 1,4-Dithiothreitol (DTT) (99%, ROTH), Thermo Scientific SnakeSkin TM Dialysis Tubing (Fisher scientific, 10K MWCO, 16 mm), Ni²⁺-Sephacrose resin (GE Healthcare), Albumin fraction V (BSA), biotin-frei, $\geq 98\%$ (Carl Roth GmbH), Imidazol, $\geq 99\%$ (Carl Roth GmbH), Isopropyl- β -D-1-thiogalactopyranoside (IPTG), $\geq 99\%$ (Carl Roth GmbH), Kanamycin (Carl Roth GmbH), L-glutamine (Merck KGaA), L-Arginine (Carl Roth GmbH), AF555 (Jena Bioscience, Germany), AF647 (Jena Bioscience, Germany), ATTO 532 (ATTO-TEC, Germany), ATTO 643 (ATTO-TEC, Germany), mPEG3400-silane (abcr, AB111226) and biotin-PEG3400-silane (Laysan Bio Inc), Biotin-NTA (Biotium), Streptavidin (Roth, Germany), Pyranose oxidase (Sigma Aldrich, Germany), Catalase (Sigma Aldrich, Germany), Glucose ($\geq 99.5\%$ GC, Sigma Aldrich, Germany), Trolox (98%, Sigma Aldrich, Germany), Potassium hydroxide ($\geq 85\%$, Honeywell, Germany), Acetone (Roth, Germany), Toluene (Roth, Germany).

Protein expression and purification

Two GlnBP double cysteine variants were generated by site-directed mutagenesis, allowing the insertion of two cysteine residues into GlnBP at positions (V111C – G192C) and (T59C – T130C), separately. *Escherichia coli* BL21-pLysS cells were freshly transformed with the plasmid carrying the coding sequence for GlnBP WT or a GlnBP variant, and grown in 2 L LB medium (100 mg/mL Kanamycin and 50 mg/mL chloramphenicol) at 37 °C under aerobic conditions. At an OD_{600nm} of 0.6-0.8, overexpression of the proteins of interest was induced upon addition of 1 mM IPTG to the culture media. The cells were further grown for 1.5-2.0 hours after induction and then harvested by centrifugation for 20 minutes at 1,529 g (Beckman, JA10) at 4 °C. All subsequent operations were carried out at 4 °C, and all solutions were stored at 4 °C. Cell pellets from 2 L culture were collected in a 50 mL falcon and resuspended in buffer A (50 mM Tris-HCl, pH 8.0, 1 M KCl, 10 mM imidazole, 10% glycerol) with 1 mM dithiothreitol (DTT). At 4 °C, the falcon was gently shaken overnight.

Cells were disrupted by sonication (Branson tip sonication; amplitude: 25%; 10 min; 0.5 s on-off pulses; temperature was kept low by the use of an ice-water bath). Centrifugation was used to fractionate the cell lysate (at 4 °C for 30 min at 4,416 g, Eppendorf, Centrifuge 5804 R) and at 4 °C for 1 hour for ultracentrifugation (70,658 g, Beckman, Type 70Ti) in vacuum, and the pellet was discarded. The protein was purified by affinity chromatography using the Ni²⁺-Sephacrose fast flow resin (GE Healthcare), pre-equilibrated with 10 column volumes of buffer A containing 1 mM DTT and gravity loaded with the supernatant from the preceding ultra-centrifugation step. The resin-bound protein was washed with 10 column volumes of buffer A containing 1 mM DTT, followed by buffer B containing 1 mM DTT (50 mM Tris-HCl, pH 8.0, KCl 50 mM, imidazole 20 mM, glycerol 10%), and finally eluted in buffer C (50 mM Tris-HCl, pH 8.0, KCl 50 mM, imidazole 250 mM, glycerol 10%) with 1 mM DTT. The eluted sample was concentrated (Vivaspin6 columns, 10 kDa MWCO, 6 mg/mL), dialyzed against PBS buffer supplemented with 1 mM DTT, and stirred gently at 4 °C overnight. SDS-PAGE was used to quantify the yield of protein overexpression and purification (Comassie staining). The absorbance at 280 nm was used to estimate the protein concentration (knowing the molar extinction coefficient of GlnBP ~25,900 M⁻¹ cm⁻¹). The protein was then split into aliquots and kept at a temperature of -20 °C. All proteins were further purified using size-exclusion chromatography (ÄKTA pure system, Superdex 75 Increase 10/300 GL, GE Healthcare). The purified protein was split into aliquots and stored at -80 °C prior to the measurements.

Unfolding and refolding process of GlnBP WT and GlnBP variants

The stock concentrations of GlnBP variants were estimated at about 6 mg/mL. Each GlnBP variant was thawed from -80 °C, then the protein was diluted to a final concentration of 3-4 μM (final volume of ~20 mL) in the unfolding buffer (PBS buffer) containing 6 M guanidine hydrochloride (GndHCl). Subsequently, the solution was incubated for 3 hours under gentle stirring at ambient temperature. Next, the unfolded GlnBP variants were centrifuged (3,046 g, 30 min at 4 °C) to remove insoluble aggregates which could act as nuclei to trigger aggregation during refolding process. A Snakeskin TM dialysis membrane was prepared (pre-cooled at 4 °C and soaked in refolding buffer - PBS buffer with 1 mM DTT, pH 7.4 - for 2 min). The GlnBP variants were transferred into the dialysis tubing, which were sealed tightly afterwards by double-knots and clips at each end. The unfolded GlnBP variant was refolded by a two-step dialysis, in the presence of a total 200-fold excess of refolding buffer. First, each protein was dialyzed against 2 L refolding buffer overnight under gentle stirring at 4 °C. Then, buffer was exchanged with additional 2 L refolding buffer for another day at 4 °C. The refolded protein was then concentrated from 20 mL to final 500 μL (Vivaspin 10 kDa MWCO; 3,000 g × 15 min at 4 °C) and further purified by size-exclusion chromatography (ÄKTA pure system, Superdex-75 Increase 10/300 GL, GE Healthcare). The unfolding and refolding process for GlnBP WT was conducted under the same conditions as described for the GlnBP variants.

Isothermal titration calorimetry (ITC) measurements

The ITC measurements were performed in a MicroCal PEAQ-ITC isothermal titration calorimeter (Malvern Instruments). The prediction ITC software “MicroCal PEAQ-ITC Control” was employed for designing and conducting the experiments. Once the K_d value and the binding stoichiometry (N) were assigned as predefined values, the concentration of both the protein and the titrant (ligand) stock solutions could be calculated by the “design-experiment” function on the software to get an optimal sigmoidal one-site binding curve. GlnBP concentration was assessed using the Nanophotometer (N60 Touch, Implen GmbH) with at least three reading repeats to get accurate determinations of concentration values. For all ITC measurements, the temperature was set at 25 °C with stirring speed at 750 rev / min. The GlnBPs solution (10 μ M in PBS buffer pH 7.4, 300 μ L) was manually loaded into the sample cell. The titrant (L-Glutamine, 100 μ M in PBS buffer, pH 7.4) was automatically loaded into the titration syringe and injected in the sample cell with a titration speed of 2 μ L every 150 second and a total of 19 injections. As a control experiment, L-Glutamine was titrated into the sample cell containing PBS buffer without GlnBPs. All the titration data were analyzed using the MicroCal PEAQ-ITC Analysis Software.

Surface plasmon resonance spectroscopy (SPR) and data analysis

SPR assays were performed on a Biacore T200 (Cytiva) using a CM5 Series S carboxymethyl dextran sensor chip coated with His-antibodies from the Biacore His-capture kit (Cytiva). Briefly, the chips were equilibrated with running buffer until the dextran matrix was swollen. Afterwards, two flow cells of the sensor chip were activated with a 1:1 mixture of N-ethyl-N-(3-dimethylaminopropyl) carbodiimide hydrochloride and N-hydroxysuccinimide according to the standard amine coupling protocol. A final concentration of 50 μ g/mL anti-histidine antibody in 10 mM acetate buffer pH 4.5 was loaded onto both flow cells using a contact time of 420 s for gaining a density of approximately 10,000 resonance units (RU) on the surface. By injection of 1 M ethanolamine/HCl pH 8.0, free binding sites of the flow cells were saturated. Preparation of chip surfaces was carried out at a flow rate of 10 μ L/min. All experiments were carried out at a constant temperature of 25 °C using PBS buffer (0.01 M phosphate buffer, 2.7 mM KCl, 0.137 M NaCl, pH 7.4) supplemented with 0.05 % (v/v) detergent P20 as running buffer.

For interaction analysis, GlnBP-6His (1.5 μ M) was captured onto one flow cell using a contact time of 240 s at a constant flow rate of 10 μ L/min. This resulted in a capture density of approximately 1,200 RU of GlnBP-6His. Eight different concentrations of glutamine (7.8, 15.6, 31.25, 62.5, 125, 250, 500 and 1,000 nM) were injected onto both flow cells using an association time of 50 s and a dissociation time of 360 s. The flow rate was kept constant at 30 μ L/min. As control, running buffer was injected. The chip was regenerated after each cycle by removing GlnBP-6His completely from the surface using 10 mM glycine pH 1.5 for 60 s at a flow rate of 30 μ L/min.

Sensorgrams were recorded using the Biacore T200 Control software 2.0.2. The surface of flow cell 1 was not coated with GlnBP-6His and used to obtain blank sensorgrams for subtraction of the bulk refractive index background with the Biacore T200 Evaluation software 3.1. The referenced sensorgrams were normalized to a baseline of 0. Peaks in the sensorgrams at the beginning and the end of the injection are due to the run-time difference between the flow cells for each chip.

In total, 26 SPR sensorgrams in three sets of measurements were recorded. To correct for remaining drift in the sensorgrams, the initial 60 s of the sensorgrams prior to Gln injection and the last 100 s of the dissociation phase were first fitted with an exponential function, which was subtracted from the sensorgrams. The drift-corrected sensorgrams were fitted to the reaction scheme of Eq. (1) based on the differential equations^{68, 101}

$$\frac{d[L]_{\text{surf}}}{dt} = k_t([L]_{\text{bulk}} - [L]_{\text{surf}}) - \frac{d[PL]}{dt}$$

$$\frac{d[PL]}{dt} = k_{\text{on}}[L]_{\text{surf}}([P]_{\text{tot}} - [PL]) - k_{\text{off}}[PL]$$

where $[L]_{\text{bulk}} = [\text{Gln}]$ and $[L]_{\text{surf}}$ are the free glutamine concentrations in the bulk flow and at the sensor surface, $[P]_{\text{tot}}$ is the total concentration of surface-immobilized protein, and $[PL]$ is the concentration of bound protein complexes. Conversion to the SPR binding response r via $[PL] = \alpha r$ and $[P]_{\text{tot}} = \alpha r_{\text{max}}$ leads to fit results for the binding rate constants that are insensitive to the (unknown) conversion factor α , which can be understood from the fact that the quasi-steady-approximation $d[L]_{\text{surf}}/dt \approx 0$ holds for SPR setups^{68, 101}. The association phases of the sensorgrams were fitted with initial conditions $[L]_{\text{surf}} = 0$ and $r = 0$ and fit parameters k_t and r_{max} at different values of k_{on} after substitution of k_{off} by $K_d k_{\text{on}}$. Prior to these fits with fixed k_{on} , a remaining small vertical off-set of the sensorgrams was determined as additional fit parameter in fits with unconstrained, large k_{on} and subtracted from the sensorgrams. The first 50 s of the dissociation phases were fitted with single fit parameter k_t for the initial conditions $[L]_{\text{surf}} = [L]_{\text{bulk}}$ and $r = r_{\text{max}}(1 + K_d[L]_{\text{bulk}})$, with r_{max} determined from fits of the association phase of the sensorgram for unconstrained, large k_{on} . Background-corrected sensorgrams that do not reach binding equilibrium in the association phase (because of small $[\text{Gln}]$), still show marked drifts in binding equilibrium, or do not resolve the initial increase of the binding signal of the association phase (because of large $[\text{Gln}]$) were discarded, which leads to the 13 sensorgrams of **Figures 6B,C** and S17 with fit results for $\alpha = 1 \mu\text{M}/\text{RU}$. Fits with e.g., $\alpha = 1 \text{ mM}/\text{RU}$ (not shown) lead to practically identical results. All fits were conducted with Mathematica 13 based on the functions ParametricNDSolveValue to obtain numerical solutions of the differential equations and NonlinearModelFit for fitting parameters of these solutions.

Protein labeling

The refolded GlnBP(111C-192C) and GlnBP(59C-130C) variants were labeled with commercial maleimide derivatives of AF555/AF647 or ATTO 532/ATTO 643⁹⁵, and then purified by SEC. The chromatogram of refolded GlnBP(111C-192C) labeled with AF555/AF647 is shown in **Figure 2B**, and those of all other variants and dye labeling combinations are displayed in **Figure S2**. First, the His-tagged protein was incubated in 10 mM DTT in PBS buffer for 30 min to reduce all oxidized cysteine residues. Subsequently, the protein was diluted 10 times with PBS buffer and immobilized on a Nickel Sepharose 6 Fast Flow resin (GE Healthcare). The resin was washed extensively with milliQ water followed by PBS buffer pH 7.4. To remove the excess of DTT, the resin was washed with PBS buffer. The protein was left on the resin and incubated overnight at 4 °C with 5-10 times molar dye excess in PBS buffer pH 7.4. Subsequently, the unreacted fluorophores were removed by washing the resin with 6 mL of PBS buffer. Bound proteins were eluted with 800 μL of elution buffer (PBS buffer, pH 7.4 400 mM Imidazole). The labeled protein was further purified by size-exclusion chromatography (ÄKTA pure, Superdex-75 Increase 10/300 GL, GE Healthcare) to eliminate remaining fluorophores and remove other contaminants and soluble aggregates. The selected elution fractions were used without further treatment for smFRET experiments as described below. In general all experiments were carried out at room temperature using 25–50 pM of double-labeled GlnBP protein in PBS buffer (pH 7.4). Titration experiments were performed by adding specific concentrations of ligand (glutamine) to the buffer.

smFRET experiments with μ sALEX

Single-molecule μ sALEX experiments were carried out at room temperature on a custom-built confocal microscope. In short, alternating excitation light (50 μ s period) was provided by two diode lasers operating at 532 nm (OBIS 532-100-LS, Coherent, USA) and 640 nm (OBIS 640-100-LX, Coherent, USA). Both lasers were combined by coupling them into a polarization maintaining single-mode fiber (P3-488PM-FC-2, Thorlabs, USA) and subsequently guided into the microscope objective (UplanSApo 60X/1.20W, Olympus, Germany) via a dual-edge dichroic mirror (ZT532/640rpc, Chroma, USA). In general, the 532 and 640 nm diode lasers operated at 60 and 25 μ W, respectively (measured at the back aperture of the objective), unless stated otherwise. Fluorescence light was collected by the same objective, focused onto a 50 μ m pinhole and separated into two spectral channels (donor and acceptor fluorescence) by a dichroic beamsplitter (H643 LPXR, AHF, Germany). Fluorescence emission was collected by two avalanche photodiodes (SPCM-AQRH-64, Excelitas) after additional filtering (donor channel: BrightLine HC 582/75 and acceptor channel: Longpass 647 LP Edge Basic, both from Semrock, USA). The detector outputs were recorded via an NI-Card (PCI-6602, National Instruments, USA) using a custom-written LabView program.

smFRET data analysis (μ sALEX)

Data analysis for μ sALEX was performed using an in-house written software package as described in¹⁶. Three relevant photon streams were extracted from the recorded data based on the alternation period, corresponding to donor-based donor emission F(DD), donor-based acceptor emission F(DA) and acceptor-based acceptor emission F(AA). Bursts from single-molecules were identified using published procedures⁶¹ based on an all-photon-burst-search algorithm with a threshold of 15, a time window of 500 μ s and a minimum total photon number (F(DD)+D(DA)+F(AA)) of 150, unless stated otherwise in the figure caption.

For each fluorescence burst, the stoichiometries S^* and apparent FRET efficiencies E^* were calculated and then presented for all bursts yielding a two-dimensional (2D) histogram. Uncorrected apparent FRET efficiency, E^* , monitors the proximity between the two fluorophores and is calculated according to $E^* = F(DA)/(F(DD)+F(DA))$. Apparent stoichiometry, S^* , is defined as the ratio between the overall fluorescence intensity during the green excitation period over the total fluorescence intensity during both green and red periods and describes the ratio of donor-to-acceptor fluorophores in the sample: $S^* = (F(DD)+F(DA))/(F(DD)+F(DA)+F(AA))$. Collecting the E^* and S^* values of all detected bursts into a 2D E^*-S^* histogram yielded subpopulations that can be separated according to their E^* - and S^* -values. The 2D histograms were fitted using a 2D gaussian function, yielding the mean apparent FRET efficiency and its standard deviation or width of the distribution. μ sALEX, assists in sorting single molecules based on their donor/acceptor dye brightness ratio (stoichiometry S^*) and uncorrected mean FRET efficiency (apparent FRET E^*), which can be related on the mean inter-dye distance^{95, 102}.

Analysis with mpH²MM was conducted as described previously by the Lerner lab⁶³. In short, the FRET Bursts software¹⁰³ was used for detecting single-molecule photon bursts using the dual channel burst search⁶¹ AND-gate algorithm with a sliding window of $m=10$ photons searching for instances with an instantaneous photon rate of at least $F=6$ times the background rate. Afterwards, bursts of such consecutive photons were filtered to have at least 50 photons originating from donor excitation and at least 50 photons originating from acceptor excitation. In the data analysis, the photon stream was then divided into photon streams of different bursts, and a time shift was applied to acceptor excitation originating photons stream so that their arrival time range overlap with that of donor excitation originating photon streams. Optimizations were conducted with state models of increasing numbers of states, and the Viterbi algorithm was employed for calculating the integrated complete likelihood (ICL). Optimizing for larger numbers of states ceased once the ICL ceased to decrease between successively larger state models.

Optimized models were manually examined, and the optimal state model selected considering the ICL and the reasonableness of the model given prior knowledge based on transition rates and the E^* and S^* values of the states. After selection of the most-likely state model, the corresponding most-likely state-path determined by the *Viterbi* algorithm was used to segment bursts into dwells and to classify burst by which states were present within each burst.

To support the idea that apo and holo state in solution match with that of the crystal structure, we performed a quantitative comparison of inter-dye distances calculated from dye accessible volumes (AV) on structural models of apo and holo protein, and those derived from the experimental smFRET results. For dye AV calculations we used the FPS method, established by the Seidel lab¹⁰⁴ (Figure S1). The experimental data were corrected for setup-dependent parameters according to refs.^{60, 96} to obtain accurate FRET values from μ sALEX data. Using a Förster distance of 5.2 nm for AF555/AF647, we found good agreement, i.e., 0.3-0.5 nm deviations (and 1.0 nm in one case) between the calculated and experimentally derived inter-dye distances for both mutants (Figure S1).

smFRET measurements with MFD-PIE and burst-wise FCS analysis

Solution-based smFRET experiments were performed on a home-built dual-color confocal microscope that combines multiparameter fluorescence detection (MFD) with pulsed interleaved excitation (PIE).⁶⁶ MFD-PIE experiments have been described in detail previously.¹⁰⁵ With MFD-PIE, it is possible extract FRET efficiency, stoichiometry, fluorescence lifetime and fluorescence anisotropy information from each single-molecule burst. Correction factors including direct acceptor excitation (α), spectral crosstalk (β) and detection correction factor (γ) are also accounted for reporting accurate the FRET efficiency values.¹⁰⁶ The accurate FRET efficiency (E) can be determined from:

$$E = \frac{F_{GR} - \alpha F_{RR} - \beta F_{GG}}{F_{GR} - \alpha F_{RR} - \beta F_{GG} + \gamma F_{GG}}$$

where F_{GG} , F_{GR} and F_{RR} are background-corrected fluorescence signals detected in green/ donor (G), red/acceptor (R) after donor excitation and acceptor channels, respectively.

Alternatively, the use of picosecond pulsed lasers and time-correlated single photon counting (TCSPC) electronics enable calculating FRET efficiencies from the quenching of the donor in presence of acceptor. According to the formula:

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

$\tau_{D(A)}$ is the fluorescence lifetime of the donor in presence of acceptor and $\tau_{D(0)}$ is the fluorescence lifetime of the donor only species. Static species can be observed on the theoretical static FRET line, which is a linear relation between E and $\tau_{D(A)}$. Sub-ms conformational dynamics can also be identified and judged by observing the right-shifted populations from the static FRET line.

For the measurements here, 100 pM of GlnBP labeled with ATTO 532 and ATTO 643 was placed on a BSA-passivated LabTek chamber and measured for 2 hours. The sample was excited with 532 and 640 nm pulsed lasers with a repetition rate of 26.6 MHz and laser powers of 45 and 23 μ W (measured at the back aperture of the objective), respectively.

Burst-wise FCS analysis is an alternative approach to observe sub-ms conformational dynamics. In this approach, donor (DD) and acceptor (AA) signals detected from single-molecule events are cross-correlated. Thus, fluctuations in the FRET efficiencies appear as an anti-correlated signal in the donor-acceptor fluorescence cross-correlation function. Burst with sufficient photons detected in both the donor and acceptor channels were selected. A time window of 50 ms was applied

around each burst. If another burst was detected within this time window, both were eliminated to ensure correlation functions that are specific to the selected bursts. All the above mentioned data analysis was done using the PIE analysis with Matlab (PAM) software package¹⁰⁷.

Surface immobilization of DNA and GlnBP(111C-192C)

Biotin-streptavidin interaction was used to immobilize tagged proteins and labeled DNA on a PEG-functionalized coverslip for single molecule studies. The protein-his-tag and a biotin-NTA chelated with Ni^{2+} were used to mark GlnBP(111C-192C) labeled with maleimide modified derivatives of ATTO 532/ATTO 643, whilst DNA labeled with Cy3B/ATTO 647N was directly tagged with a biotin. To prepare a functionalized glass surface, cover slides (1.5H Marienfeld Superior) were first sonicated in MQ water for 30 min. The slides were rinsed three times with MQ water, sonicated for 30 min in HPLC-grade acetone, rinsed three times with MQ water again. Then, the slides were sonicated with 1 M KOH for 30 min, rinsed three times with MQ water and dried with a stream of nitrogen air. To remove any organic material left on the surface, the cover slides were plasma-cleaned for 15 min with oxygen. To create a mPEG/biotin-coated surface, the slides were immediately incubated in a 99:1 solution of mPEG3400-silane (abcr, AB111226) and biotin-PEG3400-silane (Laysan Bio Inc) in a Toluene solution overnight at 55 °C. After incubation, the slides were sonicated (10 min in ethanol, 10 min in MQ water), dried under nitrogen stream, and kept under vacuum. Prior to TIRF experiments, each slide was incubated with a 0.2 mg/mL streptavidin in PBS solution for 10 min utilizing Ibidi sticky-slide (18 well) for single molecule studies. PBS buffer pH7.4 was used to wash away the unbound excess of streptavidin. For GlnBP(111C-192C) immobilization, 20 nM biotin-NTA (QIAGEN) was charged with 50 nM Ni^{2+} and incubated on the slide for 10 min before rinsing away the unbound excess biotin-NTA and Ni^{2+} with PBS (this step was omitted for the labeled DNA samples). GlnBP(111C-192C) at 0.8 nM and dsDNA at 0.04 nM were incubated for 5 and 1 min, respectively. For single-molecule data collecting, imaging buffer (PBS, pH 7.4) containing 2 mM Trolox for protein. For dsDNA we used PBS buffer in combination with an oxygen scavenging system (pyranose oxidase at 3 U/mL, catalase at final concentration of 90 U/mL, and 40 mM glucose). After that, the chambers were sealed with Silicone Isolators™ Sheet Material (Grace Bio-labs). All the single-molecule investigations were done at room temperature.

smFRET measurements with TIRF microscopy including data analysis

Single-molecule TIRF measurements were conducted on a homebuilt microscope using an Olympus iX71 inverted microscope body. Light from a 532 nm continuous wave laser (532 nm OBIS, Coherent) was transmitted off-axis onto the back-focal plane of a microscope objective (UAPON TIRF 100X 1.49NA, Olympus) via a dualband dichroic beamsplitter (TIRF Dual Line Beamsplitter zt532/640rpc, AHF Analysetechnik) to generate total internal reflection at the glass-water interface. Fluorescent emission was then split spectrally using a Dual View System (DV2, Photometrics) equipped with a dichroic beamsplitter (zt640rdc, AHF Analysetechnik). The two emission channels were then spectrally filtered using emission filters (582/75 Brightline HC and 731/137 BrightLine HC respectively, both AHF Analysetechnik). Image series were acquired using an EMCCD camera (C9100-13, Hamamatsu) in combination with the μ Manager¹⁰⁸ software. The iSMS¹⁰⁹ software was used to retrieve and calculate traces of the donor and acceptor fluorescence intensity from consecutive fluorescent images.

Supporting information

Supplementary Figures

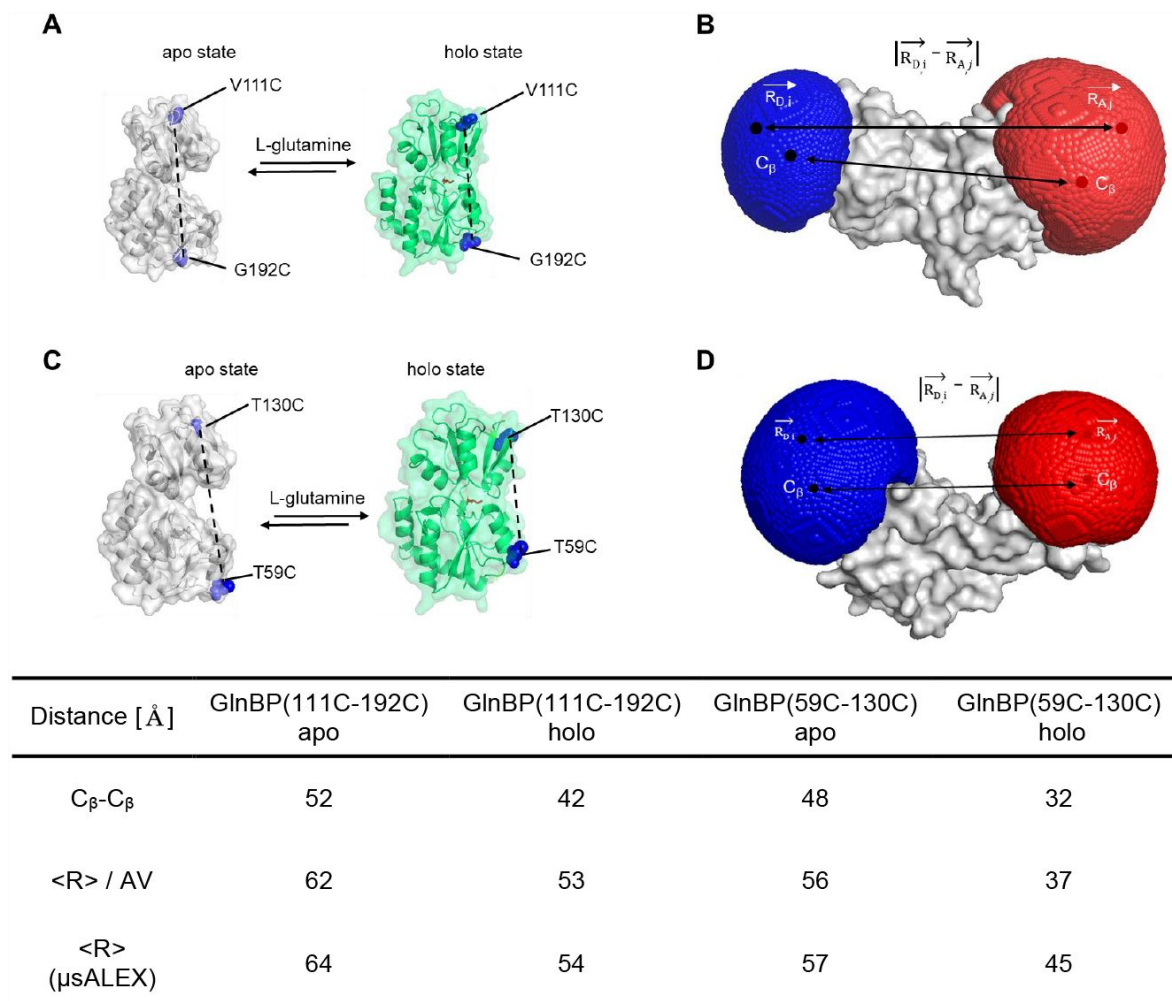


Figure S1.

Crystal structure and dye accessible volume calculations of GlnBP cysteine variants.

(A, C) Crystal structure of the ligand-free (grey structure) and ligand-bound (green structure) GlnBP with the two labeling positions of the respective variants indicated in blue. (B, D) Simulation of accessible volumes for AF555 and AF647 with values of interprobe distances based on structural predictions ($C_{\beta}-C_{\beta}'$ distances and fluorophore accessible volumes) and experimental values $\langle R \rangle$.

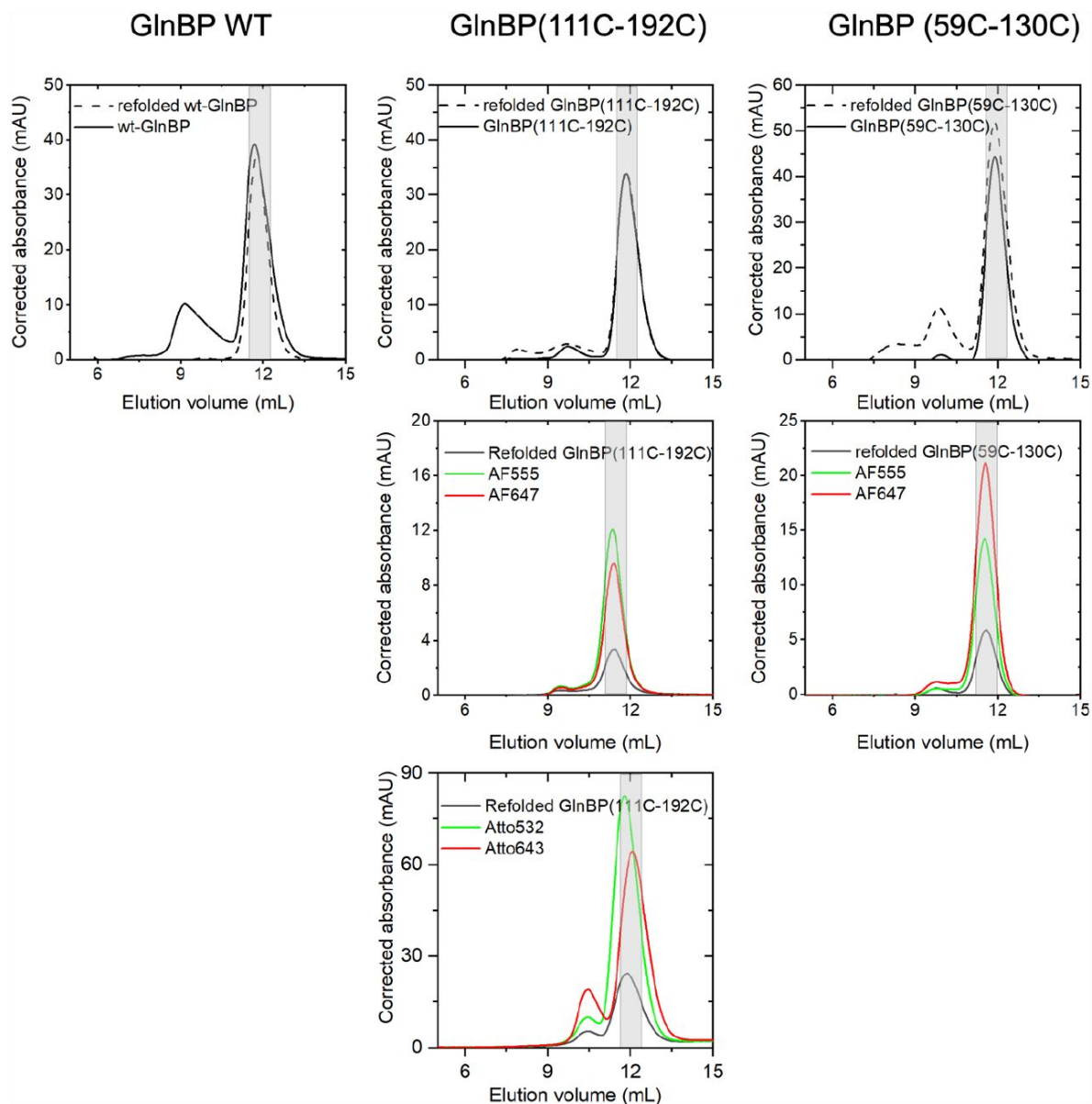


Figure S2.

Size Exclusion Chromatography (SEC) of refolded GlnBP WT and GlnBP variants.

GlnBP WT and GlnBP double-cysteine variants were unfolded with 6 M Guanidine Hydrochloride and then refolded via dialysis over two days in PBS buffer (pH 7.4, 1 mM DTT). The selected fractions (grey-shaded area) were collected and used for ITC experiments. For the solution-based smFRET measurements, the selected fractions (grey-shaded area) having the best overlap of protein, donor, and acceptor absorption were used. The protein absorption was measured at 280 nm (black curves) and the donor dye (AF555) absorption at 555 nm or donor dye (ATTO 532) absorption at 532 nm. The acceptor dye absorption (red lines) was measured at 647 nm for AF647 and 643 nm for ATTO 643.

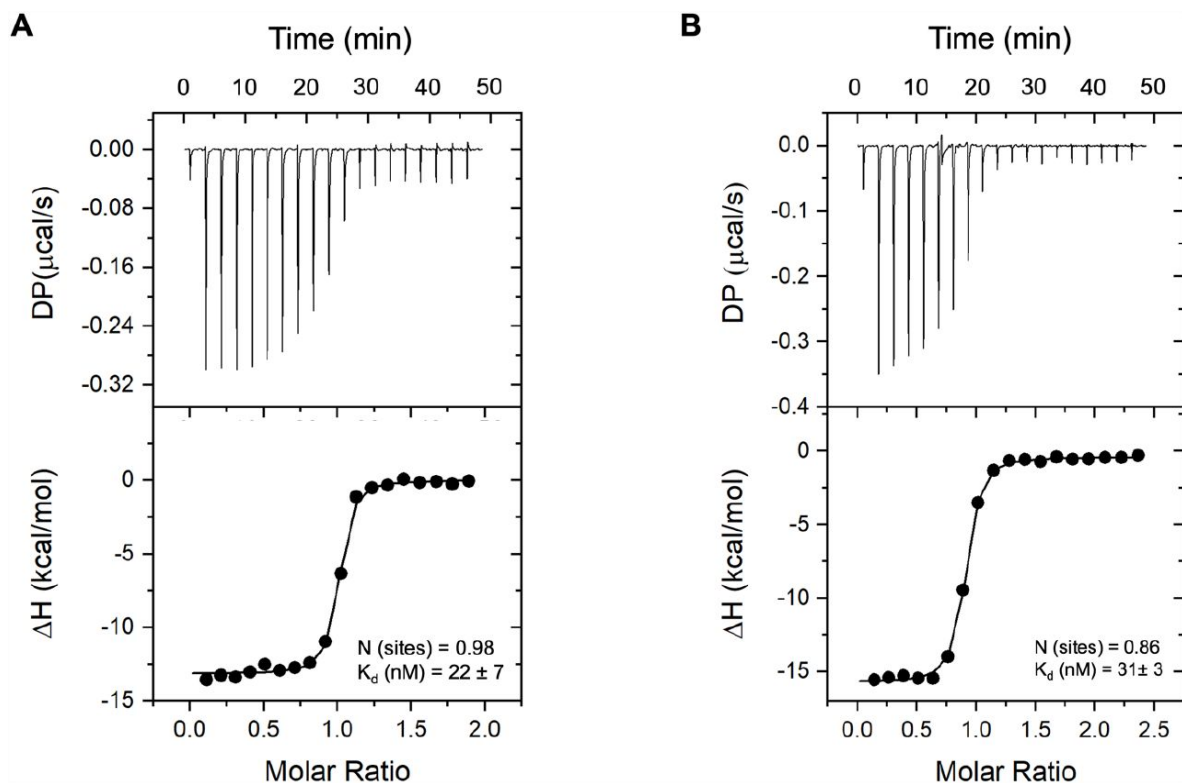


Figure S3.

Investigating binding affinity of refolded GlnBP WT and refolded GlnBP(59C-130C) using Isothermal Titration Calorimetry (ITC) measurements.

The graphs depict the changes in heat (DP, top) and enthalpy (ΔH , bottom), due to each injection of L-glutamine into the sample cell, as function of time (top x-axis of each graph) and molar ratio of refolded protein and ligand (bottom x-axis), separately. All ITC experiments were repeated three times and performed without fluorophore labeling.

(A) The mean binding affinity of the refolded GlnBP WT is 22 ± 7 nM and the binding stoichiometry is close to 1. (B) The mean binding affinity of the refolded GlnBP(59C-130C) is 31 ± 3 nM and the binding stoichiometry is close to 1.

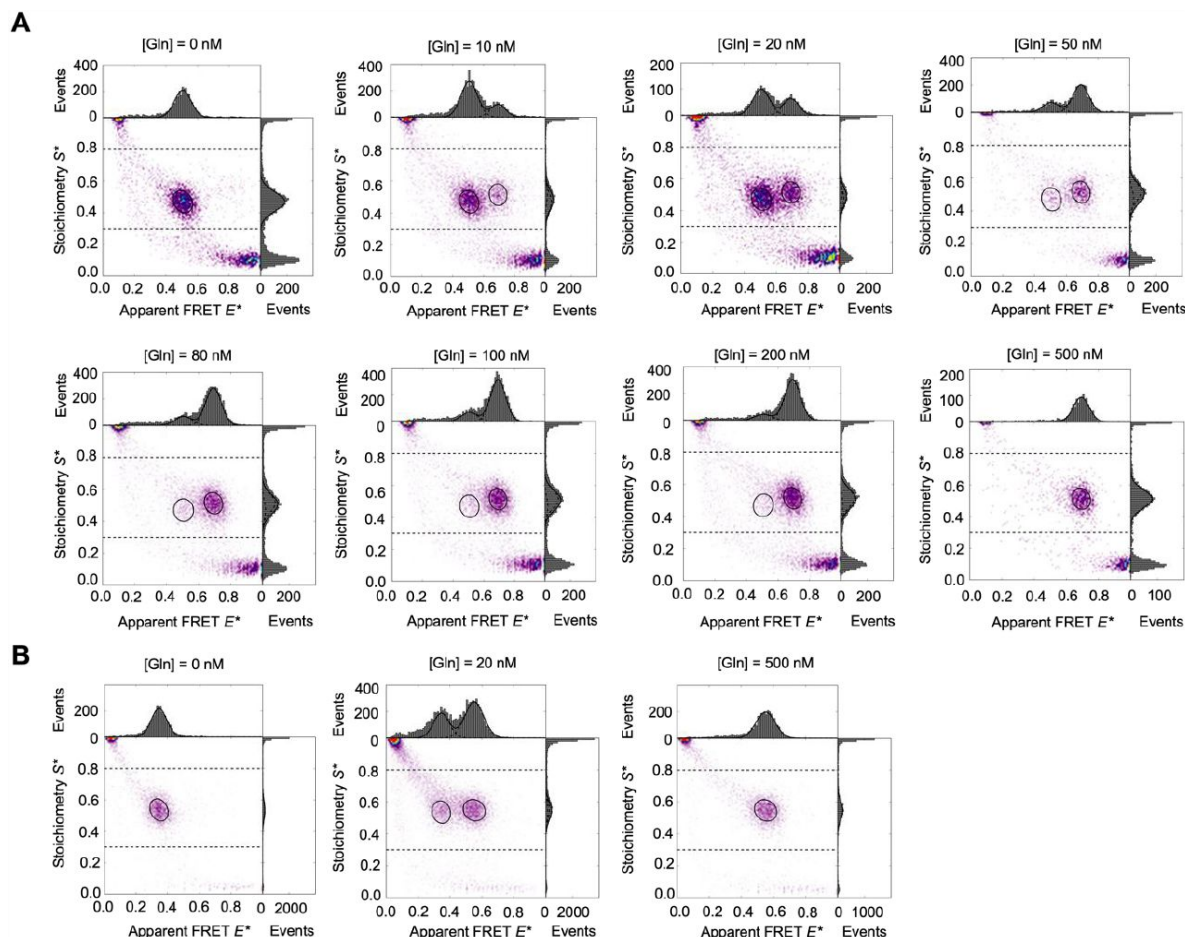


Figure S4.

L-glutamine-induced conformational changes in refolded GlnBP(111C-192C) visualized by μ sALEX measurements.

μ sALEX-based E^* - S^* histograms of the refolded GlnBP(111C-192C) double-cysteine mutants labeled with AF555/AF647 fluorophore pair (A) and labeled with ATTO 532/ATTO 643 fluorophore pair (B). First, the histograms of the apo (no L-glutamine) and holo (500 nM L-glutamine) states of the protein were fitted using a 2D gaussian distribution. Subsequently, these two distributions with variable amplitude were used to fit the intermediate ligand concentrations. Refolded GlnBP(111C-192C) labeled with AF555/AF647 shows an open state at $E^* = 0.507$ and a closed high-FRET state at $E^* = 0.694$ in the presence of a saturating concentration of L-glutamine. Refolded GlnBP(111C-192C) labeled with ATTO 532/ATTO 643 shows an open state at $E^* = 0.346$ and a closed high-FRET state at $E^* = 0.552$ in the presence of a saturating concentration of L-glutamine.

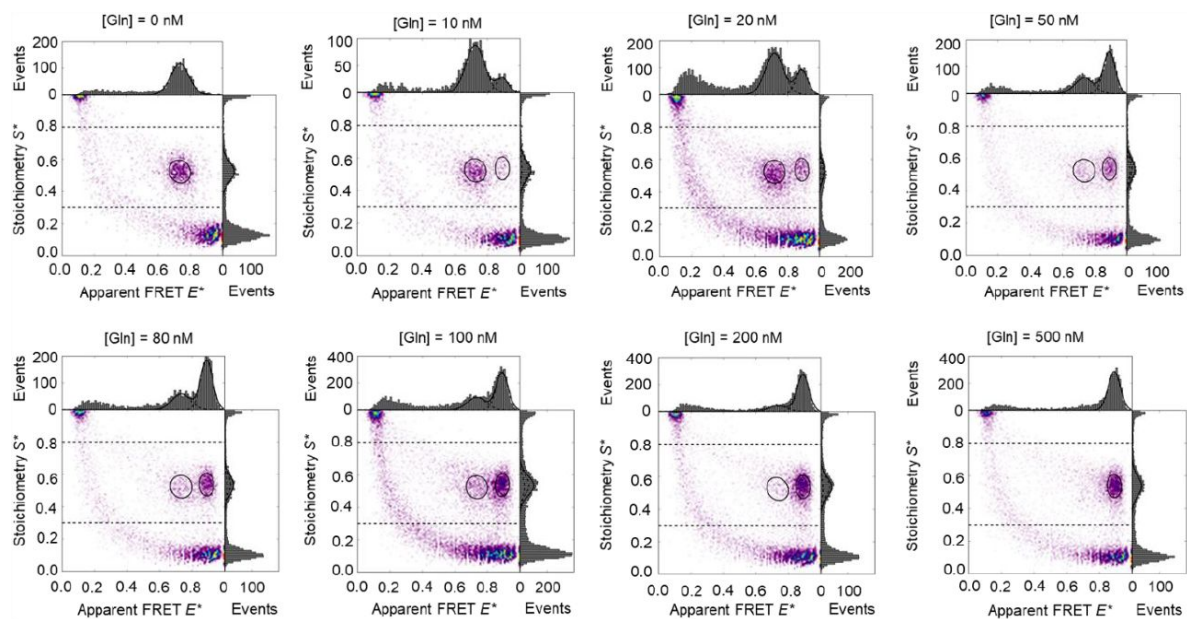


Figure S5.

L-glutamine-induced conformational changes in refolded GlnBP(59C-130C) visualized by μ sALEX measurements.

μ sALEX-based E^* - S^* histograms of the refolded GlnBP(59C-130C) double-cysteine mutants labeled with AF555/AF647 fluorophore pair. First, the histograms of the apo (no L-glutamine) and holo (500 nM L-glutamine) states of the protein were fitted using a 2D gaussian distribution. Subsequently, these two distributions were used to fit the intermediate ligand concentrations. Refolded GlnBP(59C-130C) shows an open state at $E^* = 0.735$ and a closed high-FRET state at $E^* = 0.891$ in the presence of a saturating concentration of L-glutamine.

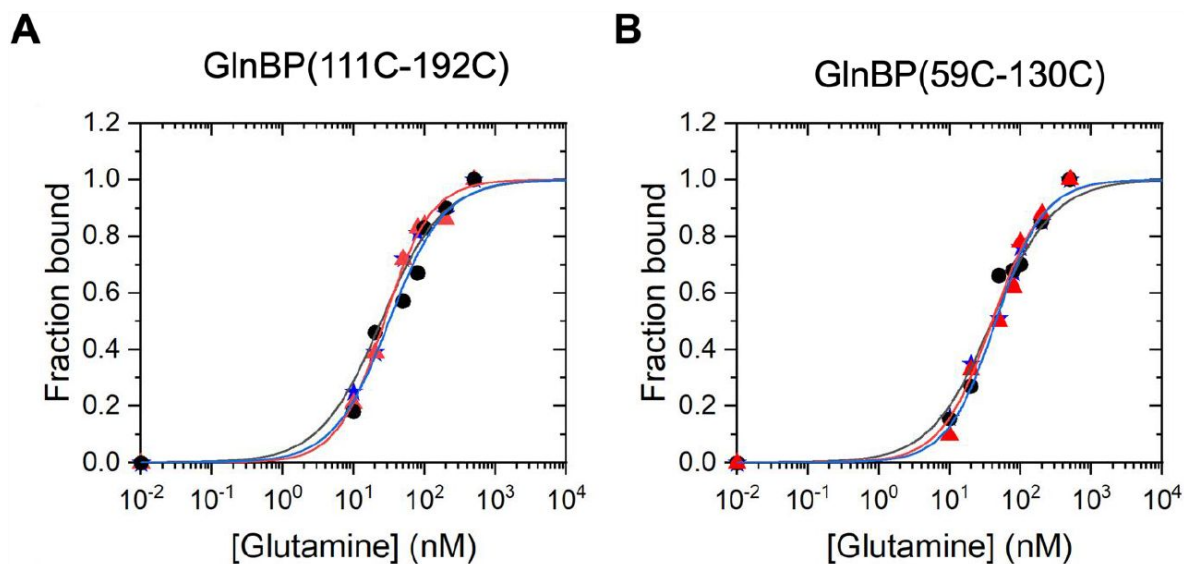


Figure S6.

Investigating binding affinities of fluorescently labeled GlnBP variants using smFRET measurements.

Binding curves of GlnBP in semilogarithmic fashion of [Gln] vs. bound fraction of protein from μ sALEX experiments using the AF555/AF647 dye pair (see [Figure S3A](#) and [Figure S4](#)). The fraction closed, i.e., the fraction of liganded protein, was determined from the ratio of the area of the high-efficiency peak and the total peak area from the projections in the apparent FRET efficiency. The fraction bound as a function of ligand (L-glutamine) concentration was fitted with the Hill equation using Origin 2016 (Origin Lab Corp, Northampton, MA), with the maximum number of binding sites fixed to 1. All the measurements were repeated three times.

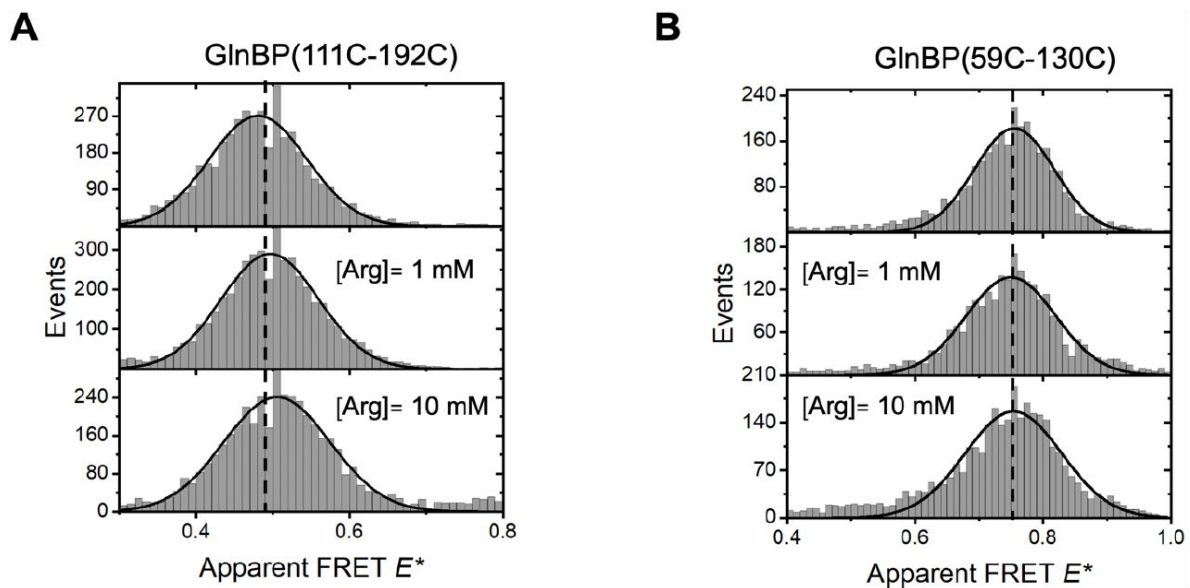


Figure S7.

Conformational states of refolded GlnBP variants probed by solution-based μ sALEX measurements reveal nearly unchanged conformations.

(A) Apparent FRET efficiency histograms of refolded GlnBP(111C-192C) labelled with AF555/647 in the absence (first row) and presence of L-arginine. (B) Apparent FRET efficiency histograms of refolded GlnBP(59C-130C) labelled with AF555/AF647 in the absence (first row) and presence of L-arginine.

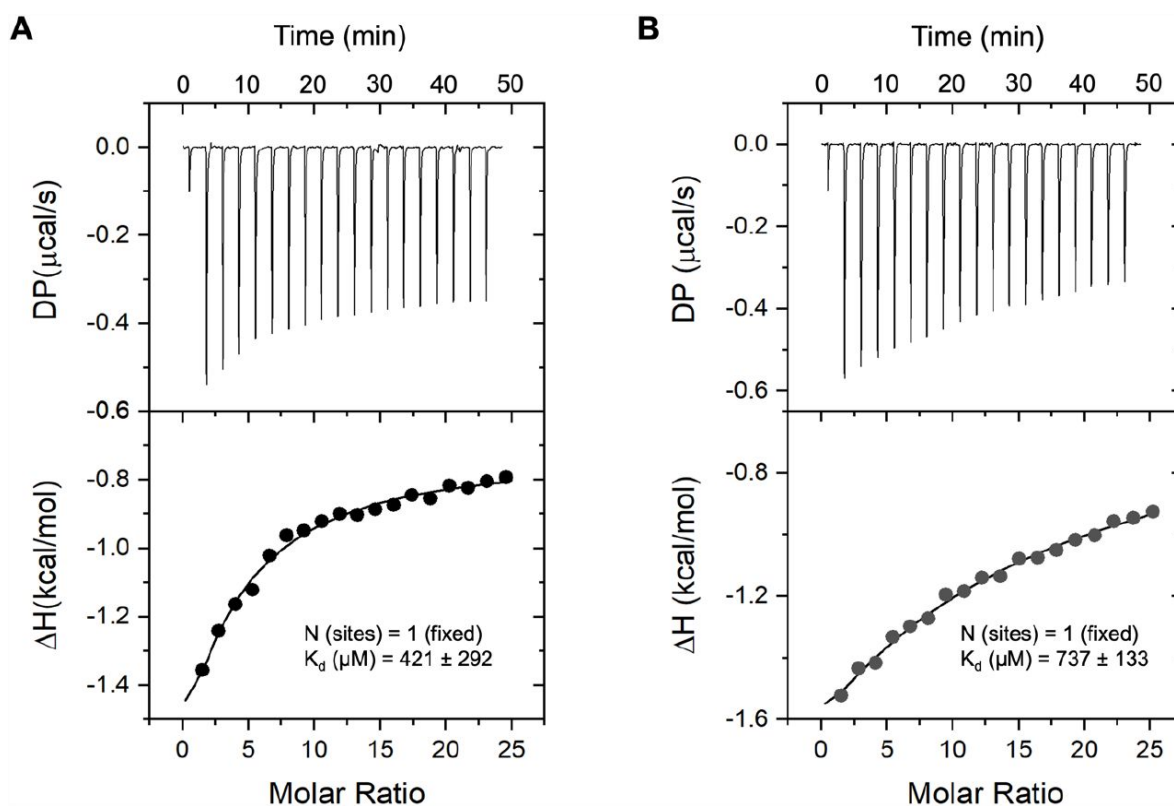


Figure S8.

Investigating L-Arginine binding affinity of refolded GlnBP(111C-192C) and GlnBP(59C-130C) variants using Isothermal Titration Calorimetry (ITC) measurements.

The graphs depict the changes in heat and enthalpy with the injection of the L-Arginine against the time and molar ratio of refolded protein and ligand, separately. All ITC experiments were repeated three times and performed without fluorophore labeling. (A) The average binding affinity of the refolded GlnBP(111C-192C) is $421 \pm 292 \mu\text{M}$. (B) The average binding affinity of the refolded GlnBP(59C-130C) is $737 \pm 133 \mu\text{M}$. The binding ratio (sites) was manually fixed to $N = 1$.

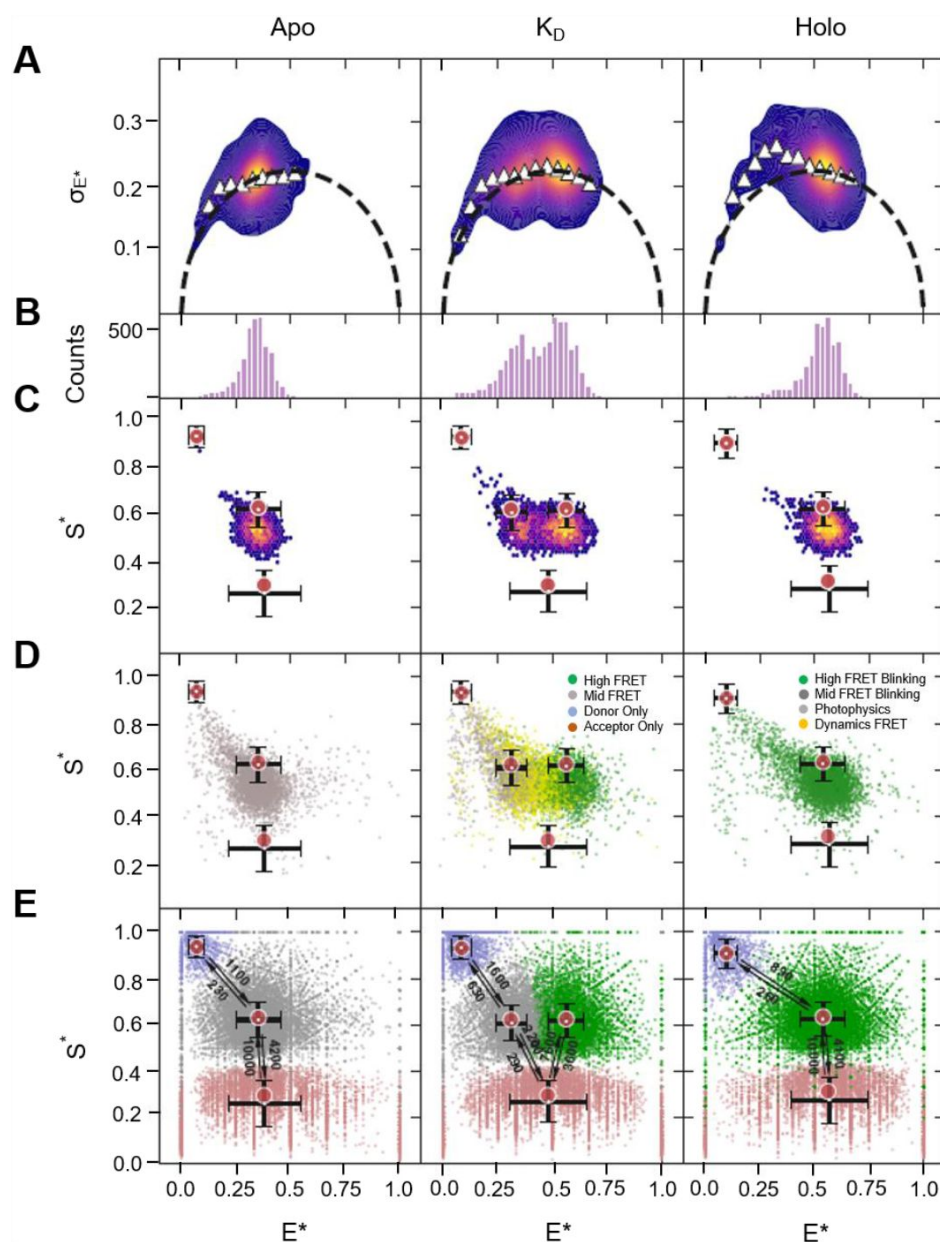


Figure S9.

Screening GlnBP(111C-192C) for rapid within-burst FRET dynamics.

Confocal-based single-molecule FRET results for GlnBP(111C-192C) doubly-labeled with ATTO 532 and ATTO 643, in the apo state (left panels), near the K_D (middle), and in the holo state (right). (A) Burst Variance Analysis (BVA) showing a weak signature of within-burst FRET dynamics in the low E^* regime. (B) Histograms of E^* values of bursts, (C) E^* versus S^* 2D histograms of bursts, (D) 2D scatter plots of bursts classified by mpH²MM, with colors corresponding to which state(s) are present within the bursts as determined with the Viterbi algorithm. Locations of states are given by red circles, and black crosses represent the SD of E^* and S^* values of dwells within each state. (E) E^* versus S^* 2D scatter plots of dwells in mpH²MM-detected states within bursts detected by the Viterbi algorithm. Red circles and black crosses are same as in (D). Arrows and adjacent numbers indicate transition rates in s^{-1} units. Transitions with rates less than $100 s^{-1}$ are omitted, since such slow transitions are improbable to occur within single-molecule bursts with durations shorter than 10 ms and are most probably a mathematical outcome of the mpH²MM framework. The dispersion of the E^* and S^* values of dwells in mpH²MM-detected states are due to the short dwell times in these states, where the shorter the dwell time in a state is, the lower the number of photons it will include, and hence the larger the uncertainty will be in the calculation of E^* and S^* values of dwells. E^* and S^* are E^* and S^* values uncorrected for background, since in mpH²MM all photons within bursts are taken into account, including ones that might be due to background.

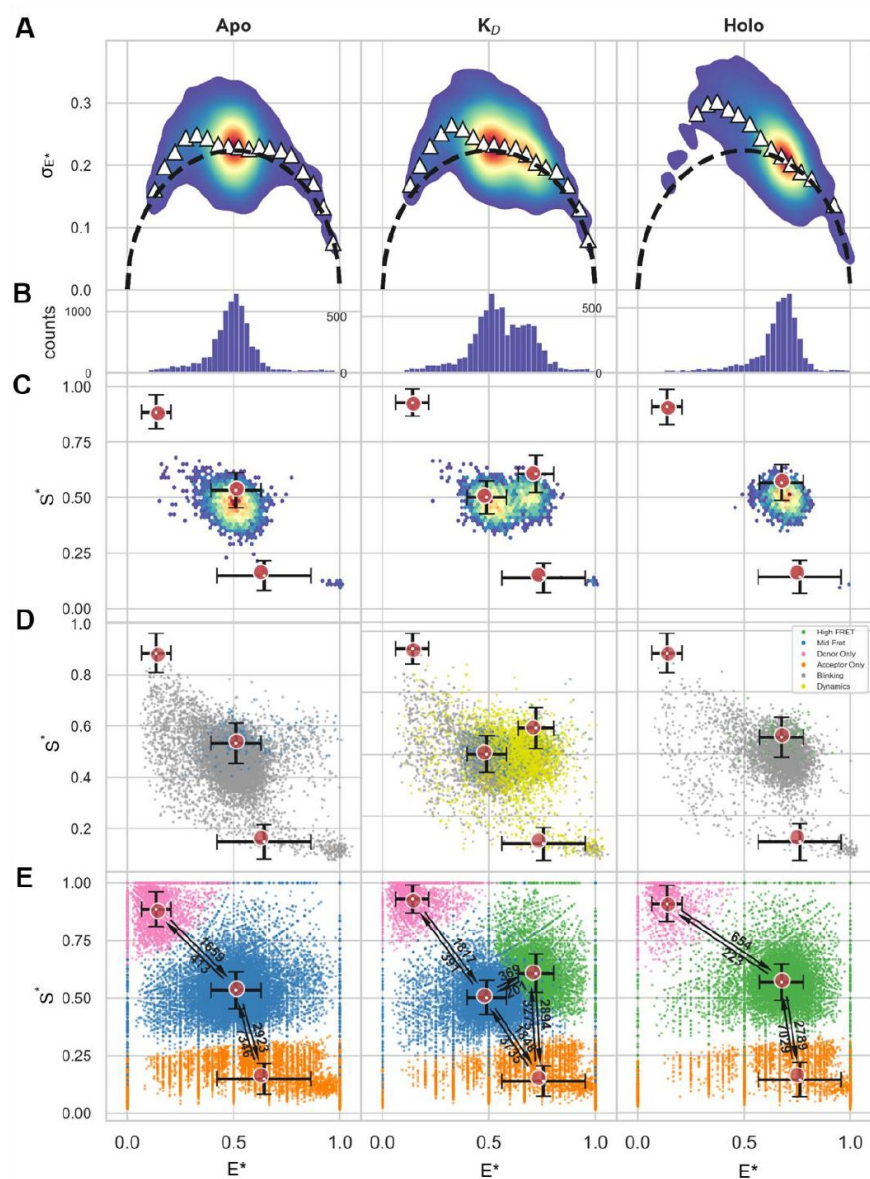


Figure S10.

Screening GlnBP(111C-192C) for rapid within-burst FRET dynamics.

Confocal-based single-molecule FRET results for GlnBP doubly-labeled at residues 111 and 192 with AF555 and AF647, in the apo state, near the K_D , and holo state. (A) Burst variance analysis showing a weak signature of within-burst FRET dynamics. (B) Histograms of E^* values of bursts, (C) E^* versus S^* 2D histograms of bursts, (D) 2D scatter plots of bursts classified by mpH²MM, with colors corresponding to which state(s) are present within the burst as determined by the Viterbi algorithm. Locations of states are given by red circles, and black crosses represent the SD of E^* and S^* values of dwells within each state. (E) E^* versus S^* 2D scatter plots of dwells in mpH²MM-detected states within bursts detected by the Viterbi algorithm. Red circles and black crosses are same as in (D). Arrows and adjacent numbers indicate transition rates in s^{-1} units. Transitions with rates less than $100 s^{-1}$ are omitted, since such slow transitions are improbable to occur within single-molecule bursts with durations shorter than 10 ms and are most probably a mathematical outcome of the mpH²MM framework. The dispersion of the E^* and S^* values of dwells in mpH²MM-detected states are due to the short dwell times in these states, where the shorter the dwell time in a state is, the lower the number of photons it will include, and hence the larger the uncertainty will be in the calculation of E^* and S^* values of dwells. E^* and S^* are E^* and S^* values uncorrected for background, since in mpH²MM all photons within bursts are taken into account, including ones that might be due to background.

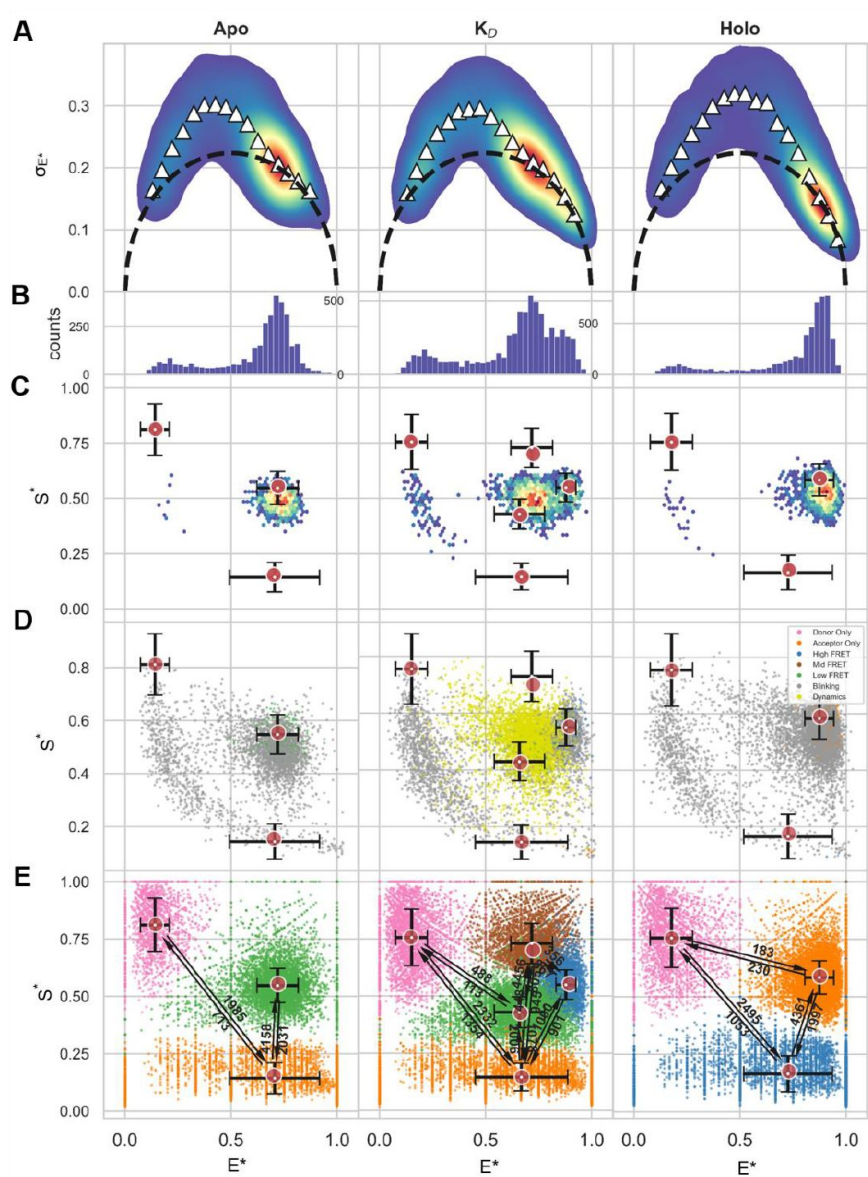


Figure 11.

Screening GlnBP(59C-130C) for rapid within-burst FRET dynamics.

Confocal-based single-molecule FRET results for GlnBP doubly-labeled at residues 59 and 130 with AF555 and AF647, in the apo state, near the K_D , and holo state. (A) Burst variance analysis showing a weak signature of within-burst FRET dynamics. (B) Histograms of E^* values of bursts, (C) E^* versus S^* 2D histograms of bursts, (D) 2D scatter plots of bursts classified by mpH2MM, with colors corresponding to which state(s) are present within the burst as determined with the Viterbi algorithm. Locations of states are given by red circles, and black crosses represent the SD of E^* and S^* values of dwells within each state. (E) E^* versus S^* 2D scatter plots of dwells in mpH2MM-detected states within bursts detected by the Viterbi algorithm. Red circles and black crosses are same as in (D). Arrows and adjacent numbers indicate transition rates in s^{-1} units. Transitions with rates less than $100 s^{-1}$ are omitted, since such slow transitions are improbable to occur within single-molecule bursts with durations shorter than 10 ms and are most probably a mathematical outcome of the mpH2MM framework. The dispersion of the E^* and S^* values of dwells in mpH2MM-detected states are due to the short dwell times in these states, where the shorter the dwell time in a state is, the lower the number of photons it will include, and hence the larger the uncertainty will be in the calculation of E^* and S^* values of dwells. E^* and S^* are E^* and S^* values uncorrected for background, since in mpH2MM all photons within bursts are taken into account, including ones that might be due to background.

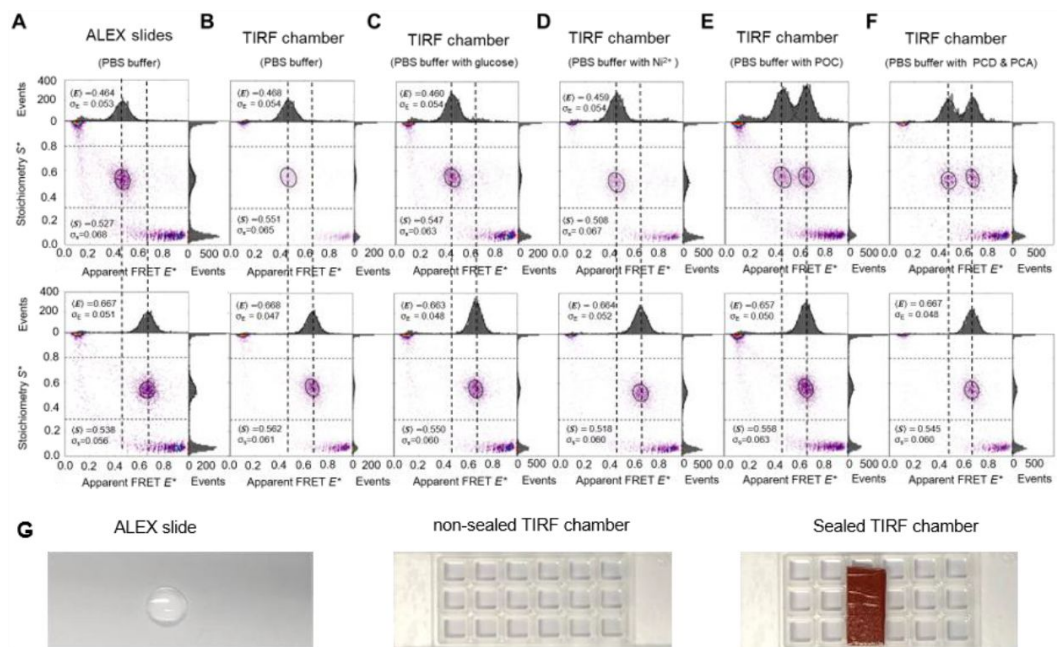


Figure S12.

Effects on conformation of GlnBP(111C-192C) under various conditions.

Due to the high binding affinity of GlnBP for L-glutamine, several control experiments under different conditions were performed to exclude artifacts induced by the reagents present in each set of experiments. The μ ALEX experiments of the refolded GlnBP(111C-192C) double-cysteine variant labeled with LD555/LD655 fluorophore pairs were measured in PBS buffer (pH 7.4) using conventional microscope glass slides (A) and using TIRF chamber (B). The PBS buffer containing (C) 40 mM glucose, (D) 50 nM Ni^{2+} , (E) pyranose oxidase/catalase (POC) and (F) protocatechuate-dioxygenase (PCD)/3,4-protocatechuic acid (PCA) was used for the ALEX measurements. (G) The conventional glass coverslips used in μ ALEX experiments (top figure) and TIRF chambers (sticky-Slide 18 well, Ibidi; non-sealed chambers: middle panel; sealed: bottom panel) glued on top of PEG-/biotin-PEG-silane microscope glass coverslips used in the TIRF experiments.

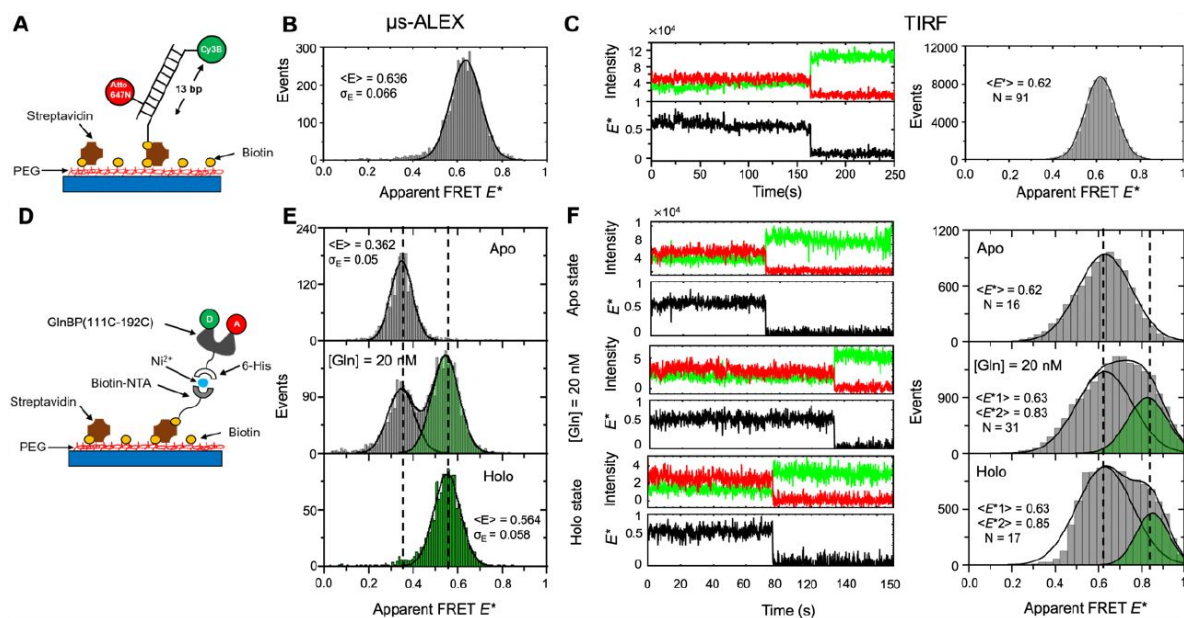


Figure S13.

Comparing smFRET measurements of biotin-modified dsDNA and GlnBP(111C-192C) using diffusion-based μ sALEX versus TIRF microscopy.

(A) Schematic view of dsDNA labeled with Cy3B and ATTO 647N for smFRET characterization on PEGylated coverslips. (B) Typical μ sALEX-based E^* - S^* histograms of the biotin-modified dsDNA labeled with Cy3B and ATTO 647N. (C) Representative fluorescence time trace of respective single emitter of the biotin-modified dsDNA sample under continuous wave excitation of $\sim 500 \mu\text{W}$ at 532 nm and the FRET histograms of all analyzed molecules and the FRET histograms of all measured molecules combined. (D) Schematic view of the refolded GlnBP(111C-192C) labeled with ATTO 532 and ATTO 643 for smFRET characterization. (E) Typical μ sALEX-based E^* - S^* histograms of the refolded GlnBP(111C-192C). (F) Representative fluorescence time trace of respective single emitter of the refolded GlnBP(111C-192C) under continuous wave excitation of $\sim 500 \mu\text{W}$ at 532 nm and the FRET histograms of all analyzed molecules. Additional data for each condition is shown in [Figures S13](#) and [S14](#).

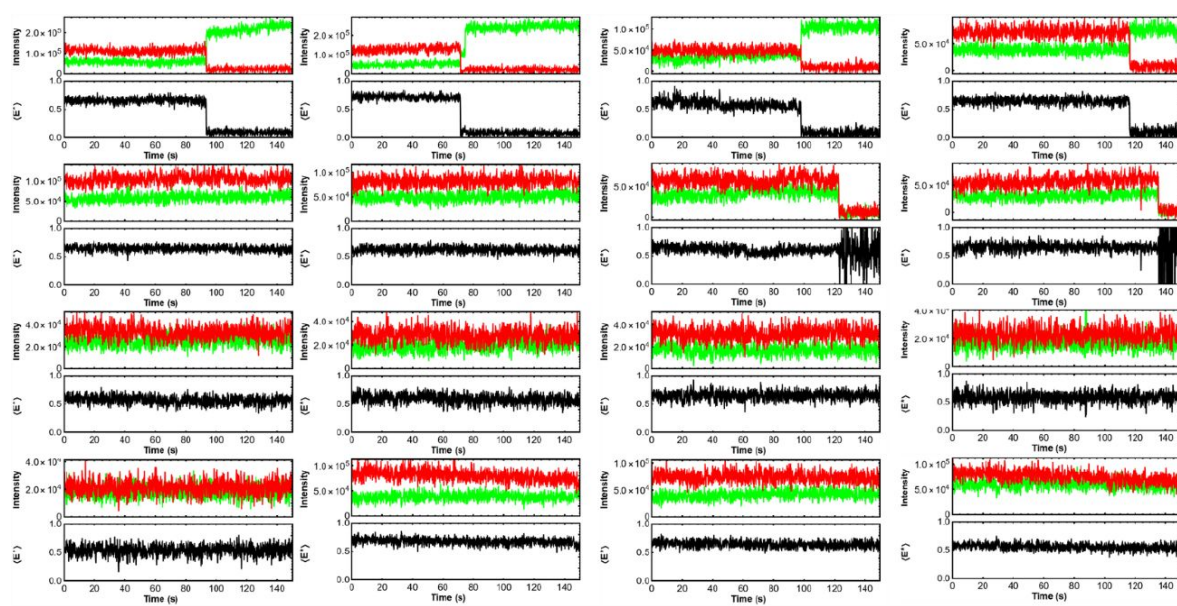


Figure S14.

Representative fluorescence time traces of respective single emitter of biotin-functionalized DNA labeled by maleimide-modified derivatives Cy3B and ATTO 647N (13 bp inter-dye distance). All measurements were done in oxygen scavenging buffer (3 U/mL of pyranose oxidase, 90 U/mL of catalase and 40 mM glucose, PBS buffer, pH 7.4). Laser power: 500 μ W.

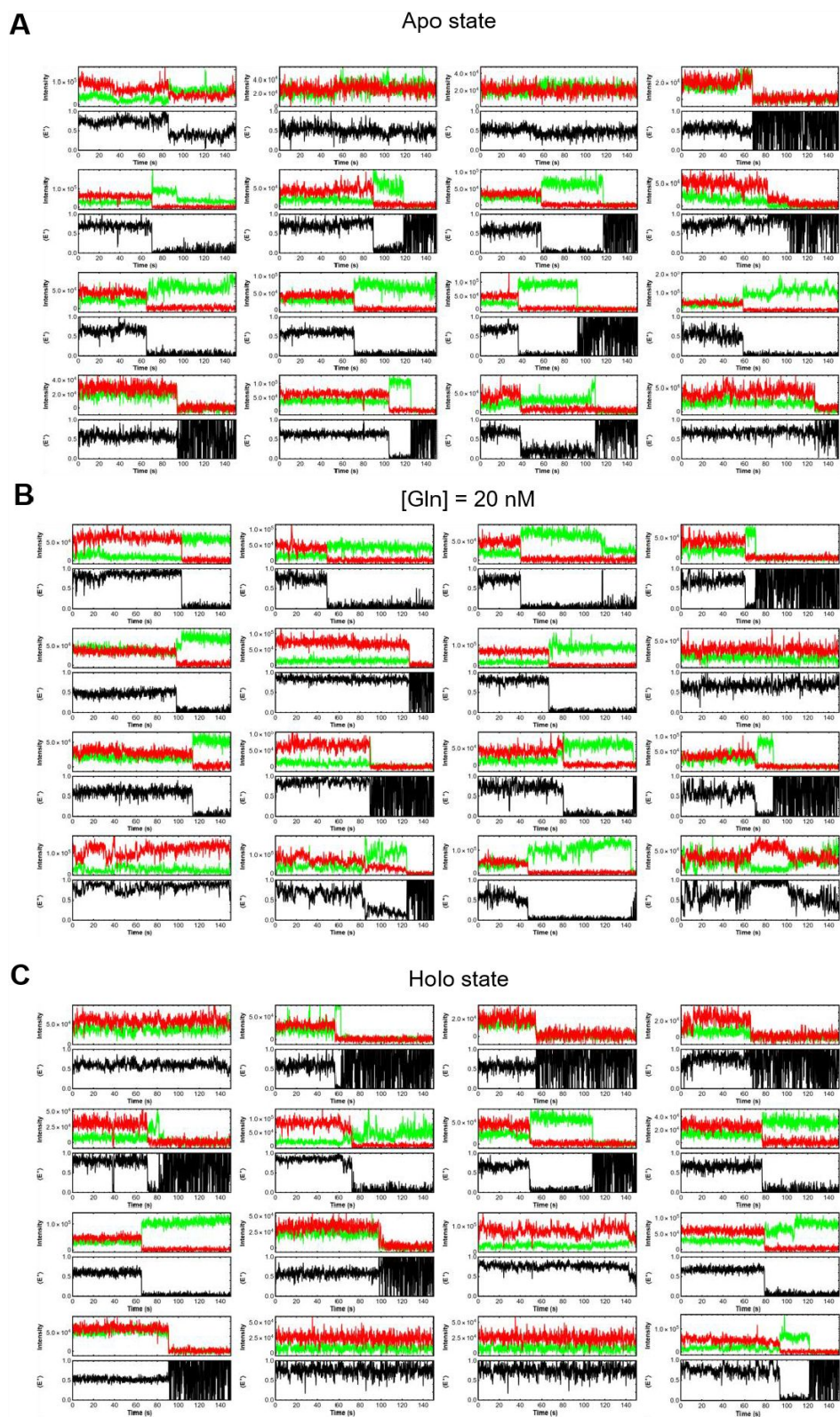


Figure S15.

Examples of fluorescence time traces of respective single emitter of refolded GlnBP(111C-192C) labeled by maleimide-modified derivatives ATTO 532 and ATTO 643. All measurements were done in PBS buffer, pH 7.4 and 2 mM Trolox. Laser power with continuous 532 nm excitation: 200 μ W.

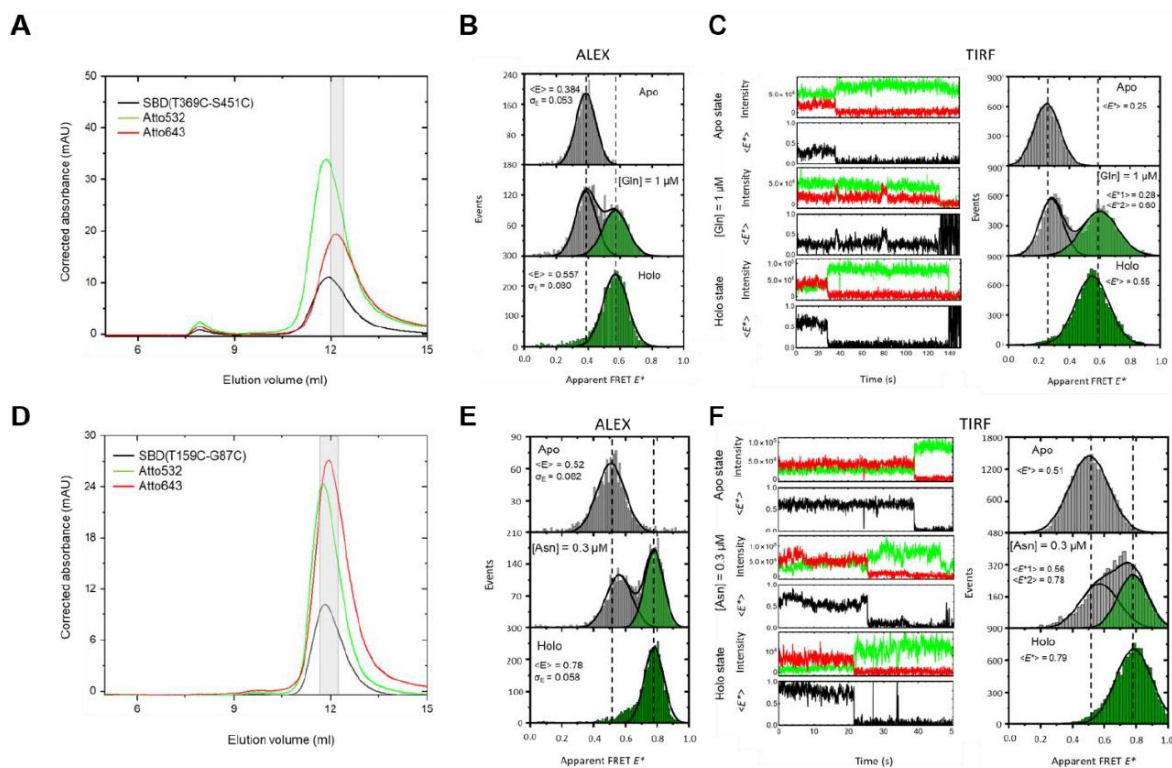


Figure S16.

(A) and (D) Size Exclusion Chromatography (SEC) of SBD(T369C-S451C) and SBD(T159C-G87C). The selected fractions (grey-shaded area) were collected and used for the solution-based smFRET measurements. The selected fractions (grey-shaded area) having the best overlap of protein, donor, and acceptor absorption were used. The protein absorption was measured at 280 nm (black curves) and the donor dye (ATTO 532) absorption at 532 nm. The acceptor dye absorption (red lines) was measured at 643 nm for ATTO 643. (B) and (E) Typical μ ALEX-based E^*-S^* histograms of the SBD(T369C-S451C) and SBD(T159C-G87C). (C) and (F) Representative fluorescence time trace of respective single emitter of the SBD(T369C-S451C) and SBD(T159C-G87C) and the FRET histograms of all measured molecules.

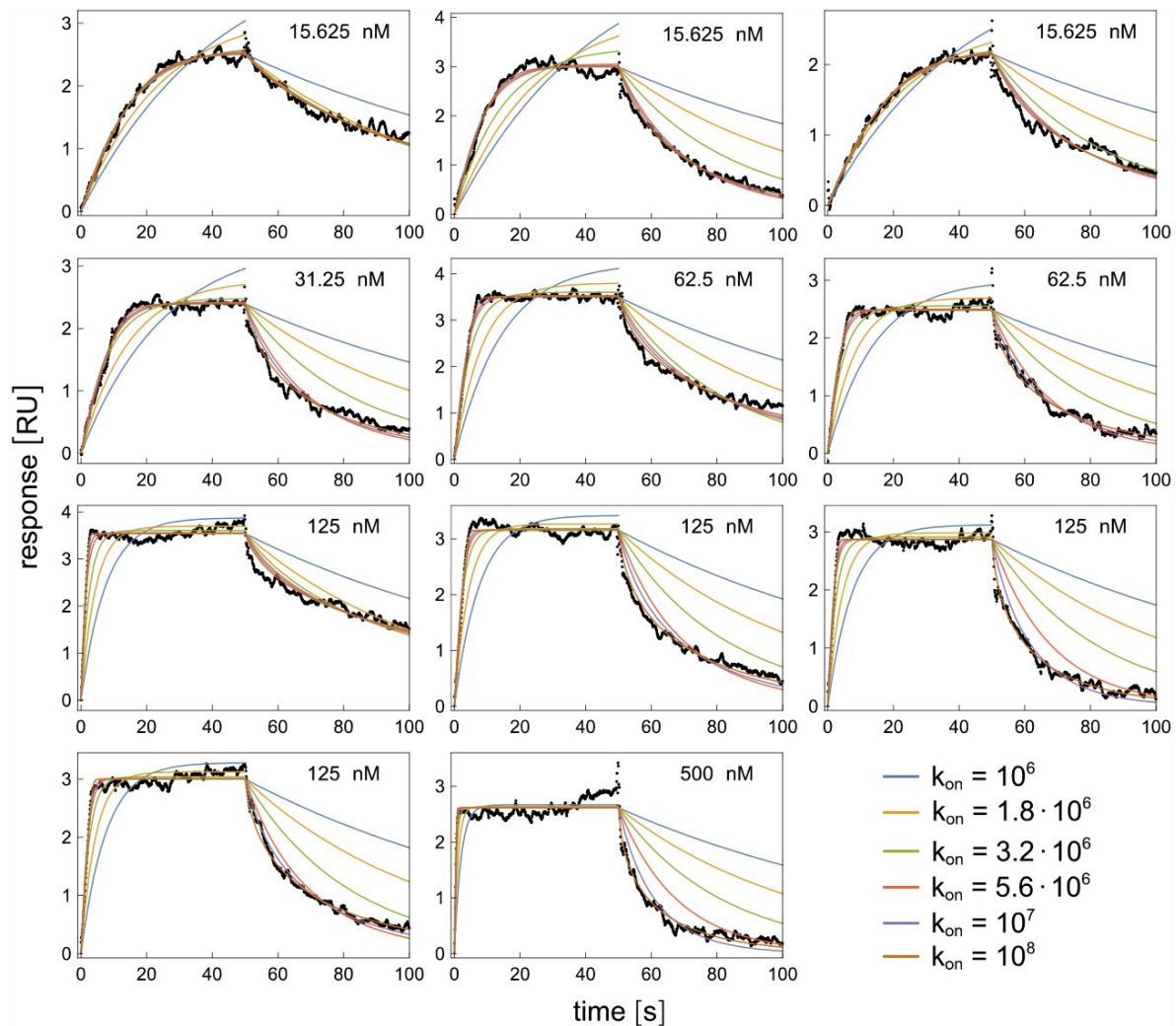


Figure S17.

SPR sensorgrams at the indicated glutamine concentrations, and fits of the sensorgrams for different values of the effective on-rate constant, k_{on} , as in [Figure 6B](#) [C](#). The rescaled sum of squared residuals versus k_{on} for these fits is shown by dashed lines in [Figure 6D](#) [C](#).

Supplementary Notes 1-4

Supplementary Note 1: Interpretation of mpH²MM analysis

For analysis of within burst dynamics, we used multi-parameter photon-by-photon hidden Markov modelling (mpH²MM) [2](#), [3](#) to identify the most-likely state model that describes the experimental results based on how E* and S* values may change within single-molecule bursts. For this analysis we (i) report the most-likely number of states and their mean E* and S* values (**Figure 4B** [2](#), red dots). (ii) We investigate whether molecules traversing the confocal excitation volume are fully static and only in the mid-FRET state or high-FRET state, or whether they undergo dynamic FRET changes including transitions of mid/high-FRET states with photo-blinking dynamics or dark donor or acceptor states (**Figure 4B** [2](#)). (iii) We finally report on E* and S* values for parts of bursts with dwells in one of the identified states and the rate constants of transitioning between them (**Figure 4B** [2](#)). These analyses confirm that among the two types of dynamic transitions that influence the burst-based E* and S* values, these are mostly donor or acceptor photo-blinking dynamics between bright and dark states of the fluorophores. Such behavior is irrelevant to understanding the conformational changes in GlnBP but does influence the mean FRET efficiency values if not decoupled. Importantly, no dynamic transitions occur between the mid-FRET and high-FRET states at timescales shorter than 10 ms (i.e., with rate constants higher than 100 s⁻¹). All measurement conditions show significant photo-blinking dynamics which occur mostly on few ms to sub-millisecond timescales most prominently for the use of AF555/AF647 and the GlnBP(59/130) variant (compare **Figure 4** [2](#) and **Figure S9** [2](#)-**S11** [2](#)). Therefore, the blinking dynamics likely account also for the signature of within-burst dynamics shown by BVA (**Figure 4** [2](#), **Figure S9** [2](#)-**S11** [2](#)).

Most importantly, mpH²MM identifies single apo and holo E*-states, which describe the open mid-FRET and closed high-FRET conformations of GlnBP. Only in the presence of low (near K_D) concentrations of glutamine two FRET states are identified which interconvert on timescales slower than 10 ms. Notably, the mean E* and S* values of the FRET states are slightly dissimilar to the centers of the burst-based E* and S* populations, owing to the effect of the rapid photo-blinking dynamics within bursts, which lead to averaging the E* and S* values of the FRET states with those of the photo-blinked states. Additionally, in the presence of near-K_D concentrations of glutamine, the FRET dynamics occur in the few ms timescale or even slower, which may contribute only slightly to the signature of FRET dynamics in BVA. In conclusion, if intrinsic conformational dynamics existed in apo GlnBP, it could only be between the highly-populated FRET conformation we identify and another conformation that is populated way below the sensitivity of our measurement and analysis (potentially <5-10% populations). Thus, we can conclude that the majority of the conformational dynamics in GlnBP is induced by glutamine, most probably as a result of its binding to GlnBP.

Supplementary Note 2: Description of TIRF data acquisition and analysis.

At first, we studied a biotin-modified double-stranded DNA (dsDNA), which was labeled with Cy3B (donor) and ATTO 647N (acceptor) in 13 bp distance, and used this as a reference sample to allow a direct comparison of μ sALEX and TIRF data (**Figure S13** [2](#)). For this, we immobilized the dsDNA on a PEG-coated glass surface via streptavidin-biotin interactions. We recorded both donor and acceptor fluorescence via a dual-view split on our EMCCD camera with 100 ms integration time per frame. With this we obtained traces that lasted multiple 10s periods. Since we did not perform ms alternation of green-and-red laser excitation, we verified that the sum-signal of the donor and acceptor channel was constant as a function of time for each molecule and discarded traces that

did not obey this condition. The dsDNA sample displays an apparent FRET efficiency E^* of ~ 0.64 for in-solution measurements, which agreed well with the analysis of surface-immobilized molecules on the TIRF microscope having a mean E^* of 0.62 (**Figure S13A** [B](#)).

Then, we investigated the conformational states and changes of GlnBP(111C-192C) with the dye pair ATTO 532/ATTO 643, since these showed least photophysical FRET-dynamics (see main text and Supplementary Note 1). To exclude the influence of buffer and other small molecules in TIRF measurements on the conformational state of GlnBP, we initially performed control experiments in μ sALEX (**Figure S12** [B](#)). We found that GlnBP was influenced by the addition of oxygen scavenger cocktails (pyranose oxidase and catalase, POC, and glucose or protocatechuate-dioxygenase, PCD, and 3,4-protocatechuic acid, PCA), resulting in the formation of artificial holo-state GlnBP molecules (**Figure S12E** [F](#)). In TIRF experiments, the effect of oxygen scavenger might have been misinterpreted as intrinsic closing. We consequently proceeded with no oxygen-removal in PBS buffer (pH 7.4) and with 2 mM Trolox as photostabilizer. GlnBP was immobilized by biotin-NTA interactions mediated by Nickel(II). To our surprise we found very different mean E^* values on TIRF in comparison to μ sALEX measurements (**Figure S13E** [F](#)). In detail, the mean E^* values were much higher on TIRF than on μ sALEX (**Figure S13E** [F](#)) in contrast to dsDNA (**Figure S13A** [B](#)). This can be interpreted as an altered conformational state of GlnBP, e.g., likely caused by protein-glass interactions due to surface-immobilization or interaction of the protein or dyes with the biotin-NTA moiety. Furthermore, addition of saturating glutamine concentrations did not show the expected behavior of a full shift of the population to a higher-FRET state (**Figure S13F** [B](#)). Instead, only a small fraction of the population is shifted for both low and saturating glutamine concentrations. At concentrations of glutamine around the K_d -value freely-diffusing GlnBP shows a mix of open- and closed state in μ sALEX experiments (**Figure 3** [B](#)). In TIRF, however, we could not identify dynamic transitions (**Figure S13** [B](#)/15). This finding indicates that a part of the immobilized fluorophore-labelled GlnBP becomes non-functional. Since our protocol deviates from that used in other studies⁴ [B](#)-⁶ [B](#), we probed whether we could reproduce published data on substrate-binding domain 1 and 2 (SBD1 and SBD2)⁷ [B](#). Again, we find a good match between biochemical properties, μ sALEX and the corresponding TIRF data for both proteins (**Figure S16** [B](#)).

Supplementary Note 3: Compatibility of IF pathway with smFRET and SPR results and estimates of kinetic rate constants.

For ligand concentrations $[L] < K_d$ and conformational excitation rates k_e that are much smaller than the relaxation rates k_r , the intermediate ligand-bound open state OL of the IF pathway has a significantly lower probability than the other two states, akin to a transition state. In this case, the relaxation rate $k_{\text{obs}}^{\text{IF}}$ of the IF pathway in **Figure 7A** [B](#) can be approximated by the relaxation rate of an effective two-state process⁸ [B](#)

$$k_{\text{obs}}^{\text{IF}} \approx k_{\text{on}}[L] + k_{\text{off}} \quad (\text{S1})$$

with effective on- and off-rate constants $k_{\text{on}} = k_+k_r/(k_- + k_r)$ and $k_{\text{off}} = k_-k_e/(k_- + k_r)$. From the equation for the effective off-rate constant k_{off} we obtain

$$k_- = k_{\text{off}}k_r/(k_e - k_{\text{off}}) \quad (\text{S2})$$

which implies

$$k_{\text{off}} < k_e \quad (\text{S3})$$

From our SPR results in **Figure 6** [B](#), we concluded a lower bound of $3 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ for k_{on} in a range of ligand concentrations $[L]$ from 15.6 to 500 nM, which likely holds also for smaller $[L]$. Based on stopped-flow mixing experiments of GlnBP and Gln more than five decades ago⁹ [B](#), an

effective on-rate constant of about $10^{8.5} \text{ M}^{-1}\text{s}^{-1}$ has been obtained from numerical fits of stopped-flow relaxation curves at concentration ratios of 1:1 and 2:1 of Gln and GlnBP. For a plausible range $3 \times 10^7 \text{ M}^{-1}\text{s}^{-1} < k_{\text{on}} < 10^{8.5} \text{ M}^{-1}\text{s}^{-1}$ of on-rate constants and K_d values of 10 – 20 nM from different methods (ITC, smFRET, SPR), we obtain $0.3 \text{ s}^{-1} < k_{\text{off}} < 2 \text{ s}^{-1}$ as range for the effective off-rate constant $k_{\text{off}} = K_d k_{\text{on}}$. Together with **Eq. (S2)**, our smFRET results with lower bounds of 100 s^{-1} for the conformational exchange rates k_e and k_r (corresponding to timescales $>10 \text{ ms}$) and an upper bound of about 10% for the relative probability $P_{\text{OL}} = k_e / (k_e + k_r)$ of conformation OL among the two bound states of GlnBP lead to

$$k_{\text{off}} = 0.3 \text{ to } 2 \text{ s}^{-1} < k_e < 9 k_r < 100 \text{ s}^{-1} \quad (\text{S4})$$

This equation shows that the IF pathway is compatible with our results. **Eq. (S4)** in turn results in a lower bound for P_{OL} of about 0.3 to 2%, and together with **Eq. (S2)**, in the lower bounds $k_{-} \approx 10 k_{\text{off}}$ for $P_{\text{OL}}=10\%$, $k_{-} \approx 20 k_{\text{off}}$ for $P_{\text{OL}}=5\%$, and $k_{-} \approx 100 k_{\text{off}}$ for $P_{\text{OL}}=2\%$. Corresponding lower bounds for the on-rate constant k_{+} of the binding-competent open conformation of the IF pathway then follow from $k_{+} = (k_{-} K_d) k_e / (k_e + k_r) = (k_{-} K_d) P_{\text{OL}}$.

Supplementary Note 4: Considerations on the accessibility of the ligand binding pocket for solvent and ligand in the closed conformation of GlnBP.

To describe the expected binding behavior of the substrate glutamine to GlnBP, we performed docking calculations of GlnBP in its open and closed conformations. The GlnBP structure that represents the open conformation is the one reported under pdb code 1GGG¹⁰. The GlnBP structure that represents the closed conformation is the one reported under pdb code 1WDN¹, with the bound ligand taken out of the file. Then, we used the 3D conformer structure of the ligand to be docked onto the structures of GlnBP. We used the SwissDock web server to perform the docking procedure^{11, 12}. The results show that (i) while glutamine can dock to many sites on GlnBP, the results that yield the lowest binding free energy are when it docks onto its cognate binding site, both in the open and closed conformation (**Figures S19/S20**). (ii) The calculated binding free energy of Gln to GlnBP in the optimized docking site leads to a dissociation constant of 20 μM in the open conformation and 230 nM in the closed conformation (**Figure S20**), about two orders of magnitude different. (iii) The higher binding free energy is due to the larger amount of GlnBP residues when the docked glutamine interacts with in the closed conformation relative to in the open conformation. (iv) The binding pocket in GlnBP seems to surround the docked glutamine from all directions (**Figure S19**), which implies that it is less probable that glutamine can access the binding pocket in the closed conformation. Instead, it is more probable that the glutamine reaches its binding site in GlnBP when it is not yet closed.

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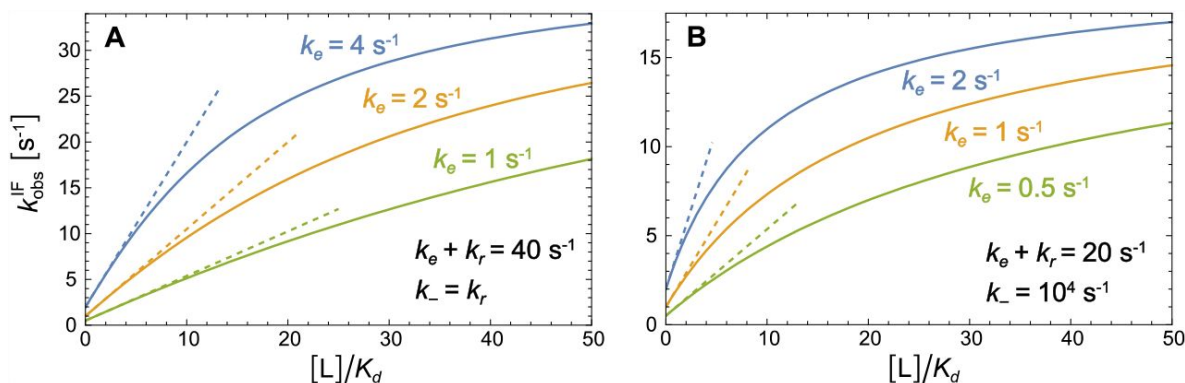


Figure S18.

Exemplary plots of the dominant relaxation rate $k_{\text{obs}}^{\text{IF}}$ versus ligand concentration $[L]$ for rate parameters consistent with Eq. (S4). Full lines represent the exact solution given in the caption of Fig. 7, and dashed lines represent the approximate solution for sufficiently small $[L]$ based on Eq. (S1). In (A), the effective off-rate resulting from the exemplary parameters is $k_{\text{off}} = 0.5 k_e$. In (B), the effective off-rate is $k_{\text{off}} \approx k_e$ because the unbinding process is dominated by the opening of the closed ligand-bound conformation with rate k_e for $k_- \gg 20 k_r$ as in this example. The limiting value of $k_{\text{obs}}^{\text{IF}}$ at large ligand concentrations is $k_e + k_r$.

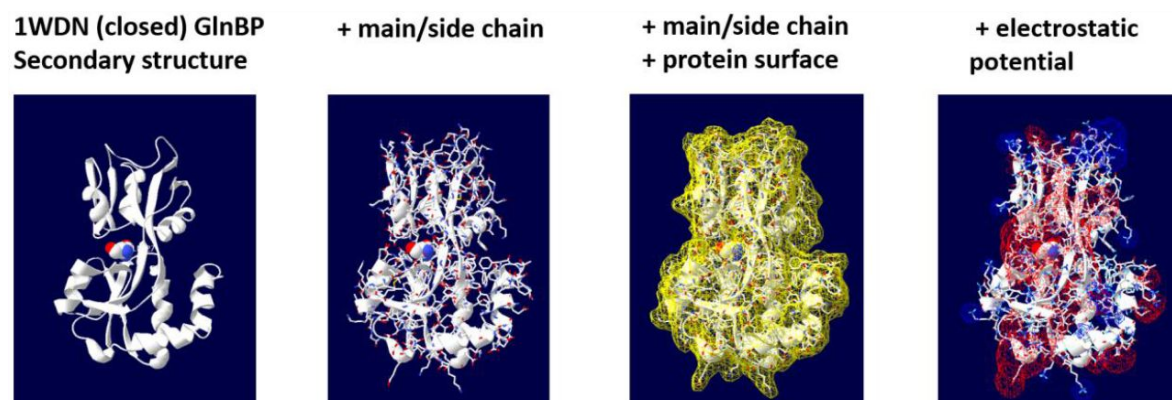


Figure S19.

The structure of holo GlnBP with optimized docking of glutamine.

The figure reports the optimized results of docking glutamine onto the crystal structure of GlnBP in holo form, after the glutamine substrate was removed from the structure, and presented back as a docking ligand using the SwissDock web server. From left to right: (i) the glutamine is docked onto the correct binding pocket within the closed conformation of GlnBP, (ii) amino acid side chains are wrapping the docked glutamine from all directions, (iii) and indeed the protein surface covers the docked glutamine, and (iv) the residues covering the docked glutamine seem to carry a net negative charge.

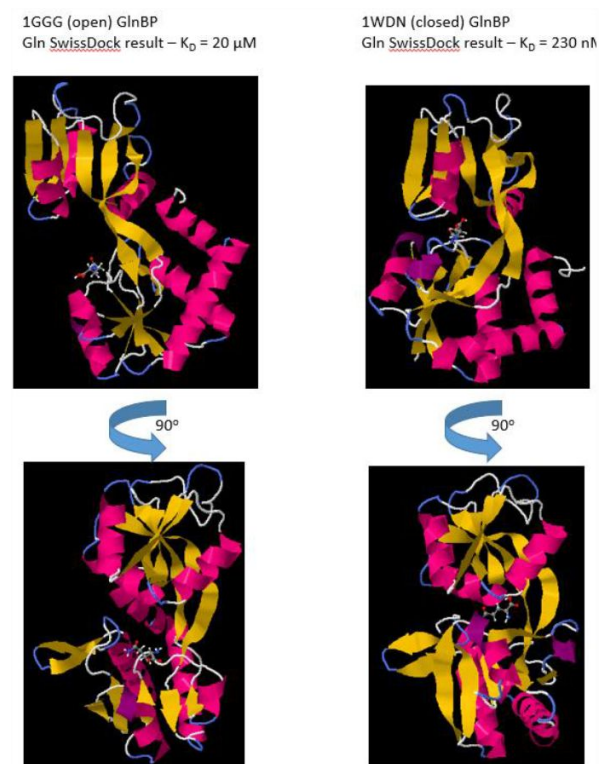


Figure S20.

Optimized docking of glutamine to GlnBP in its open and closed conformations.

Using the SwissDock web server, the molecule glutamine was docked onto the crystal structures of GlnBP in its open (pdb:1GGG) and closed (pdb:1WDN; with the glutamine substrate taken away) conformations, and the optimized docking sites as well as the calculated dissociation constant are shown (dissociation constant is calculated out of the binding energies reported in the docking results). The preferred docking of glutamine is the same site within GlnBP. The difference is that while in the open conformation glutamine binds to one domain with the other as a distant domain, in the closed conformation the other domain closes on top of the docked glutamine. Following the calculated binding energies from the optimized docking results, while the dissociation constant of glutamine to GlnBP is $20 \mu\text{M}$ in the open conformation, in the closed conformation it is 230 nM .

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Editors

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Reviewer #1 (Public Review):

Here the authors discuss mechanisms of ligand binding and conformational changes in GlnBP (a small E Coli periplasmic binding protein, which binds and carries L-glutamine to the inner membrane ATP-binding cassette (ABC) transporter). The authors have distinguished records in this area and have published seminal works. They include experimentalists and computational scientists. Accordingly, they provide comprehensive, high-quality, experimental and computational work.

They observe that apo- and holo- GlnBP does not generate detectable exchange between open and (semi-) closed conformations on timescales between 100 ns and 10 ms. Especially, the ligand binding and conformational changes in GlnBP that they observe are highly correlated. Their analysis of the results indicates a dominant induced-fit mechanism, where the ligand binds GlnBP prior to conformational rearrangements. They then suggest that an approach resembling the one they undertook can be applied to other protein systems where the coupling mechanism of conformational changes and ligand binding.

They argue that the intuitive model where ligand binding triggers a functionally relevant conformational change was challenged by structural experiments and MD simulations revealing the existence of unliganded closed or semi-closed states and their dynamic exchange with open unbound conformations, discuss alternative mechanisms that were proposed, their merits and difficulties, concluding that the findings were controversial, which, they suggest is due to insufficient availability of experimental evidence to distinguish them. As to further specific conclusions they draw from their results, they determine that a conformational selection mechanism is incompatible with their results, but induced fit is. They thus propose induced fit as the dominant pathway for GlnBP, further supported by the notion that the open conformation is much more likely to bind substrate than the closed one based on steric arguments.

Considering the landscape of substrate-free states, in my view, the closed state is likely to be the most stable and, thus most highly populated. As the authors note and I agree that state can be sterically infeasible for a deep-pocketed substrate. As indeed they also underscore, there is likely to be a range of open states. If the populations of certain states are extremely low, they may not be detected by the experimental (or computational) methods. The free energy landscape of the protein can populate all possible states, with the populations determined by their relative energies. In principle, the protein can visit all states. Whether a particular state is observed depends on the time the protein spends in that state. The frequencies, or propensities, of the visits can determine the protein function. As to a specific order of events, in my view, there isn't any. It is a matter of probabilities which depend on the populations (energies) of the states. The open conformation that is likely to bind is the most favorable, permitting substrate access, followed by minor, induced fit conformational changes. However, a key factor is the ligand concentration. Ligand binding requires overcoming barriers to sustain the equilibrium of the unliganded ensemble, thus time. If the population of the state is low, and ligand concentration is high (often the case in in vitro experiments, and high drug dosage scenarios) binding is likely to take place across a range of available states.

This is however a personal interpretation of the data. The paper here, which clearly embodies massive careful, and high-quality work, is extensive, making use of a range of experimental approaches, including isothermal titration calorimetry, single-molecule Förster resonance energy transfer, and surface-plasmon resonance spectroscopy. The problem the authors undertake is of fundamental importance.

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Reviewer #2 (Public Review):

Summary:

The manuscript by Han et al and Cordes is a tour-de-force effort to distinguish between induced fit and conformational selection in glutamine binding protein (GlnBP). It is important to say that I don't agree that a decision needs to be made between these two limiting possibilities in the sense that whether a minor population can be observed depends on the experiment and the energy difference between the states. That said, the authors make an important distinction which is that it is not sufficient to observe both states in the ligand-free solution because it is likely that the ligand will not bind to the already closed state. The ligand binds to the open state and the question then is whether the ligand sufficiently changes the energy of the open state to effectively cause it to close. The authors point out that this question requires both a kinetic and a thermodynamic answer. Their "method" combines isothermal titration calorimetry, single-molecule FRET including key results from multi-parameter photon-by-photon hidden Markov modelling (mpH2MM), and SPR. The authors present this "method" of combination of experiments as an approach to definitively differentiate between induced fit and conformational selection. I applaud the rigor with which they perform all of the experiments and agree that others who want to understand the exact mechanism of protein conformational changes connected to ligand binding need to do such a multitude of different experiments to fully characterize the process. However, the situation of GlnBP is somewhat unique in the high affinity of the Gln (slow off-rate) as compared to many small molecule binding situations such as enzyme-substrate complexes. It is therefore not surprising that the kinetics result in an induced fit situation. In the case of the E-S complexes I am familiar with, the dissociation is much more rapid because the substrate binding affinity is in the micromolar range and therefore the re-equilibration of the apo state is much faster. In this case, the rate of closing and opening doesn't change much whether ligand is present or not. Here, of course, once the ligand is bound the re-equilibration is slow. Therefore, I am not sure if the conclusions based on this single protein are transferrable to most other protein-small molecule systems. I am also not sure if they are transferrable to protein-protein systems where both molecules the ligand and the receptor are expected to have multiscale dynamics that change upon binding.

Strengths:

The authors provide beautiful ITC data and smFRET data to explore the conformational changes that occur upon Gln binding. Figure 3D and Figure 4 (mpH2MM data) provide the really critical data. The multi-parameter photon-by-photon hidden Markov modelling (mpH2MM) data. In the presence of glutamine concentrations near the K_d , two FRET-active sub-populations are identified that appear to interconvert on timescales slower than 10 ms. They then do a whole bunch of control experiments to look for faster dynamics (Figure 5). They also do TIRF smFRET to try to compare their results to those of previous publications. Here, they find several artifacts are occurring including inactivation of ~50% of the proteins. They also perform SPR experiments to measure the association rate of Gln and obtain expectedly rapid association rates on the order of $10^8 \text{ M}^{-1}\text{s}^{-1}$.

Weaknesses:

Looking at the traces presented in the supplementary figures, one can see that several of the traces have more than one molecule present. The authors should make sure that they use only traces with a single photobleaching event for each fluorophore. One can see steps in some of the green traces that indicate two green fluorophors (likely from 2 different molecules) in the traces. This is one of the frequent problems with TIRF smFRET with proteins, that only some of the spots represent single molecules and the rest need to be filtered out of the analysis.

The NMR experiments that the authors cite are not in disagreement with the work presented here. NMR is capable of detecting "invisible states" that occur in 1-5% of the population. SmFRET is not capable of detecting these very minor states. I am quite sure that if NMR spectroscopists could add very high concentrations of Gln they would also see a conversion to the closed population.

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