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journal or publication title	Photosynthesis Research
volume	130
number	1-3
page range	479-489
year	2016-12-01
URL	http://hdl.handle.net/2297/45922

doi: 10.1007/s11120-016-0285-3

Kinetics of NADP⁺/NADPH reduction-oxidation catalyzed by the ferredoxin-NAD(P)⁺ reductase from the green sulfur bacterium *Chlorobaculum tepidum*

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Keywords

Flavoenzyme; Adrenodoxin; Putidaredoxin; Stopped flow; Charge transfer complex, Kinetic isotope effect

Abbreviations

A, absorbance; *Ad*, adrenodoxin; *AdR*, adrenodoxin reductase; *Bs*, *Bacillus subtilis*; *Ct*, *Chlorobaculum tepidum*; *CT*, charge transfer; *CTC*, charge transfer complex; *Ec*, *Escherichia coli*; *FAD*, flavin adenine dinucleotide; *Fd*, ferredoxin; *FNR*, ferredoxin-NAD(P)⁺ oxidoreductase; *GR*, glutathione reductase; *HEPES*, 2-[4-(2-hydroxyethyl)-1-piperazinyl]ethanesulfonic acid; *NADPD*, (4*S*-²H)-reduced nicotinamide adenine dinucleotide phosphate; *Pd*, putidaredoxin; *PdR*, putidaredoxin reductase; *TrxR*, bacterial NADPH-thioredoxin reductase;
Suffixes: *ox*, oxidized; *red*, reduced.

Acknowledgements

We thank Dr. Hidehiro Sakurai for stimulating discussions. This work was supported in part by a grant-in-aid for Scientific Research on Innovative Areas (No. 24107004) and the Strategic Research Base Development Program for Private Universities from the Ministry of Education, Culture, Sports, Science and Technology (MEXT), Japan (to KI).

Abstract Ferredoxin-NAD(P)⁺ oxidoreductase (FNR, [EC 1.18.1.2], [EC 1.18.1.3]) from the green sulfur bacterium *Chlorobaculum tepidum* (*CtFNR*) is a homodimeric flavoprotein with significant structural homology to bacterial NADPH-thioredoxin reductases. *CtFNR* homologs have been found in many bacteria, but only in green sulfur bacteria among photoautotrophs. In this work, we examined the reactions of *CtFNR* with NADP⁺, NADPH and (4S-²H)-NADPH by stopped-flow spectrophotometry. Mixing *CtFNR*_{ox} with NADPH yielded a rapid decrease of the absorbance in flavin band I centered at 460 nm within 1 ms, then the absorbance further decreased gradually. The magnitude of the decrease increased with increasing NADPH concentration, but even with ~50-fold molar excess NADPH, the absorbance change was only ~45% of that expected for fully reduced protein. The absorbance in the charge transfer (CT) band centered around 600 nm increased rapidly within 1 ms, then slowly decreased to about 70% of the maximum. When *CtFNR*_{red} was mixed with excess NADP⁺, the absorbance in the flavin band I increased to about 70% of that of *CtFNR*_{ox} with an apparent rate of ~4 s⁻¹, whereas almost no absorption changes were observed in the CT band. Obtained data suggest that the reaction between *CtFNR* and NADP⁺/NADPH is reversible, in accordance with its physiological function.

Introduction

Ferredoxin-NAD(P)⁺ reductase ([EC 1.18.1.2], [1.18.1.3], FNR) is a member of the NAD(P)H-dependent oxidoreductases of the flavoprotein superfamily containing a non-covalently bound flavin adenine dinucleotide (FAD) or flavin mononucleotide (FMN) as a prosthetic group. FNR can reversibly catalyze either one- and two-redox-equivalent transfer with the small iron-sulfur protein ferredoxin (Fd) and NAD(P)⁺/NAD(P)H, respectively, using the redox properties of the flavin prosthetic group. In the final step of the electron transport chain in photosynthesis, Fd and FNR mediate electron transfer from a type-I photosynthetic reaction center to NAD(P)⁺ (Knaff and Hirasawa 1991; Sétif 2001; Seo and Sakurai 2002). In non-photosynthetic processes, Fd as well as small iron sulfur proteins, adrenodoxin (Ad) and putidaredoxin (Pd) participate in electron transfer in reactions involved in synthesis of a wide range of cellular materials such as fatty acid desaturation (cytochrome P450), nitrite reduction, sulfite reduction, and *S*-adenosylmethionine radical-dependent reactions in DNA and coenzyme synthesis (Munro et al. 2007; Knaff and Hirasawa 1991; Bianchi et al. 1993; Wan and Jarrett 2002). In the latter processes, FNR, adrenodoxin reductase (AdR, [EC 1.18.1.6]) and putidaredoxin reductase (PdR, [EC 1.18.1.5]) catalyze the reduction of Fd, Ad and Pd by NAD(P)H. Thus the redox coupling of Fd-FNR-NAD(P)⁺/NAD(P)H constitutes a key process of a widespread redox network in living organisms. The physiological direction of the process is assumed to be related to the redox midpoint potentials of the reaction partner(s).

Each of FNR, AdR and PdR has two nucleotide-binding domains, one for FAD/FMN and the other for NAD(P)⁺/H (Correll et al. 1993; Ziegler et al. 1999; Sevrioukova et al. 2004; Muraki et al. 2010; Medina and Gómesz-Moreno 2004; Alverti et al. 2008). This protein topology is typical of flavin-containing NAD(P)H-dependent oxidoreductases including disulfide reductases (Dym and Eisenberg 2001; Alverti et al. 2008). Based on structural and phylogenetic information, FNRs, PdR and AdR are classified into two major groups of the flavoprotein superfamily, namely the plant-type and the glutathione reductase (GR)-type FNR superfamilies and further divided into several subgroups (Alverti et al. 2008; Ceccarelli et al. 2004; Seo et al. 2004). Among FNRs, the bacterial NADPH-thioredoxin reductase-type (TrxR-type) FNR, a subgroup of the GR-type FNR superfamily, is unique in their homo-dimeric form with high structural homology to *Escherichia coli* TrxR (*EcTrxR*) (Muraki et al. 2010; Komori et al. 2010; Waksman et al. 1994). TrxR-type FNR and its homologs have been found in bacteria and archaea, including green sulfur bacteria, Firmicutes, several α -proteobacteria including Rhodospirillaceae and the archaeal genus *Sulfolobus* (Seo et al. 2002, 2004, 2009; Yan et al. 2014; Mandai et al. 2009), but it was not found in eukaryotes (Kyoto Encyclopedia of Genes and Genomes, Kanehisa et al. 2016). Among members of the TrxR-type FNR subfamily, the protein from the photoautotrophic green sulfur bacterium *Chlorobaculum tepidum* (formerly *Chlorobium tepidum*; *CtFNR*) has been isolated as a component required for NAD(P)⁺ photoreduction (Seo and Sakurai 2002). *C. tepidum* carries out non-oxygenic photosynthesis using various sulfur compounds as the

electron sources (Ogawa et al. 2008; Sakurai et al. 2010) and has a type-I photosynthetic reaction center that can reduce Fds directly (Kjær and Sheller 1996; Seo et al. 2001; Tsukatani et al. 2004). *In vitro*, purified *CtFNR* can catalyze the reduction of NAD(P)⁺ in the presence of the photosynthetic reaction center and Fd from *C. tepidum* (Seo and Sakurai 2002). The unique features of the green sulfur bacteria among phototrophs are that they use the reductive TCA cycle as the main carbon assimilation pathway (Buchanan and Arnon 1990; Tang et al. 2010) which have been found among several bacterial groups (Hügler and Sievert 2011), and that FNR and Fds are phylogenetically unrelated to their counterparts in oxygenic photosynthetic organisms/organelles but related to those from several bacterial groups described above. These would provide useful information on the origin and evolution of photosynthesis.

Although the redox reaction catalyzed by FNR is theoretically reversible, it is generally accepted that under physiological conditions, the direction of the reaction is optimized to either NAD(P)⁺ reduction in many photosynthetic organisms, or NAD(P)H oxidation in non-photosynthetic organisms and tissues. Indeed, previous studies on the TrxR-type FNR from the heterotroph *Bacillus subtilis* (*BsFNR*) revealed that *BsFNR* preferentially oxidizes NADPH to reduce Fd from the same organism at a high rate (~50 s⁻¹), rather than reducing NADP⁺ with electrons from the reduced Fd (Seo et al. 2004, 2014). Pre-steady state reaction analysis of *BsFNR* with NADP⁺/NADPH revealed that the hydride transfer rate for NADPH oxidation direction is much larger than that of NADP⁺ reduction direction (Seo et al. 2016). Therefore, research on *CtFNR* is important because green sulfur bacteria constitute a unique group among photoautotrophic organisms in that they contain TrxR-type FNRs that support NAD(P)⁺ photoreduction at a high rate (Seo and Sakurai 2002). Some differences between *CtFNR* and other TrxR-type FNRs have been revealed from deduced amino acid sequences and crystal structures of TrxR-type FNRs, for example the residue stacking on the *re*-face of the isoalloxazine ring moiety of the FAD prosthetic group is Phe337 and C-terminal extension region in *CtFNR* is longer and rich in Lys and Glu residues (Fig. 1a, b) (Muraki et al. 2010; Komori et al. 2010). It is expected that comparative studies of *CtFNR* with other members of the TrxR-type FNR family will provide valuable information on the structure-function relationships of this family and other flavin-containing NADPH-dependent oxidoreductases. In this work, we performed pre-steady-state kinetic analyses of reactions of *CtFNR* with NADP⁺ and NADPH. Our studies confirmed that the reaction between *C. tepidum* FNR and NADP⁺/NADPH is reversible as often been reported among plant-type FNR from photosynthetic organisms.

Materials and Methods

Preparation of *C. tepidum* FNR

*Ct*FNR was expressed in *E. coli* cells as described in Muraki et al. (2008). Purification of the *Ct*FNR was performed basically according to the methods used for *Bs*FNR (Seo et al. 2014), except that precipitate obtained by the addition of ammonium sulfate between 35% and 80% saturation was used for purification in place of that obtained from 0–40% saturation.

Stopped-flow spectrophotometry

Stopped-flow spectrophotometry was performed in a glove box under a nitrogen atmosphere containing hydrogen gas (ca. 5%) to decrease the residual O₂ (< 50 ppm) with a palladium catalyst as previously described (Seo et al. 2016). The reaction was initiated by mixing equal volumes of solutions in a single mixing mode at 10°C. The dead-time of the setup was estimated to be about 1 ms (Seo et al. 2016). Spectra were collected every 1 ms (Figs. 1-3, 4a-c) or 5 ms (Fig. 4e). All protein and substrate concentrations are given as the final concentrations after mixing unless otherwise stated.

Reduced *Ct*FNR (*Ct*FNR_{red}) was obtained using sodium dithionite and methyl viologen as the reductant and redox mediator, respectively, under anaerobic conditions in the glove box. Two milliliters of approximately 200 μM *Ct*FNR solution in 20 mM HEPES-NaOH buffer (pH 7.0) were reduced by the addition of an excess of sodium dithionite (0.4 mg/ml final concentration) in the presence of 10 μM methyl viologen. After a few minutes of incubation at 4°C, excess dithionite and methyl viologen were removed by size-exclusion chromatography (Bio-Gel P-4 gel, Bio-Rad Laboratories, USA) pre-equilibrated with 20 mM HEPES-NaOH buffer (pH 7.0) containing 0.32 μg/ml sodium dithionite in the glove box. The concentration of *Ct*FNR in the solution was estimated from the absorbance of air-reoxidized *Ct*FNR.

A deuterated form of NADPH, (4*S*-²H)-NADPD (referred to hereafter as NADPD), was prepared according to published methods (Viola et al. 1979, Ottolina et al. 1989, Pollock et al. 2001) as previously described (Seo et al. 2016). NADP⁺ and NADPH were purchased from Oriental Yeast Co., Ltd.

The values of A_{460} , A_{590} , ϵ_{460} and ϵ_{590} in Figs. 2b-c, 3a-c and 4c, e were obtained by subtracting A_{800} to compensate for signal drift.

Kinetic data analysis

Data collection and basic arithmetic operations on the transient spectra obtained from stopped-flow experiments were performed using Unispec (ver. 2.51, Unisoku Co., Ltd.) and Excel (ver. 14 and 15, Microsoft Corporation, Redmond, WA, USA) software. Curve fitting to the transient absorptions at a single wavelength was performed using Igor Pro ver. 6.3 software (Wavemetrics, Portland, OR, USA).

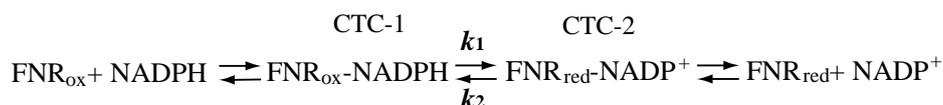
Miscellaneous methods

UV-Vis spectra were measured using a double beam spectrophotometer (UV-560, JASCO, Japan). Protein and substrate concentrations were determined from the extinction coefficients: $CtFNR_{ox}$, $\epsilon_{466} = 10.3 \text{ mM}^{-1} \text{ cm}^{-1}$ (Seo and Sakurai 2002) and NADPH and NADPD, $\epsilon_{340} = 6.2 \text{ mM}^{-1} \text{ cm}^{-1}$. NADP⁺ concentration was determined in NADPH form using glucose-6-phosphate dehydrogenase and excess amount of glucose-6-phosphate.

Results

Reduction of *CtFNR_{ox}* by NADPH

Reduction/oxidation reactions catalyzed by NAD(P)H-dependent flavoproteins are generally accompanied by the formation of charge transfer complexes (CTCs), namely $\text{FNR}_{\text{ox}}\text{-NADPH}$ (CTC-1) and $\text{FNR}_{\text{red}}\text{-NADP}^+$ (CTC-2) before the hydride transfer step, which exhibit a charge transfer (CT) absorption band in the 500–800 nm region (Scheme 1; Massey et al. 1970; Blankenhorn 1975; Batie and Kamin 1984; Lennon and Williams Jr. 1997; Tejero et al. 2007).



Scheme 1

The reaction after mixing 9 μM *CtFNR_{ox}* with 50–500 μM NADPH was measured by stopped-flow spectrophotometry (Fig. 2). Mixing *CtFNR* with NADPH resulted in a decrease in flavin absorption band I with a peak around 460 nm within 1 ms (at the end of dead time), then the absorption of this band decreased gradually with time (Fig. 2a, b). The Δ absorbance of this band region contains at least three distinctive kinetic components, namely the fast, slow and very slow phases (Fig. 2c). The fast phase observed as a rapid drop in 0 to 5 ms was too fast to estimate the rate constant. Δ absorbance of this phase was almost independent of the NADPH concentrations used (50–500 μM , inset of Fig. 2c). A_{460} in 5 to 500 ms time region can be approximated by a two-step reaction model (Fig. 2c, red continuous lines). Δ absorbance of the slow phase observed as a decay component in 5–200 ms time region increased with increasing NADPH concentration, and its observed rate decreased with increasing NADPH concentration (Fig. 2d). k_{obs} of the very slow phase was uncertain as the slight absorbance change seemed to continue beyond the measurement time scale. Although an increase in NADPH concentration decreased the total Δ absorbance of the flavin absorption band I (Fig. 2a, b), the absorbance change after 499 ms was only ~45% of that expected for fully reduced protein even with 500 μM NADPH (Fig. 2b).

The absorbance around 600 nm rapidly increased within 1 ms and reached its maximum intensity by 5 ms (Fig. 2a, b). The spectroscopic and kinetic properties of this absorption band centered at 600 nm are typical of the formation of CTCs as reported in stopped-flow works on *BsFNR*, plant-type FNRs and *EcTrxR* (Seo et al. 2016; Massey et al. 1970; Batie and Kamin 1984; Tejero et al. 2007; Lennon and Williams Jr. 1997). Subsequently, the absorption intensity of this CT absorption band decreased with time (Fig. 2a, b), but even with 500 μM NADPH, about 70% of the maximum absorption intensity remained after 2 s (Fig. 2b). No significant peak shift of the CT absorption band was observed within 2 s during *CtFNR_{ox}* reduction by NADPH (Fig. 2a, b). An increase in NADPH

concentration did not significantly affect the maximum absorption intensity of the CT band, but Δ absorbance of the decay increased with increasing NADPH concentration (inset of Fig. 2b).

Reduction of *CtFNR*_{ox} by NADPD

We studied the primary kinetic isotope effect by measuring the reactions mixing 11 μ M *CtFNR*_{ox} with 100 (Fig. 3a), 200 and 500 μ M (Fig. 3b) NADPD. Mixing *CtFNR*_{ox} with NADPD resulted in a rapid drop in the flavin absorption band I within 1 ms (Fig. 3a, b), but the Δ absorbance of the drop after 1 ms was much smaller than that with NADPH (Fig. 2a, b), and the subsequent absorption changes in 0 to 10 ms were slower than those with NADPH. These results indicated that deuterium substitution significantly affected the kinetics of the fast phase in the reduction of *CtFNR*_{ox} (inset of Fig. 3a). The k_{obs} of the fast phase in the reaction with NADPD was nearly independent of NADPD concentration; its values were 372 ± 12 , 434 ± 11 and 391 ± 11 s⁻¹ at 100, 200 and 500 μ M NADPD, respectively (inset of Fig. 3c). These values contain large errors as they are near the upper limit of the detection of the equipment. Kinetic analyses of the absorbance changes in time region 10 to 1000 ms provided similar rate constant values for the slow phase in the reaction with NADPH (36 ± 8 s⁻¹, 22.3 ± 0.4 s⁻¹ and 19.1 ± 0.3 s⁻¹ at 100, 200 and 500 μ M NADPD, respectively), which also decreased with increasing NADPD concentration.

The absorption in the CT band region increased rapidly (within 1 ms) and reached its maximum within 5 ms (Fig. 3a, b). Although the reduction of significant amount of FAD prosthetic group occurred in 1-10 ms time period (Fig. 3c), the peak wavelength and intensity of CT band were nearly stable (Fig. 3a, b). Increased NADPD concentration did not significantly affect the maximum intensity of the CT band absorption (Fig. 3a, b). Subsequently, a slight decrease in the absorbance occurred (by 100 ms), to a similar extent as was observed in the reaction with NADPH (inset of Fig. 3b). These results, almost the same as those with NADPH, indicate that the formation of CTC-1 was not affected significantly by the deuterium substitution.

Oxidation of *CtFNR*_{red} by NADP⁺

*CtFNR*_{red} was prepared with dithionite in the presence of methyl viologen followed by the removal of the latter by size-exclusion chromatography. The spectrum of the reduced *CtFNR* thus prepared is shown as the thick continuous line in Fig. 4a, b. Mixing *CtFNR*_{red} (12 μ M) with NADP⁺ resulted in a very slow increase in the flavin absorption band I (Fig. 4a, b), indicating that *CtFNR*_{red} oxidation by NADP⁺ was very slow. A_{460} in 0 to 2 sec time region can be approximated by a two-step reaction model (Fig. 4c). The k_{obs} of the faster component decreased with increasing NADP⁺ concentration (Fig. 4d). The total absorption changes at near equilibrium (i.e. at 2 s) showed a significant dependency on NADP⁺ concentration in the range 100 to 500 μ M (Fig. 4a, b). From ΔA_{460} at 2 s, the oxidation level of *CtFNR* was estimated to be about 70% (thick broken line in Fig. 4a, b).

In contrast to the reduction of $CtFNR_{ox}$ by NADPH/NADPD (Figs. 2 and 3), no significant absorption change was observed in the CT absorption band region (550–800 nm) during the reaction of $CtFNR_{red}$ with $NADP^+$ in the $NADP^+$ concentration range 100 to 500 μM (Fig. 4a, b).

As the formation of $CtFNR_{ox}$ - $NADP^+$ complex (Scheme 2) can decrease the amounts of CTCs at the equilibrium when $NADP^+$ is excess (Batie and Kamin 1986; Seo et al. 2016), the reaction mixing $CtFNR_{red}$ (24 μM) with near equimolar $NADP^+$ (25 μM) was measured (Fig. 4e). Obtained transient

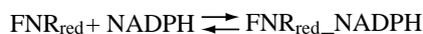


Scheme 2

spectra exhibited a gradual increase in the CT band absorption with time after 1,000 ms following accumulation of oxidized $CtFNR$ species. A_{460} predicted that the reaction was approximated by a two-step reaction model with similar k_{obs} in excess NADPH ($3.40 \pm 0.02 \text{ s}^{-1}$ and $0.426 \pm 0.001 \text{ s}^{-1}$) (inset of Fig. 4e). The change of A_{590} can be approximated by two-step reaction model with k_{obs} of $0.75 \pm 0.01 \text{ s}^{-1}$ and $0.123 \pm 0.002 \text{ s}^{-1}$ (inset of Fig. 4e).

Discussion

Previous pre-steady-state kinetic studies on flavins and other types of flavin-dependent NADP(H) oxidoreductases revealed that mixing with NADPH yielded formations of CTC-1 (FAD_{ox}-NADPH) and CTC-2 (FAD_{red}-NADP⁺) with the appearance of CT absorption bands centered at 550-750 nm (Massey et al. 1970; Blankenhorn 1975; Lennon and Williams Jr. 1997; Tejero et al. 2007). The pre-steady state kinetic study of TrxR-type *Bs*FNR catalyzing reaction with NADP⁺/NADPH revealed that formations of both CTC-1 and CTC-2 exhibited the appearance of CT absorption bands centered at approximately 600 nm (Seo et al. 2016). In the NADPH oxidation reaction by *Ct*FNR_{ox}, the CT band centered at approximately 600 nm appeared within the experimental dead-time. Its maximum intensity was not significantly affected by the NADPH concentration, indicating a formation of Michaelis complex (NADPH-FNR_{ox}, MC-1) occurred before the formation of CTC-1 as frequently reported among flavin-containing NAD(P)H-dependent dehydrogenases (Lennon and Williams Jr. 1997; Tejero et al. 2007) and 100 μM NADPH gave near saturation (Seo et al. 2016). After the rapid formation of the CTC-1, reduction of the flavin prosthetic group occurred (Figs. 2a-b, 3a-b). In the reaction, the apparent rate of the fast phase was affected by deuterium substitution (inset of Fig. 3a), indicating that the rate-limiting step in the NADPH oxidation reaction by *Ct*FNR_{ox} is the hydride transfer during CTC-2 formation (Scheme 1) in the experimental conditions used. In the course of the reaction, *Ct*FNR_{ox} reduction proceeded significantly, while the CT absorption band intensity (*A*₅₉₀) did not change in a corresponding amount. Based on Scheme 1, if CTC-2 has no absorption in the CT band region, a decrease in a CTC-1 amount reduce the absorbance in the CT band region. Thus obtained results could be explained by the presence of a CT absorption band centered at similar wavelength for CTC-2 as observed in *Bs*FNR works (Seo et al. 2016). In the NADPH oxidation reaction by *Ct*FNR_{ox}, *k*_{obs} and Δ*A*₄₆₀ of the slower phase affected by the NADPH concentration (Fig. 2c,d). As the NADPH concentration dependent second order reaction step in Scheme 1 (i.e. formation of MC-1) was rapid, such the dependency could be explained by the formation of NADPH-*Ct*FNR_{red} complex (Scheme 3) (Batie and Kamin 1986).



Scheme 3

For NADPH oxidation by *Ct*FNR_{ox}, we could not determine the rate constants of the hydride transfer step in both direction (*k*₁ and *k*₂ in Scheme 1), because the fast phase was too rapid even at 50 μM NADPH (Fig. 2c). If we assume that *Ct*FNR_{ox} and CTC-1 have a similar extinction coefficient at 460 nm (inset of Fig. 2a) and that the hydride transfer step is the rate determining, Scheme 1 can be approximated by the two-component reversible reaction model CTC-1 ⇌ CTC-2, where the rates of the forward and reverse directions corresponds to the respective hydride transfer rates in Scheme 1. According to this approximation, the ratio of CTC-1:CTC-2 at equilibrium at low NADPH concentration (Fig. 1a) corresponds to the ratio of *k*₁:*k*₂. From the spectrum at 4 ms in Fig. 1a, we can

estimate a CTC-1: CTC-2 ratio of 3:7, suggesting that k_2 is comparable to k_1 . This result is in contrast to that of *BsFNR* where the hydride transfer rate in NADPH oxidation direction was much larger than that of NADP⁺ reduction direction ($\sim 500\text{ s}^{-1}$ and $< 10\text{ s}^{-1}$, respectively, (Seo et al. 2016)). Similar reversibility has been reported on plant-type FNRs from photosynthetic organisms (Sánchez-Azqueta et al. 2012; Tejero et al. 2007).

Generally, formations of CTCs are considered to lead the stacking of the two-ring system, the isoalloxazine ring and the nicotinamide ring moieties of FAD and NAD(P)⁺/NAD(P)H, respectively, which enables efficient hydride transfer between C4 of nicotinamide ring portion of NAD(P)⁺/NAD(P)H and N5 of isoalloxazine ring portion of FAD in the reaction catalyzed by flavin-dependent NAD(P)H-oxidoreductases (Blankenhorn 1975; Tejero et al. 2007; Peregrina et al. 2010; Sánchez-Azqueta et al. 2014). The pre-steady state reaction analyses of WT and mutated plant-type *Anabaena* FNRs with NADP⁺/NADPH revealed a significant correlation between the absorbance of CT bands and hydride transfer rates (Peregrina et al. 2010). In present work, NADPH oxidation by *CtFNR*_{ox} with larger k_{obs} (Fig. 2) was accompanied by an appearance of a CT absorption band, whereas NADP⁺ reduction reaction with smaller k_{obs} did no obvious CT absorption band (Fig. 4). In the NADP⁺ reduction by *CtFNR*_{red}, as the slope of the initial increasing phase in A_{460} was not affected by the NADP⁺ concentrations used (Fig. 4c), it seemed that the rate of the complex formation with NADP⁺ was saturated with respect to NADP⁺ concentration. Accordingly, the rate limiting step in NADP⁺ reduction by *CtFNR*_{red} would be present after the formation of *CtFNR*_{red}-NADP⁺ complex. If the process of NADP⁺ reduction by *CtFNR*_{red} occurred along the same conformations in NADPH oxidation by *CtFNR*_{ox}, i.e. kinetically reversible, the hydride transfer process in NADP⁺ reduction by *CtFNR*_{red} would be rapid the same as in the NADPH oxidation by *CtFNR*_{ox} ($k_{\text{obs}} = k_1 + k_2$). According to this scenario, the transition of *CtFNR*_{red}-NADP⁺ complex to CTC-2 would be the rate limiting and CTC-2 would not accumulate during the reaction, but the rapid formation of CTC-1 would give an absorption in CT band region in the NADP⁺ reduction by *CtFNR*_{red}. Obtained results, however, provided almost no CT absorption bands during the NADP⁺ reduction by *CtFNR*_{red}. One can consider that rapid and strong *CtFNR*_{ox}-NADP⁺ complex formation will decrease the absorbance of CT band region (Scheme 2). But mixing *CtFNR*_{red} with near equimolar of NADP⁺ exhibited a faster formation of *CtFNR*_{ox} followed by a slower formation of CTCs (Fig. 3e). These results seem to indicate a difference in the conformation of CTCs between in the NADP⁺ reduction by *CtFNR*_{red} and in the NADPH oxidation by *CtFNR*_{ox}. In the NADPH oxidation reaction, the nicotinamide ring reacts with the oxidized isoalloxazine ring, whereas in the NADP⁺ reduction reaction, the nicotinamide ring reacts with the reduced isoalloxazine ring. The crystal structure analysis of *BsFNR* with NADP⁺ revealed that the nicotinamide ring portion of NADP⁺ attached on the surface of the NADPH-binding domain with substantial electron density and within a hydrogen-bonding distance to Asp177 (Fig. 1a). In this conformation, S⁻²H at C4 of the nicotinamide moiety of NADPD can face to the N5 of isoalloxazine

ring portion by the domain motion, which is consistent with the results of the primarily kinetic isotope effect (Seo et al. 2016). Similarly, structurally related *Ec*TrxR also provided a substantial isotope effect in steady-state assay with *S*-²H NADPD (Williams Jr. 1995). The crystal structure analyses of *Ec*TrxR revealed that the isoalloxazine ring portion of the reduced FAD hydroquinone was in a bent conformation whereas that in the oxidized FAD quinone was in a planar one (Lennon et al. 1999), suggesting that a conformational change of the isoalloxazine ring portion of FAD prosthetic group can occur during the redox transition. In the crystal structure of *Bs*FNR_{ox} and *Ct*FNR_{ox}, the isoalloxazine ring portion was in a planer conformation (Komori et al. 2010; Muraki et al. 2010), but we have no structural information about the conformation of isoalloxazine ring portion in the reduced TrxR-type FNR. On the UV-vis spectra, *Bs*FNR_{red} had an apparent small absorption band with a peak at 430 nm, which red-shifted in the presence of NADP⁺ (Seo et al. 2016) whereas in this work, *Ct*FNR_{red} had only a small shoulder at 410 nm (Fig. 4a, b). According to the previous reports discussing the relation between the spectroscopic properties and the conformation of the isoalloxazine ring portion of the reduced flavin hydroquinone (Ghisla et al. 1974; Blankenhorn 1975), it is likely that the isoalloxazine ring moiety in *Bs*FNR_{red} was in a more planer conformation whereas that in *Ct*FNR_{red} was in a bent one (Ghisla et al 1974). Actually *Bs*FNR demonstrated the rapid formation of CTC-2 with an apparent CT absorption band in the NADP⁺ reduction reaction (Seo et al. 2016), indicating that the conformation of the isoalloxazine ring portion would affect the accessibility and /or distance of the nicotinamide ring portion to the isoalloxazine ring moiety. In addition, the positioning of the C-terminal extension region might play an important role during the reaction as this region hamper the stacking of the rings (Fig. 1a). Further works especially focused on the C-terminal region should be performed to reveal the mechanism of the redox reactions between TrxR-type FNRs and NADP⁺/H.

Redox equilibrium between NADP⁺/NADPH and *Ct*FNR shows that *Ct*FNR can catalyze NADP⁺/NADPH reduction/oxidation reversibly, which is in accordance with previous report (Seo and Sakurai 2002) and its physiological requirements. However, in our *in vitro* experiments in the absence of Fd, the rate of NADP⁺ reduction was rather slow compared with those by plant-type FNRs (Batie and Kamin 1984; Tejero et al. 2007) and steady-state assay (Seo and Sakurai 2002). In the report by Batie and Kamin (1984), the rate of NADP⁺ reduction was enhanced by the presence of Fd, which led to the conclusion that ternary complex formation is involved in the reaction. The effects of Fd on the kinetics of *Ct*FNR reactions remain to be investigated.

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Figure legends

Fig. 1. (a) Close-up view around the isoalloxazine ring moiety of the FAD prosthetic group and NADP⁺-binding domain in the crystal structure of *CtFNR* (PDB code: 3AB1). Main chains of subunit A which includes the FAD prosthetic group, and C-terminal region of subunit B (I327 to A348) in *CtFNR* are colored in green and purple, respectively. NADP⁺ is superimposed based on the superposition of NADPH-binding domain of *BsFNR* which binds NADP⁺ (Komori et al 2010, PDB code 3LZW) against *CtFNR* on UCFS Chimera package (Pettersen et al. 2004). Side chains of Tyr57, Asp64 and Asp177 in subunit A, and Phe337 and Ser339 in subunit B are depicted as a ball and stick model. Hydrogen-bondings between Tyr57 and O2' hydroxyl group of ribitol moiety of FAD, Asp64 and N3 of isoalloxazine ring portion, Asp177 and N7 of nicotinamide ring portion of NADP⁺, and Ser339 and N5 of isoalloxazine ring portion of FAD are depicted as red dotted lines. The figure was prepared using Discovery Studio 4.0 Visualizer (Accelrys Inc., USA). **(b)** Partially aligned amino acid sequences around the *si*-face and *re*-face residues in TrxR-type FNRs. The number of the first amino acid residue in each row is indicated at the left. The positions of the Tyr57, Asp64, Phe337 and Ser339 residues in *CtFNR* sequence are indicated by arrows at upside. *bsu_BSU32110*: *Bacillus subtilis* subsp. *subtilis* str. 168 FNR, *rpa_RPA3954*: *Rhodopseudomonas palustris* CGA009 FNR, *ttj_TTHA0465*: *Thermus thermophilus* HB27 FNR, *cte_CT1512*: *Chlorobaculum tepidum* FNR, *cli_Clim_1719*: *Chlorobium limicola* DSM245 FNR.

Fig. 2. (a, b) Transient spectra induced by mixing 9.0 μM *CtFNR*_{ox} with a. 100 μM or b. 500 μM NADPH. The reaction was performed in 20 mM HEPES-NaOH buffer (pH 7.0) at 10°C. The spectra at 0 ms and 2,000 ms are shown as thin continuous lines. The spectra shown by thin dotted lines from the top to the bottom at 450 nm correspond to those at 2, 4, 9, 19, 49, 99, 199 and 499 ms, respectively. The spectrum of *CtFNR*_{ox} in the absence of NADPH is shown as a thick continuous line, and that of *CtFNR*_{red} (see Fig. 4) as a thick dashed line, respectively. Arrows indicate the directions of the absorbance change at the respective wavelengths; the dashed parts of the arrows indicate change that occurred within the first data acquisition period (1 ms). The inset in Fig. 2b shows the time course of A_{590} at a. 100 μM and b. 500 μM NADPH. The data of the inset are the average of five measurements. **(c)** The time course of A_{460} after mixing *CtFNR*_{ox} with NADPH. The measurement conditions were the same as those in Fig. 2a, except the NADPH concentrations of a: 0 μM , b: 50 μM , c: 100 μM d: 200 μM and e: 500 μM . The data are an average of four to five measurements. The time course in the short time range is shown in the inset. Double exponential decay curves based on the data for each NADPH concentration in the time range 5 to 500 ms are indicated by continuous red lines. **(d)** NADPH concentration dependency of k_{obs} of the slow phase. The observed rate constant at each NADPH concentration was obtained as shown in Fig. 2c. One standard deviation of each observed rate constant was less than 0.6 s⁻¹.

Fig. 3. (a, b) Transient spectra induced by mixing 11 μM $CtFNR_{\text{ox}}$ with a. 100 μM or b. 500 μM NADPD. The reaction was performed in 20 mM HEPES-NaOH buffer (pH 7.0) at 10°C. The spectra at 0 ms and 1,999 ms are shown as thin continuous lines. The spectra shown by thin dotted lines from the top to the bottom at 450 nm correspond to those at 2, 4, 9, 19, 49, 99, 199 and 499 ms, respectively. The spectra of 11 μM $CtFNR_{\text{ox}}$ in the absence of NADPD and $CtFNR_{\text{red}}$ in 20 mM HEPES-NaOH buffer (pH 7.0) are shown as a thick continuous line and a thick dashed line, respectively. Arrows indicate the directions of change in absorbance at the respective wavelengths; the dashed parts of arrows indicate change that occurred within the first acquisition period (1 ms). The insets in Fig. 3a and b show time courses of ϵ_{460} (Fig. 3a) and ϵ_{590} (Fig. 3b) after mixing $CtFNR_{\text{ox}}$ with 100 μM NADPH (\circ) and NADPD (Δ) shown in Fig. 2a and 3a, respectively. The absorbance was normalized by $CtFNR$ concentration. The dotted line in the inset of Fig. 3a represents the absorption of $CtFNR_{\text{ox}}$. (c) The time course of A_{460} after mixing $CtFNR_{\text{ox}}$ with NADPD. The measurement conditions were the same as those in Fig. 3a, except the NADPD concentrations of 0 μM (closed circle), 100 μM (open circle), 200 μM (open triangle) and 500 μM (open square). The data are an average of four to five measurements. The time course in the short time range is shown in the inset.

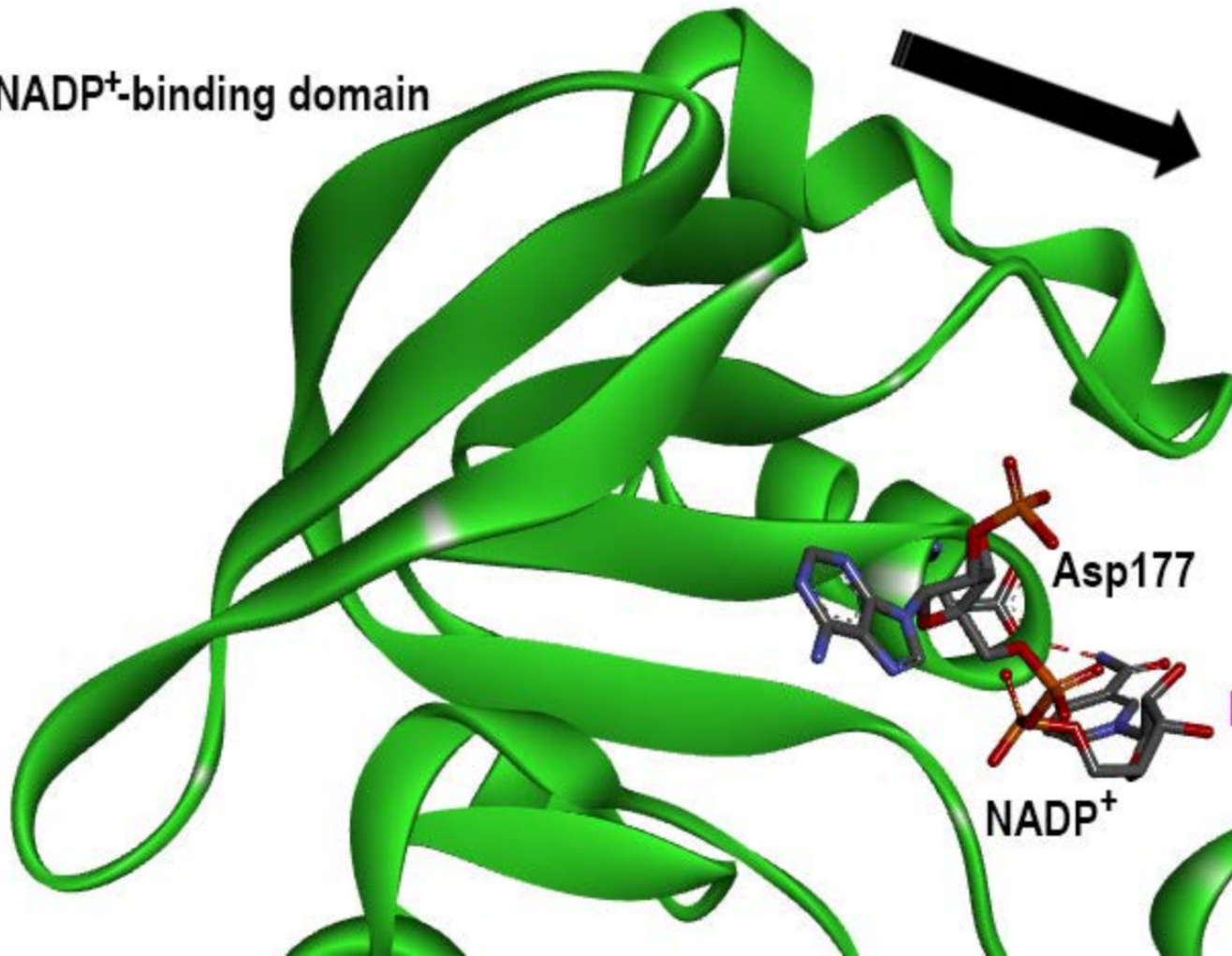
Fig. 4. (a, b) Transient spectra induced by mixing chemically reduced 12 μM $CtFNR$ with a. 100 μM and b. 500 μM NADP^+ . The reaction was performed in 20 mM HEPES-NaOH buffer (pH 7.0) at 10°C. The spectra of 12 μM of $CtFNR_{\text{ox}}$ and chemically reduced $CtFNR_{\text{red}}$ in 20 mM HEPES-NaOH buffer (pH 7.0) are represented by thick dashed and thick continuous lines, respectively. Spectra for 0 ms and 3,999 ms are indicated by thin continuous lines, and for 9, 19, 49, 99, 199, 499, 999 and 1,999 ms, respectively, by thin dotted lines from the bottom to the top at 450 nm. Arrows indicate the direction of change in absorbance. (c) The time course of A_{460} after mixing $CtFNR_{\text{red}}$ with NADP^+ . The measurement conditions were the same as those in Fig. 4a, except the NADP^+ concentrations of a: 0 μM , b: 50 μM , c: 100 μM d: 200 μM and e: 500 μM . The data are an average of four to five measurements. The time course in the short time range is shown in the inset. Double exponential decay curves based on the data for each NADP^+ concentration are indicated by continuous red lines. (d) NADP^+ concentration dependency of the observed rate of the faster phase. The observed rate constant at each NADP^+ concentration was obtained as shown in Fig. 4c. One standard deviation of each observed rate constant was less than 0.2 s^{-1} . (e) Transient spectra induced by mixing chemically reduced 24 μM $CtFNR$ with 25 μM NADP^+ . The reaction was performed in 20 mM HEPES-NaOH buffer (pH 7.0) at 10°C. The spectra of 24 μM of $CtFNR_{\text{ox}}$ and chemically reduced $CtFNR_{\text{red}}$ in 20 mM HEPES-NaOH buffer (pH 7.0) are represented by thick dashed and thick continuous lines, respectively. Spectra for 0 ms, 1 s and 20 s are indicated by thin continuous lines, and for 9 ms, 19 ms, 49 ms, 99 ms, 199 ms, 499 ms, 2 s, 5 s and 10 s, respectively, by thin dotted lines from the bottom to

the top at 450 nm. Arrows indicate the direction of change in absorbance. Inset of Fig. 4e represent the time course of A_{460} (\circ) and A_{590} (\square). The data are average of four measurements. Double exponential decay curves based on the data for both wavelengths are indicated by continuous red lines.

NADP⁺-binding domain

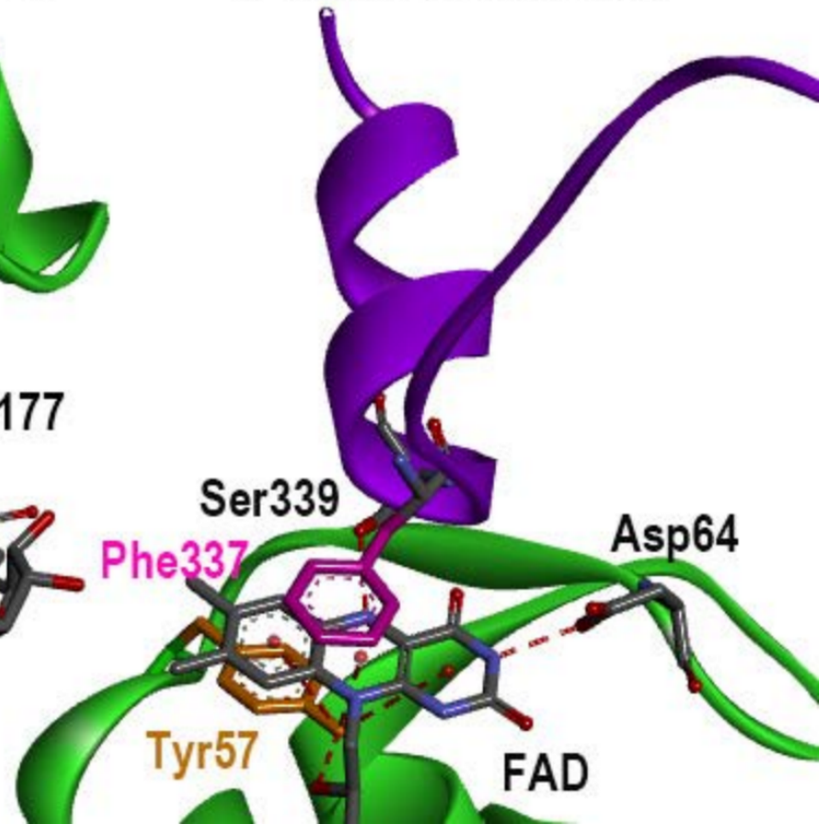


C-terminal extension



Asp177

NADP⁺



Ser339

Phe337

Tyr57

Asp64

FAD

Figure 1b

si-face

```
bsu_BSU32110      43...GGQLSALYPEKYIYDVAG...
rpa_RPA3954       42...GGQCAELYPEKPIYDIPG...
ttj_TTHA0465      41...GGQLTALYPEKYIYDVAG...
cte_CT1512        50...GGQLAALYPEKHIYDVAG...
cli_Clim_1719     50...GGQLAALYPEKHIYDVAG...
                  *** : ***** ***.*
```

re-face

```
bsu_BSU32110      313...YMDPKARVQPLHSTSLFENK
rpa_RPA3954       317...YVYPDKRVVFQYITTSSTNLQKKLGVN
ttj_TTHA0465      315...YANPALKVNPGHSSSEKAAPGT
cte_CT1512        326...YIKPGEKIRNVFSSVKMAKEKKAEEAGNATENKAE
cli_Clim_1719     326...YIRPGEKIRHTFSSVKMAKQKKNEQ
                  * * :: .::
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Figure 2a

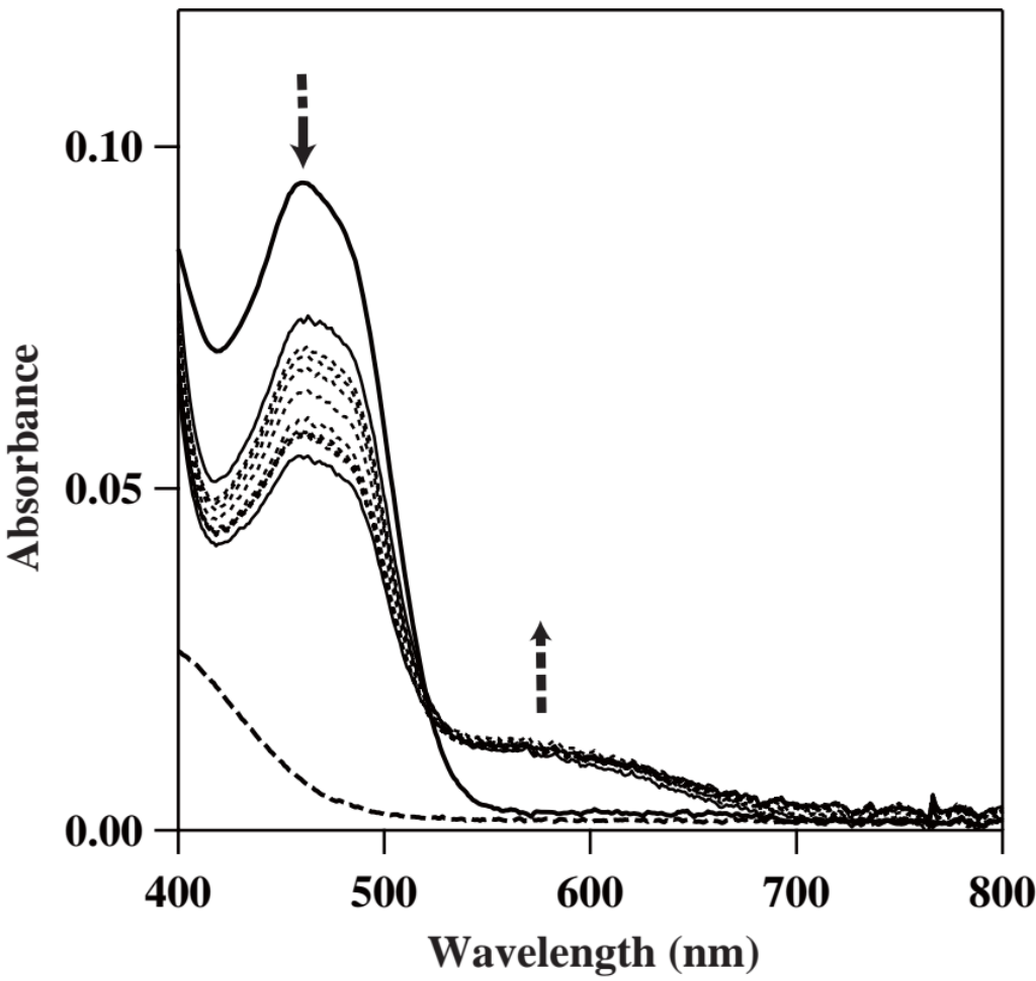


Figure 2b

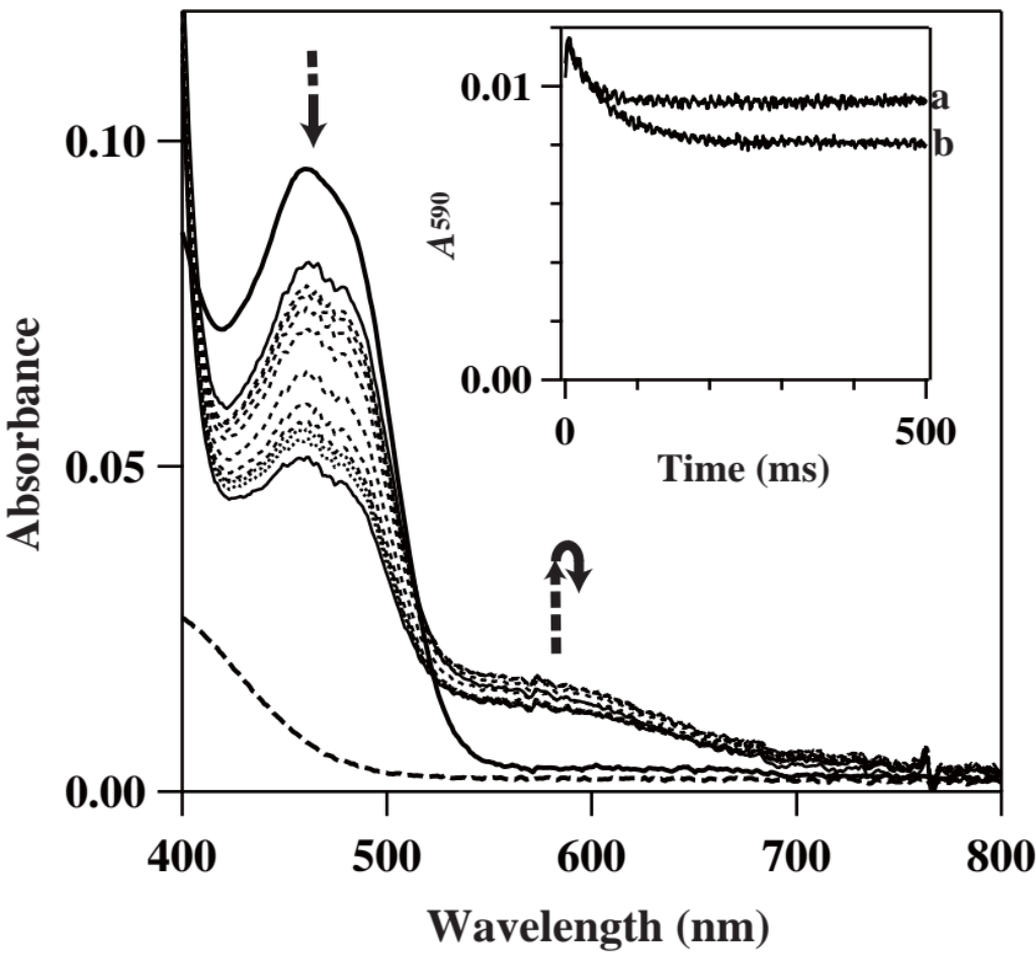


Figure 2c

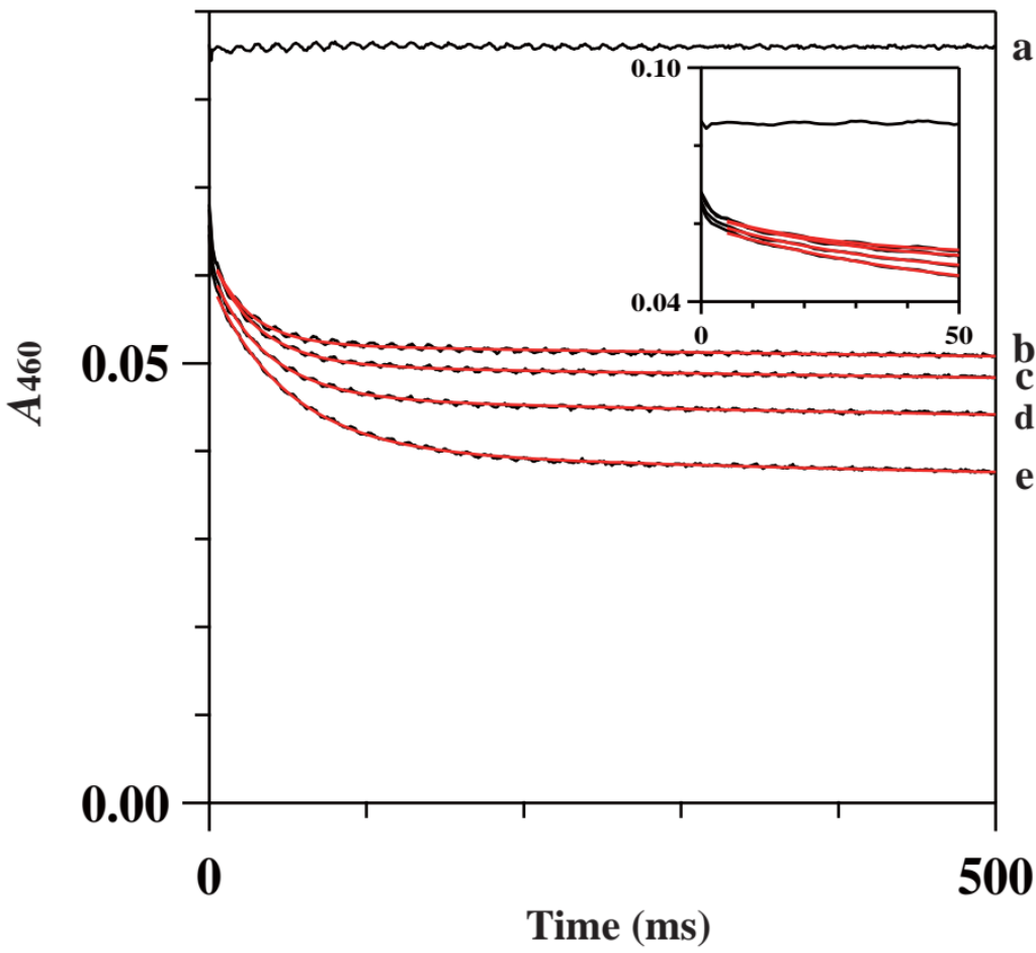


Figure 2d

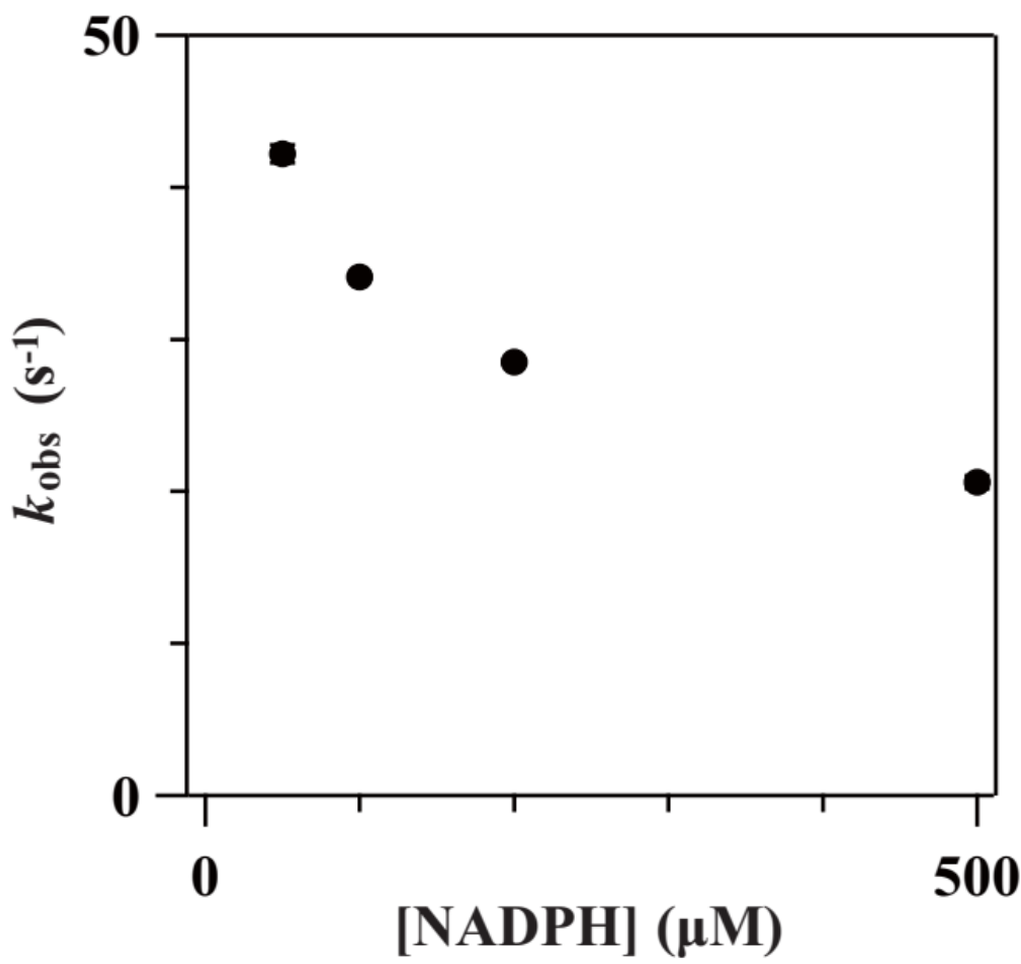


Figure 3a

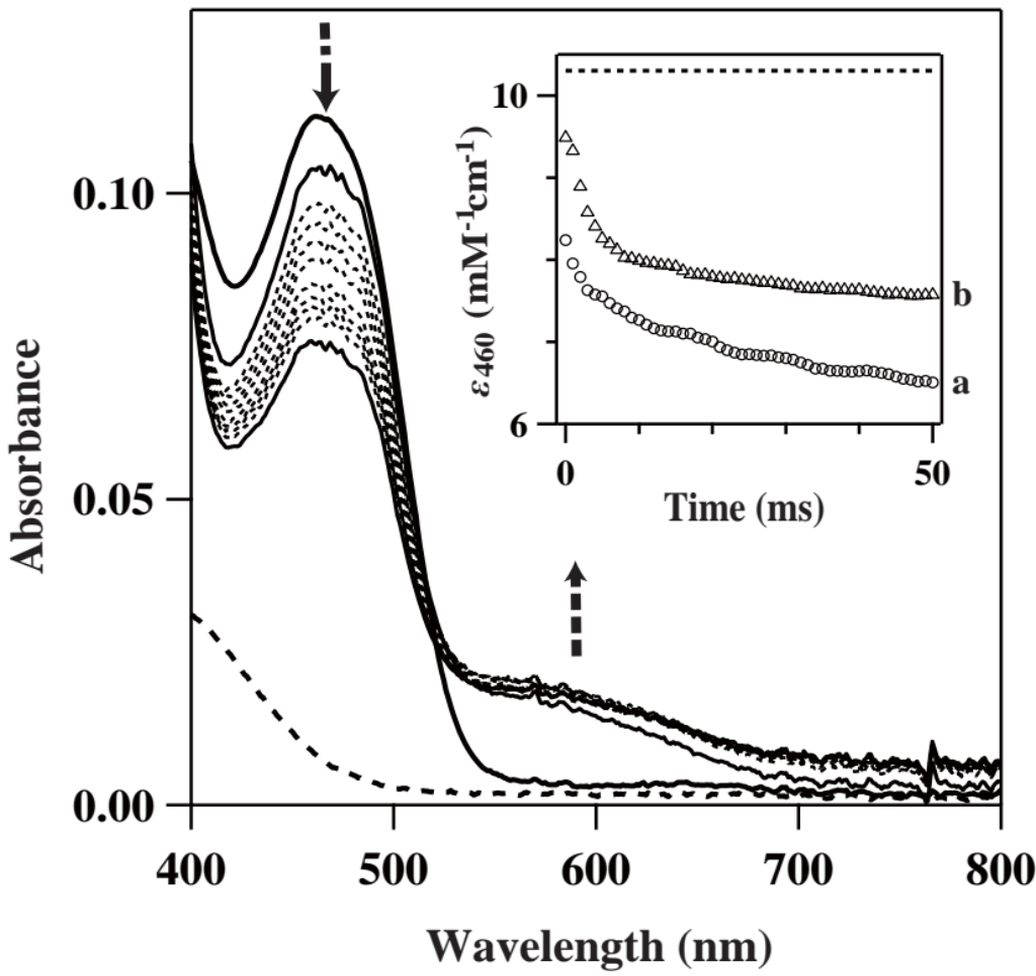


Figure 3b

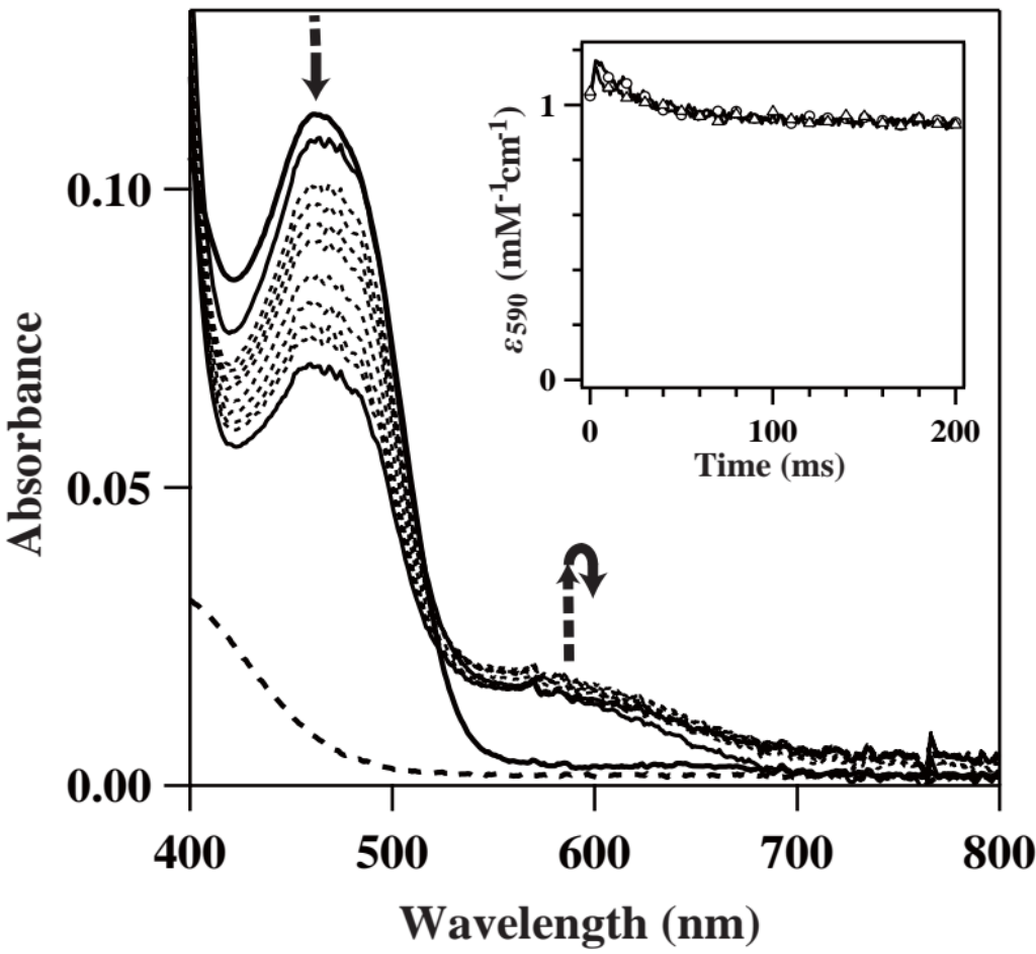


Figure 3c

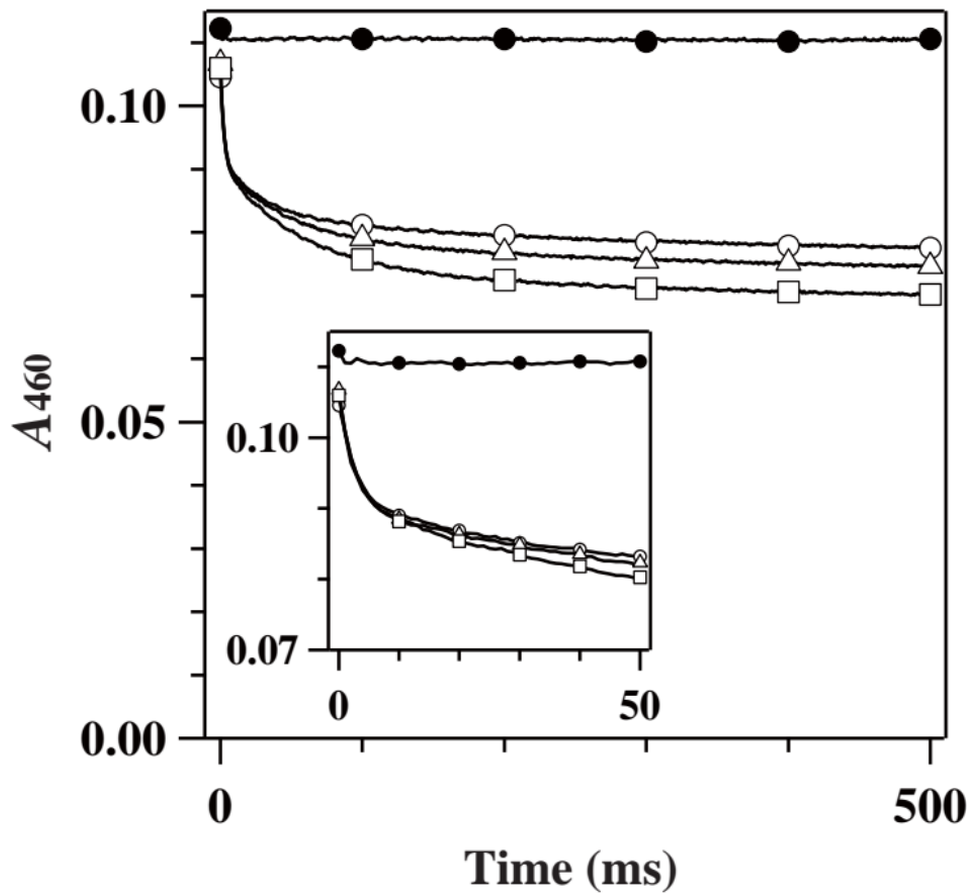


Figure 4a

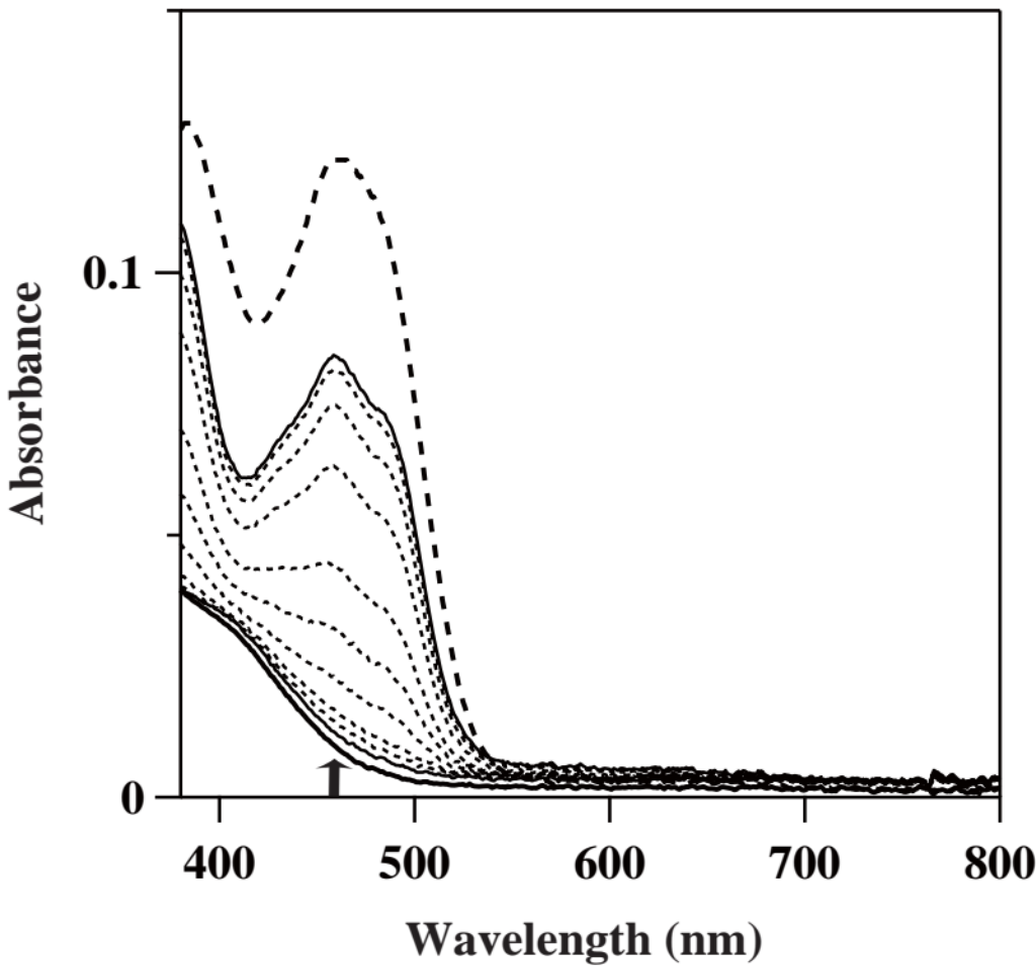


Figure 4b

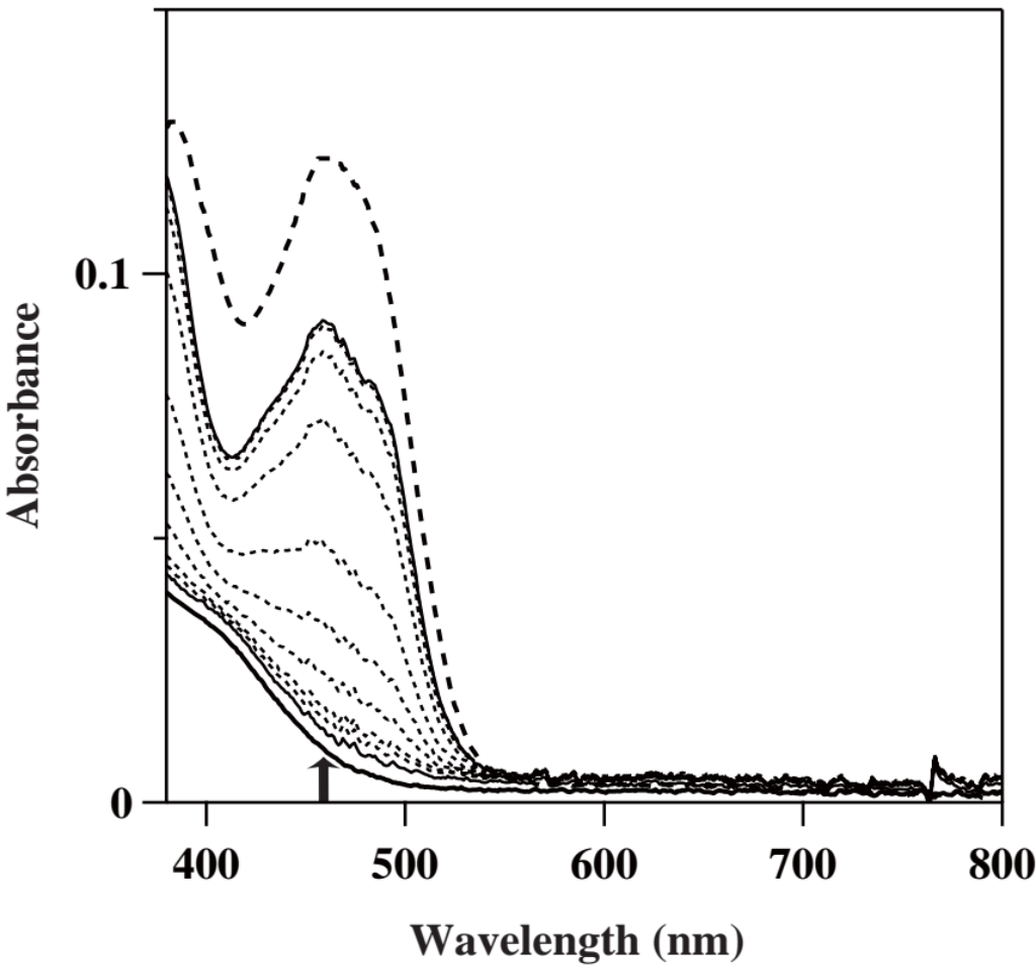


Figure 4c

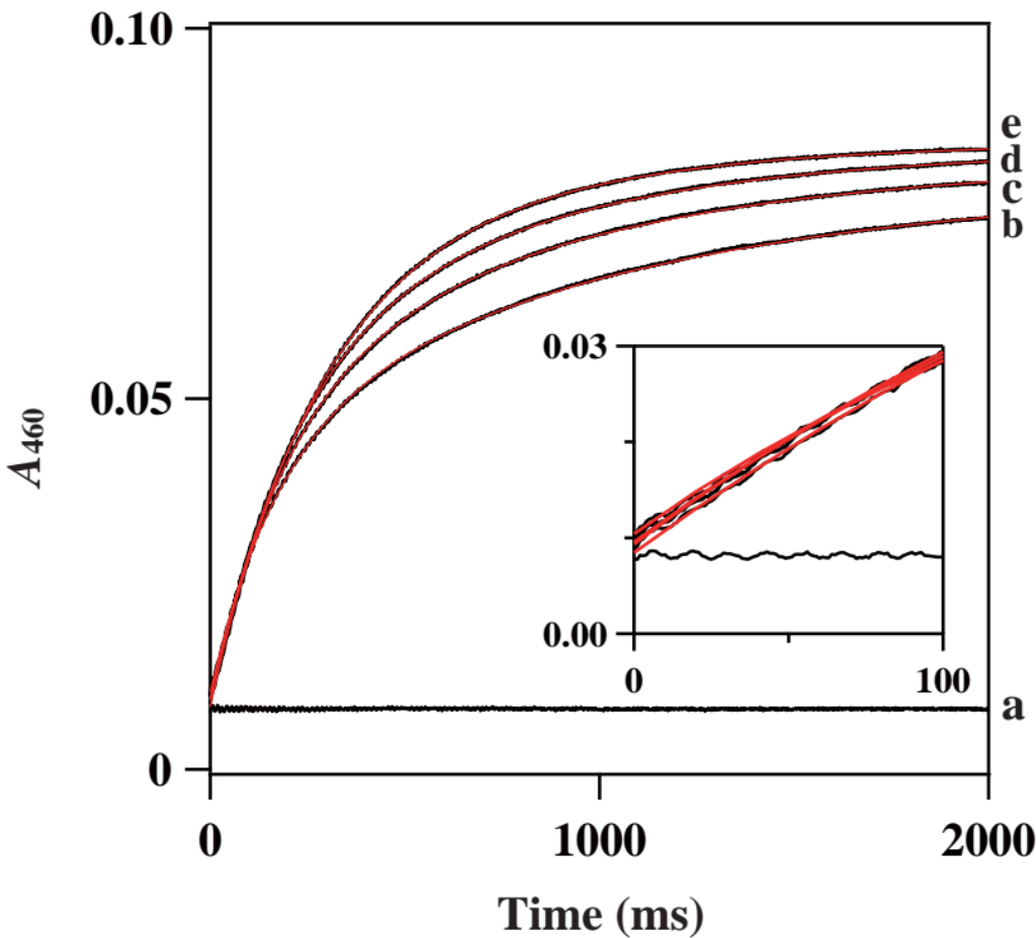


Figure 4d

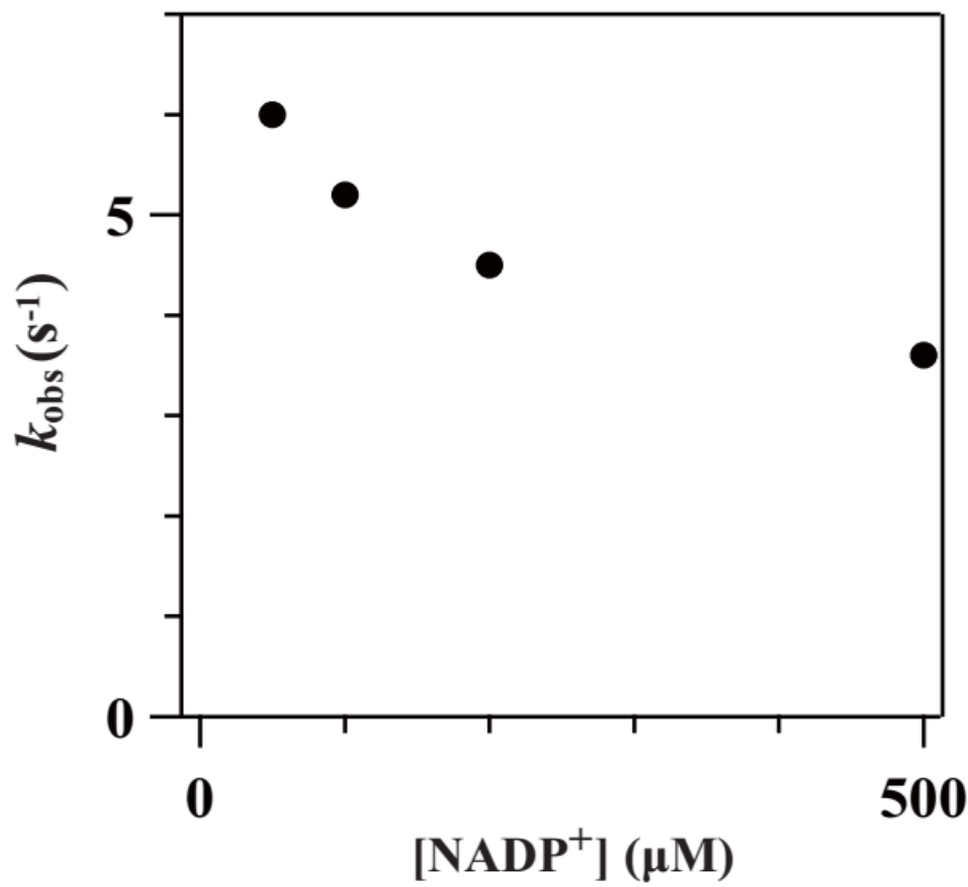


Figure 4e

