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Redox cycling and kinetic analysis of single molecules of solution-phase nitrite reductase

Randall H. Goldsmith^{a,2}, Leandro C. Tabares^{b,c}, Dorota Kostrz^d, Christopher Dennison^d, Thijs J. Aartsma^b, G. W. Canters^{b,1}, and W. E. Moerner^{a,1}

^aDepartment of Chemistry, Stanford University, Stanford, CA 94305; ^bHuygens Laboratory, Leiden Institute of Physics, Leiden University, Niels Bohrweg 2, 2333CA, Leiden, The Netherlands; ^cCommissariat à l'Énergie Atomique, Institut de Biologie et de Technologies de Saclay, Service de Bioénergétique, Biologie Structurale et Mécanismes, 91191 Gif-sur-Yvette Cedex, France; and ^dInstitute for Cell and Molecular Biosciences, Medical School, Newcastle University, Newcastle upon Tyne NE2 4HH, United Kingdom

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Single-molecule measurements are a valuable tool for revealing details of enzyme mechanisms by enabling observation of unsynchronized behavior. However, this approach often requires immobilizing the enzyme on a substrate, a process which may alter enzyme behavior. We apply a microfluidic trapping device to allow, for the first time, prolonged solution-phase measurement of single enzymes in solution. Individual redox events are observed for single molecules of a blue nitrite reductase and are used to extract the microscopic kinetic parameters of the proposed catalytic cycle. Changes in parameters as a function of substrate concentration are consistent with a random sequential substrate binding mechanism.

ABEL trap | enzymology | electron transfer | fluorescence | catalysis

Measurements made on the activity of single enzymes have revolutionized the picture of these molecular machines' mode of operation by providing evidence for static and dynamic heterogeneity, enabling the observation of transient intermediates, and allowing determination of the microscopic rate constants comprising complex catalytic cycles under turnover conditions (1–4). However, conclusions regarding enzyme dynamics require the observation of multiple turnovers of a single molecule with a requisite observation window of several seconds or more. Because this window is several orders of magnitude longer than the typical dwell time of a solution-phase molecule diffusing through a confocal detection volume, molecules that naturally exist in solution are often immobilized on a solid support (1–3, 5). Although gross comparison between properties measured on immobilized and freely diffusing biomolecules tends to show good agreement (1, 3), more subtle aspects of biomolecule behavior such as property distribution widths and shapes (6, 7) and conformational dynamics (6, 8) have been shown to be affected by immobilization, and in some cases, immobilization has been suspected of contributing to observed enzyme behavioral heterogeneity (4). In this report we take an important step toward viewing the richness of single-enzyme dynamics without immobilization by making observations of multiple turnover events of single solution-phase molecules of a blue nitrite reductase (bNiR), an important redox-active enzyme in bacterial denitrification (9).

Prolonged solution-phase measurements are enabled through use of a specialized microfluidic trapping device that cancels the Brownian motion of a single emissive solution-phase object through the use of directed electroosmotic forces, the anti-Brownian electrokinetic (ABEL) trap (10, 11). The ABEL trap estimates particle position via a revolving laser spot phase-locked in a homebuilt analogue circuit to a restoring force gated by photon detection and applied via two orthogonal pairs of electrodes (11). This scheme results in the confinement of the object in a spatially homogeneous excitation volume for multiple seconds without immobilization and can be applied to single biomolecules substantially smaller than the large dielectric beads that are trapped with optical tweezers. We have previously used the ABEL trap to study

conformational dynamics of single photosynthetic antenna proteins (6) and fluorescently labeled DNA molecules (12).

bNiR is a multicopper enzyme that carries out the one-electron reduction of nitrite, NO_2^- , to nitric oxide, NO, in the presence of a suitable reductant (9). It is a trimer, 10 nm in diameter, and contains two different Cu sites (13). The type 1 (T1) Cu, coordinated by two His, a Cys, and a Met residue (13), is the initial site of reduction by an electron donor. Intramolecular electron transfer (ET) (14) over 12.6 Å results in reduction of the proximal type 2 (T2) Cu, which is located at the monomer-monomer interface and is coordinated by three His residues (13). One of these T2 Cu ligands is adjacent to the Cys residue at the T1 site, and this covalent network has been proposed as the pathway for ET (15–17). Nitrite reduction occurs at the T2 Cu with protons for water formation being provided by a hydrogen bond network composed of a bound water, and the second coordination sphere Asp92 and His249 residues (18). Recently, it has been suggested that the intramolecular ET event is coupled to proton transfer (19). Considerable mechanistic interest on nitrite reductases (NiRs) has focused on whether intramolecular ET proceeds consistently before or after nitrite binding (ordered mechanisms) or via a mixture of both pathways (random-sequential mechanism) (20, 21). Much seemingly conflicting evidence has appeared for both mechanisms, with ET without nitrite binding occurring via pulse radiolysis (17), ET shown to be blocked in the absence of nitrite via X-ray absorption spectroscopy (22), and both pathways shown to be operating simultaneously (a random-sequential mechanism) via activity assays and protein film voltammetry (21).

The T2 Cu possesses no significant absorption in the visible range (23). However, the oxidized T1 Cu possesses an intense $\text{S}(\text{Cys}) \rightarrow \text{Cu}(\text{II})$ charge transfer transition giving rise to strong visible absorption (9). This absorption vanishes upon reduction of the T1 Cu (23). This spectral change was recently employed to make a proximal dye molecule a fluorescent reporter of the T1 Cu oxidation state (24). The oxidized Cu significantly quenches the dye fluorescence via Förster resonant energy transfer (FRET) due to overlap between the dye's emission and the colored Cu's absorption, whereas the reduced colorless Cu leaves the dye unquenched and maximally emissive (24), Fig. 1. Single green NiR enzymes (gNiR) (25) from *Alcaligenes faecalis* were immobilized and fluorescence intensity fluctuations were shown to be indicative of enzyme turnovers (24), although individual turnovers

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¹To whom correspondence may be addressed. E-mail: wmoerner@stanford.edu or canters@chem.leidenuniv.nl.

²Present address: Department of Chemistry, University of Wisconsin, Madison, Wisconsin 53706.

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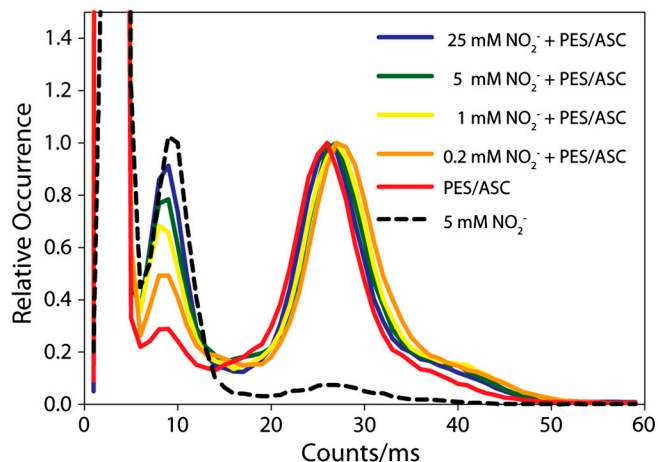


Fig. 3. Histogram of intensities compiled from many molecules under varying conditions at 10 ms binning normalized to the central peak near 25 counts/ms. The amplitude of the lower intensity peak is seen to rise as a function of substrate concentration. The lower peak is seen to nearly disappear when no substrate is added, whereas the upper intensity peak is seen to nearly disappear when no reductant is added. The off-scale peak at lowest intensity is from background.

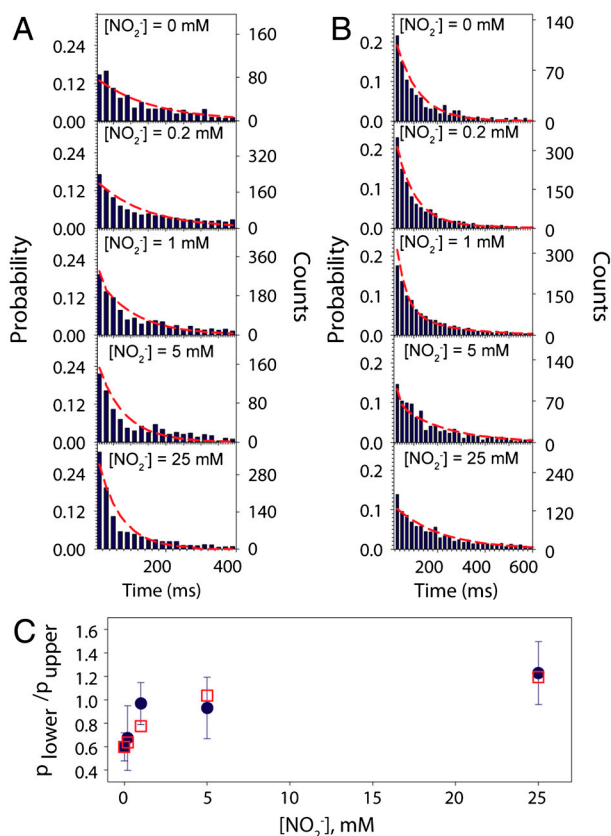


Fig. 4. Fits for dwell times of the upper intensity state (A), lower intensity state (B), and population ratio (C), at multiple substrate concentrations for one representative dataset. In A and B, dwell times in 20-ms bins (dark blue bars) are fit with red lines that follow the function forms derived from Fig. 6A. In C, population ratios calculated from the determined parameters (red squares) are seen to fall within the error bars of the measured ratios (blue circles). Increasing substrate concentration results in a shift toward shorter dwell times in the upper state, a shift to longer dwell times in the lower state, and a larger fraction of time spent in the lower state.

Discussion

To model changes in dwell time distributions, we began with an ordered kinetic scheme (Fig. 5). Kinetic equations for single-molecule dynamics are expressed in terms of probabilities rather than concentrations. Expected dwell time distributions are determined by differentiating the integrated rate law with respect to time (*SI Appendix*) (28, 29).

Assuming the validity of the scheme in Fig. 5, dwell times in the upper state, RO (where the first letter, R, describes the oxidation state of the T1 Cu and the second letter, O, describes the oxidation state of the T2 Cu, see Fig. 5), can be modeled as a simple monoexponential decay to directly extract the rate constant of intramolecular ET, k_{23} . These fits are shown in *SI Appendix*, Fig. 3. Surprisingly, k_{23} is not a simple rate constant, as it is observed to increase as a function of nitrite concentration (Fig. 5B). Previously, conflicting effects of nitrite binding on intramolecular ET have been reported. ET was inhibited by the absence of nitrite binding in crystallized gNiR variants at 100 K (22), although in our data, state shifts in singly reduced bNiR molecules at zero nitrite concentration, summarized in Fig. 4A and B, suggest that k_{23} is not blocked by the absence of substrate under these conditions. Electron-nuclear double resonance measurements (30) and crystallographic studies (31) on the T2 Cu suggest a greater thermodynamic driving force for ET after nitrite bonding due to weaker coordination of His ligands, and a consequent increase in rate constant. In contrast, pulse radiolysis measurements showed a sharp decrease in rate upon nitrite binding (16). Bulk activity assays and electrochemical measurements have suggested that an increase in ET rate constant upon nitrite binding is expected at higher pH values, whereas a decrease upon binding is expected at lower pH, depending on whether the nitrite displaces a water molecule or hydroxide ion (21). Our direct observation of an ET rate constant increase at pH 7.0 is consistent with a shift between a mechanism where ET precedes nitrite binding to one where ET occurs after nitrite binding with a consequent increase in ET rate (21). These changes are in agreement with a random-sequential mechanism whereby ET can occur before or after binding with the relative contributions of each determined by environmental variables such as substrate concentration and pH.

If the intramolecular ET rate constant is affected by nitrite binding via allosteric regulation, where the nitrite interacts with the protein at some site other than the active site, the scheme in Fig. 5A remains valid. However, if as suggested, the ET rate constant is altered by conformational and energetic changes as a result of nitrite binding to the active site, then the scheme in Fig. 5A is too simplistic and a more complex scheme is required that explicitly considers substrate binding. Such an alternative kinetic scheme (24) is shown in Fig. 6A, with the upper right

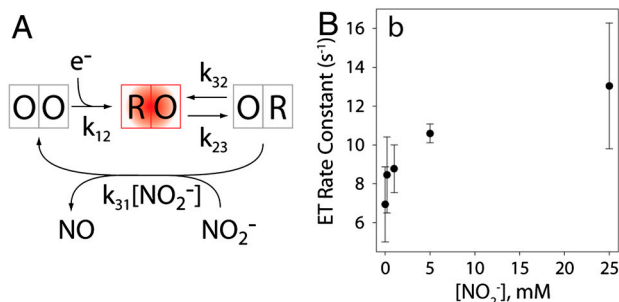


Fig. 5. A simple kinetic scheme and extracted intramolecular ET rate constants. (A) A kinetic scheme where the left side box represents the oxidation state of the T1 Cu (O/O, oxidized/reduced), whereas the right side describes the T2 Cu. Only molecules with a reduced T1 Cu are fluorescent (red highlight). (B) The rate constant of intramolecular ET, k_{23} , is seen to increase as a function of substrate concentration. Error bars are from four independent measurements, each comprised of between one thousand and several thousand individual intensity traces.

Population in the lower state manifold is initially concentrated in the T1 Cu oxidized, T2 Cu reduced (OR) state at vanishing substrate concentration, Fig. 6B. However, this population rapidly decreases with increasing substrate concentration, with the majority of the population in the manifold residing in the OO state, before the bottleneck of slow reduction via the solution phase electron donor. As shown in Fig. 6D, population can enter the lower state manifold at either OR or ORs (both show nonzero values at time zero). As the population in the ROs state increases, the majority of the population is shown to enter via ORs. However, population is quickly shifted to the OO state before leaving the manifold after reduction, k_A .

The scheme in Fig. 6 and parameters in Table 1 describe a system that proceeds with an ordered mechanism at low substrate concentrations where intramolecular ET precedes substrate binding. However, at intermediate concentrations both pathways in the scheme in Fig. 6 are active. In this regime, individual molecules will stochastically follow one or the other pathway with some characteristic partitioning probability, i.e., a random-sequential mechanism. At high substrate concentrations, the system approaches the opposite ordered mechanism, where substrate binding precedes intramolecular ET. In this picture, only the microscopic rate constants in the kinetic scheme determine the partitioning between the two ordered mechanisms.

In contrast to some other single-enzyme experiments (2, 3), product formation was not directly measured. The rate of substrate consumption, $d[\text{NO}_2^-]/dt$, can be calculated from the kinetic scheme. The complex scheme in Fig. 6A will exhibit a slight deviation from Michaelis-Menten (MM) kinetics at high substrate concentration (32–34) (*SI Appendix*, Fig. 5). Nevertheless, approximation by the MM functional form yields k_{cat} , 2.9 s^{-1} and K_m , $860 \text{ }\mu\text{M}$. This value of k_{cat} is well within the range of k_{cat} values established from multiple bulk experiments. A more detailed comparison is presented in *SI Appendix*.

Though measured from observations on individual bNiR molecules, the rate constants shown in Table 1 are ensemble-averaged values. Determination of these parameters for a single enzyme requires a substantially longer viewing window than can be achieved with the current implementation of the ABEL trap and future designs hold promise of extending this window (35). Molecule-to-molecule variability is suggested by the diversity of behavior seen in Fig. 2 but at the moment is not required to explain the observed behavior. Molecule-to-molecule variability has begun to be quantified in immobilized NiR molecules (24, 26).

Individual kinetic parameters can be measured via bulk experiments by tuning reaction conditions to contrived configurations so as to make a specific step rate limiting and thus experimentally accessible for measurement. However, single-molecule measurements allow observation of unsynchronized reaction intermediates, the analysis of which enables simultaneous determination of many kinetic parameters under relevant turnover conditions. The use of the ABEL trap to make these measurements on single solution-phase enzymes opens a window of unprecedented detail into the dynamics of single enzymes in solution.

In conclusion, we extracted kinetic parameters from single molecules of bNiR probed individually in solution. Molecules trapped for multiple seconds and discrete intensity levels corresponding to oxidized and reduced states of the T1 Cu were observed. Extracted parameters lend further support to a random-sequential mechanism and offer direct evidence that an increase in substrate concentration leads to higher rate constants of intramolecular ET under these conditions. Further experiments are ongoing to include effects of pH and to enable long enough trapping windows to allow parameter estimation for individual enzymes.

Materials and Methods

Protein Isolation and Labeling. The Lys329Cys mutant of the bNiR from *Alcaligenes xylosoxidans*, which introduces a Cys residue 2 nm from the T1 Cu within the same monomer, was isolated and purified as described previously (25). The Cys was modified with Atto647N-maleimide yielding protein that was 9% labeled (26).

ABEL Trap Loading Procedure. A 700-nm deep quartz microfluidic cell (11) was cleaned in a UV-ozone cleaner (Jelight) for 30 min, sonicated for 10 min in 1 M potassium hydroxide, thoroughly rinsed with water, and dried with a stream of dry N_2 . Following a previously published procedure (6, 36), a drop of positively charged polyethylene imine (PEI, 1 g/L, pH 8.0, passed through a 0.02- μm filter), was deposited at the microfluidic input port and allowed to enter and coat the cell via capillary action. The cell was allowed to sit for 10 min with the fluid inside. The cell was then blown dry with N_2 , and filled with ultrapure water the same way. The cell was then blown dry, and a drop of negatively charged polyacrylic acid (PAA, 1 g/L, pH 8.0, passed through a 0.02- μm filter) was allowed to coat the inside in the same manner as PEI. The cell was allowed to sit for 10 min, blown dry, and rinsed with water as above. The entire coating procedure was repeated one more time, leaving the cell coated with two alternating pairs of layers, $\text{SiO}_2/\text{PEI}/\text{PAA}/\text{PEI}/\text{PAA}$. If a bare silica cell is used, rampant sticking of bNiR to the surface was observed. With the above coating procedure, sticking was nearly completely eliminated.

Following deposition of the polyelectrolyte multilayer, the dried cell was filled via capillary action with freshly prepared buffer, 1:1 20 mM Hepes, pH 7.0:glycerol with sacrificial electron donor, 5 mM sodium ascorbate (ASC), electron shuttle, 100 nM phenazine ethosulfate (PES) (37), and varying amounts of substrate, sodium nitrite. This solution was allowed to sit for 5 min and then a freshly prepared solution of the same composition but including $<0.1 \text{ nM}$ labeled bNiR was added via capillary action. The loaded ABEL trap cell was then allowed to stand for 5 min before beginning optical measurements.

Optical Setup and Trapping. Excitation was provided by a continuous-wave 638-nm excitation laser (FiberTEC, Blue Sky), which was passed through a 515–640-band-pass filter to eliminate red spectral sidebands. The laser light was steered into the epifoot of a Nikon TE-300 microscope and focused with an oil objective (N.A. = 1.3) to provide an average power of $15 \text{ kW}/\text{cm}^2$ at the sample. Beam deflection to produce the revolving spot was achieved through the same apparatus described previously (11). The excitation spot is $0.6 \text{ }\mu\text{m}$ in diameter in the sample plane, as measured by scanning a fluorescent bead through the focal volume, and is revolved at 40 kHz with a radius of $0.4 \text{ }\mu\text{m}$ in the sample plane. Sample fluorescence is collected back through the same objective and passed through dichroic (645DRLP) and long pass (640ALP) filters, a 200- μm pinhole, and focused onto an Si avalanche photodiode detector (Perkin Elmer, SPCM-CD 2801). Time-correlated single photon counting is achieved using the PicoHarp 300 (Picoquant) timing module. Feedback direction and magnitude are calculated for each 25- μs cycle by phase-sensitive integration of photons detected in the previous cycle. A single detected photon produces a feedback pulse of 12-V vector magnitude lasting 25 μs , which is applied across the 20- μm transverse width of the shallow region of the cell. Voltages are produced by four PA-83 high-voltage operational amplifiers (Apex Microtechnologies) to generate forces in the appropriate direction to cancel Brownian motion. Only objects that maintained residency in the trap for over 200 ms were included in the analysis.

Data Analysis. Time-tagged photon detection data was analyzed in Matlab 7.7. A change-point-finding algorithm (27) was applied with a low critical value and then a thresholding algorithm was used to identify shifts between the upper and lower intensity states and background. A full description of the analysis procedure can be found in *SI Appendix*.

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