Roles of Closed- and Open-Loop Conformations in Large-Scale Structural Transitions of L-Lactate Dehydrogenase

Kimichi Suzuki,*†‡ Satoshi Maeda,*†§∥ and Keiji Morokuma‡

†Department of Chemistry, Faculty of Science, Hokkaido University, Sapporo 060-0810, Japan
‡Fukui Institute for Fundamental Chemistry, Kyoto University, Kyoto 606-8103, Japan
§Research and Services Division of Materials Data and Integrated System (MaDIS), National Institute for Materials Science (NIMS), Tsukuba 305-0044, Japan
∥Institute for Chemical Reaction Design and Discovery (WPI-ICReDD), Hokkaido University, Hokkaido 001-0021, Japan

Supporting Information

ABSTRACT: The mechanism of l-lactate generation from pyruvate by l-lactate dehydrogenase (LDH) from the rabbit muscle was studied theoretically by the multistructural microiteration (MSM) method combined with the quantum mechanics/molecular mechanics (QM/MM)−ONIOM method, where the MSM method describes the MM environment as a weighted average of multiple different structures that are fully relaxed during geometry optimization or a reaction path calculation for the QM part. The results showed that the substrate binding and product states were stabilized only in the open-loop conformation of LDH and the reaction occurred in the closed-loop conformation. In other words, before and after the chemical reaction, a large-scale structural transition from the open-loop conformation to the closed-loop conformation and vice versa occurred. The closed-loop conformation stabilized the transition state of the reaction. In contrast, the open-loop conformation stabilized the substrate binding and final states. In other words, the closed- to open-loop transition at the substrate binding state urges capture of the substrate molecule, the subsequent open- to closed-loop transition promotes the product generation, and the final closed- to open-loop transition at the final state prevents the reverse reaction going back to the substrate binding state. It is thus suggested that the exchange of stability between the closed- and open-loop conformations at different states promotes the catalytic cycle.

1. INTRODUCTION

Enzyme catalysis generally proceeds under moderate conditions and is fast and highly selective.1,2 Enzyme catalysis has therefore been applied to syntheses of various organic compounds.3−4 In general, the entire process of an enzyme-catalyzed reaction consists of three steps; that is, substrate binding, catalytic reaction in the active site, and product release.5 It has been shown experimentally that some catalytic reactions proceed with a large-scale structural transition in the surrounding protein.6−9 In such cases, one needs to take into account the large structural change in the surrounding protein before and after the chemical reaction that occurs at a local active site. However, revealing the roles of such a large structural change either experimentally or theoretically is challenging.2,4

L-Lactate dehydrogenase (LDH) exists in various organisms, and its catalytic mechanism has been the topic of many experimental and computational studies.10−23 LDH catalyzes a reversible transformation of pyruvate to l-lactate in the presence of NADH/NAD⁺ as a cofactor. Holbrook and co-workers proposed that the mechanism of this transformation consists of the five steps shown in Figure 1.7 In this

Figure 1. Schematic illustration of the full LDH catalytic process.7

Received: October 15, 2018
Accepted: December 28, 2018
Published: January 14, 2019
mechanism, substrate binding occurs while LDH is in the open-loop conformation. Then, LDH changes its form to the closed-loop conformation. The transformation of pyruvate to L-lactate occurs while LDH is in the closed-loop conformation. The product is released with LDH in the open-loop conformation. This proposed mechanism involves large-scale structural transitions from the closed-loop conformation to the open-loop conformation and vice versa. The closure of the open loop over the active site after substrate binding is considered to be the rate-limiting step in the LDH catalytic process of wild-type Bacillus stearothermophilus.18

Figure 2 illustrates the transformation of pyruvate to L-lactate in the LDH active site. In early theoretical studies, quantum mechanics/molecular mechanics (QM/MM) molecular dynamics (MD) calculations were performed to examine whether these events occur in a concerted or stepwise manner. Recently, free energy surfaces were calculated for two collective variables through QM/MM MD simulations using the AM1 and RM1 methods for the QM part. After making a correction based on density functional theory calculations, these free energy surfaces showed that proton and hydride transfers were concerted in LDH from the rabbit muscle. This theoretical result is supported experimentally by a recent spectroscopic study.10

The roles of closed- and open-loop conformations of LDH are not yet well understood theoretically. In principle, their behavior can be studied if MD trajectories over a long timescale can be computed at a QM or QM/MM level. The QM/MM free energy perturbation method coupled with geometry optimization calculations is another approach to study this topic. However, these calculations could be practically difficult because of the high computational cost to achieve converged statistical sampling describing the slow structural transition between open- and closed-loop conformations. We therefore use an alternative approach in which both open- and closed-loop conformations are prepared beforehand and then the entire structure is expressed as a weighted sum of these conformations. The weight changes depending on the stage of the reaction cycle and the transition between the two conformations are described as changes in weight. Such a calculation can be performed easily in the framework of our recently proposed multistructural microiteration (MSM) method.

2. THEORETICAL BACKGROUND

In this study, the structures of two conformations of LDH are prepared by replica-exchange MD simulations. Then, twelve initial structures are prepared. Among the twelve structures, six have an open-loop conformation and the other six have a closed-loop conformation. To distinguish between the two conformations, we introduced four distance parameters which are Ca-Glu103 and Ca-Tyr238, O3-pyruvate and CZ-Arg105, O1-pyruvate and CZ-Arg105, and CZ-Arg105 and CZ-Arg168 (see Figure S1). The structures with distances between Ca-Glu103 and Ca-Tyr238, O3-pyruvate and CZ-Arg105, O1-pyruvate and CZ-Arg105, and CZ-Arg105 and CZ-Arg168 of longer than 13, 8, 10, and 13 Å are regarded as open-loop conformation. Experimental values for these distances for open-loop conformations are 17.9, 12.0, 15.2, and 18.6 Å, while those for closed-loop ones are 8.0, 3.9, 4.5, and 7.2 Å. Details of the structural preparation are described in the Computational Details section. In the conventional QM/MM, our own N-layer integrated molecular orbital molecular mechanics (ONIOM) method33,34 the total energy E is expressed as

$$E = E_{\text{model-QM}} - E_{\text{model-MM}} + E_{\text{real-MM}}$$

where $E_{\text{model-QM}}$, $E_{\text{model-MM}}$, and $E_{\text{real-MM}}$ correspond to the QM energy of the model system, MM energy of the model system, and MM energy of the real system, respectively. In contrast, in the QM/MM–ONIOM–MSM method, E is expressed as

$$E = E_{\text{model-QM}} - E_{\text{model-MM}} + \sum_{i}^{N} \alpha_i E_{\text{real-MM}(i)}$$

where N is the number of structures considered and $E_{\text{real-MM}(i)}$ corresponds to the energy of the ith structure. The weight $\alpha_i$ is given as

$$\alpha_i = \frac{\epsilon^{-\beta \Delta E_{\text{real-MM}(i)}}}{\sum_{j}^{N} \epsilon^{-\beta \Delta E_{\text{real-MM}(j)}}}$$

where $\beta = 1/k_B T_{\text{MSM}}$, in which $k_B$ is the Boltzmann constant and $T_{\text{MSM}}$ is a model temperature parameter (see Computational Details section). During geometry optimization, all the N structures in the real system were optimized one-by-one with the atoms fixed in the model system (reaction-center region) before changing the positions of atoms in the model system. In other words, the optimization of the model system was performed in the geometrical subspace in which the gradient of the region not involved in the reaction (part of the real system excluding the model system) was zero.

The minimum energy path (MEP) for the abovementioned reaction process was computed. The MEP was obtained by optimizing an initial path using the locally updated plane (LUP) method.35 Initial paths were obtained as described below. During the path optimization, energy peaks (maxima) were directly optimized to transition states (TSs) using the quasi-Newton method. Three different calculations were performed: (i) a mixed calculation in which all twelve structures were used, (ii) an open-only calculation in which only the six structures with the open-loop conformation were used, and (iii) a closed-only calculation in which only the six structures with the closed-loop conformation were used. In calculations (ii) and (iii), the system adopts either an open- or closed-loop conformation throughout the calculation. The open- to closed-loop conformational transition or vice versa could only take place in calculation (i). The results of these three calculations are compared below. In calculation (i), the initial path was generated by the artificial force-induced
reaction (AFIR) method. In contrast, in calculations (ii) and (iii), the MEP of the mixed calculation was used as the initial path, and then the six surrounding structures with the non-target conformation were deleted before performing the LUP path optimization.

3. RESULTS AND DISCUSSION

An energy profile and variations of $\omega_i$ along the MEP obtained by the mixed calculation are shown in Figure 3a,b, respectively.

Three energy peaks were observed in the MEP, where the highest peak corresponds to the TS of concerted proton and hydride transfers. In Figure 3b, $\omega_i$ changed around the first and third peaks. The first and third peaks therefore describe the open- to closed-loop and closed- to open-loop structural transitions, respectively. Among the three peaks, the highest peak corresponds to the TS. The other two peaks were arisen through the dramatical change in $\omega_i$; we therefore call these energy maxima as surrounding structural TSs (SSTSs). For coordinates of the reaction-center atoms (atoms in the QM model system plus those bonded to the QM model system), each SSTS is a local maximum along the MEP of the potential energy surface calculated by the MSM method. We note that each SSTS also corresponds to a local minimum in the geometrical subspace in which the two main contributing surrounding structures have the same energy. Therefore, assuming immediate thermal equilibration for the surrounding structure, each SSTS corresponds to the energetically most preferred structure at which the population moves from one surrounding structure to the other. Figure 3b shows that the system adopts the open-loop conformation before SSTS1, the closed-loop conformation between SSTS1 and SSTS3, and the open-loop conformation again after SSTS3. This behavior is consistent with the mechanism in Figure 1 proposed from experimental results.

Figure 4 compares energy profiles of the corresponding paths obtained from the open-only (denoted o) and closed-only (denoted c) calculations with the path of the mixed (denoted m) calculation. On the energy profile, surrounding structural features in open- or closed-loop conformations are clearly distinguishable (see Table S1). At the substrate binding equilibrium state EQ1, the open-loop conformation m(o)EQ1 is more stable than the closed-loop conformation cEQ1. Therefore, the substrate binding is likely to take place in the open-loop conformation. The mSSTS1, which describes the structural transition from the open-loop conformation to the closed-loop conformation, leads to intermediate state m(c)EQ2. Then, the concerted proton and hydride transfers occur from m(c)EQ2 through m(c)TS2. Meanwhile, oEQ1 is directly connected to oTS2. After passing oTS2, the system again directly reaches the most stable state m(o)EQ4. In contrast, the Eqs in the closed-loop conformation, m(c)EQ3 and cEQ4, are both less stable than m(o)EQ4. From m(c)EQ3, the system can also reach m(o)EQ4 accompanying the surrounding structural transition through mSSTS3. Figure 4 clearly shows that the path obtained by the mixed calculation is the most energetically favorable path, and the surrounding structural transitions through mSSTS1 and mSSTS3 are necessary for the system to follow this path. In addition, the energy landscape was not largely affected (only $\sim$2.0 kcal mol$^{-1}$) by the polarization effects taken by the electronic embedding scheme as shown in Figure 4. The calculated activation energy 22.3 kcal mol$^{-1}$ is larger than an experimental value 14.0 kcal mol$^{-1}$ obtained by taking a logarithm of observed $k_{cat}$ (= 245 s$^{-1}$) at 295 K. Taking account of the
free-energy correction at 295 K estimated based on the harmonic vibrational analysis, the activation energy for this step decreased from 22.3 to 19.4 kcal mol\(^{-1}\) showing a better agreement with the experimental value, although the value still is larger than the experimental one. One reason would be because initial surrounding structures were prepared at the reactant state. This treatment may significantly overestimate relative stability of the reactant state and consequently the barrier height. However, we believe that the relative stability between open- and closed-loop conformations at each state, which is the main focus of this study, can be discussed from the results in Figure 4. There are many other factors that may affect the computational barrier height such as QM computational level, size of the QM part, parameters of the MM force field, efficiency of quantum tunneling, and dynamical fluctuation in the surrounding part, although taking these into account is beyond the scope of this study.

Interestingly, the profile in Figure 4 reveals that the reaction does not only proceed in the closed-loop conformation. Without the surrounding structural transition to the open-loop conformation through \(m\)SSTS3, the reaction is reversible because the stability of \(c\)EQ4 is similar to that of \(c\)EQ1. The surrounding structural transition from the open-loop conformation to the closed-loop conformation through \(m\)SSTS1 shifts the reaction bottleneck from \(o\)TS2 to the lower energy \(m\)TS2 and also promotes the reaction. We therefore conclude that the exchanges between the two conformations play roles in facilitating the reaction in the present system.

Next, we discuss reaction-center structures along the MEPs. At all the EQs and TSs in Figure 4, the volume which is estimated with assuming that each atom is a sphere with 1.5 times of the corresponding van der Waals radius is smaller in the closed-loop conformation than in the open-loop conformation throughout the process. This indicates that the open-loop conformation provides a larger reaction space than the closed-loop conformation. At TS2, the substrate must approach both His192 and NADH to transfer the proton and hydride. It is known in hydrogen-bonded cluster systems that a tight reaction space accelerates proton transfers.\(^{39,40}\) Consequently, the volume is the smallest at TS2 in the entire energy profile. Figure 5a–c compares structures of \(o\)TS2 and \(c\)TS2. The distances between the proton/hydride donor and acceptor are shorter in \(c\)TS2 than in \(o\)TS2 because of the tighter reaction space in the closed-loop conformation than in the open-loop conformation. In contrast, the greater space is available in the substrate binding state EQ1 and the final product state EQ4 than in TS2. In EQ4, the open-loop conformation allows the product to rotate in the reaction space and adopt a more stable configuration, as shown in Figure 5d–f. Because the substrate can move easily in the open-loop conformation, the system can reach the more stable configurations EQ1 and EQ4 directly without being captured at the tighter configurations EQ2 and EQ3. In short, the volume is small in the closed-loop conformation but is large in the open-loop conformation, while the volume is small at TS2 but is large at EQ1 and EQ4. In other words, the closed-loop conformation fits well to TS2, while the open-loop conformation fits well to EQ1 and EQ4. We thus propose that the volume matching is one factor controlling the variation of relative stability between the two conformations along the reaction path. The volume matching would strengthen some attractive interactions such as electrostatic interactions between the QM and MM parts and enhance stability of one of conformations at each state.

Figure 6 shows structural differences between EQ1 and EQ2 and between EQ3 and EQ4. The difference between \(c\)EQ1 and

![Figure 5. Reaction-center structures: (a) superposition of \(o\)TS2 (red) and \(c\)TS2 (blue), (b) \(o\)TS2, (c) \(c\)TS2, (d) superposition of \(o\)EQ4 (red) and \(c\)EQ4 (blue), (e) \(o\)EQ4, and (f) \(c\)EQ4. Distances (in Å) between the proton/hydride donor and acceptor at TS2 are shown in parentheses.](image)

![Figure 6. Reaction-center structures: (a) superposition of \(m\)EQ1 (red), \(c\)EQ1 (blue), and \(c\)EQ2 (yellow), and (b) superposition of \(m\)EQ3 (red), \(c\)TS3 (blue), and \(c\)EQ4 (yellow).](image)
m(\(\sigma\))EQ1 is in the interaction form between pyruvate and His192. The \(\epsilon\)EQ1 becomes m(\(\sigma\))EQ2 via \(\epsilon\)TS1 through the reorientation of His192 as highlighted in Figure 6a by an arrow. Although structures of m(\(\sigma\))EQ3 and \(\epsilon\)EQ4 resemble each other as shown in Figure 6b, changes occur in the surrounding structure through the movement of acetamide group in NAD via \(\epsilon\)TS3 as highlighted in Figure 6b by an arrow. The energy difference between m(\(\sigma\))EQ3 and \(\epsilon\)EQ4 is 13.4 kcal mol\(^{-1}\). The energy lowering of \(\epsilon\)EQ4 is due to the surrounding structural difference between m(\(\sigma\))EQ3 and \(\epsilon\)EQ4, where QM and MM components of the energy difference in m(\(\sigma\))EQ3 and \(\epsilon\)EQ4 (\(\eta\)-lactate) were 3.7 and \(-17.1\) kcal mol\(^{-1}\), respectively.

Finally, we discuss about four issues in the model adopted in this study. First, a previous study showed that formation of the tetramer structure of LDH affects the active site structure and the binding energy of the substrate.\(^{32}\) Because interactions between different chains were not considered in our model, it is expected that the energy profile in Figure 4 changes when the tetramer structure is used. Second, the dynamic fluctuation of the entire molecule that affects the entropic term of the free energy barrier is not taken into account by the MSM method. Third, it was shown that the transition between open- and closed-loop configurations was the slowest process among several reaction steps in the entire reaction.\(^{22}\) On the other hand, the MSM method describes the open to closed transition just as the weight change occurring along the reaction coordinate for the reaction-center part. In other words, actual atomic motions as well as the time required in the transition are not considered explicitly by the MSM method. Nevertheless, the MSM method is useful for predicting the sequence of stability changes in the surrounding structure part occurring along the reaction coordinate for the reaction-center part. Beyond this model, couplings between motions in the reaction-center part and the surrounding structure must be taken into account through QM/MM-MD simulations in the future. Fourth, we used twelve surrounding structures with six open- and six closed-loop conformations in this study. As shown in Figure 3b, the entire catalytic process was represented only with four surrounding structures, that is, “open 1”, “open 3”, “closed 1”, and “closed 4”. From this fact, we concluded that twelve were enough in this system. It should be noted that the number of structures required to describe an entire process would change depending on systems. It is thus recommended to prepare twice or more of structures than those required to describe the system.

4. CONCLUSIONS

We studied the roles of the open- and closed-loop conformations in the transformation reaction of pyruvate to \(\eta\)-lactate in LDH. The substrate binding state and final product binding state are more stable in the open-loop conformation than in the closed-loop conformation. In contrast, the TS for the concerted proton and hydride transfers is more stable in the closed-loop conformation. These stability differences at different states are found to correlate with the variation of the volume of the reaction center along the MEP, where the initial and final binding states are stabilized in the larger reaction space of the open-loop conformation and the TS is stabilized in the smaller reaction space of the closed-loop conformation. We suggest that the exchanges of the stability between the open- and closed-loop conformations at the different states play the central role in promoting the present reaction in one direction.

5. COMPUTATIONAL DETAILS

The initial structure was obtained from the protein data bank (PDB code: 3H3F).\(^{32}\) The structure of LDH from the rabbit muscle consists of two independent tetramers, each of which has four active sites; that is, two with the open-loop conformation and the other two with the closed-loop conformation. The model structure was built using chain C (closed loop) and D (open loop), where oxamate in the PDB structure was replaced by the reactant pyruvate. Hydrogen atoms were added assuming the normal protonation states at a pH of 7.0 using the PROPKA program,\(^{42,43}\) and His192 was manually protonated. The charged protein was neutralized with chloride ions and fully solvated in the octahedron water box. To prepare the surrounding structures in the open- and closed-loop conformations, two replica exchange MD simulations with fixing of reaction-center atoms were performed: one for the open-loop conformation and the other for the closed-loop conformation. In each MD simulation, 20 replicas for the temperature range of 300 to 400 K were used and 5 000 000 steps were computed for each replica.\(^{31}\) Time step size and cutoff radius were set to 0.1 (fs) and 12.0 Å, respectively. Structures were printed every 10 000 MD steps, and among the 10 000 printed structures (500 for each replica), the six with the lowest energies were chosen as the initial surrounding structures of the open- and closed-loop conformations. Distance parameters defined at the theoretical back ground and distances between CZ-Arg168 and O1-pyruvate, and CZ-Arg168 and O3-pyruvate during MD simulations were shown in Figure S2. Afterward, aqueous environment molecules more than 50 Å away from the center of protein coordinates and counterions were removed for the reaction path calculations, where total charge of the protein was +2. From open- and closed-loop conformations, both six initial structures were merged for the mixed-loop calculations. Among the twelve structures, six have an open-loop conformation and the other six have a closed-loop conformation. It should be noted that our model structure constructs from a single reaction-center and twelve different surrounding structures.

The B3LYP/6-31+G(d,p) level of theory and general AMBER force field\(^{44}\) were used as the high-level QM and low-level MM methods, respectively. The mechanical embedding scheme\(^{33,34}\) was employed in all calculations, while the electronic embedding scheme\(^{35,46}\) was used for single-point calculations. AMBER force field parameters of NADH cofactor and pyruvate were obtained from generalized AMBER force field files (gaff.dat). The reaction center consisted of 46 atoms including the three link atoms placed at the QM/MM boundary indicated by wavy lines in Figure 2. All molecular structures were depicted by the UCSF Chimera package.\(^{47}\) The MEP calculations were performed by the MSM method using the GRRM program combined with the Gaussian 09 program.\(^{30,48,49}\) The initial path of the mixed calculation was obtained by the AFIR method with the model collision energy parameter of the AFIR method set to \(\gamma = 150\) kJ/mol and the model temperature parameter of the MSM method set to \(T_{\text{MSM}} = 50\) 000 K; these settings are discussed in our previous papers.\(^{30,36}\) The artificial force was applied to two atom pairs: between the shifting proton and acceptor oxygen and between the shifting hydride and acceptor carbon. The initial path was further optimized by the LUP path optimization method.\(^{35}\)
where the model temperature parameter of the MSM method was set to $T^{\text{MSM}} = 300$ K. Three different LUP calculations were performed (as discussed in the main text): (i) a mixed calculation, (ii) an open-only calculation, and (iii) a closed-only calculation. During the LUP calculations, all energy maxima were directly optimized to actual TSs using the quasi-Newton algorithm.

### ASSOCIATED CONTENT

#### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.8b02813.

Illustration of distances between key residues defined in the section of the theoretical background, variations in MD simulations, and optimized structures, full reference of the Gaussian 09 program and Cartesian coordinates of the reaction center for all structures described in Figure 4 (PDF)

### AUTHOR INFORMATION

#### Corresponding Authors

*E-mail: ki_suzuki@eis.hokudai.ac.jp* (K.S.).

*E-mail: smaeda@eis.hokudai.ac.jp* (S.M.).

**ORCID**

Satoshi Maeda: 0000-0001-8822-1147

#### Notes

The authors declare no competing financial interest.

### ACKNOWLEDGMENTS

This work was supported by a grant from the Japan Science and Technology Agency with a Core Research for Evolutional Science and Technology (CREST) grant in the area of “Establishment of Molecular Technology towards the Creation of New Functions” at Hokkaido University (grant no. JPMJCR14LS). Calculations were performed at the Research Center Computational Science (RCCS), Okazaki, Japan. We thank Natasha Lundin, PhD, from Edanz Group ([www.edanzediting.com/ac](http://www.edanzediting.com/ac)) for editing a draft of this manuscript.

### REFERENCES

(1) Wolffen, R.; Snider, M. J. The Depth of Chemical Time and the Power of Enzymes as Catalysts. *Acc. Chem. Res.* 2001, 34, 938–945.

(2) Nesi, B. M.; Hammer, S. C.; Nebel, B. A.; Hauer, B. New Generation of Biocatalysts for Organic Synthesis. *Angew. Chem., Int. Ed. 2014*, 53, 3070–3095.

(3) Raso, J. P.; Voss, E. Enzyme-Catalyzed Processes in Pharmaceutical Industry. *Appl. Catal., A* 2001, 221, 145–158.

(4) Laurent, N.; Haddoub, R.; Flitsch, S. L. Enzyme Catalysis on Solid Surfaces. *Trends Biotechnol.* 2008, 26, 328–337.

(5) Fersht, A. *Structure and Mechanism in Protein Science: A Guide to Enzyme Catalysis and Protein Folding*; W.H. Freeman: New York, 1999.

(6) Yasuda, R.; Noji, H.; Yoshida, M.; Kinoshita, K.; Itoh, H. Resolution of Distinct Rotational Substeps by Submillisecond Kinetic Analysis of F1-ATPase. *Nature* 2001, 410, 898–904.

(7) Kędzierski, P.; Moreton, K.; Clarke, A. R.; Holbrook, J. J. The A245K Mutation Exposes Another Stage of the Bacterial L-Labelate Dehydrogenase Reaction Mechanism. *Biochemistry* 2001, 40, 7247–7252.

(8) Boehr, D. D.; McElheny, D.; Dyson, H. J.; Wright, P. E. The Dynamic Energy Landscape of Dihydropyruvate Reductase Catalysis. *Science* 2006, 313, 1638–1642.

(9) Henzler-Wildman, K. A.; Thai, V.; Lei, M.; Ott, M.; Wolf-Watz, M.; Fenn, T.; Pozharski, E.; Wilson, M. A.; Petsko, G. A.; Karplus, M.; et al. Intrinsic Motions along an Enzymatic Reaction Trajectory. *Nature* 2007, 450, 838–844.

(10) Wang, Z.; Chang, E. P.; Schramm, V. L. Triple Isotope Effects Support Concerted Hydride and Proton Transfer and Promoting Vibrations in Human Heart Lactate Dehydrogenase. *J. Am. Chem. Soc.* 2016, 138, 15004–15010.

(11) Basner, J. E.; Schwartz, S. D. How Enzyme Dynamics Helps Catalyze a Reaction in Atomic Detail: A Transition Path Sampling Study. *J. Am. Chem. Soc.* 2005, 127, 13822–13831.

(12) Quaytman, S. L.; Schwartz, S. D. Reaction Coordinate of an Enzymatic Reaction Revealed by Transition Path Sampling. *Proc. Natl. Acad. Sci. U.S.A.* 2007, 104, 12253–12258.

(13) Deng, H.; Vu, D. V.; Clinch, K.; Desamero, R.; Dyer, R. B.; Callender, R. Conformational Heterogeneity within the Michaelis Complex of Lactate Dehydrogenase. *J. Phys. Chem. B* 2011, 115, 7670–7678.

(14) Quaytman, S. L.; Schwartz, S. D. Comparison Studies of the Human Heart and Bacillus stearothermophilus Lactate Dehydrogenase by Transition Path Sampling¹. *J. Phys. Chem. A* 2009, 113, 1892–1897.

(15) Pineda, J. R. E. T.; Antoniou, D.; Schwartz, S. D. Slow Conformational Motions That Favor Sub-Picosecond Motions Important for Catalysis. *J. Phys. Chem. B* 2010, 114, 15985–15990.

(16) Pan, X.; Schwartz, S. D. Free Energy Surface of the Michaelis Complex of Lactate Dehydrogenase: A Network Analysis of Microsecond Simulations. *J. Phys. Chem. B* 2015, 119, 5430–5436.

(17) Pan, X.; Schwartz, S. D. Conformational Heterogeneity in the Michaelis Complex of Lactate Dehydrogenase: An Analysis of Vibrational Spectroscopy Using Markov and Hidden Markov Models. *J. Phys. Chem. B* 2016, 120, 6612–6620.

(18) Yadav, A.; Jackson, R. M.; Holbrook, J. J.; Warshel, A. Role of Solvent Reorganization Energies in the Catalytic Activity of Enzymes. *J. Am. Chem. Soc.* 1991, 113, 4800–4805.

(19) Ranganathan, S.; Gready, J. E. Hybrid Quantum and Molecular Mechanical (QM/MM) Studies on the Pyruvate toL-Lactate Interconversion inL-Lactate Dehydrogenase. *J. Phys. Chem. B* 1997, 101, 5614–5618.

(20) Moliner, V.; Williams, I. H. Flexible QM/MM Modelling Embraces Alternative Mechanisms for Lactate Dehydrogenase. *Chem. Commun.* 2000, 19, 1843–1844.

(21) Swiderer, K.; Tuñón, I.; Marti, S.; Moliner, V. Protein Conformational Landscapes and Catalysis. Influence of Active Site Conformations in the Reaction Catalyzed by L-Lactate Dehydrogenase. *ACS Catal.* 2015, 5, 1172–1185.

(22) Dunn, C. R.; Wilks, H. M.; Halsall, D. J.; Atkinson, T.; Clarke, A. R.; Muirhead, H.; Holbrook, J. J. Design and synthesis of new enzymes based on the lactate dehydrogenase framework. *Philos Trans R Soc Lond, B, Biol Sci* 1991, 332, 177–84.

(23) Ferrer, S.; Tunón, I.; Martí, S.; Moliner, V.; García-Viloca, M.; Gonzalez-Lafont, A.; Lluch, J. M. A Theoretical Analysis of Rate Constants and Kinetic Isotope Effects Corresponding to Different Reactant Valleys in Lactate Dehydrogenase. *J. Am. Chem. Soc.* 2006, 128, 16851–16863.

(24) Nishihara, Y.; Kato, S.; Hayashi, M.; Hachiya, Y.; Atkinson, T.; Clarke, A. R.; Muirhead, H.; Holbrook, J. J. Design and synthesis of new enzymes based on the lactate dehydrogenase framework. *Philos Trans R Soc Lond, B, Biol Sci* 1991, 332, 177–84.

(25) Jorgensen, W. L. Free Energy Calculations: A Breakthrough for Modeling Organic Chemistry in Solution. *Acc. Chem. Res.* 1989, 22, 184–189.

(26) Ståhlb, M.; Hong, G.; Warshel, A. Ab Initio QM/MM Simulation with Proper Sampling: “First Principle” Calculations of the Free Energy of the Autodissociation of Water in Aqueous Solution. *J. Phys. Chem. B* 2002, 106, 13333–13343.
(27) Rod, T. H.; Ryde, U. Accurate QM/MM Free Energy Calculations of Enzyme Reactions: Methylation by CatecholO-Methyltransferase. *J. Chem. Theory Comput.* 2005, 1, 1240–1251.

(28) Hu, H.; Yang, W. Free Energies of Chemical Reactions in Solution and in Enzymes with Ab Initio Quantum Mechanics/Molecular Mechanics Methods. *Annu. Rev. Phys. Chem.* 2008, 59, 573–601.

(29) Kosugi, T.; Hayashi, S. QM/MM Reweighting Free Energy SCF for Geometry Optimization on Extensive Free Energy Surface of Enzymatic Reaction. *J. Chem. Theory Comput.* 2011, 8, 322–334.

(30) Suzuki, K.; Morokuma, K.; Maeda, S. Multistuctural Microiteration Technique for Geometry Optimization and Reaction Path Calculation in Large Systems. *J. Comput. Chem.* 2017, 38, 2213–2221.

(31) Case, D. A.; Cheatham, T. E.; Darden, T.; Gohlke, H.; Luo, R.; Merz, K. M., Jr.; Onufriev, A.; Simmerling, C.; Wang, B.; Woods, R. J. The Amber Biomolecular Simulation Programs. *J. Comput. Chem.* 2005, 26, 1668–1688.

(32) Świderek, K.; Panczakiewicz, A.; Bujacz, A.; Bujacz, G.; Paneth, P. Modeling of Isotope Effects on Binding Oxamate to Lactic Dehydrogenase. *J. Phys. Chem. B* 2009, 113, 12782–12789.

(33) Vreven, T.; Morokuma, K.; Farkas, O.; Schlegel, H. B.; Frisch, M. J. Geometry optimization with QM/MM, ONIOM, and other combined methods. I. Microiterations and constraints. *J. Comput. Chem.* 2003, 24, 760–769.

(34) Chung, L. W.; Sameera, W. M. C.; Ramozzi, R.; Page, A. J.; Hatanaka, M.; Petrova, G. P.; Harris, T. V.; Li, X.; Ke, Z.; Liu, F.; et al. The ONIOM Method and Its Applications. *Chem. Rev.* 2015, 115, 5678–5796.

(35) Choi, C.; Elber, R. Reaction Path Study of Helix Formation in Tetrapeptides: Effect of Side Chains. *J. Chem. Phys.* 1991, 94, 751–760.

(36) Maeda, S.; Morokuma, K. Communications: A systematic method for locating transition structures of A+X→B−X type reactions. *J. Chem. Phys.* 2010, 132, 241102.

(37) Peng, H.-L.; Deng, H.; Dyer, R. B.; Callender, R. Energy Landscape of the Michaelis Complex of Lactate Dehydrogenase: Relationship to Catalytic Mechanism. *Biochemistry* 2014, 53, 1849–1857.

(38) Peng, H.-L.; Egawa, T.; Chang, E.; Deng, H.; Callender, R. Mechanism of Thermal Adaptation in the Lactate Dehydrogenases. *J. Phys. Chem. B* 2015, 119, 15256–15262.

(39) Marx, D. Proton Transfer 200 Years after Von Grotthuss: Insights from Ab Initio Simulations. *ChemPhysChem* 2006, 7, 1849–1870.

(40) Suzuki, K.; Shiga, M.; Tachikawa, M. Temperature and Isotope Effects on Water Cluster Ions with Path Integral Molecular Dynamics Based on the Fourth Order Trotter Expansion. *J. Chem. Phys.* 2008, 129, 144310.

(41) Świderek, K.; Paneth, P. Importance of the Lactate Dehydrogenase Quaternary Structure in Theoretical Calculations. *J. Phys. Chem. B* 2010, 114, 3393–3397.

(42) Li, H.; Robertson, A. D.; Jensen, J. H. Very fast empirical prediction and rationalization of protein pKa values. *Proteins: Struct., Funct., Genet.* 2005, 61, 704–721.

(43) Bas, D. C.; Rogers, D. M.; Jensen, J. H. Very fast prediction and rationalization of pKa values for protein-ligand complexes. *Proteins: Struct., Funct., Genet.* 2008, 73, 765–783.

(44) Wang, J.; Wolf, R. M.; Caldwell, J. W.; Kollman, P. A.; Case, D. A. Development and Testing of a General Amber Force Field. *J. Comput. Chem.* 2004, 25, 1157–1174.

(45) Bakowies, D.; Thiel, W. Hybrid Models for Combined Quantum Mechanical and Molecular Mechanical Approaches. *J. Phys. Chem.* 1996, 100, 10580–10594.

(46) Vereven, T.; Byun, K. S.; Komáromi, I.; Dapprich, S.; Montgomery, J. A., Jr.; Morokuma, K.; Frisch, M. J. Combining Quantum Mechanics Methods with Molecular Mechanics Methods in ONIOM. *J. Chem. Theory Comput.* 2006, 2, 815–826.