

Programming xenon diffusion in maltose-binding protein

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ABSTRACT Protein interiors contain void space that can bind small gas molecules. Determination of gas pathways and kinetics in proteins has been an intriguing and challenging task. Here, we combined computational methods and the hyperpolarized xenon-129 chemical exchange saturation transfer (hyper-CEST) NMR technique to investigate xenon (Xe) exchange kinetics in maltose-binding protein (MBP). A salt bridge ~ 9 Å from the Xe-binding site formed upon maltose binding and slowed the Xe exchange rate, leading to a hyper-CEST ^{129}Xe signal from maltose-bound MBP. Xe dissociation occurred faster than dissociation of the salt bridge, as shown by ^{13}C NMR spectroscopy and variable- B_1 hyper-CEST experiments. “Xe flooding” molecular dynamics simulations identified a surface hydrophobic site, V23, that has good Xe binding affinity. Mutations at this site confirmed its role as a secondary exchange pathway in modulating Xe diffusion. This shows the possibility for site-specifically controlling xenon protein-solvent exchange. Analysis of the available MBP structures suggests a biological role of MBP’s large hydrophobic cavity to accommodate structural changes associated with ligand binding and protein-protein interactions.

SIGNIFICANCE It is possible to program the ^{129}Xe NMR signal in maltose-binding protein by engineering two sites—a salt bridge (normally stabilized by maltose binding) and a surface hydrophobic site, V23—which are each separated by 1 nm from a central xenon-binding cavity. The ability to control the linear flow of xenon in a protein has applications for magnetic resonance sensors employing hyperpolarized Xe-129 and is relevant to understanding naturally occurring gas-binding proteins.

INTRODUCTION

Proteins are porous materials with internal cavities of various size and duration. Calculation of high-resolution protein crystal structures found that the packing fraction of protein cores is only $\phi \approx 0.56$, significantly lower than the ideal packing fraction of solid spheres ($\phi \approx 0.74$ for face-centered cubic crystals) (1). This reveals that the interiors of most natural proteins contain considerable empty space between amino acids. These “packing defects” play an important role in controlling protein dynamics and functions, especially in the case of gas-binding proteins. Internal hydrophobic cavities and pores define pathways for small gas molecules and allow efficient diffusion to the protein active site. For example, X-ray crystallography and photodissociation experiments confirmed that myoglobin possesses multiple preexisting internal cavities that are

involved in oxygen diffusion and reactivity (2). Computational methods also provide insights into gas pathways that are difficult to probe by experiments (3). Hydrogenases are another class of proteins where kinetics of gas diffusion is of great interest owing to their bioenergetic metabolism involving H_2 conversion (4). One challenge in studying gas diffusion in hydrogenase is that H_2 is invisible in X-ray crystallography and difficult to model accurately by computational methods. Heavy noble gases such as xenon and krypton are therefore frequently used as a surrogate for small gaseous molecules such as H_2 (5,6) and CO (7,8) to probe protein interiors and gas dynamics given their comparable molecular size (4.3 Å diameter for xenon [Xe], 4.0 Å for krypton [Kr], and 3.4–3.8 Å for H_2) (9,10). Analysis on the basis of krypton-pressurized crystal structures and molecular dynamics (MD) simulations predicted possible tunnels for both H_2 and the inhibitor O_2 (5,11–13). However, gas diffusion rates were still unknown without input from experimental approaches, such as protein film voltammetry and isotope exchange assays (14). Moreover, gas delivery in proteins is important in various biological processes that

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are fundamental to all life, including carbon and nitrogen fixation and noble gas narcosis (15–20). Investigation of gas migration pathways largely relied on computational methods, but kinetic information and the mechanisms of gas exchange remain poorly studied. Knowing the kinetics of gas exchange processes in proteins is useful for deciphering catalytic or inhibitory mechanisms and advancing biotechnology applications.

Interestingly, large internal cavities and channels are found not only in gas-reactive proteins and gas vesicle nanostructures (21,22) but also in many other monomeric proteins whose functions are completely unrelated to gas permeability and interactions, such as periplasmic binding proteins, T4 lysozyme, and TEM-1 β -lactamase (23–29). These proteins can host Xe using a large hydrophobic cavity, thus allowing the spin-1/2 nucleus, ^{129}Xe , to serve as a biocompatible probe for the study of protein-protein interactions (30) and small-molecule sensing (23,24). For example, maltose-binding protein (MBP) in its wild-type form has a cavity volume of 75–95 \AA^3 (26), which is surprising because an intramolecular cavity of this large size is rarely found in naturally occurring proteins (31) and void space is usually associated with protein instability despite the high stability of MBP. The use of ^{129}Xe NMR spectroscopy enabled detection of the protein cavity and conformational changes in MBP (32). In combination with X-ray crystallography and heteronuclear single-quantum coherence spectroscopy, an Xe-binding cavity was identified in the N domain, and Xe binding affinity was estimated ($K_a \sim 20 \pm 10 \text{ M}^{-1}$) (26). Using hyperpolarized ^{129}Xe chemical exchange saturation transfer (hyper-CEST) (33), our laboratory observed a distinctive chemical shift for ^{129}Xe bound to maltose-saturated MBP, shifted ~ 90 ppm downfield relative to free aqueous ^{129}Xe (23). However, it remains unexplained how the maltose-bound state, instead of the maltose-free state, produces the unique ^{129}Xe signal. This prompted us to investigate the structural basis for Xe-MBP interactions, which could guide future design of novel Xe-based biosensors.

In this paper, we employ MD simulations to find Xe exchange pathways in MBP and the hyper-CEST technique to characterize Xe dissociation rate. Maltose exchange kinetics and mutations made to a surface hydrophobic site provide insights into additional Xe exchange mechanisms. Cavity volume analysis reveals the possible biological role of the large hydrophobic pocket.

MATERIALS AND METHODS

MD simulations

MD simulations were performed with NAMD 2.14 software (34) using the Bridges-2 Regular Memory system at the Pittsburgh Supercomputing Center (35,36). Initial structures were obtained by adding a Xe atom to MBP crystal structures (PDB: 1LLS for maltose-free form (26) and PDB: 1ANF for maltose-bound form (37)). Each protein was solvated in a TIP3P water box (38), and 150 mM NaCl was used to neutralize total

charge. Each system was minimized with 1,000 steps, after which the system was equilibrated at a temperature of 300 K and a pressure of 1 atm using the Langevin thermostat and barostat methods. The CHARMM36 force field was used to perform all MD simulations (39). The Xe atom was treated with only nonbonded interactions (40). Periodic boundary conditions were employed, and the SHAKE algorithm was applied to constrain the lengths of all bonds that involve a hydrogen (41). At least a 43 ns MD trajectory was obtained from each simulation. Salt-bridge distances and root-mean-square deviation data were calculated using VMD 1.9.3. (42).

Cavity volume and tunnel analysis

Cavity volume was calculated using the Fpocket algorithm (43), which detects protein cavity based on Voronoi tessellation. Input structures were obtained from PDB. Tunnel analysis was carried out on MD trajectories of MBP using Caver 3.0 software (44).

Site-directed mutagenesis

K15D, D14K/K15D, V23A, and V23L mutations were introduced to the pET His6 MBP TEV LIC cloning vector, a gift from Scott Gradia acquired via Addgene (#29656), using the forward and reverse primers listed in Table S1. The mutated plasmids were amplified in NEB 5 α competent *E. coli* cells (New England Biolabs, Ipswich, MA, USA) and purified using a miniprep kit (Qiagen, Germantown, MD, USA). All mutated sequences were verified at the University of Pennsylvania DNA Sequencing Facility.

Protein expression and purification

Preparation of MBP mutants uses the same procedures as wild-type MBP, which has been described before (23). Briefly, the plasmid encoding the MBP sequence was transformed into *E. coli* BL21(DE3)-competent cells (New England Biolabs), which were then grown in 1 L lysogeny broth media supplemented with 50 $\mu\text{g}/\text{mL}$ kanamycin. Protein expression was induced by addition of 1 mM isopropyl- β -D-thiogalactoside, followed by overnight incubation at 25°C. Cell lysis was carried out using hen egg white lysozyme (Alfa Aesar, Ward Hill, MA, USA) in 20 mM sodium phosphate (pH 7.4) in the presence of benzonase nuclease (Sigma-Aldrich, Burlington, MA, USA). The lysate was subjected to fast protein liquid chromatography. Pure protein was eluted from HisTrap HP Ni column (Cytiva, Marlborough, MA, USA) and HiLoad 16/600 Superdex 75 pg column (GE Life Sciences, Chicago, IL, USA). Protein concentrations were determined from the absorbance at 280 nm using the extinction coefficient 67,840 $\text{M}^{-1} \text{ cm}^{-1}$, as calculated by PROTPRAM (45).

^{129}Xe hyper-CEST NMR

HP ^{129}Xe was generated using the spin-exchange optical pumping method (46) with a home-built ^{129}Xe polarizer based on the IGLXe.2000 commercial model by GE. A Shark 65 W tunable ultranarrow band diode laser (OptiGrate, Oviedo, FL, USA) set to 794.770 nm was used for optical pumping of Rb vapor. A gas mixture of 88% helium, 10% nitrogen, and 2% natural abundance Xe (Linde Group, Bridgewater, NJ, USA) was used as the hyperpolarizer input. ^{129}Xe hyperpolarization level was roughly 10%–15%. For each data point in the hyper-CEST z -spectra, hyperpolarized (hp) ^{129}Xe was bubbled into the NMR tube through capillaries for 20 s, followed by a 3 s delay to allow the bubbles to collapse. A d-SNOB saturation pulse was used. The pulse length was $\tau_{\text{pulse}} = 3.80$ ms, number of pulses was $n_{\text{pulse}} = 600$, and saturation time was $T_{\text{sat}} = 2.28$ s. NMR experiments were performed using a Bruker BioDRX 500 MHz NMR spectrometer and 10 mm PABBO probe at 300 K. A 90° hard pulse of this probe has a pulse length of 22 μs . Protein samples were 80 μM in PBS (pH 7.4),

with 0.1% (v/v) Pluronic L81 (Aldrich) added to mitigate foaming. All results were averaged over at least three trials.

¹²⁹Xe exchange rate determination by variable- B_1 measurements

Estimation of $k_{\text{off,Xe}}$ is based on an approximation to the full hyper-CEST solution (47). As described previously (48), the Xe dissociation rate $k_{\text{off,Xe}}$ can be obtained by measuring the CEST effect with various saturation strengths, B_1 . Exponential Lorentzian fits were applied to model the CEST peaks of protein-bound and free ¹²⁹Xe. A simplified expression for hyper-CEST is

$$\text{CEST} = 1 - e^{-\lambda t},$$

where $\lambda = \frac{fk_{\text{off}}(\gamma B_1)^2}{(\gamma B_1)^2 + k_{\text{off}}}$ and k_{off} represents the ¹²⁹Xe dissociation rate. Rearrangement of the above equations gives

$$\ln^{-1}(1 - \text{CEST}) = -\frac{k_{\text{off}}}{tf} \times \frac{1}{(\gamma B_1)^2} - \frac{1}{tfk_{\text{off}}}.$$

Therefore, the linear plot of $\ln^{-1}(1 - \text{CEST})$ versus $(\gamma B_1)^{-2}$ gives $k_{\text{off,Xe}} = (\text{slope}/y\text{-intercept})^{1/2}$. Protein samples were prepared at 80 μM concentration in PBS (pH 7.4) with 0.1% (v/v) Pluronic L81. Each z-spectrum was averaged over at least three trials.

¹³C NMR spectroscopy

0.6 mM protein was mixed with 2 mM D-maltose in PBS (pH 7.4) with 10% D₂O. Data collection was performed on a Bruker NEO600 NMR spectrometer. Sample temperature was 300 K. Maltose dissociation rate, $k_{\text{off, maltose}}$, was estimated based on the line shape of ¹³C signal of the bound maltose. The apparent transverse relaxation of the bound signal results from both intrinsic relaxation and chemical exchange. As a rough estimation, intrinsic relaxation can be neglected given the broad width of the bound signal. Therefore, $k_{\text{off, maltose}} \approx 1/T_2^* = \pi \times w_{1/2}$.(49).

Fluorescence spectroscopy

For fluorescence quenching of MBP mutants, 1 μM protein was prepared in PBS buffer for measurement. A stock of 100 mM maltose in PBS buffer was added to the protein solution to a final concentration of 1 mM. The excitation wavelength was 280 nm, and fluorescence emission was recorded over a range of 300 to 420 nm. All measurements were obtained at 298 K using a Varian Cary Eclipse fluorescence spectrophotometer.

RESULTS

Salt bridge K15-E111 controls Xe exchange

The different ¹²⁹Xe hyper-CEST response in the presence and absence of maltose indicates a difference in either the Xe binding environment or exchange kinetics. In order to characterize the differences in dynamics, we performed MD simulations on MBP in both the maltose-bound “closed” conformation and maltose-free “open” conformation, initiated with a single Xe atom in the well-defined Xe-binding cavity (23). In the closed conformation, the salt bridge K15-E111 forms an H-bond network with hydroxyl groups

of maltose, whereas the same side chains show no interactions in the open conformation (Fig. 1). Interestingly, the Xe-binding pocket resides underneath this salt bridge at a distance of ~ 9 Å (Fig. 2), which leads us to postulate that the K15-E111 salt bridge forms a barrier between the bound Xe atom and solvent. Superposition of the cavity-containing N domains of both MBP conformations shows almost identical structure, with an average root-mean-square deviation of only 0.33 Å, indicative of no change in the Xe binding environment in different conformations of MBP. In addition to protein dynamics, we also analyzed Xe trajectories. Notably, in the closed state, Xe is constrained by the protein matrix during the 43 ns simulation with a root-mean-square fluctuation of 1.6 Å, only $\sim 38\%$ of the van der Waals diameter of Xe. In contrast, during the same simulation length of the open state, Xe can escape from the initial hydrophobic cavity through the crevice between K15 and E111 and finally become completely solvated. Caver tunnel analysis estimates the average width of this crevice to be at least 2.2 ± 0.3 Å in the open state and 1.8 ± 0.4 Å in the closed state during the entire simulation length (50). Despite this narrow bottleneck, Xe can pass the tunnel probably due to transient rearrangement of its surrounding amino acids. Further analysis revealed the most frequently involved bottleneck residues include I11, K15, and L262.

We set out to test experimentally our hypothesis about the effect of the K15-E111 salt bridge on Xe motion. To this end, we attempted to destroy the salt-bridge interaction with a K15D mutation and then restore it with a second mutation, D14K. Salt-bridge analysis using MD trajectories of both mutants showed distinct results. Even though both mutants can bind maltose as observed in MD simulations and fluorescence quenching experiments (Fig. S1), the salt bridge exists only in the D14K/K15D double mutant, with

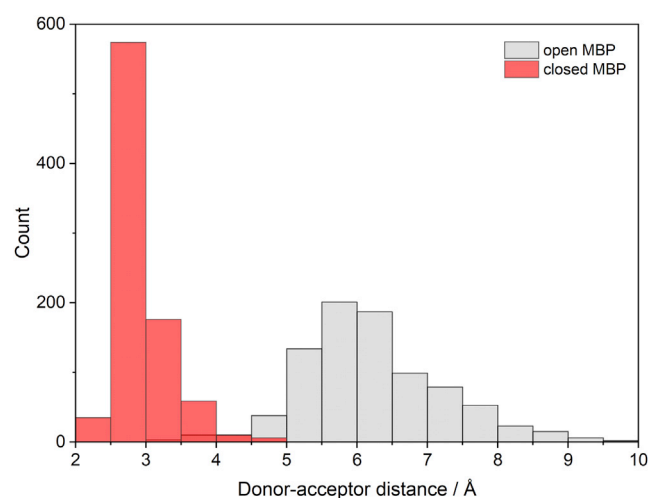


FIGURE 1 Donor-to-acceptor distance between K15 and E111 in maltose-free open MBP (gray) and maltose-bound closed MBP (red). To see this figure in color, go online.

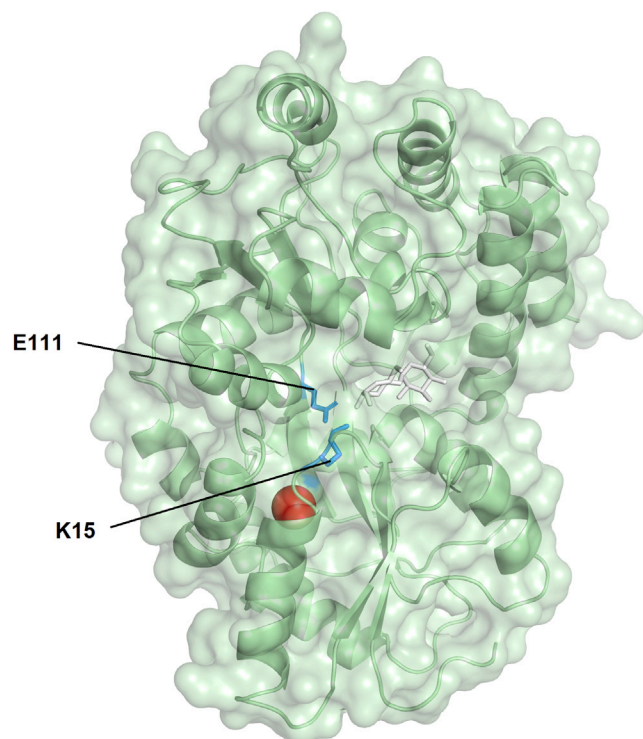


FIGURE 2 Cartoon representation of closed MBP with a bound Xe atom (red sphere). Maltose is shown as white sticks, and side chains of K15 and E111 are shown as blue sticks. To see this figure in color, go online.

an occupancy of 87% (Fig. S2). As a result, K15D protein exhibits no bound ^{129}Xe NMR signal, similar to the open state, whereas D14K/K15D is able to restore the bound ^{129}Xe NMR signal at +80 ppm (Fig. 3). Our results confirmed the important role of this proximal salt bridge in controlling Xe exchange.

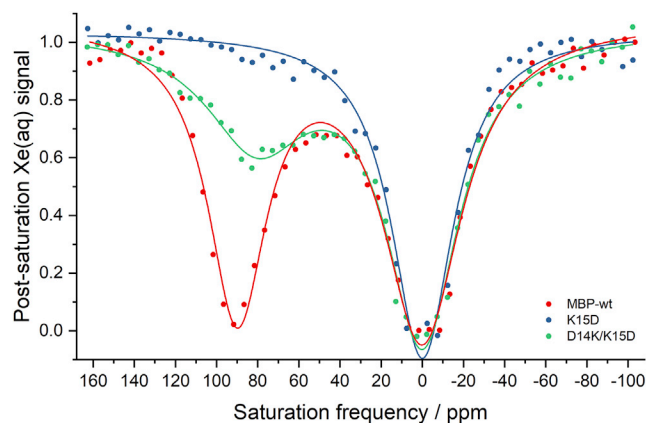


FIGURE 3 Hyper-CEST z-spectra of wild-type MBP (red), K15D (blue), and D14K/K15D (green) in presence of 1 mM maltose. 80 μM protein was dissolved in PBS (pH 7.4) at 298 K. Wild-type MBP gives a hyper-CEST signal at +90 ppm, referenced to the free ^{129}Xe in PBS. The K15D mutant completely lost its ^{129}Xe @protein signal due to faster Xe exchange, while D14K/K15D was able to restore the hyper-CEST signal at +80 ppm. To see this figure in color, go online.

Alternative Xe exchange pathways

It remains unresolved whether the Xe egress pathway in the closed MBP structure is the same salt-bridge site because it is possible that other pathways that are minor in the open MBP structure could become significant upon closure of the K15-E111 salt bridge. If there are no alternative pathways, the rate of Xe exchange should be comparable to or slower than the dynamics of salt-bridge formation, which in turn depends on maltose dissociation rate. Therefore, we characterized the maltose dissociation rate using $[1\text{-}^{13}\text{C}]$ -labeled maltose and the Xe dissociation rate via variable- B_1 hyper-CEST method (47,48). From the line fitting and peak-width analysis ($k_{\text{off}} \sim \pi \times w_{1/2}$) of the ^{13}C NMR signal of bound maltose, we can estimate the maltose off rate, $k_{\text{off, maltose}}$, to be $1.2 \times 10^2 \text{ s}^{-1}$ for β -maltose and $1.3 \times 10^2 \text{ s}^{-1}$ for α -maltose (Fig. 4). This value agrees with the result of the stopped-flow fluorescence measurement (51) and is larger than other reported values by one order of magnitude (52–54). This is probably because ^{13}C NMR signal reflects the H-bonding of the hydroxyl group directly attached to $1\text{-}^{13}\text{C}$ and intrinsic fluorescence is affected by the local environment of nearby tryptophan, whereas other techniques measured the rate of overall protein conformational change. In this aspect, our result better represents the local dynamics of the salt bridge as the hydroxyl group of $1\text{-}^{13}\text{C}$ forms a direct H-bond with the K15 side chain. Next, we estimated the ^{129}Xe dissociation rate via the variable- B_1 hyper-CEST method (47,48). A series of z-spectra using varying saturation pulse strengths was acquired, and the bound ^{129}Xe signal was analyzed to obtain a Xe dissociation rate, $k_{\text{off, Xe}} = (8.6 \pm 2.1) \times 10^2 \text{ s}^{-1}$ (Fig. 4, b and c). The fact that $k_{\text{off, Xe}}$ is roughly four times greater than $k_{\text{off, maltose}}$ suggests that alternative Xe pathways contribute significantly to gas diffusion in the closed MBP structure.

Tunnel prediction using the hard sphere model may not correctly identify all Xe exchange pathways since tunnel hydrophobicity is not considered. Also, evidence for alternative pathways might be hidden in the single-Xe simulations due to insufficient sampling. Therefore, the “Xe flooding” approach, in which Xe atoms were initially placed in the surrounding solvent, was applied, and the concentration of Xe was artificially increased to $\sim 150 \text{ mM}$ to enhance Xe diffusion through the protein. (This is equivalent to nearly 30 atm Xe at room temperature (55)). Using this computational method, we identified several protein surface sites that associate with Xe exchange due to transient Xe affinity, defined by the average number of bound Xe atoms, n , at each residue of the protein (Fig. 5 a). These sites include V23, F92, and the hinge region of MBP, all of which consist of nonpolar residues. Meanwhile, Xe has relatively low affinity ($n < 0.05$) toward the K15-E111 region, while maltose is stably bound during the simulation. F92 and the hinge region residues were not taken into further consideration

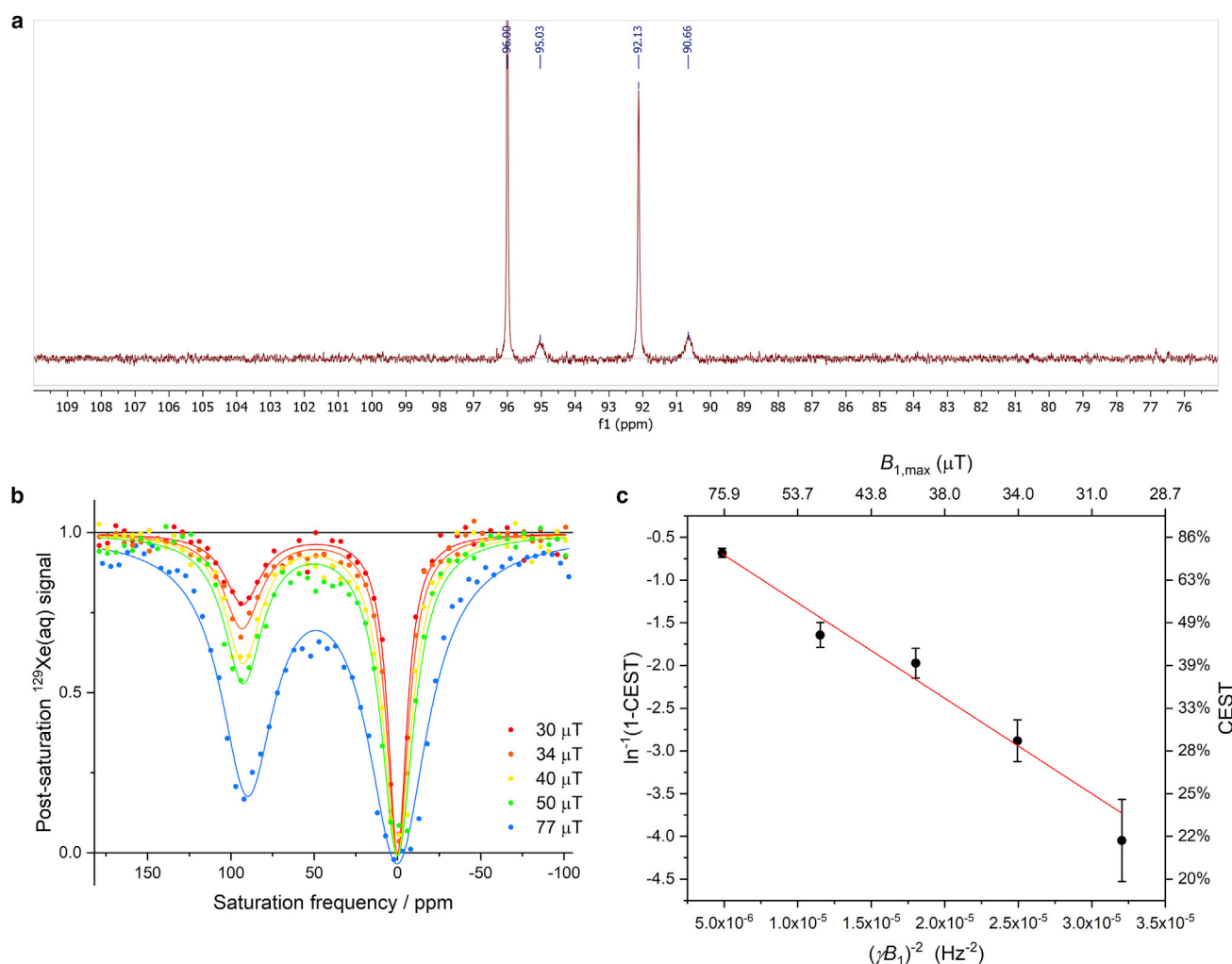


FIGURE 4 (a) ^{13}C NMR spectrum of 2 mM $[1-^{13}\text{C}]$ -maltose and 0.6 mM MBP in PBS (pH 7.4). From high frequency to low frequency are free β -maltose ($\delta = 96.00$ ppm), bound β -maltose ($\delta = 95.03$ ppm), free α -maltose ($\delta = 92.13$ ppm), and bound α -maltose ($\delta = 90.66$ ppm). The peak width (full width at half maximum) is 37.7 Hz for bound β -maltose and 40.4 Hz for bound α -maltose. (b) Hyper-CEST z-spectra of wild-type MBP taken in a series of saturation pulse strengths from 30 to 77 μT at 300 K. (c) Linear fitting of $\ln^{-1}(1-\text{CEST})$ versus $(\gamma B_1)^{-2}$ yields the slope = $(-1.1 \pm 0.1) \times 10^5 \text{ s}^{-2}$, y-intercept = -0.15 ± 0.09 , and $k_{\text{off,Xe}} = (\text{slope}/\text{y-intercept})^{1/2} = (8.6 \pm 2.1) \times 10^2 \text{ s}^{-1}$. To see this figure in color, go online.

because of their long distance ($>18 \text{ \AA}$) from the Xe cavity. Importantly, the solvent-exposed residue V23 is only $\sim 10 \text{ \AA}$ from the Xe binding pocket with a relatively high Xe affinity ($n = 1.0$) (Fig. 5 b). Xe accumulation at this site and its close proximity to the cavity may lead to a considerable rate of Xe exchange through V23. In order to examine the contribution of the V23 pathway, we mutated V23 to the smaller alanine and the bulkier leucine and performed variable- B_1 measurements. Both mutations caused less than 5 ppm difference in chemical shift of the bound ^{129}Xe NMR signal, meaning that the Xe binding pocket was minimally perturbed. Moreover, V23A and V23L mutations do not affect maltose-MBP interactions or salt-bridge dynamics, as confirmed by ^{13}C NMR spectroscopy (Figs. S3 and S4). Consequently, the Xe dissociation rate, $k_{\text{off,Xe}}$, increased to $(1.2 \pm 0.5) \times 10^3 \text{ s}^{-1}$ with V23A mutation (Fig. S5) and decreased to $(5.4 \pm 0.3) \times 10^2 \text{ s}^{-1}$ with the introduction of V23L

(Fig. S6). We then inspected the “Xe flooding” MD trajectories of V23A and V23L proteins. Surprisingly, V23L no longer exhibits high Xe affinity ($n = 0.4$) at this site, although other affinity sites are mostly unchanged, whereas V23A still has a similar affinity ($n = 0.8$) compared with wild-type (Fig. S7). These results confirm that V23 is one of the exchange pathways, which can be explained by a hydrophobic cryptic site between the V23 side chain and the aliphatic part of K26.

Analysis of cavity size in different states of MBP

In addition to gas exchange rate and pathways in MBP, there is also curiosity regarding the origin of gas binding. Xe binding to MBP’s unusually large internal cavity serves no obvious function, and it is unclear about its biological role in different states of MBP. It has been well characterized

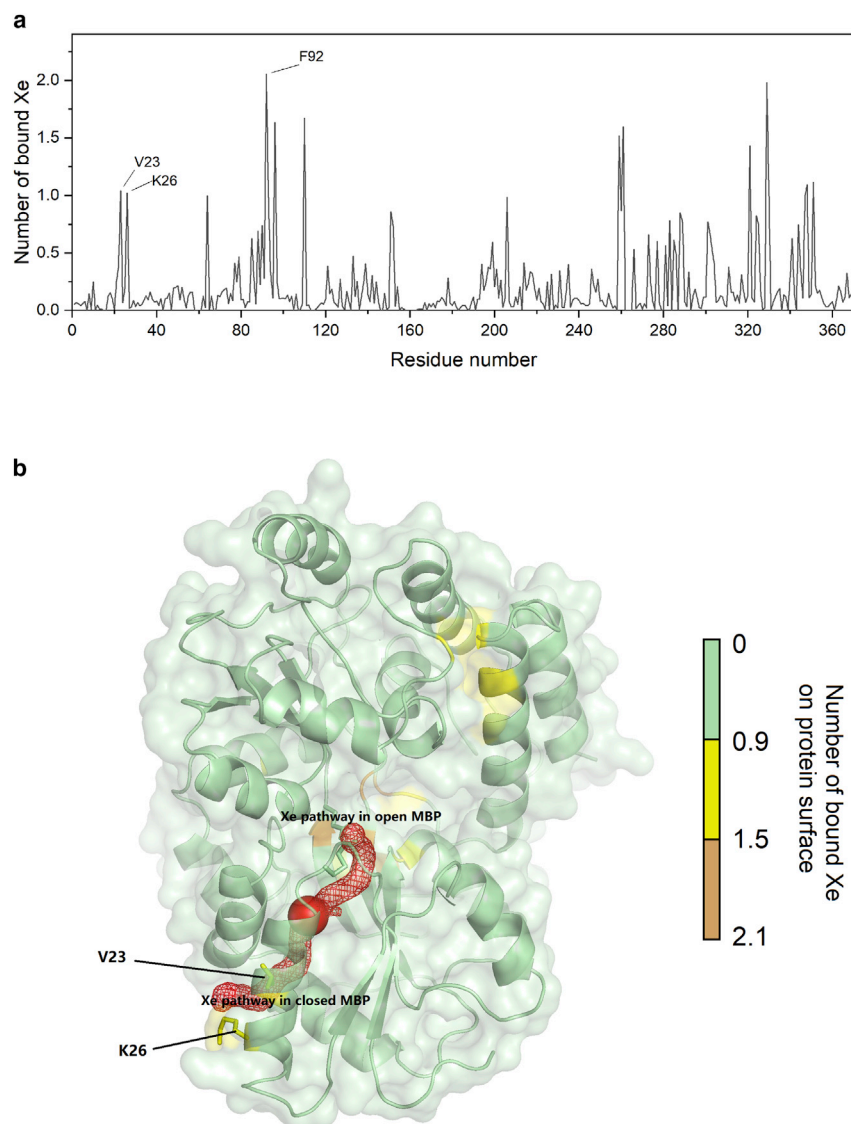


FIGURE 5 (a) Average number of Xe atoms bound to each residue of MBP during the “Xe flooding” simulation. Xe affinity for the protein surface is represented by peaks, which indicate the number of bound Xe atoms. (b) Visualization of Xe affinity sites on MBP surface. Interior Xe binding site is depicted as the red sphere. Xe exchange pathway switches from the site of K15-E111 salt bridge to surface hydrophobic site V23 upon formation of the K15-E111 salt bridge. To see this figure in color, go online.

by crystallography that MBP interacts with a variety of binding substrates, including maltodextrins of various lengths and other oligosaccharides, in different conformations (37,56,57), as well as the maltose transporter complex, MalFGK₂ (58). Cavity volume was then calculated using Fpocket (43) based on the crystal structures of MBP in complex with various biomolecules. The largest cavity found in each structure corresponds to the same site as the Xe pocket. As listed in Table 1, the unliganded MBP harbors a cavity of 97 Å³, very similar to the cavity volume found in Xe-occupied cryptophane-A and its derivatives (85–89 Å³) (59). Binding of maltooligosaccharides causes only minor volume change (10%–20%). However, the cavity shrinks by 40% when maltodextrin-bound MBP interacts with MalFGK₂, which cannot be explained merely by crystal packing effects. Instead, the extensive protein-protein interaction probably requires MBP to arrange itself into a more compact structure.

DISCUSSION

Our previous work has discussed the possibility of different Xe exchange rates in open and closed MBP conformations, but the structural basis was elusive, and no kinetic information was obtained (23). The current study reveals that the salt bridge K15-E111 is pivotal to Xe exchange rate by

TABLE 1 Cavity analysis of MBP in different structures.

Description	PDB code	Cavity size, Å ³
Unliganded MBP	1OMP	97
Maltose bound	1ANF	101
Maltotriose bound	3MBP	92
Maltotetraose bound	4MBP	121
Unliganded MBP in complex with MalFGK ₂	4K10	87
Maltotetraose bound, in complex with MalFGK ₂	4KHZ	58

combining computational analysis and hyper-CEST experiments. In the maltose-bound closed conformation, this salt bridge helps to constrain Xe in the hydrophobic cavity, resulting in a $k_{\text{off,Xe}}$ of $(8.6 \pm 2.1) \times 10^2 \text{ s}^{-1}$. However, in the absence of the K15-E111 salt bridge, MBP in the open conformation can release Xe at a much faster rate. Xe escape from the protein cavity adopts a strategy similar to the “breathing motion” of gas tunnels in myoglobin. Although the bottleneck of the Xe tunnel is only about half of its van der Waals diameter, the dynamics of the bottleneck are highly dependent on the salt bridge, as K15 participates in the formation of a bottleneck. Transient rearrangement of the local environment promoted by the lack of the salt-bridge interaction allows faster Xe diffusion. Notably, the bound maltose is barely involved in the bottleneck, suggesting that the salt bridge itself, instead of the bound maltose, determines Xe exchange. This conclusion may be applied to study the rate-determining step of general gas-protein interactions. According to the quantitative hyper-CEST model (47,60), the CEST signal is optimal when Xe dissociation rate $k_{\text{off,Xe}}$ is close to the saturation pulse strength. Considering the broad peak width of protein hyper-CEST signal and the potential heating effect from strong pulses, the saturation pulse strength γB_1 used in hyper-CEST experiments is usually limited to $\sim 5 \times 10^2 \text{ Hz}$ ($B_{1,\text{max}} \approx 77 \mu\text{T}$ using dSNOB pulses). Therefore, the $k_{\text{off,Xe}}$ in the closed conformation is nearly optimal to allow efficient depolarization of ^{129}Xe , giving a strong hyper-CEST signal of the maltose-MBP-bound ^{129}Xe . The fast off rate, $k_{\text{off,Xe}}$, in the open conformation reaches the “weak saturation” condition, leading to inadequate depolarization of the fast exchanging ^{129}Xe by the same pulse strength. Consequently, the bound ^{129}Xe signal can only be observed in the presence of maltose even though the Xe-binding cavity remains the same without maltose. We also showed that the ^{129}Xe CEST signal could be manipulated by programming the salt bridge. Switching off the salt bridge by K15D mutation can completely turn off the bound ^{129}Xe signal, whereas a reconstructed salt bridge D14K-K15D was able to restore the bound ^{129}Xe NMR signal.

It is conceivable that the salt bridge can be functionalized by rational protein engineering. Metal coordination and organic-covalent interactions can form a switchable and controllable bridge at this site to regulate ^{129}Xe exchange and generate different ^{129}Xe NMR responses, which holds great promise for ^{129}Xe NMR/magnetic resonance imaging applications, such as genetically encoded metal biosensors and light/electro-controlled contrast agents. Design and characterization of novel MBP-based ^{129}Xe NMR biosensors will be elaborated in future publications.

Formation of the K15-E111 salt bridge blocks the fastest Xe exchange pathway with an opening rate of $\sim 10^2 \text{ s}^{-1}$, as estimated by NMR line shape analysis. This method measures ligand exchange at a millisecond-to-second scale

with reasonable approximations, and 1- ^{13}C -labeled maltose represents the local dynamics of the K15-E111 salt bridge. Xe residence time ($1/k_{\text{off,Xe}} = 1.2 \pm 0.3 \times 10^{-3} \text{ s}$) is shorter than the salt-bridge lifetime ($\sim 10^{-2} \text{ s}$), and prior studies showed that the conformational transition rate of maltose-bound MBP is even slower ($\sim 1 \text{ s}$), suggesting that Xe can dissociate without opening the K15-E111 salt bridge or switching protein conformation. Our “Xe flooding” simulation found that Xe has relatively high affinity for several hydrophobic sites on the protein surface but almost no affinity toward the polar environment of the K15-E111 salt bridge and maltose-binding site. In addition, solvent water and salt ions may form a shell at the maltose-binding site via H-bonding and electrostatic interactions, preventing Xe access and egress through this site. On the other hand, surface hydrophobic residues such as V23 can increase local Xe concentration by providing a transient binding site and help to desolvate Xe atoms. Similar surface aggregation of inert gas molecules near hydrophobic regions was also observed in pepsin (61). As a result, surface hydrophobic residues reduce the energy barrier for Xe migration, thus enhancing exchange between solvent and a proximal Xe-binding cavity. This action may be analogous to the “breathing motion” as well (62). Our variable- B_1 experiments confirmed that V23 impacts the Xe dissociation rate without affecting the Xe-binding cavity and salt bridge. Importantly, the V23L mutation can inhibit Xe exchange, probably by decreasing local Xe concentration at the surface hydrophobic site. If we hypothetically inhibited all hydrophobic Xe pathways, the low limit of Xe exchange rate can approximate the dynamics of salt-bridge opening. Because the optimal saturation power is found to be $\sqrt{2}$ times the Xe dissociation rate (60), a narrower and stronger hyper-CEST signal will be observed with a slower Xe exchange. It is meaningful to constrain Xe in protein to a greater extent because protein-bound ^{129}Xe is usually in rapid exchange and its chemical shift is not directly observable in a simple ^{129}Xe NMR measurement. It has been reported that $k_{\text{off,Xe}}$ values in myoglobin and hemoglobin are on the order of $10^4\text{--}10^5 \text{ s}^{-1}$ (63). MBP in its closed conformation has a significantly slower $k_{\text{off,Xe}}$ by more than one order of magnitude, which is still not slow enough to give a directly detectable ^{129}Xe NMR signal. Identification and reshaping gas pathways in these proteins could potentially afford protein-based biosensors that are useful in direct-detection ^{129}Xe NMR/magnetic resonance imaging applications.

Investigation of Xe-MBP interactions not only improves our understanding of gas exchange but also raises interesting questions as to the origin of the large packing defect in MBP, which fundamentally determines gas-protein interactions. An internal cavity is often considered to destabilize protein (64,65), but our prior study found that the cavity-filling mutation V293L in unliganded MBP does not increase its thermal stability (melting temperature [T_m] = $53.9^\circ\text{C} \pm 0.7^\circ\text{C}$ for wild-type and $53.8^\circ\text{C} \pm 0.8^\circ\text{C}$ for V293L) (23). Hence, the

internal cavity volume may not correlate with protein stability. More interestingly, although ligand binding to MBP does not change the cavity as evidenced by MD simulations and volume analysis (Table 1), filling the cavity in maltose-bound MBP has a destabilizing effect ($T_m = 61.0^\circ\text{C} \pm 0.8^\circ\text{C}$ for wild-type and $59.5^\circ\text{C} \pm 0.9^\circ\text{C}$ for V293L), suggesting that the cavity may facilitate ligand binding. Presumably, void space strengthens salt-bridge and H-bond interactions, e.g., K15-E111, due to its low dielectric constant and separation from solvent. Furthermore, the N domain of MBP can undergo considerable compression when interacting with the transmembrane complex MalFGK₂ as the cavity becomes 40% smaller. This supports the functions of the cavity in providing structural flexibility to engage with MalFGK₂ and prepare for the release of maltodextrins. Also, compression of the cavity may be detectable using hyper-CEST and push ¹²⁹Xe chemical shift further downfield because it has been noted that smaller cavities in proteins are correlated with larger downfield ¹²⁹Xe NMR shifts (26).

In summary, we have demonstrated the structural details behind the turn on ¹²⁹Xe hyper-CEST signal in MBP and verified the role of the K15-E111 salt bridge in regulating Xe exchange. Dissociation rate of Xe in maltose-bound MBP exceeds the opening rate of the salt bridge, which led to the discovery that Xe exchange can also occur near a surface hydrophobic site, V23. Finally, we discuss the role of the hydrophobic cavity in strengthening ligand binding and protein-protein interactions and its flexibility in different states of the protein. These findings will foster future development and applications of ¹²⁹Xe-based bio-reporters and contrast agents and provide general insights into mechanisms of gas-protein interactions.

SUPPORTING MATERIAL

Supporting material can be found online at <https://doi.org/10.1016/j.bpj.2022.10.025>.

AUTHOR CONTRIBUTIONS

Z.Z. and I.J.D. designed the study. Z.Z., N.A.R., and J.H. performed all experiments and analyzed the data. Z.Z. and I.J.D. wrote the manuscript.

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DECLARATION OF INTERESTS

The authors declare no competing interests.

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