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Solvent dynamics play a decisive role in the complex formation of biologically relevant redox proteins†

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Electron transfer processes between proteins are vital in many biological systems. Yet, the role of the solvent in influencing these redox reactions remains largely unknown. In this study, terahertz-time domain spectroscopy (THz-TDS) is used to probe the collective hydration dynamics of flavoenzyme ferredoxin-NADP+-reductase (FNR), electron transfer protein ferredoxin-1 (PetF), and the transient complex that results from their interaction. Results reveal changes in the sub-picosecond hydration dynamics that are dependent upon the surface electrostatic properties of the individual proteins and the transient complex. Retarded solvent dynamics of 8-9 ps are observed for FNR, PetF, and the FNR:PetF transient complex. Binding of the FNR:PetF complex to the substrate NADP⁺ results in bulk-like solvent dynamics of 7 ps, showing that formation of the ternary complex is entropically favored. Our THz measurements reveal that the electrostatic interaction of the protein surface with water results in charge sensitive changes in the solvent dynamics. Complex formation between the positively charged FNR:NADP+ pre-complex and the negatively charged PetF is not only entropically favored, but in addition the solvent reorganization into more bulk-like water assists the molecular recognition process. The change in hydration dynamics observed here suggests that the interaction with the solvent plays a significant role in mediating electron transfer processes between proteins.

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Introduction

In the center of photosynthetic metabolic redox-reactions of plants and other photosynthetic microorganisms, ferredoxins (FDXs) serve as universal electron shuttles by distributing electrons originating from photosynthetic water oxidation and fermentation to specific FDX-dependent enzymes in the chloroplast. 1-4 Plant-type FDXs are small metalloproteins that usually have low redox potentials (-300 to -460 mV) and harbor [2Fe-2S]-clusters. 1-8

In the photosynthetic electron transport chain of the well characterized unicellular green algae Chlamydomonas reinhardtii, FDX1 (also known as PetF) transfers electrons from photosystem I (PSI) to the flavoenzyme ferredoxin-NADP+-reductase (FNR) to generate nicotinamide adenine dinucleotide phosphate (NADPH), an important energy carrier in most biological systems. Regardless of whether occurring in vivo under physiological conditions or in vitro, electron transfer reactions between two or more proteins require the formation of transient protein complexes. A reversible two-electron transfer between PetF and FNR initiates the hydride (H⁻)-transfer to generate NADPH; ^{10,11} protein crystallography revealed that this process takes place as a ternary complex between FNR, NADP⁺ and PetF, in which FNR and NADP⁺ form a pre-complex. 12-14 The general interaction pattern of this redoxcouple has already been well characterized by protein crystallography, biochemical methods and nuclear magnetic resonance (NMR) experiments. These studies revealed that the intermolecular recognition and subsequent stabilization of this complex is mainly driven by electrostatic charge-charge interactions between basic residues of FNR and acidic residues of PetF (Fig. 1a and Fig. S1 in the ESI†). 13,15-17 This transient complex is further stabilized by several amino acids from both proteins that form a hydrophobic environment near the two redox centers which were reported to be in close proximity to each other (\sim 6 Å) to ensure fast electron transfer (Fig. S2, ESI†).15

Interestingly, despite the presence of at least 13 FDXisoforms in C. reinhardtii, which are similar in their sequence and share structural homology (~40% sequence similarity), only a few of these isoforms (including PetF, FDX2, and FDX7) show an interaction with FNR.^{6,8} Most importantly, the relevant acidic amino acids (including D58 of PetF) are strictly conserved in FDX-isoforms 1-3. However, the fact that FDX3 is not

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Paper

A B PetF and FNI (THz-TDS). B with the non-

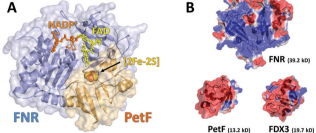


Fig. 1 (a) Structural model of the transient FNR-NADP*-PetF complex from *Chlamydomonas reinhardtii* as a cartoon and surface model (based on PDB ID 1GAQ). The redox-centers ([2Fe-2S]-cluster, FAD and NADP*) are displayed as sticks and spheres. (b) Visualization of the surface electrostatic potential of FNR (upper panel, structural model, based on PDB ID 1EWY), PetF (left, PDB ID: 2MH7) and FDX3 (right, structural model, based on PDB ID 2MH7). Surface charge distribution has been calculated using APBS. Red represents regions of negative charges and blue represents regions of positive charge.

able to interact with FNR indicates that the processes of molecular recognition and complex stabilization cannot be solely attributed to the function of individual amino acids as these patterns are mainly conserved in FDX isoforms.

In recent years the importance of the aqueous solvent in influencing protein structure, function, and dynamics has been recognized. 18,19 Coupling of water molecules to the protein surface results in an interfacial region in which water molecules within this region have distinctly different properties than bulk water. However, the structure and dynamics within this interfacial region are still not easy to access experimentally. Studies of protein solutions by surface sensitive techniques such as sum frequency generation (SFG) spectroscopy have found that organization of water in the interfacial region largely depends on the structure and net charge of the protein.²⁰⁻²² Furthermore, probing of water near specific molecular groups relevant to proteins such as carboxylic acids, alcohols, and amines have found that the hydrogen-bonding network is largely influenced by the local electric field (i.e. charge), hydrophobicity, and the ability of the molecular group to form hydrogen bonds. 23-25 This technique, however, is restricted to water near air/liquid or liquid/solid interfaces.

Terahertz (THz) spectroscopy has been shown to be a powerful tool to investigate solvent dynamics in bulk solutions. Radiation in the THz regime is directly sensitive to the low frequency collective intermolecular hydrogen-bonding vibrations of water (0.3–6 THz or 10–200 cm⁻¹), and thus to any changes in the hydrogen-bonding network. Changes in these sub-picosecond collective motions, such as protein–water interactions, can be probed by changes in the measured THz absorption. Individual hydration shells of proteins have been shown to contribute largely to structure–function relationships and ultimately modulate the binding properties of proteins. However, solvation dynamics are still poorly characterized in protein–protein complexes in spite of the fact that they may play a key role in unraveling the molecular recognition of redox-protein couples. In this study, the transient protein complex between

PetF and FNR was studied by THz-time domain spectroscopy (THz-TDS). By comparing the native redox couple PetF:FNR with the non-functional couple FDX3:FNR at different protein concentrations, compelling evidence is provided that solvation water plays a crucial role for the molecular recognition and electron transfer processes of redox proteins. Investigation of the ternary complex PetF:FNR:NADP⁺ reveals a change in solvent dynamics upon substrate binding which could promote release of the reduced substrate after the electron transfer process.

Experimental

Protein expression and purification

FNR, PetF, and FDX3 proteins were expressed and purified according to previously established protocols (see ESI† for details). All protein samples were stored at -80 °C until further use. Specific concentrations were prepared from dilution of the stock solution. All mixtures were freshly prepared on the day of measurement prior to injection into the sample cell.

Terahertz-time domain spectroscopy

THz-TDS measurements were carried out with a custom-built spectrometer setup similar to that previously described. 31,32 Briefly, the 800 nm output of a Ti:sapphire oscillator (Mai Tai EHP, Spectra Physics, United States) with a repetition rate of 80 MHz and pulse duration of 70 fs is split into pump and probe beams. The pump beam propagates to a photoconductive antenna (Tera-SED 3, Gigaoptics, Germany) that is biased at 20 V and modulated at 92 kHz to produce THz radiation. The generated THz radiation is collimated and focused by two off-axis parabolic mirrors to the sample cell. The THz beam transmitted through the cell is collected and focused by two off-axis parabolic mirrors to a 0.5 mm ZnTe(110) crystal for electro-optical sampling (EOS). A hole in the last parabolic mirror allows a collinear sampling geometry with the probe beam. Based on the principles of EOS, the electric field of the THz pulse generates a birefringence in the ZnTe crystal which induces a rotation in the polarization state of the probe beam that is directly proportional to the electric field of the THz pulse. The probe beam is split into orthogonal polarization components with a Wollaston prism and directed to a balanced photodetector (Nirvana Auto-Balanced Optical Receiver, Model 2007, New Focus, United States), where the intensity difference of the two components is measured. The signal of the photodetector is processed by a lock-in amplifier (HF2-LI, Zurich Instruments, Switzerland) with a 6 dB filter and 5 µs time constant, and the phase of the lock-in amplifier is set such that the signal in the X-component is maximized. A time window of approximately 20 ps is probed through the use of a rapid delay stage (ScanDelay, Model 150, APE, Germany) in the pump beam path which operates at 2 Hz. All measurements were conducted at 30 °C, where the temperature was held constant by an external chiller (±0.1 °C). Humidity within the sampling chamber was fixed at $\leq 5\%$.

The sample cell was a commercially available demountable cell (Bruker, Germany), in which two z-cut quartz windows were used. Sample thickness of 100 µm was achieved with a Teflon spacer inserted between the windows. The front window had two holes bored in it, allowing for the sample to be injected. An overall solution volume of 200 µL was needed to fill the cell. The frequency dependent refractive index $n(\nu)$ and absorption

coefficient $\alpha(\nu)$ were calculated with eqn (1) and (2)

$$n(\nu) = 1 + \frac{c[\varphi - \varphi_i]}{2\pi\nu d} \tag{1}$$

$$\alpha(\nu) \text{ (cm}^{-1}) = -\frac{2}{d} \log \left| \frac{A (n(\nu) + n_0)^2}{A_i n(\nu) (1 + n_0)^2} \right|$$
 (2)

where c is the speed of light, d is the sample thickness, v is the frequency, ϕ and ϕ_i are the phases of the Fourier transforms (FT) of the fields transmitted through the filled and empty cell, A and A_i are the FT amplitudes, and n_0 is the refractive index of z-cut quartz. Reiterative calculations were done to reduce artifacts from multiple reflections in the cell.³³ The electric field transmitted through the reference and sample was measured alternatively 10 times, in which each scan was integrated for 30 seconds, and resulted in an average error of the calculated refractive index and absorption coefficient of 0.01 and 0.6 cm⁻¹, respectively. Measurements were repeated at least two times to ensure reproducibility.

Results and discussion

Fig. 2 shows the THz-TDS absorption spectra of solutions containing FNR and PetF with concentrations ranging from 0 to 2 mM. The absorption coefficient increases nearly linearly over the measured frequency range for all spectra. With increasing protein concentration a decrease in the absorption coefficient occurs, consistent with a decrease in absorption due to a decrease in bulk water concentration as similarly observed in

previous measurements of other proteins. 34-37 If the decrease in absorption is occurring solely due to a lowered water concentration, a linear decrease at every measured frequency as a function of protein concentration would be observed. In the case of PetF, it can be clearly seen that changes in the absorption coefficient are frequency dependent, where the absorption coefficient is largely independent of concentration below 1 THz, but decreases significantly at higher frequencies. Based on this observation, the average absorption coefficient (α_{avo}) for frequency ranges of 0.3–1, 1–1.75, and 1.75–2.5 THz were determined, and provides the same information as an individual frequency within these ranges. Results shown here will focus on the range 1.75-2.5 THz (the other two regions are shown in Fig. S3, ESI†). The larger absorption coefficient at higher frequencies means that changes in the α_{avg} due to water exclusion will be more pronounced in this regime. Any deviations from the expected water exclusion value report on the hydrodynamic solvation shell of the protein. However, similar effects can be observed at other frequency ranges and will be discussed where pertinent. Fig. 3 shows α_{avg} of FNR and PetF solutions from 1.75-2.5 THz for protein concentrations ranging from 0 to 2 mm. The black line in Fig. 3 represents the expected decrease in α_{avg} of the buffer solution by excluding a volume of bulk water equal to the volume of the protein in which spherical volume of the protein was assumed to be R_{FNR} = 29 Å and $R_{\text{PetF}} = 17$ Å from the radii measured with dynamic light scattering in previous literature.³⁸ The α_{avg} of both individual FNR and PetF solutions deviates from this black line which takes into account solely bulk water exclusion. At all volume fractions FNR has an increased absorption relative to the black line while PetF features a decreased absorption. The nonlinear behavior in α_{avg} observed here for both proteins indicates that the water molecules directly interacting with the protein surface, i.e. the hydration shell, have a distinct absorption which differs from bulk water, consistent with previous studies of proteins in solution. 18,36,37,39 However, the manner in which water interacts with the two proteins is

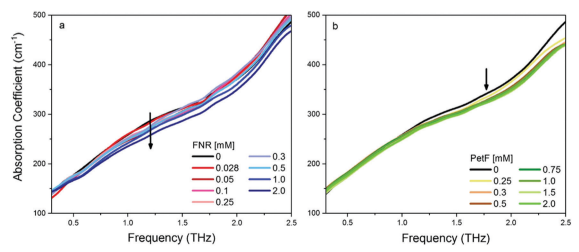


Fig. 2 THz-TDS absorption spectra of aqueous solutions at pH 8 of (a) FNR and (b) PetF. The error of each measurement is on average 0.6 cm⁻¹. The absorption coefficient decreases with increasing protein concentration, consistent with a decreasing water content.

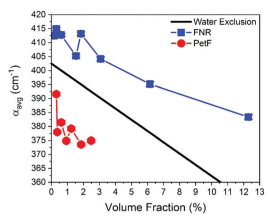


Fig. 3 The average absorption coefficient (α_{avg}) in the 1.75–2.5 THz range of aqueous solutions containing either FNR (blue) or PetF (red) as a function of volume fraction of protein in solution. The black line shows the expected decrease in α_{avg} based on exclusion of water equal to the volume of protein in solution. Error bars are on average 0.6 cm $^{-1}$.

different, evidenced by the opposite behavior in α_{avg} . Coupling of water to the protein depends upon its surface properties, in which the local topography influences interactions with water. For instance, water molecules within hydrophobic pockets or grooves form strong hydrogen-bonds mostly due to sterical constraints, 40 whereas coordination of water molecules to hydrophilic sites appears to be correlated with charge. 41 The resulting dynamical solvation shell emerges from the cooperative effect of water solvating the heterogeneous surface of the protein. 42

Mapping of the surface electrostatic potential of FNR and PetF at pH 8 (Fig. 1b, based on available PDB structures) shows that the surface of FNR has two distinct regions of negative and positive charges while the surface of PetF is largely negatively charged. The opposite surface electrostatic potentials of these proteins influence water to different extents, and is reflected in their corresponding THz spectra. *Ab initio* MD (AIMD) simulations and THz spectra of glycine and valine have shown that positively charged groups, such as aminos (NH₃⁺), result in an increased THz intensity in the low frequency regime (below 100 cm⁻¹, 3 THz). This increased intensity was attributed to cage rattling modes of the solvating water

molecules at 65 and 70 cm $^{-1}$ (1.8 and 2.0 THz). 45-47 While the surface of FNR has both positively and negatively charged regions, the positively charged region covers a larger surface area, and the results obtained here show that the contribution of the positively charged groups dominates the THz response. In contrast, PetF has a negatively charged surface, which is mostly comprised of aspartic acid and glutamic acid residues. MD and AIMD simulations on glycine have shown that water molecules solvating carboxylate groups (COO⁻) are well ordered and have a cage librational mode, i.e. a correlated solute-water mode, in the THz regime centered at 2.4 THz (80 cm⁻¹). The THz intensity below 150 cm⁻¹ is predicted to be suppressed for the negative form of the carboxylic acid group compared to the neutral form. 45,47,48 Measurements of glycine confirm this, where it was observed that the negatively charged molecule has a larger intensity than the zwitterionic molecule, 49 and is consistent with results shown in Fig. 2 for PetF. This observation demonstrates how solvent dynamics are sensitive to differences in the surface properties of individual proteins, and can be used to report on changes in the electrostatics of protein surfaces.

THz absorption spectra for solutions of the native redox couple FNR:PetF, the non-functional couple FNR:FDX3, and the ternary complex FNR:PetF:NADP⁺ (NADP⁺ purchased from Roth GmbH, >97% purity) are shown in Fig. 4. All solutions were prepared with equimolar amounts of each component. Fig. 5 shows the corresponding α_{avg} determined for each solution in the 1.75-2.5 THz range (see Fig. S4, ESI† for other frequency regions). The α_{avg} follows the expected water exclusion for FNR:PetF solutions (assuming R = 30 Å for the transient complex),38 while increased values are observed for FNR:FDX3 solutions (assuming R = 17 Å for FDX3). FDX3 alone has decreased α_{avg} similar to PetF as shown in Fig. S5 and S6 (ESI†). This result demonstrates differences in the collective water dynamics in the hydration shell of proteins that remain unbound in solution (FNR and FDX3) compared to those that form a complex (FNR:PetF). Formation of site-specific salt bridges is imperative for the complexation of FNR and PetF. 14,43,50,51 Positively charged lysine and arginine residues on the surface of FNR couple to negatively charged glutamic acid and aspartic acid residues on PetF (Fig. S1, ESI†). 16,17,44

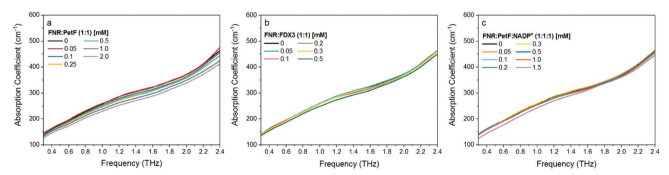


Fig. 4 THz-TDS absorption spectra of aqueous solutions at pH 8 of equimolar mixtures of (a) FNR:PetF (b) FNR:FDX3 and (c) FNR:PetF:NADP⁺. The error of each measurement is on average 0.6 cm⁻¹. The absorption coefficient decreases with increasing protein concentration, consistent with decreasing water content.

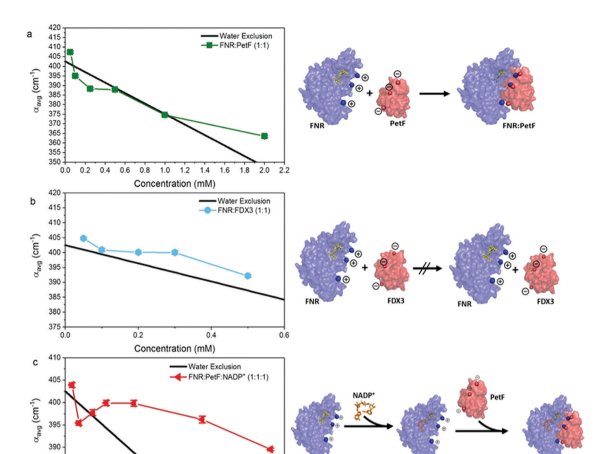


Fig. 5 The average absorption coefficient (α_{avg}) in the 1.75–2.5 THz range of aqueous solutions of (a) FNR:PetF (b) FNR:FDX3 and (c) FNR:PetF:NADP+ as a function of concentration. See Fig. S4 in ESI† for other frequency ranges. The black line in each panel shows the expected decrease in α_{avg} based on exclusion of water equal to the volume of protein in solution. Error bars are on average 0.6 cm⁻¹. The complexation state of proteins in solution is illustrated on the right hand side.

Neutralization of these surface charges upon complexation results in a change in the surface electrostatic potential such that the solute is overall neutral. Our results demonstrate that the THz responses of FNR, PetF, and the FNR:PetF complex are clearly distinguishable.

1.0

Concentration (mM)

Fig. 5c shows that the α_{avg} of the FNR:PetF:NADP⁺ ternary complex in aqueous solutions deviates significantly from the expected water exclusion, where an increased α_{avg} is observed for concentrations greater than 0.3 mM. This result is surprising as it is well known that no major changes in surface morphology of the complex occur upon NADP binding (Fig. S2, ESI†), thus yielding valuable insight. NADP binds within the interior of FNR prior to complexation with PetF (Fig. 5c and Fig. S2, ESI†) and is consequentially solvent inaccessible. Measurements of NADP+ show that it has no influence on the measured THz absorption (Fig. S7 and S8, ESI†), and therefore cannot explain the observed increase in absorption. Thus, we propose binding of the NADP⁺ alters the charge distribution at the surface of the complex, which consequently influences hydration water.

Evaluation of the complex dielectric response of the protein solutions (see ESI† for details) reports on the water solvent dynamics, specifically the time scale of fluctuations in the hydrogen-bonding network. Here, the complex dielectric response was modeled with a double Debye function, 52 which reveals a slow relaxation process, τ_1 , associated to the cooperative rearrangement of the hydrogen bonded network, and a fast relaxation process, τ_2 , assigned to the rotational modes of individual polar water molecules.53,54 Fig. 6 shows values of τ_1 determined for all protein solutions (see Fig. S9, ESI† for τ_2). For solutions of FNR and PetF (Fig. 6a), τ_1 is observed to increase from the initial 7.0 ps value at 0 mM up to 9 ps and 8 ps, respectively, with increasing protein concentration. This trend is consistent with MD simulations that found water molecules in the hydration shell of proteins have significantly slower reorientation times than pure, bulk water. 26,40,42,52 The slower relaxation times observed for FNR relative to PetF are indicative of its predominantly hydrophobic nature, in which water molecules near the hydrophobic surface will form

FNR:NADP

FNR:NADP*:PetF

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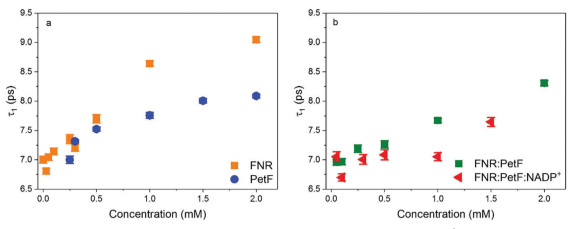


Fig. 6 Relaxation time constants τ_1 of aqueous solutions of (a) FNR and PetF (b) FNR:PetF and FNR:PetF:NADP⁺.

stronger hydrogen-bonds with neighboring water molecules to minimize interaction with the hydrophobic surface which ultimately results in retarded dynamics.

Similarly, an increase in τ_1 with increasing protein concentration is observed for the transient FNR:PetF complex (Fig. 6b). In contrast, the presence of the substrate NADP⁺ does not initially affect the relaxation dynamics (up to 0.5 mM). For the FNR:PetF:NADP⁺ complex the values of τ_1 remain relatively constant around 7.0 ps until a concentration of 1.5 mM, where a sudden increase to 7.6 ps occurs. The relaxation dynamics for the ternary complex are similar to pure bulk water while the binary complex has retarded dynamics.

Changes in the solvent dynamics upon complex formation indicate the solvent contributes to changes in the free energy of the system ($\Delta G = \Delta H - T\Delta S$). Water molecules tightly bound to the protein surfaces have the potential to act as an entropic reservoir to offset the enthalpic and entropic costs of forming the complex. FNR and PetF solutions show retarded water dynamics stemming from water molecules tightly bound to charged surface residues. As illustrated in Fig. 7, these tightly bound water molecules (shown in blue) must be displaced and released into the bulk phase in order to favor binding of PetF to the pre-formed FNR:NADP⁺ complex. The bulk-like solvent dynamics observed for the ternary complex indicate that water molecules within its hydration shell have a low frequency response which is similar to that of pure water (*i.e.* bulk-like water),

meaning that binding of the complex in the presence of the NADP⁺ is entropically favorable. The increased THz absorption upon embedding of NADP⁺ into its binding pocket is on the other hand an indication of a change of the surface charge distribution due to binding. For a binding reaction the overall change of $\Delta G = \Delta H - T\Delta S$ is decisive. Whereas most of the time there is a focus on ΔH , the magnitude of $T\Delta S$ is mostly of the same size. Here, we show indications for a change in solvent dynamics which favors ternary complex formation. Our results suggest that the solvent is a vital participant in the electron transfer process, where changed solvent dynamics upon binding of the substrate NADP⁺ will most likely serve as a prompt for the following step in the redox reaction. In electron transfer processes, dissociation of the transient complex after electron transfer to the substrate is usually the rate-limiting step. 55-57 The bulk-like solvent dynamics observed here upon NADP+ binding is proposed to help facilitate dissociation of the complex so that the formed NADPH molecule can be released after the electron transfer.

Conclusions

These results demonstrate that THz-spectroscopy is a powerful technique to study the role of water in redox protein couples in solution that are of biological and biotechnological relevance. While the transient FNR:PetF complex has already been

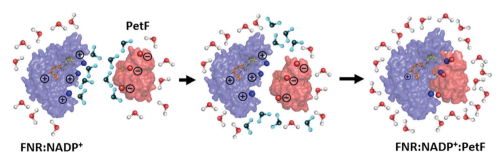


Fig. 7 Formation of the FNR:NADP*:PetF ternary complex is entropically favorable. Tightly bound, retarded water molecules (shown in blue) near the charged surface residues on the pre-formed FNR:NADP* complex and PetF protein are released as the charged surfaces approach each other, resulting in a hydration shell that is more bulk-like in nature.

characterized extensively by several biochemical and spectroscopic methods, results here shed light on the role of the solvent. We show the THz response of the solvent is sensitive to surface properties of the individual proteins, with a change in the THz response observed upon protein complexation. Our results further reveal that the interaction of proteins does not solely account to individual amino acids on the protein surface as PetF and FDX3 share a conserved amino acid pattern, which were shown to be relevant for a mutual interaction with FNR. Furthermore, we find that formation of the ternary complex FNR:NADP+:PetF, in which the substrate NADP+ binds first to FNR, results in bulk-like hydration dynamics. The change in solvent dynamics from a more retarded to a bulk-like nature is entropically favorable and facilitates complex formation to trigger electron transfer. In general, solvation dynamics play a decisive factor for the molecular recognition and interaction of redox proteins.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

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