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Effect of Occluded Ligand Migration on the Kinetics and Structural Dynamics of Homodimeric Hemoglobin

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Abstract

Small molecules such as molecular oxygen, nitric oxide, and carbon monoxide play important roles in life, and many proteins require the transport of small molecules to and from the bulk solvent for their function. Ligand migration within a protein molecule is expected to be closely related to the overall structural changes of the protein, but the detailed and quantitative connection remains elusive. For example, despite numerous studies, how occluded ligand migration affects the kinetics and structural dynamics of the R-T transition remains unclear. To shed light on this issue, we chose homodimeric hemoglobin (HbI) with the I114F mutation (I114F), which is known to interfere with ligand migration between the primary and the secondary docking sites, and studied its kinetics and structural dynamics using time-resolved X-ray solution scattering. The kinetic analysis shows that I114F has three structurally distinct intermediates (I₁, I₂, and I₃) as in the wild type (WT), but its geminate CO recombination occurs directly from I₁ without the path via I₂ observed in WT. Moreover, the structural transitions, which involve ligand migration (the transitions from I₁ to I₂ and from I₃ to the initial state) are decelerated compared to WT. The structural analysis revealed that I114F involves generally smaller structural changes in all three intermediates compared to WT.

Graphical Abstract

Supporting Information.

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Sample preparation; Data collection; Data processing; SVD analysis; Kinetic model; kinetic analysis; Origin of the small amplitude in the SADS curve of I114F; Relation between the amplitude of the SADS curve and the magnitude of the structural change; Structure refinement; Photoconversion yield; and Table and Figures (PDF)



Keywords

picosecond time-resolved X-ray solution scattering; homodimeric hemoglobin; ligand migration; protein structural dynamics

INTRODUCTION

Ligand binding to a protein is a fundamental process for the protein function in biology. The active site for ligand binding is usually located inside of the protein and surrounded by residues that stabilize the bound ligand or the intermediate structure of the substrate. Ligand binding to a protein requires the translocation of the ligand from the bulk solvent to the active site inside the protein, which is often facilitated by conformational changes of the protein.^{1–2} Therefore, the ligand migration pathway within the protein and its connection to protein conformational changes have been the topic of extensive studies.^{1–15} Heme proteins such as myoglobin (Mb), hemoglobin (Hb), and homodimeric hemoglobin (HbI), which play crucial roles in life by storing and delivering simple diatomic molecules such as molecular oxygen, nitric oxide, and carbon monoxide, have been targeted for such studies.^{16–17}

HbI is an excellent model system to investigate the ligand-linked structural changes during allosteric regulation.^{18–20} Unlike Mb, which is a monomeric protein, the binding of the ligand in a multimeric protein is more complicated due to the additional effects of allosteric regulation, which generally proceeds by a series of structural changes.²¹ For example, the binding of the ligand (effector molecule) to one active site triggers subsequent structural changes that affect the reactivity of a distant active site. Hb is a textbook example, but its tetrameric structure naturally involves highly complex reaction dynamics.^{21–27} In this regard, HbI offers a simpler model system due to its homodimeric structure, and it has been studied to observe ligand migration during allosteric regulation with various site-directed mutations that can alter the ligand migration pathways.^{28–31}

X-ray crystallography^{3–9, 29, 31} and optical and infrared spectroscopic tools^{10–15, 28, 30} have been used to investigate the ligand migration pathways in heme proteins. To gain better insight into ligand migration, a series of mutants, which can modify protein-ligand interactions and the ligand migration pathways by introducing a voluminous residue to block the putative transient ligand docking sites or to occlude possible migration pathways between these sites, have been made and investigated.^{15, 30–31} As a result of these comprehensive studies of the ligand migration pathways of heme proteins, for example, it is known that for Mb the ligand migrates between the cavities inside the protein, such as the

primary docking site and Xe sites (secondary docking sites).^{6–8} Those cavities act as transient ligand docking sites, thereby reducing the possibility of the immediate rebinding to the heme group and consequently facilitating the easy escape of the ligand from the protein to the bulk solvent. It is also known that the migration of ligands requires structural changes in the protein that facilitate the movement of the ligand between internal cavities and the binding site.^{1–2} However, despite numerous studies, it remains elusive how occluded ligand migration affects the kinetics and structural dynamics of the R-T transition. To address this issue, information about both ligand migration and the global structural changes should be obtained, though this is not a trivial task. Here, we applied time-resolved X-ray solution scattering (TRXSS) to the I114F mutant of HbI (I114F) and compared the results with those from the wild type (WT) to investigate the relationship between ligand migration and the allosteric structural changes of HbI.

I114F is known to modify protein-ligand interaction by occluding possible ligand migration pathways between the heme pocket and the putative transient ligand docking sites.^{30, 32–33} As shown in Figure 1, the 114th residue, together with the 36th and 73rd residues, forms a channel for the migration of a photo-dissociated ligand between the primary docking site (distal pocket) and the secondary docking sites (Xe4 and Xe2 sites). Whereas this channel is relatively free from obstruction in WT, Phe114 in I114F makes this channel narrower.³² As a result, the geminate recombination would occur only from the primary docking site in I114F whereas it also occurs from the secondary docking sites in WT.³⁰ In addition, since the steric hindrance imposed by Phe114 of the mutant restricts the ligand-linked heme movement deep into the subunit in the liganded state of I114F, as a result, the structure of the liganded state of I114F is mainly affected by the mutation. More specifically, the heme-heme distance and the subunit rotation angle in the liganded state of I114F are similar to those in the T state of WT, that is, the unliganded state of WT. Accordingly, the heme movement and the intersubunit rotation involved in the R-T transition are reduced in I114F.³²

The static structures of I114F have been studied by X-ray crystallography³² and the ligand migration in I114F has been studied over wide time and temperature ranges using optical and infrared spectroscopy.^{30, 33} An X-ray crystallographic study has revealed detailed structural differences between the R and T states with the atomic resolutions,³² but could not access the intermediate structures and relevant kinetics. Fourier-transform infrared (FTIR) cryospectroscopy can discern the kinetics of CO migration,³⁰ but it lacks information about global structural changes. TRXSS, also known as time-resolved X-ray liquidography (TRXL), is a relevant tool for probing the global structural changes linked to the CO migration and relevant kinetics.^{35–37} Due to its relatively low sensitivity to position of the CO molecule, the kinetics observed by TRXSS data reflect the kinetics associated with the overall conformation changes rather than the CO movement itself. However, since it is known that I114F interferes with the ligand migration pathway by obstructing the movement of the ligand migration between the primary and the secondary docking sites,³⁰ it serves as an excellent model system to study the effect of an occluded ligand migration pathway on the kinetics of ligand migration and the structural dynamics of a protein. For example, ligand migration is expected to be decelerated in I114F compared to WT, but there has been no experimental evidence supporting this prediction.

EXPERIMENTAL METHODS

TRXSS data of I114F were collected at the 14-ID-B beamline of the Advanced Photon Source (APS). The solution sample of CO-bound I114F HbI(CO)₂ sealed in a capillary with 3.5 mM concentration was excited by 35 ps long laser pulse at 532 nm. Subsequently, a ~100 ps long X-ray pulse probed the sample, and the scattered X-ray photons were recorded as a function of the time delay between the laser pump pulse and the X-ray probe pulse. Time-resolved scattering curves were collected at 36 pump-probe time delays ranging from 100 ps to 56.2 ms (four time points per decade). By taking the difference between the scattering curve measured at each time delay point and the reference scattering curve measured at -5μ s, we obtained time-resolved difference scattering curves, S(q,t). The sample preparation, experiment, and the data processing are provided more in detail in the Supporting Information.

RESULTS AND DISCUSSION

The time-resolved difference X-ray solution scattering curves, S(q, t), for I114F as shown in Figure 2 were measured after the photodissociation of CO ligands from the initial I114F HbI(CO)₂ for time delays from 100 ps to 56.2 ms. To obtain the number of reaction intermediates and time constants of the reaction, we performed a singular value decomposition (SVD) analysis on the TRXSS data. The analysis revealed that the same kinetics framework involving three structurally distinct intermediates (termed I_1, I_2 , and I_3) according to their appearance time), which were previously identified for WT,35 can also be applied for I114F on the basis of the number of significant singular components (Figure S4). We fit the first three right singular vectors multiplied by their singular values with the sum of the exponentials, as shown in Figure S4d, and obtained six time constants of 640 (\pm 96) ps, $23 (\pm 1.5)$ ns, 990 (± 190) ns, 7.6 (± 1.0) µs, 3.2 (± 0.41) ms, and 29 (± 24) ms, whereas seven time constants were found for WT. In order to find the optimal kinetic model based on the observed time constants, the number of intermediates at each time delay should be determined. The number of intermediates in a certain time range can be obtained accurately through SVD analysis of reduced time ranges (see the Supporting Information for the detailed procedures). As a result, the optimal kinetic model for I114F (Figure 3a) is found and compared with that of WT (Figure 3b).³⁵ Note that the latest two time constants are replaced by a bimolecular rate constant in the kinetic model. The main difference between I114F and WT is the fact that I114F has a different geminate CO recombination pathway, which is responsible for the lack of one time constant in I114F. As shown in Figure 2, the theoretical difference scattering curves (red curves) obtained from the kinetic model are in excellent agreement with the experimental difference scattering curves (black curves) at all time delays. From the kinetic analysis, we obtained the time-dependent population changes (Figure 3c) and the species-associated difference scattering (SADS) curves of each individual intermediate (Figure 3d). To distinguish the intermediates for WT and I114F, we labeled I₁, I₂, and I₃ of WT as I₁^{WT}, I₂^{WT}, and I₃^{WT}, respectively, and likewise labeled those of I114F as I1^{I114F}, I2^{I114F}, and I3^{I114F}, respectively.

The kinetic models for I114F and WT are shown in Figures 3a and 3b,³⁵ respectively. Both have three structurally distinct intermediates. The earliest intermediate, I_1 , is formed by laser

excitation from the initial state within our experimental time resolution of ~100 ps. In I114F, 16% of I₁ goes back to the initial I114F HbI(CO)₂ by the geminate recombination of CO with a time constant of 640 ps, and the rest transforms into the second intermediate, I₂, with a time constant of 23 ns, which is substantially (about 7 times) slower than that of WT (3.2 ns). The geminate recombination pathway has the smaller fraction (16 %) in spite of its faster time constant, compared to the I₁-to-I₂ transition pathway, indicating that the geminate recombination for details). The subsequent biphasic I₂-to-I₃ transitions, which are interpreted as the R-T transitions of two substates of fully and partially photolyzed forms, occur with time constants of 990 ns and 7.6 µs in I114F, similar to those of WT (730 ns and 5.6 µs). The last intermediate, I₃, goes back to the initial I114F HbI(CO)₂ state via the bimolecular CO recombination with a bimolecular rate constant of 47 mM⁻¹s⁻¹, which is two times slower than that of WT (95 mM⁻¹s⁻¹).

The unveiled kinetics shows that there are two significantly decelerated transitions in I114F. Because the occluded ligand migration pathway in I114F^{30, 32} is expected to decelerate the ligand migration between the primary and the secondary docking sites and TRXSS measures global structural changes rather than the ligand migration process itself, we can infer that the decelerated structural transitions manifested in the TRXSS data of I114F would involve CO migration between the primary and the secondary docking sites. The first decelerated process is the I₁-to-I₂ transition. Since the I₁ intermediate is formed directly after photodissociation within 100 ps, CO should still likely be in the primary docking site. Thus, the delayed I_1 -to- I_2 transition by the mutation means that the structural transition from I_1 to I₂ involves the migration of CO from the primary to the secondary docking sites. The significant deceleration of the I1-to-I2 transition observed in I114F was not observed in TRXSS studies on other mutants such as F97Y35 and T72V37 where their ligand migration pathways are not expected to be affected by the mutations, supporting that the I_1 -to- I_2 transition is related to ligand migration from the primary to the secondary docking sites. The second decelerated process is the bimolecular CO rebinding of I₃. As Phe114 makes the access of CO to the active site difficult, the bimolecular CO recombination can be slowed as well, which is consistent with the results of flash photolysis and oxygen equilibrium binding studies.^{30, 32–33} On the other hand, other major processes, the biphasic structural transitions from I₂ to I₃, in I114F are not delayed and have rates similar to those of WT. This means that the I₂-to-I₃ transition is not likely to involve the movement of CO between the primary and the secondary docking sites. According to a previous structural analysis of WT, it is known that I_1^{WT} and I_2^{WT} have structures close to the R state of WT, whereas I_3^{WT} has a structure close to the T state of WT.³⁵ Therefore, we expect that the CO migrates from the inside of the protein to the bulk during the I₂-to-I₃ transition.

In addition, I114F has a different geminate recombination pathway compared to WT, which supports our interpretation of the CO migration during the transition between I₁ and I₂. In WT, the geminate recombination occurs from the I₂ intermediate. A part of I₂ transforms into a ligated form of I₁ (the two red octagons in Figure 3b), and then subsequently returns to the initial liganded state. In contrast, in I114F, the geminate recombination occurs from I₁ not I₂. According to a FTIR cryospectroscopic study which is sensitive to the CO position, the geminate recombination occurs only from the primary docking site in I114F, in contrast

in WT, it occurs from both of the primary and the secondary docking sites.³⁰ By comparing the previous FTIR study and our TRXSS results, we can infer the location of the ligand (CO molecule) in the I_1 and I_2 intermediates. The geminate recombination in I114F is likely to occur directly from rather the primary docking site than the secondary docking sites, since Phe114 residue disturbs the movement of the ligand from the secondary to the primary docking sites, thereby blocking the I_2 -to- I_1 transition. These differences in the geminate recombination kinetics between the I114F and WT suggest that the ligand in the I_1 and I_2 intermediates resides in the primary and the secondary docking sites, respectively.

The SADS curves shown in Figure 3d, extracted from the kinetic analysis, contain the structural information of each intermediate. The difference in the amplitude between the SADS curves of I114F and WT is attributed to the difference in the magnitude between the structural change occurring in I114F and WT (see the Supporting Information for details). As shown in the upper panel of Figure 3d, the SADS curves of I114F have smaller amplitudes than those of the WT for all intermediates, suggesting that the degree of structural change is smaller in I114F. Notably, in the case of I₃, which was identified as the intermediate arising from the R-T transition, the SADS curves of I114F and WT have nearly identical shapes, whereas the magnitude of the SADS curve of I114F is 2.8 times smaller than that of WT (the lower panel of Figure 3d). The identical shapes with different magnitudes suggest that the trend of structural changes between the liganded state and I₃ in WT is maintained in I114F similarly, while the amount of the structural change is less pronounced in I114F (see the Supporting Information for details). This observation is consistent with the result expected from the previous crystallographic study.³² Due to the restricted heme movement, the liganded state of I114F has closer structure to rather the T state of WT than the R state of WT.

We conducted a structure refinement analysis on the SADS curves to extract key structural parameters of each intermediate structure according to the protocol established in our previous study on WT.³⁵ The detailed procedures are given in the SI. The refined candidate structures are obtained from the analysis and their theoretical scattering patterns show excellent agreement with the experimental SADS curves (Figure S9). The three key structural parameters, the displacement of the residues, the heme-heme distance, and the subunit rotation angle, are extracted from the sets of the refined candidate structures (Figure 4). The first and the other two are relevant to the tertiary and quaternary structural changes, respectively.

The absolute displacements of the residues for the three intermediates, related to the tertiary structural changes, are shown in Figure 4a. This figure visualizes the degree of movement of the residues with respect to the reference structure. Here, we defined (i) the position of a residue as the distance between the C α atom and the iron atom of the heme in the same subunit, and (ii) the displacement of a residue as the difference in the positions between the candidate and reference structures. The reference structures of WT and I114F are their own static crystal structures of the liganded state (pdb codes WT: 3SDH,³⁴ and I114F: 1JWN³²). For all three intermediates, the area under the absolute displacement curve of I114F is smaller than that of WT, meaning that I114F undergoes less structural changes during the transition to each intermediate. In the case of I₃, the magnitudes of the displacements are

much smaller in I114F than in WT, whereas the directions of the displacements in I114F are similar to those in WT. Hence, the magnitude of the structural changes involved in the transition to I₃ is greatly reduced in I114F, whereas the direction of the structural changes remains the same in WT and I114F. This is fully consistent with the fact that the shape of the SADS curve of I_3^{I114F} is identical to that of I_3^{WT} but with a smaller magnitude by 2.8 times.

The occurrence distribution of the heme-heme distance and the subunit rotation angle of the intermediates for I114F and WT are also obtained and compared in Figures 4b and 4c, respectively. They are the key structural parameters about the quaternary structural transition as the cooperative ligand binding of HbI is modulated by the hydrogen-bonding network between two hemes³² and the subunit rotation.^{34, 38–41} In WT, the transitions from the liganded state (heme-heme distance = 18.4 Å, subunit rotation angle = 0°) to I_1^{WT} (hemeheme distance = 18.0 ± 0.2 Å, subunit rotation angle = $-0.1 \pm 0.5^{\circ}$) and to I_2^{WT} (17.9 ± 0.3) Å and $0.1 \pm 0.5^{\circ}$) involve very small changes in the heme-heme distance and subunit rotation angle. The major structural changes during the photoreaction of WT occur in I_3^{WT} (16.6 ± 0.2 Å and 3.4 \pm 0.6°), whose structure is close to the T state of WT (16.6 Å and 3.5°). In I114F, the heme-heme distances of both I_1^{I114F} (17.4 ± 0.3 Å) and I_2^{I114F} (17.4 ± 0.3 Å) are identical to that of the liganded state of I114F (17.4 Å) but much smaller than that of the R state of WT (18.4 Å). The heme-heme distance of I_3^{1114F} (17.0 ± 0.3 Å) is similar to that of the unliganded state of I114F (16.9 Å) and the T state of WT (16.6 Å). The subunit rotation angles of I_1^{114F} (2.9 ± 0.6°), I_2^{1114F} (3.1 ± 0.6°), and I_3^{1114F} (3.3 ± 0.6°) are also close to that of the T state of WT (3.5°). As can be seen in Figures 4b and 4c, the extent of quaternary structural changes such as the heme-heme distance change and intersubunit rotation are smaller in I114F, compared to WT. The smaller quaternary structural changes in I114F are attributed to the quaternary structure of the liganded state of I114F. Specifically, the starting structure of the R-T transition (the initial I114F HbI(CO)₂) in I114F is already shifted toward the T state of WT, whereas the end structure (I₃^{1114F}) is still similar to the T state of WT. Accordingly, the amount of structural changes involved in the R-T transition of I114F is smaller than that of WT (Figure S10). The restricted heme movement by Phe114 reduces the degree of quaternary structural change, which is consistent with conclusions drawn from the crystal structures.³²

CONCLUSIONS

In this work, we harnessed the power of TRXSS to investigate the detailed reaction kinetics and intermediate structures. The effects of the occluded ligand migration pathway on the kinetics and structural dynamics were elucidated by comparing the TRXSS data from WT and I114F, which is known to occlude ligand migration between the primary and the secondary docking sites. The kinetic analysis of the TRXSS data revealed that I114F has a kinetic framework with three structurally distinct intermediates (I₁, I₂, and I₃), as in WT. From a comparative study with WT, we found that two structural transitions (the I₁-to-I₂ transition and the bimolecular recombination of I₃) are significantly delayed in I114F. Therefore, we suggest that they involve ligand migration between the primary and the secondary docking sites. We also found that the geminate CO recombination occurs from the I₁ intermediate instead of the I₂ intermediate in I114F. A structural analysis using the SADS curves of the three intermediates revealed that the restriction of heme movement by Phe114

reduces the degree of allosteric structural change, which is consistent with results from the crystal structures. The delayed and different structural transitions caused by the obstructed ligand migration indicate that the mutation causes both the deceleration of ligand migration and the alteration of the conformation change during the protein dynamics.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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ABBREVIATIONS

HbI	homodimeric hemoglobin			
I114F	I114F mutant of HbI			
WT	wild type of HbI			
Mb	myoglobin			
Hb	hemoglobin			
TRXSS	time-resolved X-ray solution scattering			
FTIR	Fourier-transform infrared			
TRXL	time-resolved liquidography			
SVD	singular value decomposition			
SADS	species-associated difference scattering.			

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Figure 1.

Close-up views of the heme environment of crystal structures of HbI(CO)₂ for WT (left, pdb code: $3SDH^{34}$) and I114F (right, pdb code: $1JWN^{32}$). Heme and some residues forming secondary docking sites (Xe4 and Xe2 sites) are represented by the gray sticks, and hemebounded CO molecules are shown as spheres. The 36th, 73rd, and 114th residues, which separate the primary and the secondary docking sites, are depicted with their van der Waals spheres (gray for the 36th and 73rd residues and blue for the 114th residue). Phe97 is shown as the magenta stick, which faces the subunit interface in WT but faces the proximal site in I114F. The positions of the Xe4 and Xe2 sites are labeled based on the crystal structure of Xe-exposed WT HbI(CO)₂ (pdb code: $3G46^{31}$).

[— Exp	eriment				
$\Delta S(q, t)$	— The	ory	100 ps		\frown	3.16 µs
	Comment		178 ps			5.62 µs
	1 million	magniture and	316 ps			10 µs
		······	562 ps			17.8 µs
	- marine - m		1 ns			31.6 µs
	/	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1.78 ns			56.2 µs
	m		3.16 ns			100 µs
			5.62 ns			178 µs
			10 ns			316 µs
	Participant		17.8 ns	-		562 µs
			31.6 ns			1 ms
	Junio		56.2 ns	-		1.78 ms
	June		100 ns			3.16 ms
			178 ns			5.62 ms
			316 ns			10 ms
			562 ns			17.8 ms
	h		1 µs			31.6 ms
			1.78 µs			56.2 ms
ľ	0.3	0.6	0.9	0.3	0.6	0.9
		α (Å-1)			α (Å ⁻¹)	
		ч (°)			9.(*)	

Figure 2.

Time-resolved difference X-ray solution scattering curves, S(q, t), measured for a solution sample of I114F. Experimental curves (black curves) are compared with theoretical curves (red curves) generated by linear combinations of left singular vectors based on the optimized kinetics model.



Figure 3.

(a, b) Kinetic models for I114F and WT. The red (with "CO") and white symbols represent ligated and photolyzed subunits, respectively. To indicate the three structurally distinguishable intermediates with different tertiary structures, the subunit of each intermediate is presented with a different shape. For all three intermediates and HbI(CO)₂ of I114F, two subunits are described as rotating with respect to the other, reflecting the quaternary structural difference with respect to the R state of WT, that is, the liganded state of WT. For WT, only I_3 is represented in such a way, indicating that the quaternary transition involving subunit rotation occurs only in I₃. In the kinetic model for WT, two connected red octagons represent a ligated form of I₁, which is formed by the geminate recombination of CO with I_2 and structurally indistinguishable from the photolyzed forms (two connected white octagons and the white-red mixed ones) of I_1 . (c) Population changes of the three intermediates and initial HbI(CO)₂ as a function of time for I114F (red) and WT (black). The lines correspond to the populations obtained from the kinetics analysis. The open circles correspond to the optimized populations obtained by fitting the experimental curve at each time point with a linear combination of SADS curves for the three intermediates. (d) SADS curves of the three intermediates for I114F (red) and WT (black). The scaled curve of I3^{I114F} (blue) has a shape nearly identical to that of I_3^{WT} , but with a smaller magnitude by 2.8 times.

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Figure 4.

(a) Absolute displacement as a function of the residue for the candidate structures of three intermediates for I114F (red) and WT (black). For the calculation of the displacement, the reference structure for I114F is its liganded crystal structure, whereas for WT it is its liganded crystal structure. (b, c) The occurrence distribution of (b) the heme-heme distance and (c) the subunit rotation angle for the candidate structures of I114F (red) and WT (black) with respect to the R state of WT. The values from the crystal structures for the liganded state of WT and I114F are indicated by the blue and green vertical solid lines, respectively, and those for the unliganded state of WT and I114F are likewise indicated by the blue and green vertical dotted lines, respectively. The pdb codes for liganded WT, unliganded WT, liganded I114F, and unliganded I114F are 3SDH,³⁴ 4SDH,³⁴ 1JWN,³² and 1JZK,³² respectively.