



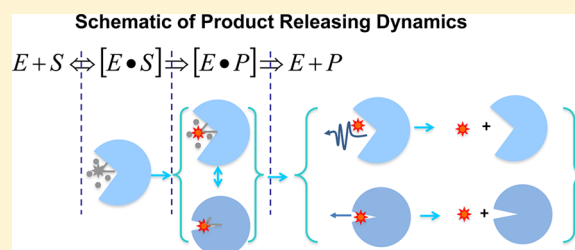
Single-Molecule Enzymatic Conformational Dynamics: Spilling Out the Product Molecules

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Supporting Information

ABSTRACT: Product releasing is an essential step of an enzymatic reaction, and a mechanistic understanding primarily depends on the active-site conformational changes and molecular interactions that are involved in this step of the enzymatic reaction. Here we report our work on the enzymatic product releasing dynamics and mechanism of an enzyme, horseradish peroxidase (HRP), using combined single-molecule time-resolved fluorescence intensity, anisotropy, and lifetime measurements. Our results have shown a wide distribution of the multiple conformational states involved in active-site interacting with the product molecules during the product releasing. We have identified that there is a significant pathway in which the product molecules are spilled out from the enzymatic active site, driven by a squeezing effect from a tight active-site conformational state, although the conventional pathway of releasing a product molecule from an open active-site conformational state is still a primary pathway. Our study provides new insight into the enzymatic reaction dynamics and mechanism, and the information is uniquely obtainable from our combined time-resolved single-molecule spectroscopic measurements and analyses.



INTRODUCTION

Conformational motions of enzymes often play critical roles in enzymatic reactions. A fundamental understanding of enzymatic reaction dynamics and mechanisms relies on the experimental characterization of the specific steps of the enzymatic reactions. Beyond the well-known Michaelis–Menten mechanism, there are a number of fundamental questions that are critical but have not been experimentally answered: for example, how is a product molecule released from an enzymatic active site? Typically, an enzymatic reaction involves multiple kinetic steps, such as substrate binding to form an enzyme–substrate complex, $[E \cdot S]$, nascent product generation, $[E \cdot P]$, and products releasing from the enzyme active site to complete an enzymatic productive turnover, as illustrated in the Michaelis–Menten mechanism, $E + S \rightleftharpoons [E \cdot S] \rightarrow [E \cdot P] \rightarrow E + P$, where E , S , and P represent enzyme, substrate, and product, respectively.^{1–3} The intrinsically dynamic and inhomogeneous processes of enzymatic reactions involve conformational fluctuations ranging over 10^{-15} s to >1 s and 10^{-2} Å to >10 Å, in time and space, respectively.^{4,5} These spatial and temporal dynamics play critical roles in defining the enzymatic free energy potential surfaces, reactive coordinates, and rate processes.^{6,7} Particularly, enzymatic conformational dynamics is essential for enzyme specificity and selectivity. Multiple reaction coordinates of the conformational dynamics in an enzyme complicate the understanding of their structure and function.^{8,9} Therefore, a large variety of experimental and theoretical techniques have been developed to probe internal

dynamics of enzymes, particularly at the picosecond to second time scale.^{10–12}

Single-molecule fluorescence spectroscopy and imaging have been proved as a powerful technique in characterizing enzymatic conformational dynamics and mechanisms.^{4,6,13} Photophysical properties of fluorophores, such as fluorescence intensities, fluorescence quantum yield, and fluorescence decay rate constants, are sensitive to the properties of the local environment.^{14,15} Fluorescence anisotropy provides information about the motions of the protein fragments to which the fluorescent probe is attached.^{16,17} For example, time-resolved anisotropy is capable of analyzing the conformational change fluctuations and the protein flexibility.¹⁶ Furthermore, single-molecule fluorescence lifetime fluctuation measurements are sensitive in characterizing the electrostatic and hydrophobic interactions between the probe molecules and their fluctuating local environment.^{18–21} The intrinsic ability to sense the molecular surrounding of the probe molecule makes the single-molecule spectroscopy an ideal tool to study conformational dynamics of enzymes. Complementary to conventional ensemble-averaged measurements that average over a population, single-molecule measurements are able to detect conformational heterogeneities, identify transient or specific conformations, follow conformational changes, and reveal parallel reaction pathways.^{2,6,22} Generally, three categories of

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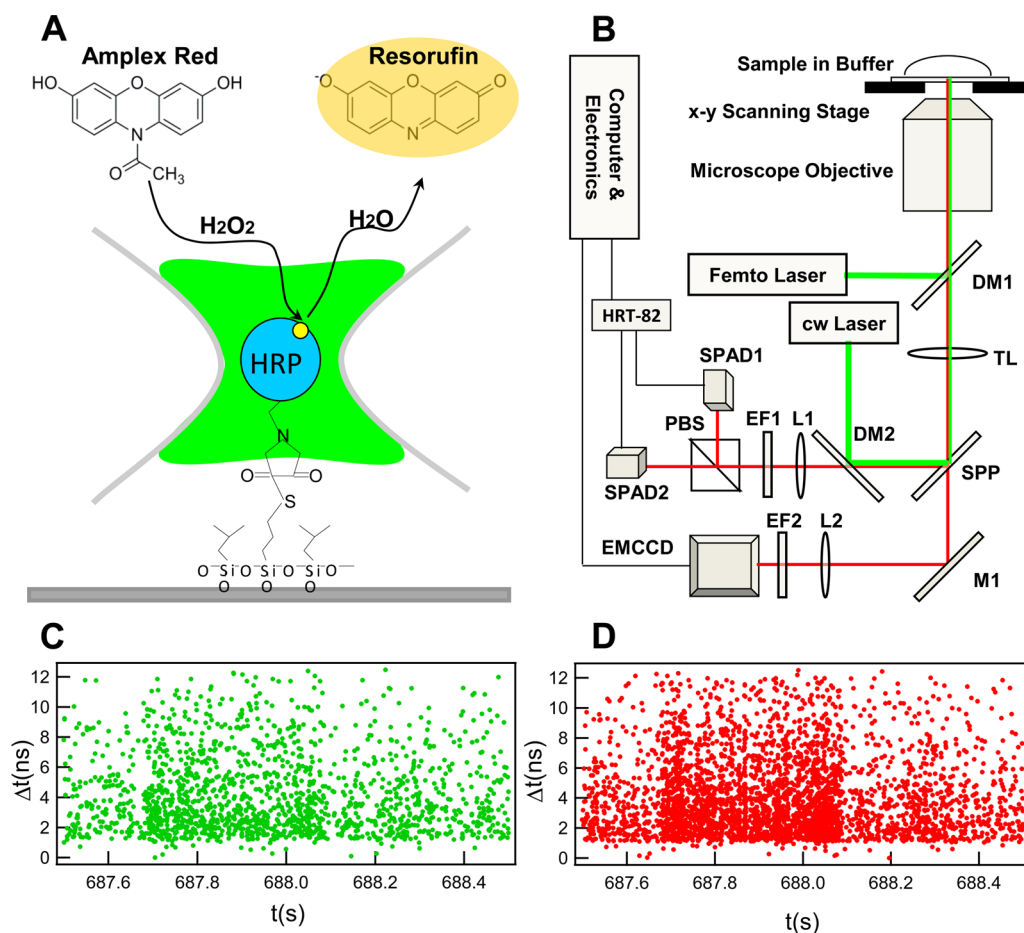


Figure 1. Single-molecule fluorescence experimental scheme. (A) Schematic representation of the enzymatic reaction on (3-aminopropyl) trimethoxysilane modified cover glass. Maleimide-activated HRP is linked to the sulfhydryl ($-SH$) group of 3-mercaptopropyl-trimethoxysilane. Nonfluorescent substrate Amplex red in PBS buffer is converted to fluorescent resorufin product by a single HRP molecule in the presence of hydrogen peroxide initiator, the single-molecule fluorogenic assay. (B) Schematic representation of the total internal reflection fluorescence microscopy imaging-guided confocal fluorescence spectroscopy (TIRFM-CFS). M1: reflection mirror. DM1–DM2: dichroic mirror beam splitters. SPP: side port prism for left/vis obs. TL: tube lens. EF1–EF2: emission filters. L1–L2: lens. PBS: Polarization beam splitter. SPAD1–SPAD2: single photon avalanche photodiode. (C), (D) The typical raw data of single-molecule photon time-stamping spectroscopy of each detector channel with the perpendicular and parallel polarization components, respectively. Each data point represents a detected photon plotted by its arrival time (t) and delay time (Δt).

fluorophores have been used in single-molecule studies of enzymatic reaction systems, which include intrinsic fluorescence from tryptophan or tyrosine residues, site-specific fluorescent labeling, and fluorescent products.¹³ Comparing with others, fluorescent product molecules are generated constantly during the course of an enzymatic reaction, which eliminates the photobleaching problem and provides a chance to record and analyze a long time trajectory of the enzymatic reaction from a single enzyme. In a fluorogenic enzymatic assay measurement, the recorded trajectories are sufficiently long to represent a statistically relevant sampling of all possible conformational states.

Horseradish peroxidase (HRP), a monomeric enzyme 44K Dalton, containing a heme prosthetic group, catalyzes the oxidation of a broad range of substrates such as aromatic amines, indoles, phenols, and sulfonates in the presence of hydrogen peroxide as an oxidizing agent (H_2O_2).²³ The crystal structure and the catalytic mechanism of HRP were described previously.^{24–26} The HRP catalytic dynamics of turning over the fluorogenic substrate has been investigated by single-molecule fluorescence spectroscopy techniques.^{9,27–30} How-

ever, previously reported results are primarily limited to the reaction rate and the correlation analysis of the dynamic disorder. Specifically, there is still a lack of experimental understanding of the molecular mechanism and dynamics of the interactions between the enzymatic active site and the nascent product and the conformational dynamics associated with the releasing of the product from the enzyme. Nevertheless, various fluorescence characteristics, such as fluorescence intensity, spectrum, lifetime, and anisotropy, deserve to be exploited as they are potentially capable of probing enzyme–substrate and enzyme–product molecular interactions under complex local environments.^{31–35} Our unique technical approach of multiparameter single-molecule spectroscopic investigation of enzymatic reactions provides information beyond the reaction rates and correlation analysis of dynamic disorder.^{36,37}

In this work, we use HRP-catalyzed oxidation of non-fluorescent substrate Amplex Red (10-acetyl-3,7-dihydroxyphenoxazine) to fluorescent resorufin as a model system to study the conformational dynamics of the enzyme during the fluorescent products releasing from the enzymatic active site

into the solution, i.e., the last step of the enzymatic reaction turnover cycle. We have used a home-built total internal imaging guided confocal single molecular spectroscopy technique to probe the consecutive fluorescent product releasing from the enzymatic active site. Conformational dynamics of the single-molecule HRP is characterized by measuring real-time photophysical properties of the nascent fluorescent enzymatic products. We have identified a wide distribution of the multiple conformational states involved in active-site interacting with the nascent product molecule during the product releasing, and there is a significant conformational fluctuation between tight states and loose states of the active site of the enzyme involved in the product releasing process. Furthermore, we have experimentally revealed a new pathway in which the product molecules are spilled out from the enzymatic active site, possibly from the squeezing effect, besides the typical pathway of product releasing from an open active-site conformational state through diffusion.

■ EXPERIMENTAL METHODS

1. Chemicals and Sample Preparation. Horseradish peroxidase immobilized on the cover glass was used in our experiments. The cover glass (Gold seal, 3419) was first washed in fresh prepared sulfuric acid dichromate cleaning solution for 1 h to eliminate grease and possible fluorescent spots. After washing with water and drying with nitrogen gas, the cover glass was treated overnight with a mixture solution of 3-mercaptopropyl-trimethoxysilane (Fluka, 09324), isobutyltrimethoxysilane (Sigma, 444065), and dimethyl sulfoxide (Sigma, D4540) with a volume ratio of 1:300:6000. After baking at 110 °C for 10 min, the silanated cover glass was washed with methanol and water. Then the coverslips were incubated in 50 mM PBS buffer (pH 8.0) with about 1 nM maleimide-activated HRP (Thermo scientific, 31485) for 2 h and followed by rinsing with water and PBS buffer. Phosphate buffer (PBS) was prepared with potassium phosphate monobasic solution (Sigma-Aldrich, P8709) and potassium phosphate dibasic solution (Sigma-Aldrich, P8584). Maleimide-activated HRP was linked to the sulfhydryl (–SH) group of 3-mercaptopropyl-trimethoxysilane on the cover glass as shown in Figure 1A. Substrate Amplex Red (Invitrogen, A12222) was dissolved in dimethyl sulfoxide (DMSO) at 5 mg/mL and stored at –20 °C in the dark before use. The reaction solution was prepared just prior to experimentation, with 200 nM Amplex Red and 2 mM H₂O₂ in PBS buffer (pH 7.4). All chemicals were used without further purification. In our experiment, about 0.5 mL of reaction solution was filled in a home-built magnetic chamber that is composed with the coverglass tethered with HRP at the bottom and a lid on the top of the chamber to eliminate the evaporation.

2. Experimental Setup. The single-molecule enzymatic reaction assay was performed using a total internal reflection fluorescence microscopy imaging-guided confocal fluorescence spectroscopy (TIRFM-CFS) system built in our lab.³⁸ The system is based on an inverted microscope (Axiovert 200M, Carl Zeiss) in an epi-illumination configuration combining the TIRFM mode and the confocal mode, facilitated by a home-developed software for recording images from TIRFM mode and moving the pinpoints of interest from TIRFM imaging to confocal single-molecule spectroscopy measurement.³⁸ The TIRFM mode is used to locate spatially random distributed enzymes from the stochastic on–off bursts of fluorescent signals, and the confocal mode is used for the time-resolved

dynamic measurements of the targeted single enzyme.³⁸ Figure 1B shows the experimental setup schematically. Briefly, in the TIRFM mode, a cw laser (GCL-050-L, CrystaLaser) is aligned to the side port by a dichroic mirror beam splitter (DM2) then reflected by the side port prism (SPP) and focused by the tube lens (TL) on the back focal plane of the objective (Plan-Fluar, 1.45 NA, 100×, Carl Zeiss). By slightly deviating DM2 from 45° relative to the incident direction of the beam, the incident angle at the cover glass/solution interface can be adjusted to exceed the critical angle of total internal reflection. The fluorescence emission is collected by the same microscope objective, passing SSP, reflected by M1 to the EMCCD (EMCCD; Photomax 512B, Princeton Instruments), and further the emission signal is purified by an emission filter (EF2). In the confocal mode, a femtosecond pulse laser (Ti:sapphire Mira 900F/P, Coherent Inc.) is used. After the optical parametric oscillator (OPO BASIC, Coherent Inc.) and frequency doubling by a BBO crystal, the linear-polarized pulse laser is aligned to the back port of the microscope stand, reflected by the dichroic mirror beam splitter (DM1) in the filter cube, and focused on the cover glass–solution interface by the objective. The magnetic chamber filled with reaction solution is fixed on the piezoelectric scanning stage (Nano-H100, MCL). Fluorescence emission from the excitation focal volume is collected by the same objective. After transmitting DM1, the signal is reflected by the side port prism (SPP) and transmitting DM2 and refocused by the remagnification lens (L1) to the SPAD1 and SPAD2 (PDM50ct, MPPD) entrances. The polarization beam splitter (PBS) is used to separate the signal into parallel and perpendicular polarization components. Here the sensor target diameters of SPADs are 50 μm, which also serve as pinholes to reject stray and ambient light noise. The signals from SPADs are routed by the HRT-82 module and then connected with the single-photon counting module (SPC-830, Becker & Hickl GmbH) to record both photon arrival time (*t*) and delay time (Δt) between the excitation pulse and excited state emission with high time resolution for each single-photon event. The typical raw data of single-molecule photon time-stamping spectroscopy of each detector channel with the perpendicular and parallel components have been shown as Figure 1 C and D. The arrival times of the photons provide the information about the photon flux, and the histograms of arrival times yield fluorescence intensity trajectories with a given time-bin resolution. Also, the histograms of the delay times give the fluorescence lifetime.^{39,40} The control experiment has been carried out with the same conditions except no enzyme in the focus volume of laser illumination.

The efficiencies of two detection channels are typically different, primarily because of the difference of the quantum efficiencies of the two SPAD detectors and the bias of the dichroic beam splitter, mirrors, and polarizing beam splitter. To correct the efficiency imbalance of the two detection channels, the overall weighting factor, so-called *G* factor, is determined by fluorescent intensity response of the two channels with different excitation polarization. In our experiments, the measured weighting factor *G* is 1.48.

3. Fluorescence Intensity and Anisotropy. The single-molecule fluorescence intensity decays of the perpendicular and parallel polarized components are typically distinct: The fluorophores that have their transition moment aligned parallel to the electric vector of the excitation laser field preferentially absorb and emit photons. Using the polarization beam splitter,

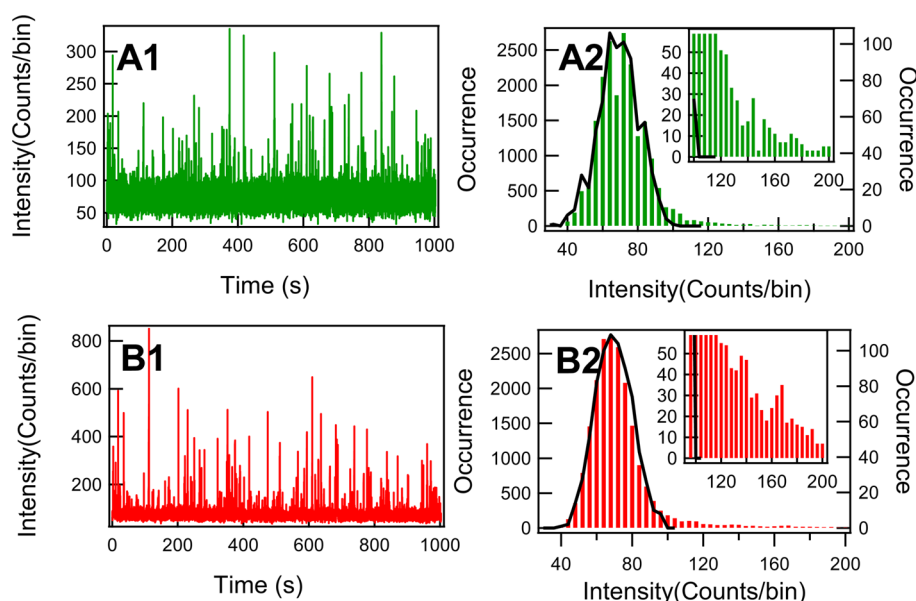


Figure 2. Single-molecule fluorescence intensity trajectories and the intensity distributions of HRP-catalyzed oxidation of Amplex Red, binning with 50 ms. (A1) Intensity trajectory of the perpendicular polarization component relative to the excitation polarization after the compensation by the G factor of photon detection. (B1) Intensity trajectory of the parallel polarization component relative to the excitation polarization. (A2) and (B2) The distributions of fluorescence intensity of the trajectories in A1 and B1, respectively. The black curve denotes the background intensity distribution from control experiment, and the inset is the zoom-in near the threshold of the background.

the emission intensity signal is divided into perpendicular and parallel components relative to the polarization of the excitation light. After the signal compensation by the G factor, these intensity trajectories of perpendicular ($I_{\perp}(t)$) and parallel ($I_{\parallel}(t)$) components are used to calculate total fluorescence intensity $I_T(t)$ and anisotropy $r(t)$ from the nascent single-molecule product at the enzymatic reaction active site, following eq 1 and eq 2.

$$I_T(t) = I_{\parallel}(t) + 2GI_{\perp}(t) \quad (1)$$

$$r(t) = \frac{I_{\parallel}(t) - GI_{\perp}(t)}{I_{\parallel}(t) + 2GI_{\perp}(t)} \quad (2)$$

4. Fluorescence Lifetime. Taking convolution with the instrument response function into consideration, the time-resolved polarization intensities $I_{\perp}(t)$ and $I_{\parallel}(t)$ are determined using eqs 3 and 4, respectively:⁴¹

$$I_{\parallel}(t) = \text{IRF} \otimes \left[\sum e^{-t/\tau_i} (1 + 2r_0 e^{-t/\tau_r}) \right] \quad (3)$$

$$I_{\perp}(t) = \text{IRF} \otimes \left[\sum e^{-t/\tau_i} (1 - r_0 e^{-t/\tau_r}) \right] \quad (4)$$

Here, IRF is the instrument response function; τ_r is the nascent single-molecule product rotational correlation time; τ_i is fluorescence lifetime; and \sum sums over all possible terms with different lifetime or rotational correlation time. To simplify the data analysis, the depolarization due to the curvature of the spherical wavefront of the focused field is ignored, due to the low influence even at high apertures.⁴² We assumed the excited-state population decay as single-exponential functions with lifetime.³⁹ So the total fluorescence intensity could be described by the following equations

$$I_T(t) = \text{IRF} \otimes Ae^{-t/\tau_i} \quad (5)$$

Although different methods have been described and applied to extract the lifetime from low count rates,^{43–47} we use the least-

squares (LS) method to get the monoexponential lifetime of the nascent single-molecule enzymatic product since this method is widely applied and converges to maximum likelihood estimation (MLE) at high count rates. The IRF and the time offset are determined from a bulk fluorescence measurement of the resorufin with the same time binning. Therefore, we have three free parameters to be extracted from the fluorescence data analysis: the fluorescence lifetime, fluorescence amplitude, and constant noise term.

5. Rotational Correlation Time. The rotational correlation time of a fluorescent molecule characterizes the molecule rotation diffusion, being dependent on the size, shape, and hydrodynamics of the molecule, as well as the bulk physical characteristics of the solvent. Therefore, for a fluorescent molecule buried in an enzyme active site, the variation of the rotation correlation time is directly related to the molecular confinement status and its local environment in the enzymatic active site. The rotation correlation time of the rotation fluctuation can be derived through the Perrin equation

$$\frac{1}{r} = \frac{1}{r_0} + \frac{\tau_f}{r_0\tau_r} \quad (6)$$

Here, r is steady-state anisotropy; r_0 is the fundamental anisotropy in the absence of molecular rotation; τ_f is the fluorescence lifetime; and τ_r is rotational correlation time. In the current experiment, the fundamental anisotropy of the resorufin is 0.318, which was measured in 80% glycerol solution.^{48,49}

RESULTS AND DISCUSSION

Fluorescence intensity is the most straightforward parameter to characterize fluorescence signals of fluorogenic enzymatic reactions. Fluorescence from a single-molecule product can only be detected and identified while it is still confined in the enzyme. After being released from the enzymatic active site, the fluorescent product molecule diffuses out of the laser excitation

confocal volume within submilliseconds.⁵⁰ By the single-molecule photon time stamping spectroscopy technique, we record the intensity trajectories of different polarization channels. Figures 2 A1 and 2 B1 show perpendicular channel and parallel channel fluorescence intensity trajectories, respectively, with a binning time of 50 ms, of the fluorescent products from a single HRP enzyme tethered to the coverglass in PBS buffer solution at pH 7.4. The fluctuations of fluorescence bursts recorded from both parallel and perpendicular polarization channels show similar behaviors and photon count distributions. On average, the intensity of the parallel component is higher than that of the perpendicular component even after the G-factor compensation. The corresponding intensity distributions of different polarizations are shown in Figures 2 A2 and 2 B2, respectively. The black curve is from the intensity distribution of the background, which was taken at the same conditions but without enzyme in the focus volume. This curve clearly shows the profile of the photon time stamping measurement without fluorescence turnovers. The insets show the intensity distributions above the edge of the background. The intensity distribution of the background shows a narrow band below 120 photons, whereas the distributions of intensity beyond the background are asymmetric and elongated toward the higher intensity side. For the distribution portion of intensity beyond the background, it is evident that the distribution likely contains more than one component since the distribution is significantly wider than the highest intensity shot noise.

Generally, the fluorescence intensity of a single fluorophore is proportional to the photon absorption and quantum yield, which is the ratio of emitted photons to the number of absorbed photons. Besides the absorption coefficient, the angle between the excitation polarization and the electron transition dipole also determines the photon absorption probability, which is revealed by the anisotropy measurement. Quantum yields are generally more sensitive to the environment than photon absorption.⁵¹ Typical local environmental factors that influence the quantum yield include local refractive index, solvent polarity, proximity and concentrations of quenching species,¹⁴ ultimately related to the nonradiative relaxation of the dye molecules, which could be revealed by fluorescence lifetime measurement. Single-molecule fluorescence lifetime analysis is often the choice of probing and characterizing the local environment interactions and fluctuations.^{18,52,53} Therefore, to identify the relationship between the local environment of a single enzyme's active site and the fluorescence properties of the fluorescence single-molecule product, it is necessary to perform the anisotropy and lifetime analyses.

Anisotropy provides a critical analysis of the rotational dynamics of a fluorescent molecule; furthermore, specific physical parameters of the local environment influencing the probe molecule rotational motions can also be either qualitatively or semiquantitatively characterized, such as the free space, force field, electric field, hydrophobicity, hydrodynamics, solvation effect, and hydrodynamic volume changes. Figure 3A shows the anisotropy fluctuation trajectory and the corresponding polarization intensity trajectory of a portion of typical single-molecule photon stamping trajectories (Figure 2). Here the intensity is calculated from 300 photons divided by the time intervals for accumulating the consecutive 300 photons to control the shot noise at the same level. The anisotropy shows as constant around zero between the fluorescence photon bursting. During the bursting, the

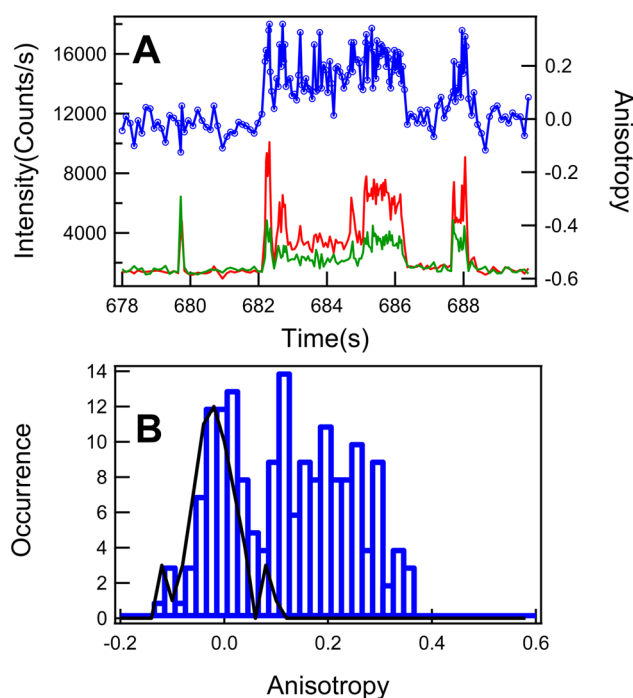


Figure 3. Single-molecule fluorescence anisotropy fluctuation and distribution of single HRP-catalyzed oxidation of Amplex Red fluorogenic assay. (A) Single-molecule fluorescence anisotropy fluctuation. Intensity trajectory from the perpendicular polarization component relative to the polarization of excitation (green) and the simultaneously recorded intensity trajectory from the parallel polarization component relative to the polarization of excitation (red); the calculated anisotropy trajectory (blue) using eq 2 from the pair of polarization components (red and green). (B) Anisotropy distribution from the trajectory (blue) in A. The anisotropy distribution from the background (black) with mean of -0.020 and standard deviation of 0.054 . It is evident that the signal anisotropy distribution (blue) is identifiable, beyond the background distribution.

anisotropy shows a dramatic fluctuation. Figure 3B shows the corresponding anisotropy distribution with two distinct peaks. The anisotropy distribution around the peak near zero has a narrow Gaussian-like shape and is dominated by the background, as compared to the control background with mean at -0.02 and with standard deviation as 0.054 , as shown in the black curve. The anisotropy distribution at higher anisotropy beyond the background is asymmetric and elongated toward higher values. This broad distribution of the high anisotropy spans from 0.1 and 0.36 , which is larger than the standard deviation of the background. To identify the specific interactions of the enzyme–substrate and enzyme–product at the enzymatic active site, it is reasonable to assume that the substrate amplex red and the nascent fluorescent product bind to the active site in a similar configuration as a typical aromatic substrate does, such as a reported binding structure of a HRP–acetate complex.^{23,25} Aromatic substrates form stable, reversible 1:1 complexes with HRP through both hydrogen-bonded and hydrophobic interactions at the distal side of the heme plane. The amino acid residues Arg38 and His42 play the roles in binding and stabilization of aromatic substrates (see Supporting Information). At the enzymatic active site of the HRP–resorufin complex, the rigid hydrogen bonding of resorufin to the HRP enzyme ensures that the HRP product active site rotates relatively confined, which in turn makes the measured fluorescence anisotropy trajectories likely to yield information

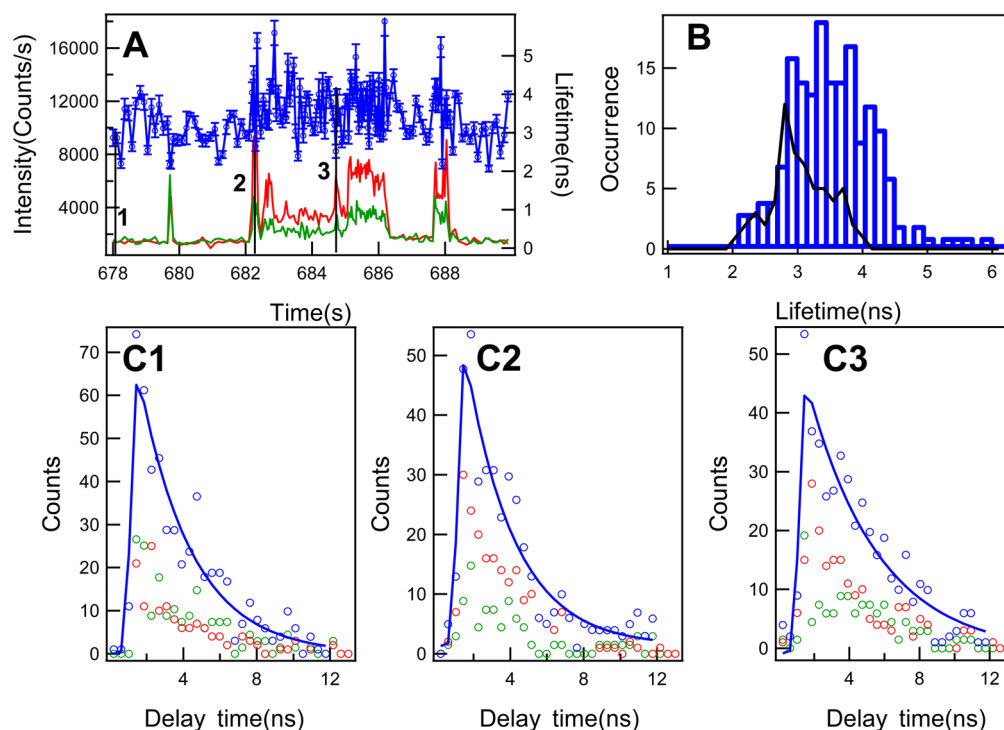


Figure 4. Single-molecule fluorescence lifetime fluctuation and distribution of single HRP-catalyzed amplex red fluorogenic assay. (A) Single-molecule fluorescence lifetime fluctuation trajectory. Intensity trajectory from the parallel polarization component relative to the polarization of excitation (red) and the simultaneously recorded intensity trajectory from the perpendicular polarization component relative to the polarization of excitation (green). Lifetime fluctuation trajectory (blue) calculated using eq 5 from the pair of polarization component trajectories (red and green). (B) Single-molecule enzymatic reaction product fluorescence lifetime distribution and the lifetime background distribution (black) from the trajectory (blue) in A. The lifetime background distribution is deduced from the lifetime trajectory from 678 to 681.5 s in (A), and the background distribution gives the mean of 3.01 ns and standard deviation of 0.60 ns. (C1), (C2), and (C3) Fluorescence decays at point 1 at 678.01 s, point 2 at 682.29 s, and point 3 at 684.75 s, respectively. The parallel polarization component decays (red), the perpendicular polarization component decays (green), and the fluorescence decays with fit (blue). The lifetimes at point 1, point 2, and point 3 are 2.9 ± 0.2 ns, 2.7 ± 0.1 ns, and 4.1 ± 0.1 ns, respectively.

about the rotational and conformational dynamics of not only the fluorescent product but also the HRP protein, particularly of the solvent exposed heme prosthetic group at the active site, where the oxidations of aromatic substrates happen. The broad distribution of the anisotropy in Figure 3B also indicates that likely multiple conformational states are involved in active-site interaction with the product molecules during the product releasing, which is consistent with the binding site being a relatively flexible structural region.

Fluorescence lifetime can be determined independently from molecular rotation through the linear combination of eqs 3 and 4. Figure 4A shows the lifetime fluctuation trajectory and the corresponding polarization intensity trajectories from a portion of the photon stamping trajectories in Figure 2. Each data point in the lifetime trajectory is calculated from the consecutive 300 photons of the two time-resolved photon stamping polarization channels. Figure 4B shows the lifetime distribution of the fluctuation lifetime trajectory in Figure 4A. There are no evident multi-peaks in the lifetime distribution as that in the anisotropy distribution. The lifetime background distribution (Figure 4B) shows the mean at 3.0 ± 0.6 ns, which is consistent with the lifetime of free resorufin molecules in pH 7.4 buffer. Compared to the lifetime of the background, the average lifetime of the nascent enzymatic reaction product, resorufin, is much longer than that of free resorufin molecules (Figure 4B). Since the lifetime of the resorufin is sensitive to the pH of the solution,⁵⁴ the longer average lifetime of the confined resorufin

is likely influenced by the local basicity of the active site. This is consistent with the fact that the hydrogen bond from amide oxygen of amino acid residue Asn70 to imidazole NH of amino acid residue His42 contributes to the basicity of the local environment.²³ Since the fluorescence lifetime is sensitive to the fluctuation of the local environment and the movements of macromolecules,^{52,55–59} the lifetime fluctuations of the confined resorufin most likely reflect the fluctuations of the active-site conformation of the HRP enzyme as well as the enzyme–product molecular interactions. Figure 4 C1 shows the typical fluorescence decay behavior of the free resorufin molecules from point 1 at 678.01 s in Figure 4A. The fluorescence intensity of parallel polarization shows similar decay as that of the perpendicular polarization component, which is consistent with the rotation of the free resorufin molecules being faster than the fluorescence lifetime.^{60,61} Figures 4 C2 and 4 C3 have shown the fluorescence decay behaviors during the bursting from point 2 at 682.29 s and point 3 at 684.75 s, respectively, in the trajectory 4A. At point 2, the lifetime is shorter than that of the free resorufin molecules in buffer solution, and the shorter lifetime is likely due to the quenching effect from the heme group at the enzymatic active site, since there exists the energy transfer between the enzymatic product, resorufin, and the heme group.^{56,57,62,63} At point 3, the fluorescence decay shows a longer lifetime than that in point 2. The significant increase of the lifetime suggests a significant change of the enzyme–product molecular

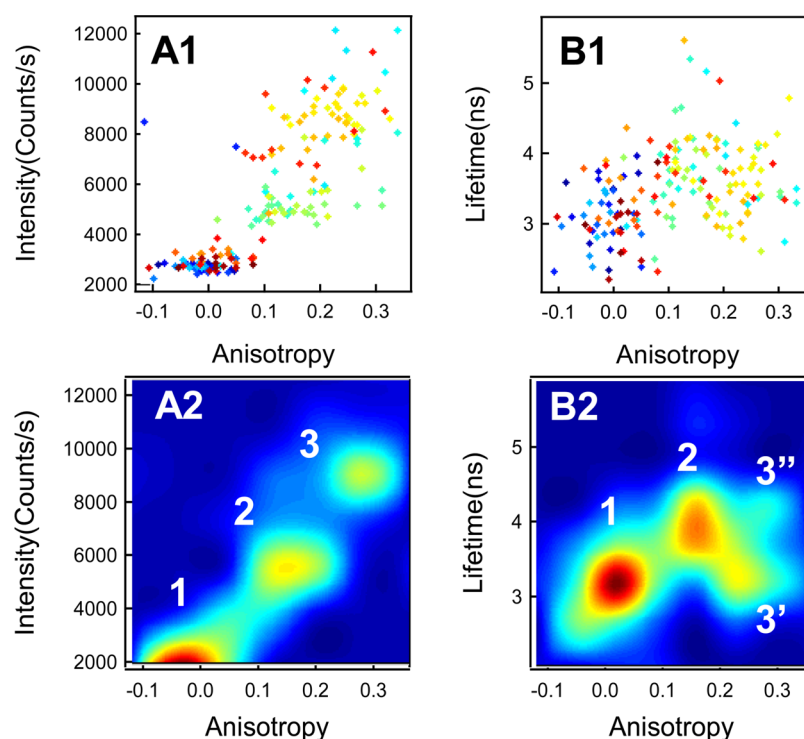


Figure 5. Correlation plots of the lifetime, anisotropy, and intensity on the single HRP-catalyzed amplex red fluorogenic assay. (A1) Correlation of fluorescence intensity and anisotropy and (B1) correlation of lifetime and anisotropy. For A1 and B1, the color (from cold color to warm color, i.e., from blue to red) of the data points represents the time sequence from the start to the end of the fluorogenic enzymatic turnover event. (A2) The correlation plots of fluorescence intensity and anisotropy with identified rotational correlation times. (B2) Correlation of lifetime and anisotropy with identified rotational correlation times.

interaction at the active site change, presumably making the nascent product molecule more efficiently prevented by the Phe41 residue to access the heme iron,²⁵ which decreases the energy transfer between the product and the heme group. At the enzymatic active site and at the particular time of point 3 the product most likely exists as the resorufin anion (R^-) state since it is dramatically different in lifetime compared to that of the protonated form of resorufin (RH).^{54,64}

Performing correlation analysis of the lifetime, anisotropy, and intensity on the single-molecule photon-by-photon trajectories allows us to characterize the complexity and heterogeneity of enzymatic conformational dynamics and mechanism.^{37,65,66} Figures 5 A1 and 5 B1 show the correlated distributions of intensity vs anisotropy and fluorescence lifetime vs anisotropy deduced from the trajectories shown in Figure 3 and Figure 4, respectively. Figures 5 A2 and 5 B2 show the correlated counter plots of Figures 5 A1 and 5 B1. There are distinctly three domains in the intensity–anisotropy 2D plot (Figure 5A2), while in the lifetime–anisotropy 2D plot (Figure 5 B2), there are two domains 3' and 3'' separated at the high anisotropy value, except the same domains 1 and 2 as in Figure 5 A2. These domains reflect the different local environments of the nascent fluorescent enzymatic reaction product, resorufin, at the enzymatic active site. To assign the origin of each domain in the correlation distribution 2D plots, the rotational correlation times (τ_r) are needed as shown in eq 6.

In Figure 5 B2, the regions 1, 2, 3', and 3'' are centered at $(r, \tau_r) = (0.01, 3.1 \text{ ns}), (0.15, 3.7 \text{ ns}), (0.24, 3.1 \text{ ns}),$ and $(0.26, 4.0 \text{ ns})$, respectively. On the basis of eq 6, the corresponding rotational correlation times, τ_r , of the regions 1, 2, 3', and 3'' are calculated to be 0.10, 3.30, 9.54, and 17.93 ns, respectively. The rotation correlation time of 0.10 ns from region 1 is consistent

with that of the freely rotating resorufin in aqueous solution.^{61,67} Region 3'' has the longest rotation correlation time, which is close to the rotation correlation time of HRP in solution, 18.3 ns, since the τ_r is proportional to the molecular weight and increases about 1 ns for each 2400 Da increase in molecular weight.⁶⁸ In region 3'', the resorufin is most likely bound with HRP tightly in the form of an enzyme–product complex, and there is essentially no measurable relative rotation between resorufin and HRP. In regions 2 and 3', the calculated rotational correlation times are shorter than the rotational correlation time of HRP in the solution. The shorter rotation correlation time most likely comes from the less tight spatial confinement of the unbounded or loosely bound nascent resorufin at the HRP enzymatic active site. Therefore, the apparently different τ_r can be associated with the enzyme conformational dynamics associated with the enzyme–product interactions. Here, the relatively shorter rotation correlation time in region 2 can be ascribed to a relatively loose state of HRP, which makes more free space for the rotation of nascent resorufin. The relatively longer rotation correlation time in region 3' could be ascribed to a relatively tight state of the HRP, which makes the rotation time of the resorufin molecule much closer to that of the enzyme HRP. In other words, when the confinement space of the active site becomes larger, the enzyme–product interaction is looser, and the product molecule has higher flexibility of movements, which constitutes a shorter τ_r , and vice versa. The assignment of loose state and tight state of HRP from the rotation correlation time of nascent resorufin molecules is also consistent with the quenching effect of the heme group to the fluorescence lifetime of resorufin.^{54,69,70} As in the loose states, the average distance from the nascent resorufin to the heme group is larger than that in tight

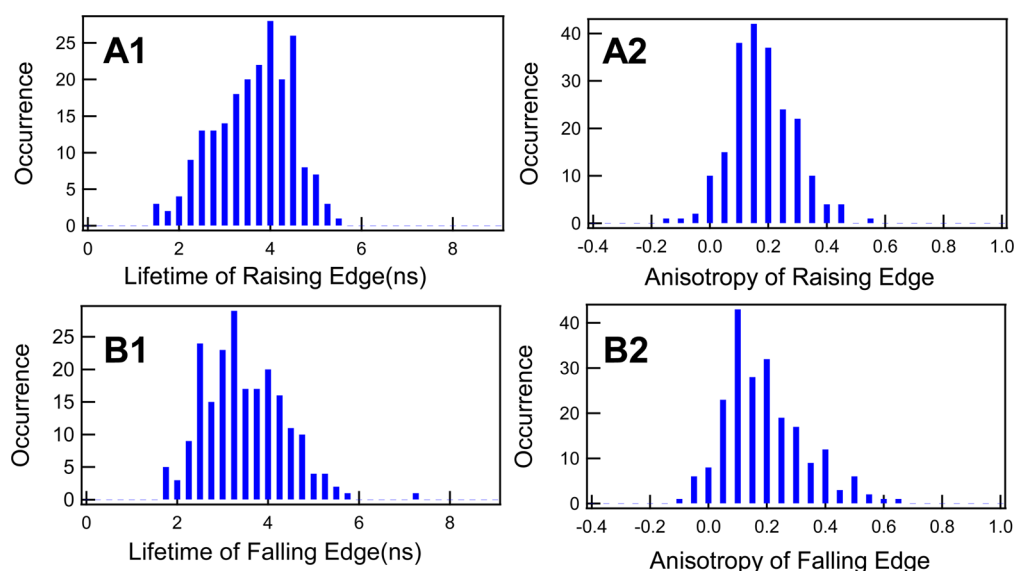


Figure 6. Single-molecule fluorescence lifetime and anisotropy distributions of the rising edges and falling edges of single HRP-catalyzed oxidation of amplex red reaction fluorescence turnover events. (A1) The lifetime distribution of rising edges with mean at 3.75 ns and standard deviation 1.26 ns. (A2) The anisotropy distribution of rising edges with mean at 0.17 and standard deviation 0.14 ns. (B1) The lifetime distribution of falling edges with mean at 3.37 ns and standard deviation 1.3. (B2) The anisotropy distribution of falling edges with mean at 0.15 and standard deviation 0.16.

states, which results in the decrease of the quenching effect from the heme and thus makes the lifetime longer than that in tight states. Furthermore, the local dielectric constant variation also influences the fluorescence lifetime of the enzymatic product that is embedded in the active site of the HRP enzyme. Because the average refractive index of proteins and amino acids is larger than that of water,^{71,72} the tight state of the conformation gives a larger local dielectric constant. The increase in local dielectric constant at the enzymatic active site induces a shorter lifetime since the spontaneous emission rate scales with the square of the refractive index based on a simple density-of-states argument.^{73–75} Therefore, we note that HRP has breathing-type conformation motions between the tight states and loose states of the enzyme active site under enzyme–product interactions before the nascent product molecule is released from the enzyme into the solution.

To further identify the conformational dynamics of the product releasing from the enzyme active site, we have analyzed the distributions of the fluorescence lifetime and anisotropy at the rising edges and falling edges of about 211 fluorescence bursting spikes from the trajectories shown in Figure 2. Here, the signal rising edge corresponds to the fluorescent product generated at a single-molecule reactant-to-product turnover, and the signal falling edge corresponds to the product releasing from the active site into the solution and diffusing away from the confocal volume. Figure 6 shows the distributions of the lifetimes and anisotropies of the products from the rising edges and the falling edges of each fluorescence spike. Comparing the lifetime distributions from the rising and falling edges of the fluorescence spikes (Figure 6 A1 and Figure 6 B1), we notice that the mean lifetime of the falling part is about 0.38 ns shorter than that of the rising part, while the anisotropy distributions are almost the same. On the basis of the influence of the quenching effect and local dielectric constant on the fluorescence lifetime of the nascent product as discussed above, the most possible explanation for the shorter lifetime of the single resorufin molecules at the falling edges is that the

product molecules are buried in tight states compared to the rising edge.

Rotation correlation time of the enzymatic reaction product, resorufin, at the falling edges of the turnover fluorogenic signals is a sensitive parameter to reveal the enzyme–product interactions right at the product releasing events. Figure 7

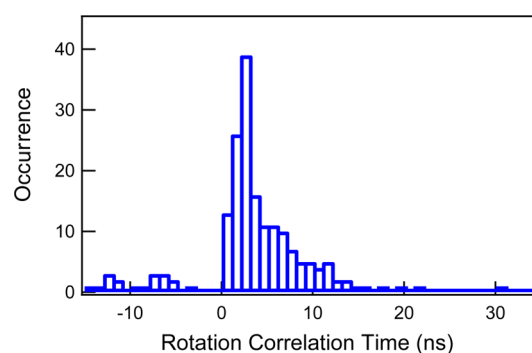
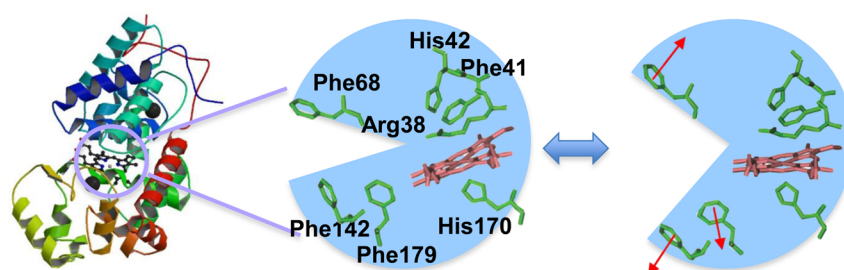


Figure 7. Distribution of rotation correlation time of the product at the signal falling edges that correspond to the moments of releasing the product molecules from the enzyme into the solution.

shows the distribution of the rotation correlation time of the resorufin at the falling edges and the distribution peaks at 2 ns that are close to the rotation correlation time of the loose enzyme–product states in region 2 of Figure SB2, and these product events are most likely related to the enzyme active site opening up to release the fluorescent product into the solution. Nevertheless, about 20% of the product releasing events are associated with the rotation correlation time larger than 9 ns, which suggests that the product releasing events occurred at a strong enzyme–product interaction so that the fluorescent product rotational motions are at the protein motion time scale. Furthermore, the fluorescence lifetime of the product releasing events is also associated with shorter fluorescence lifetime, as we have discussed above, which also suggests a tight enzyme–substrate interaction at the events of product releasing. The

A. Breathing motion of active site



B. Schematic of Product releasing dynamics

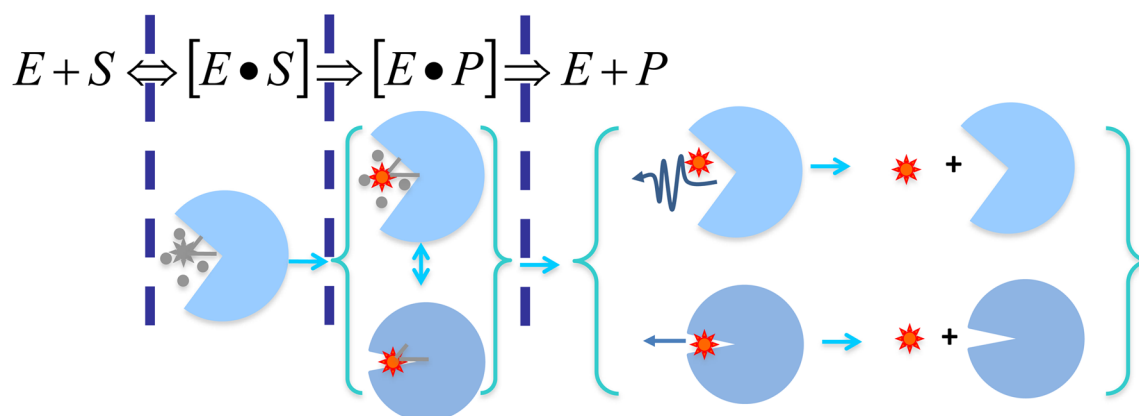


Figure 8. Proposed dynamics of single HRP-catalyzed oxidation of amplex red reaction. (A) The breathing motions of the enzymatic reaction active site. (B) The schematic of the two pathways of product release. Our experimental observation suggests that about 20% product releasing is involved in the spilling pathways.

correlated shorter lifetime and longer rotation correlation time of the products at the falling edges support our attribution that there is a significant pathway in which the product molecules are spilled out from the enzymatic active site, driven by a squeezing effect by the breath-type motions between loose and tight active-site conformation states; i.e., there is a significant portion of product molecules that are released from a tight active-site conformational state rather than from an open active-site conformational state. This most remarkable phenomenon provides a new insight into the enzymatic reaction dynamics and mechanism—specifically on the product releasing mechanism and dynamics.

It is well-known that enzymatic reactions involve multiple kinetic steps, and each step involves complex molecular interactions, inhomogeneous conformational changes, and fluctuations of confined local environment.^{6–9} Both the overall dynamics and mechanism of the enzymatic reaction have been well investigated in ensemble-averaged experiments about the HRP enzyme, although there are different fluorogenic enzymatic reaction mechanisms proposed in the recent literature (see Supporting Information).^{23–26,30,85–89} Here, the catalytic networks of the HRP-catalyzed amplex red have been further characterized by our single-molecule fluorogenic assay. Rotational correlation times of the product during the releasing process have shown the enzyme undergoing breathing-type conformational motions fluctuating between loose states and the tight states. On the basis of the crystal structure of HRP, these breathing-type configurational motions may come from the active-site segment motion as shown in Figure 8A. As the enzyme changes to the loose states, the water-

soluble substrate is facilitated to transport into the active site. Conformation fluctuations between the loose states and tight states occur during the products releasing. Nevertheless, we propose a new mechanism of the HRP enzymatic reaction (Figure 8B): There are multiple conformations involved in the enzymatic reaction enzyme–substrate and enzyme–product interactions as well as the product releasing from the active site, and the conformational fluctuations occur in consecutive and parallel pathways in the mechanism with both the active-site open releasing product pathway and the active-site close spilling product pathway.⁷ Overall, the active-site close spilling product pathway reported in this work is the most significant new knowledge for the HRP enzymatic reaction dynamics and mechanism. We anticipated that the “spilling” product releasing pathway widely existed in other enzymatic reaction product releases, especially for the enzyme active-site associated with large conformational motions in enzymatic reaction turnovers.

The enzymatic reaction rate could be influenced by different factors, such as temperature, pH, enzyme concentration, substrate concentration, local environment, and the presence of any inhibitors or activators.^{1,3} Beyond the ensemble-averaged reaction kinetic scope, the enzymatic reaction turnover rate fluctuation for a single enzyme is often observed, which is characterized as the dynamic disorder.^{2,6,34,76–81} In our experiment, only the slow product releasing events, giving more than 600 detected photons, were used for analyzing the lifetimes, anisotropy, and rotational correlation times of the rising edges and the falling edges. The detected turnover rate in our single-molecule fluorogenic enzymatic assay is typically about 0.2 events/s. Compared to the turnover rate of the

conventional enzymatic reaction assay at mM or μ M substrate concentrations,^{9,30,82} our measured turnover rates are slow, mostly due to the low concentration of 200 nM substrate used in our experiments and also due to that only the high fluorescence photon counts events were detected and analyzed. We used the low concentration for two reasons: (1) This low concentration of substrate effectively controls the background increasing from laser-induced and autocatalytically enhanced photo-oxidation of amplex red to resorufin, and (2) in this work, we focus on studying the enzymatic reaction product releasing event analysis but not the enzymatic activity or enzymatic reaction rates; therefore, isolated single events are ideal for our detailed analysis of lifetime, anisotropy, and rotational correlation time for both the rising edge and falling edge of the fluorogenic turnover event. Nevertheless, the substrate concentration is the dominate factor in the diffusion-controlled reaction processes, and we further note that we cannot completely rule out the effect of the local steric hindrance around the tethered enzyme,⁸³ which may play a role in the influence of the turnover rate. The local steric hindrance can affect the motions of the enzymes, both the substrate accessing to the active site and product releasing from the active site may be slowed, although the statistics of analyzing hundreds of single fluorogenic turnover events should essentially remove the significance of the steric hindrance effect for our examination on the fluorescence lifetime, anisotropy, and rotational correlation time distributions.

One of the highest aims of studying enzymology is to obtain a fundamental understanding of why the enzymes are capable of catalyzing a biochemical reaction by millions or even billions of times.^{84,85} Probing and characterizing the enzymatic reaction active site has been one of the central efforts in modern enzymology. In this work, we focus on resolving the product releasing pathway analysis as it is one of the critical processes in an enzymatic reaction, as described by the Michaelis–Menten mechanism; however, the significance of being able to probe the nascent fluorescent product formation at an enzymatic reaction site is much more profound and beyond just illuminating the product releasing pathways because the nascent fluorescent product molecule at the active site serve as an ideal and sensitive probe at the fluctuations and distributions of the chemical bonding configuration, the local static electric field, local force field, solvation local environment, and the fluctuation dynamics of the enzymatic reaction transition state.^{8,10,12,86–99} Clearly, there are significant challenges for single-molecule spectroscopy to probe the details of these important physical properties at the active site and at the exact time of enzymatic reaction of forming the product, and the recent developments of high time-resolved and spatial resolved single-molecule imaging and spectroscopy with the multiple physical parameter selectivity and sensitivity are definitely promising for carrying out these high efforts.^{34,36,37,100–120}

CONCLUSIONS

We have used an enzymatic reaction, horseradish peroxidase-catalyzed oxidation of amplex red, as a model system to carry out single-molecule enzymatic conformational dynamics studies. Using our combined and correlated single-molecule fluorescence intensity, anisotropy, and lifetime measurements, we are able to identify the specific pathways with distinctive specifications on the conformational dynamics. Broad distributions of fluorescence intensity, anisotropy, and lifetime beyond

their standard deviation values identify the presence of intermediate conformational states associated with the product releasing step. Rotation correlation time and correlation between the fluorescence lifetime and anisotropy confirm the existence of a tightly confined product–enzyme complex prior to the releasing step. The resorufin anion state is responsible for the increased lifetime value of resorufin confined in the enzyme's active site. The lifetime fluctuations in the active site result from conformational fluctuations of the resorufin–enzyme complex between loose and tight states. Although only the enzyme–product complexes are investigated in this experiment, the conclusion could also be used to understand the conformational dynamics of enzymes during catalysis. All the conformational fluctuations between the loose states and tight states happen before the product releasing. The lifetime and the rotation correlation times of the products at the falling edges have shown that there is a significant pathway during which the product molecules are spilled out from the enzymatic active site, driven by a squeezing effect; i.e., a significant portion of product molecules are released from a tight active-site conformational state beside an open active-site conformational state. This most interesting phenomenon provides a new insight into the enzymatic reaction dynamics and mechanism, and the information is uniquely obtainable from our combined time-resolved single-molecule spectroscopic measurements and analyses. This multiple conformations occur in consecutive and parallel in the mechanism, which would help to understand the rugged multidimensional standard free energy.

ASSOCIATED CONTENT

Supporting Information

Additional information about the catalytic mechanism of horseradish peroxidase with hydrogen peroxide and the crystal structure of the active site; movie about single-molecule fluorogenic turnovers from the tethered HRP enzyme; complementary fluorogenic assay results of tethered single-molecule HRP enzymes. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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Notes

The authors declare no competing financial interest.

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