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ALTERED STRUCTURE OF THE Mn₄Ca CLUSTER IN THE OXYGEN EVOLVING COMPLEX OF PHOTOSYSTEM II BY A HISTIDINE LIGAND MUTATION*


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Running head: Altered Mn₄Ca cluster in a PSII mutant

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The effect of replacing a histidine ligand on the properties of the oxygen-evolving complex (OEC) and the structure of the Mn₄Ca cluster in Photosystem II (PSII) is studied by X-ray absorption spectroscopy using PSII core complexes from the Synechocystis sp. PCC 6803 D1 polypeptide mutant H332E. In the X-ray crystallographic structures of PSII, D1-His332 has been assigned as a direct ligand of a Mn ion, and the mutation of this histidine ligand to glutamate has been reported to prevent the advancement of the OEC beyond the S2YZ’ intermediate state. The Mn K-edge absorption spectrum of D1-H332E shifts to a lower energy compared to that of the native wild type (WT) samples, suggesting that the electronic structure of the Mn cluster is affected by the presence of the additional negative charge on the OEC of the mutant. The EXAFS (Extended X-ray Absorption Spectrum) shows that the geometric structure of the cluster is altered substantially from that of the native WT state, resulting in an elongation of Mn-ligand and Mn-Mn interactions in the mutant. The Sr-H332E mutant, in which Ca is substituted by Sr, confirms that Sr(Ca) is a part of the altered cluster. The structural perturbations caused by the D1-H332E mutation are much larger than those produced by any biochemical treatment or mutation examined previously with X-ray absorption spectroscopy. The substantial structural changes provide an explanation not only for the altered properties of the D1-H332E mutant, but also the importance of the histidine ligand for proper assembly of the Mn₄Ca cluster.

The oxygen-evolving complex (OEC) located in the Photosystem II (PSII) membrane-bound protein in plant, algae, and cyanobacteria catalyzes the water-oxidation reaction (1-6). The OEC couples the four-electron chemistry of water oxidation with the one-electron photochemistry of the reaction center by sequentially storing oxidizing equivalents through five intermediate S-states (Sᵢ, i = 0 to 4), before one molecule of dioxygen is evolved. The heart of the OEC consists of four Mn atoms and one Ca atom; the Mn₄Ca cluster provides a high degree of redox and chemical flexibility so that several oxidizing equivalents can be stored during the S-state cycle. Unlike inorganic catalysts, however, the uniqueness of catalytic centers in metalloenzymes arises from their protein environment. This environment provides (i) direct ligands to the cluster for maintaining the metal structure, while giving structural flexibility to the cluster at the same time, (ii) controls the cluster’s redox potential by participating in charge distribution during the catalytic reaction, and (iii) provides a hydrogen-bonding network. Thus, nature has taken advantage of the special properties of the metal ions and tuned them by protein encapsulation to perform a wide variety of specific functions associated with life processes.

The assignment of the protein ligands to the Mn₄Ca cluster has been attempted by a wide range of mutational studies. These studies have revealed the potential ligands to the Mn₄Ca cluster. Considerable structural information has also been obtained through extensive studies by X-ray crystallography and various spectroscopic techniques over the past several years (1-5,7). The X-ray crystallography (XRD) studies up to the current resolution of 2.9 Å (8-11) have located the electron density associated with the water-oxidizing Mn₄Ca cluster within the large complex of PS II peptides. Structural ambiguity remains,
however, because of the limited resolution and radiation damage specifically to the Mn₄Ca cluster (12,13). The ligand environment of the Mn₄Ca cluster in the 3.5 and 3.0 Å structural models also differ in a number of aspects, including the binding mode of the carboxylate ligands (i.e., bidentate versus monodentate) and the orientations of the backbone residues. These differences are likely due to the limited resolution of the diffraction data, different extents of radiation damage, and different interpretations of the electron density. Polarized X-ray spectroscopy with PSII single crystals also provides Mn₄Ca models based on the accurate distances and the polarization characteristics of metal-metal or metal-ligand vectors within the protein environment (14). However, the assignment of the ligand environment remains tentative in this case because the proposed OEC environment obtained by this method is a simple combination of the Mn₄Ca cluster model obtained from the polarized EXAFS and the ligand environment from the crystal structure (Fig. 1).

Despite the structural ambiguities mentioned above, all the experimental evidence so far points that His332 residue is an important direct ligand to the Mn₄Ca cluster. In the recent X-ray crystallographic structural models of PSII, the Mn₄Ca cluster is ligated by a single histidine residue, His332 of the D1 polypeptide (8-11). This assignment is supported by recent 31 – 34 GHz ESEEM studies of the Mn₄Ca cluster in its S₂ state (15,16). These studies show that the nitrogen couplings observed in earlier ESEEM studies conducted at lower frequencies (9 – 12 GHz) (17-20) originate from a single ¹⁴N nucleus. The higher frequency studies permit the quantitative determination of the hyperfine and nuclear quadrupolar couplings of the coupled nucleus; these are consistent with direct coordination of the ¹⁴N atom to a Mn ion in the S₂ state (15,16). On the basis of earlier 9 GHz ESEEM studies conducted with unlabeled and [¹⁵N]histidine, this ¹⁴N atom is known to originate from the nitrogen of a histidine imidazole group (19,21). Because the ¹⁴N coupling essentially disappears in PSII core complexes from the D1-H332E mutant of *Synechocystis* sp. PCC 6803 (20), the ¹⁴N nucleus has long been assigned to D1-His332, consistent with the X-ray structural models (e.g., (5)).

However, in a recent study (22) of a different mutant, D1-H332S, of *Synechococcus elongatus* the nitrogen coupling observed in the three-pulse 9 GHz ESEEM spectrum of wild-type PSII core complexes was largely unchanged by the D1-H332S mutation. It was proposed that the nitrogen coupling observed in all the ESEEM studies originates from D1-His337 rather than D1-His332 and that the loss of the nitrogen coupling in the D1-H332E mutant (20) was caused by secondary structural perturbations introduced into PSII by the mutation. These same structural perturbations were proposed (22) to give rise to the altered S₂ state multiline EPR signal that is observed in the D1-H332E mutant (20,23) and the inability of D1-H332E cells (24) and PSII core complexes (23) to advance beyond the S₂Y' state.

To study whether the failure of the catalytic cycle of D1-H332E is due to the structural changes of the OEC itself, or to the secondary effect of the mutation, we applied X-ray spectroscopy in the current study. Manganese X-ray absorption spectroscopy is exquisitely sensitive to changes in Mn-Mn and Mn-Ca vectors in the Mn₄Ca cluster of PSII and to changes in the charge density on the Mn ions. Moreover, XAS can detect the signals from Mn or Ca/Sr even when there are no other spectroscopic signatures from the cluster, as the X-ray spectroscopic signals are never silent. This property is useful in studying whether the cluster assembles in the mutant species. For example, it is possible that the cluster assembles in a modified geometry which makes it inactive and or incapable of exhibiting other spectroscopic signals. Determining the geometric and electronic structure of such altered clusters in mutants is particularly important because the changes may give us a rationale for identifying the properties of the native cluster that make it unique for catalyzing the difficult water oxidation reaction. XAS is also useful for studying the changes in the geometric and electronic structures as the altered cluster advances through the S-state cycle, and in determining the state at which the cycle is impaired. These structure-function correlations in the altered structures that may assemble in some mutants can be critical for understanding the native, active catalytic cluster. Accordingly, to characterize structural perturbations that may be caused by the D1-H332E mutation and to further investigate the structural role of D1-His332 in the
Mn$_2$Ca cluster of PSII, we conducted a Mn X-ray absorption study of the D1-H332E mutant of *Synechocystis* sp. PCC 6803.

**Experimental Procedures**

**Construction of Mutant and Propagation of Cultures.** The construction of the D1-H332E mutation was described previously (20). Briefly, the mutation was introduced into the *psbA*-2 gene of *Synechocystis* sp. PCC 6803 and transformed into a host strain of *Synechocystis* that lacks all three *psbA* genes and that contains a hexahistidine-tag (His-tag) fused to the C-terminus of CP47. Solid media contained 5 mM glucose, 10 μM DCMU, and 5 _g/mL_ kanamycin monosulphate. The DCMU and antibiotic were omitted from the liquid cultures. Large-scale liquid cultures (each consisting of three 7 L cultures held in glass carboys) were propagated as described previously (25). For the isolation of Sr-containing PSII core complexes, the CaCl$_2$ in the liquid media was replaced with an equivalent concentration of SrCl$_2$ (26,27). To verify the integrity of the mutant cultures that were harvested for the purification of PSII core complexes, an aliquot of each culture was set aside and the sequence of the relevant portion of the *psbA*-2 gene was obtained after PCR amplification of genomic DNA (28).

**Purification of PSII core complexes.** PSII core complexes of the wild type (WT) and D1-H332E mutant of *Synechocystis* sp. PCC 6803 retaining the Mn cluster were purified under dim green light at 4 °C with Ni-NTA superflow affinity resin (Qiagen, Valencia, CA) as described previously (27). The purification buffer consisted of 1.2 M betaine, 10% (v/v) glycerol, 50 mM MES-NaOH (pH 6.0), 20 mM CaCl$_2$, 5 mM MgCl$_2$, and 0.03% (w/v) _n_-dodecyl _D_-maltoside by concentrating them to ~ 9 mg of Chl/mL, diluting them 20-fold with a buffer containing 42% (v/v) glycerol, then again concentrating them to ~ 9 mg of Chl/mL. The resulting concentrated PSII core complexes were transferred to epoxy sample holders (40 μL each) designed to fit into both EPR and X-ray liquid He cryostats. After dark-adaptation for one hour at room temperature, the samples were predominantly in the S$_1$ state. These samples were stored in liquid nitrogen until used. Half of the S$_1$ sample holders were taken out later and illuminated with a 400 W tungsten lamp at 200 K for 5 minutes in order to generate the S$_2$ state for the wild type S$_2$ state and at 273 K for 2 minutes for the H332E mutant sample. The illuminated samples were then stored in liquid nitrogen.

**EPR measurements.** Samples were characterized by EPR spectroscopy at 8 K in order to ensure the quality of the samples before and after the illumination. Spectra were collected using a Varian E-109 EPR spectrometer equipped with an Air Products Helitrak liquid helium cryostat. The generation of the S$_2$ state after the illumination was checked by the intensity of EPR multiline signal characteristic to the S$_2$ state. All the samples were also checked with EPR prior to XAS measurement to see whether free Mn(II) was produced during any biochemical and sample handling processes. No detectable Mn(II) signal was observed for any of the mutant and wild type samples.

**XAS measurements.** X-ray absorption spectroscopy (XAS) was performed at the Stanford Synchrotron Radiation Laboratory (SSRL) on beamline 9-3 at an electron energy of 3.0 GeV with an average current of 85-100 mA. The radiation was monochromatized by a Si(220) double-crystal monochromator. The intensity of the incident X-ray beam was monitored by a N$_2$-filled ion chamber (I$_0$) in front of the sample. The monochromator energy was calibrated using a pre-edge peak of KMnO$_4$ (6543.3 eV) for Mn XAS, and using an edge peak of strontium acetate (16120 eV) for Sr XAS. These standards were placed between two N$_2$-filled ionization chambers (I$_1$ and I$_2$) after the sample. The total number of X-ray photons used for the measurements was ~ 4 x 10$^{13}$ photons mm$^{-2}$ for the Sr XAS (16-17 KeV) and ~ 5 x 10$^{12}$ photons mm$^{-2}$ for the Mn XAS (6-7
keV), and are well below the X-ray damage threshold as previously established (12). The Mn K-edge was also closely monitored for any reduction of Mn as seen by a shift in the K-edge inflection point energy. The monochromator was detuned at 6600 eV (16200 eV for Sr) to 50% (or greater) of maximal flux to attenuate the X-ray 2nd harmonic. Samples were kept at 10 K in a liquid helium flow cryostat to minimize radiation damage. Data reduction of the XAFS spectra was performed using EXAFSPAK (29). Pre-edge and post-edge background were subtracted from the XAS spectra and the results were normalized with respect to edge height. After correction of background-corrected spectra from energy space to photoelectron wave vector (k) space (E0 = 6561.3 eV for Mn and 16120 eV for Sr), and weighted by k3, a four-domain spline function was subtracted. Curve fitting was performed using ab initio calculated phases and amplitudes from the program FEFF 8.2. (30,31). The details of the curve fitting analysis is described in the supplementary information.

Results

Mn XANES of D1-H332E. The XAS spectra of *Synechocystis* sp. PCC 6803 wild type (WT) and D1-H332E PSII core complexes in the dark state (S1) and illuminated (S2) state are compared in Figure 2. The WT S1 and S2 XANES spectra are similar to those observed in spinach PSII membrane preparations (32), in which the formal oxidation states have been assigned as Mn(III)2Mn(IV)2 for the S1 state and Mn(III)Mn(IV)3 for the S2 state. During the S1 to S2 transition, the rising edge energy shifts ~0.7 eV to higher energy in the WT (Fig. 2a). In its dark state (S1), the D1-H332E mutant has a lower rising edge energy and a slightly different shape as seen in the second derivatives than the WT S1 state (Fig. 2b). This is not due to the release of free Mn(II) in the mutant samples as no Mn(II) signal was observed in the EPR spectra taken prior to the XAS measurement. One possible explanation for this low energy shift in D1-H332E is that the formal oxidation state of the D1-H332E dark state is not Mn(III)2(IV)2, but lower such as (III)2(IV) or (II)(III)(IV)2. The energy shift between the WT and the D1-H332E is 1.69 eV. This is much larger than the energy shift between the S1 to S2 transition of the WT, but within the range of one-oxidation state differences (for example, shifts of ~2.1 eV for the S0 to S1 transition and ~1.1 eV for the S1 to S2 transition have been observed in spinach PSII membrane preparations (32)). This variation in the magnitude of the energy shift corresponding to a single oxidation event demonstrates the difficulties of making formal oxidation state assignments on the basis of the XANES edge shift: the edge shape is also highly sensitive to structural changes.

In H332E PSII core complexes, only ~60% of the PSII centers contain Mn clusters, although all of these advance to the S2 state in response to continuous illumination at 273 K (23). Upon illumination at 273K, the H332E XANES spectrum shifts ~0.6 eV towards higher energy (Fig. 2(c)), indicating that Mn is oxidized during this transition in the mutant similar to the WT. The dark state of D1-H332E is EPR silent, and upon illumination, exhibits an altered multilane EPR spectrum that superficially resembles that in Ca-depleted, Sr-substituted, and NH4-inhibited WT PSII preparations (20,23). It is therefore likely that the formal oxidation state of the D1-H332E mutant in its dark state is the same as that of the WT, i.e. Mn(III)2Mn(IV)2.

Mn EXAFS of D1-H332E. Figure 3 shows the Fourier transforms of k3-weighted Mn EXAFS spectra of *Synechocystis* sp. PCC 6803 WT and D1-H332E PSII core complexes in the dark (S1) and illuminated (S2) states (k-space data shown in Figure S1 in Suppl. Info.). The WT spectra (Fig. 3a) are almost identical to those of the S1 and S2 spectra of spinach PSII membrane preparations, suggesting that the structure of the Mn6Ca cluster in the S1 and S2 states is the same in *Synechocystis* as in spinach. The D1-H332E mutant dark (S1) state spectrum (Fig. 3b) differs significantly from that of the WT S1 state spectrum (Fig. 3a). The first FT peak (labeled I in Figs. 3a and 3b), which arises primarily from Mn-O interactions with a possible small contribution from one or two Mn-N interaction(s) in the WT, becomes more intense and shifted towards longer distance in the mutant. In addition, the second FT peak (labeled II in Figs. 3a and 3b), which largely arises from the di-µ-oxo bridged Mn-Mn interactions in the WT, decreases and merges with the third peak (labeled III in Fig. 3a and 3b, and assigned predominantly to oxo-
bridged Mn-Mn and Mn-Ca distances at ~3.3 and ~3.4 Å in the WT) in the mutant. These changes indicate that the D1-H332E mutation substantially alters the structure of the Mn$_4$Ca cluster. The spectra obtained with three different preparations of D1-H332E PSII core complexes were identical. This reproducibility provides proof that the structure of the OEC cluster in the mutant is homogeneous, and that it does not change as a function of the preparation. Additionally, we estimated that the Mn stoichiometry in the D1-H332E mutant is the same as the native PSII, i.e., 4 Mn ions per PSII, based on previous EPR studies (20, 23) and the analysis of Mn X-ray fluorescence data (see Suppl. Info.).

Upon illumination, the first FT peak intensity decreases in the mutant (Fig. 3b, gray trace). Because the first FT peak of the WT is essentially unchanged upon illumination (Fig. 3a, gray trace), the substantially decreased amplitude of the first FT peak in the mutant indicates that substantial structural changes (e.g., ligand reorganization) take place during the S$_1$ to S$_2$ transition in the mutant, whereas little or no structural change takes place during this transition in WT (33).

**Sr XAS on Sr-substituted H332E.** It has been suggested on the basis of biochemical studies that mutations of D1-His332 diminish the affinity of Ca for the Mn$_4$Ca cluster (34). Because the X-ray crystallographic structural models place the Ca ion and His332 ligand on opposite sides of the Mn$_4$Ca cluster, the apparently diminished affinity of Ca has been hypothesized to be a consequence of mutation-induced perturbations to the Mn$_4$Ca cluster. In order to investigate whether the Ca ion is present in the Mn cluster of D1-H332E PSII core complexes, samples were prepared having Sr substituted for Ca and measurements were carried out using Sr XAS (instead of Ca XAS). Many studies have shown that the Ca ion can be replaced with Sr while retaining O$_2$-evolving activity in PSII (26,27,35). Additionally, Sr XAS (absorption energy: ~16120 eV) has several experimental advantages over Ca XAS (absorption energy: ~4500 eV), such as (i) the X-ray radiation damage is less for the same number of photons incident on the sample, (ii) the attenuation of the incident and scattered X-rays is far less at the Sr X-ray absorption energy, and (iii) the fluorescence yield of Sr is higher than that of Ca (see references (36,37)).

Figure 4 shows the Sr XANES (a) and EXAFS (b) spectra of Sr-substituted H332E PSII core complexes. For a comparison, the published data of the Sr-WT PSII core complexes from *Synechococcus elongatus* in the S$_1$ state is shown in the figure. The Sr EXAFS of the mutant (Fig. 4b, gray trace) shows a distinctive second FT peak (arising primarily from Mn-Sr interactions) that closely resembles the second FT peak in the WT. These data show that the Mn-Sr interaction exists in the H332E mutant and that the Mn-Sr interaction(s) is(are) at a similar distance and amplitude to the WT. On the other hand, the first FT peak in the mutant (corresponding to Sr-O interactions) is much stronger than in the WT. We also collected Mn XAS on this sample, in order to see if the substitution of Sr for Ca caused any structural perturbations to the Mn$_4$ cluster (Figure S2 in Suppl. Info.). Unlike the situation in *Synechococcus elongatus*, where biosynthetically incorporated Sr does not exchange with exogenous Ca ions (26), Sr/Ca exchange occurs when *Synechocystis* PSII core complexes containing biosynthetically incorporated Sr are purified in buffers containing Ca (27). Because Sr XAS requires the elimination of exogenous Sr ions, our PSII core complexes containing biosynthetically-incorporated Sr were purified in Ca-containing buffers. Therefore, only about 75% of the Sr-H332E PSII core complexes contained Sr, while the remainder contained Ca (27). This heterogeneity is not an issue for carrying out Sr XAS measurements, because only the Sr-H332E fraction is probed. For Mn XAS on this sample, on the other hand, both Sr-H332E and Ca-H332E fractions contribute to the spectrum, and we cannot distinguish their individual contributions (Fig. S2 in Suppl. Info.). In the Mn EXAFS shown in Fig. S2, the spectra of the Ca-H332E and Sr(Ca)-H332E PSII samples are almost identical up to the FT peak II region, but the FT peak III region differs significantly. This difference might be explained as a consequence of a larger contribution of the Sr H332E fraction which results in a distance elongation (typically, actual distance of 3.4 Å for Mn-Ca and 3.5 Å for Mn-Sr) and increase in the intensity (due to heavier atom contribution) around the peak III region. Although further detailed structural information cannot be extracted from the Mn XAS on the Sr-H332E sample, the geometry of the Mn$_4$ cluster in the Sr-
containing H332E sample seems to be very similar to that in the Ca-containing H332E sample.

\textbf{Mn EXAFS Curve fitting of the WT S\textsubscript{i} state spectrum.} Mn EXAFS curve fits were carried out for the k\(^3\)-weighted EXAFS spectra of the WT S\textsubscript{i} state and the D1-H332E mutant dark state (S\textsubscript{i}). Fitting results for the WT S\textsubscript{i} state for the Mn-nearest neighbor interactions are given in Table 1, in which R, N, and \(\sigma^2\) show actual distance (Å) (note the Fourier transforms only show apparent distances), coordination number, and EXAFS Debye-Waller factor (Å\(^2\)), respectively. N values are defined as the total number of absorber-backscatterer vectors divided by the number of absorber atoms per OEC. R factor \((R_\text{fit}, \%)\) shows the goodness of the fit.

The predominant contribution to the FT peak I is from Mn-oxygens at \(R\approx 1.87\) Å. Although one or two Mn-nitrogen interaction(s) from the histidine residue(s) contributes to this region, it cannot be distinguished in the data because of the minor contribution (one out of \(\sim 24\) Mn-ligand interactions at similar distances). Therefore, Mn-O and Mn-N interactions were treated together in the current curve fitting.

The second and third peak region (Fig. 3a) are due mainly to the di-\(\mu\)-oxo bridged Mn-Mn \((\sim 2.7\) Å), and mono-\(\mu\)-oxo-bridged Mn-Mn \((\sim 3.3\) Å) interactions. The presence of Mn-Ca interactions \((3.4\) Å) in addition to the predominant mono-\(\mu\)-oxo bridged Mn-Mn interaction in the third peak have previously been demonstrated on the basis of Ca XAS experiments conducted with spinach PSII preparations (Sr XAS experiments on Sr-substituted PSII show four Mn-Sr interactions between \(\sim 3.5\) and 4.0 Å, see below) (37). Therefore, Mn-Ca interactions are included in the fit. In Table 1, the most preferable structural motif that supports the vector components of the recent EXAFS studies (the ratio of di-\(\mu\)-oxo bridged Mn-Mn : mono-\(\mu\)-oxo bridged Mn-Mn : Mn-O-Ca bridged interactions = 3:1:x, x=2 or 3, Figure S3, A-D in Suppl. Info.) was used for fitting the WT S\textsubscript{i} state spectrum. The contribution of three Mn-Mn interactions and their distance heterogeneities \((2.7\) to \(2.8\) Å) becomes clear only when the polarized spectra or the range-extended EXAFS data are used for the fit (38,39). Therefore, this heterogeneity was not considered in this study. Including Mn•••C interactions, which arise from the second shell carbons at \(\sim 3.2\) Å from Mn, slightly improves the fitting quality. Although the coordination number, \(N=2.75\) for the Mn•••C interactions was used through the fits which is based on the ligand model reported in the 2.9 Å crystal structure (11), this number remains highly ambiguous due to the uncertainly in the ligand binding modes (for example, whether the binding modes of the carboxylate ligands are bidentate or monodentate). Nevertheless, the Debye-Waller factor \((\sigma^2)\) of this absorber-backscatterer path is large \((\sim 0.018\) Å\(^2\)), indicating that the distance distribution of the Mn•••C interactions is high, and its contribution to the EXAFS spectrum has only a minor effect. In addition, the inclusion of this path does not noticeably change the fitting parameters for the other paths.

We have also considered two shell contributions to the first FT peak region, one part from the Mn-bridging oxygen and the other from the Mn-terminal oxygen. We assumed that the Mn-bridging oxygens have shorter distances \((\sim 1.8\) Å) than those of the Mn-terminal oxygens \((1.9\) Å) on the basis of model compound studies (40,41). No obvious improvement was observed in the two shell fit (not shown). The Fit#2 and 4 (Table 1) are shown in Figure 5a.

\textbf{Mn EXAFS Curve fitting of the D1-H332E spectra.} Table 2 summarizes the curve fitting results of the D1-H332E mutant in its dark-stable state (S\textsubscript{i}). In Fig. 3b, the increased intensity and the longer distance shift of the first FT peak of the D1-H332E sample compared to that of the WT indicates that the Mn-ligand distances are more uniform in the mutant. The one shell fit of the first peak gives a Mn-ligand distance of 1.89 Å in the mutant while it is \(\sim 1.86\) Å in the WT, suggesting that an elongation of the Mn-ligand interactions occurs in the mutant (see also Table S1 in Suppl. Info.).

The averaged Mn-Mn distance (peak II) is also longer \((\sim 2.77\) Å) in the mutant compared to the WT \((\sim 2.72\) Å). On the other hand, the intensity of the peak II region is decreased significantly in the mutant. This could be either due to the increased distance heterogeneity in the \(\sim 2.7\) Å region, or due to the reduced number of di-\(\mu\)-oxo bridged Mn-Mn interactions in the mutant. In order to test these two hypothesis, N of the Mn-Mn \(\sim 2.7\) Å interaction was fixed either to \(N=1\) (two Mn-Mn interactions,) or to \(N=1.5\) (three Mn-Mn interactions) in the fits.
interactions) in the fit (3:1:x and 2:1:x, or 3:2:x and 2:2:x). The result between two or three Mn-Mn 2.7 Å interactions was not conclusive as the fitting quality is similar within the error of the method (for example, Fit#1 compared to Fit#7 in Table 2). The result, however, shows that the averaged di-µ-oxo-bridged Mn-Mn distance is about 0.05 Å elongated in the mutant. If there are three di-µ-oxo bridged Mn-Mn interactions, the distance heterogeneity is much higher than that of the WT based on the decreased intensity of peak II.

The presence of the Mn-Ca(Sr) interactions was confirmed by the Sr XAS data obtained with the Sr-substituted H332E PSII core complexes, described below. Therefore, the Mn-Ca interaction was included in all the fits. The mono-µ-oxo bridged Mn-Mn interaction (~3.2 Å) seems to be necessary for maintaining a reasonable fit quality, and elimination of this interaction made fit quality worse (not shown). This implies that the unclear peak III region in the mutant spectrum (Fig. 3b) is a consequence of the elongation of peak II components.

Fits corresponding to an increase in the Mn-Mn 3.3 Å interactions to two as compared to one in the WT, namely 3:2:x and 3:1:x, or 2:2:x and 2:1:x, were also considered without much improvement in the quality of the fit. The addition of the second shell Mn•••C(ligand) ~ 3.1 Å interactions slightly improved the R factor. However, the contribution of this absorber-backscatter path has only a minor impact on the modeling of the Mn-Ca cluster due to its high Debye-Waller factor and it is not included in the Table. The representative fits (Fit# 2 and 8) are shown in Fig. 5b.

**Sr EXAFS Curve fitting of the D1-H332E spectra.** Fitting results for the Sr XAS on Sr-H332E are given in Table 3. Previously, Ca EXAFS of native PS II and Sr EXAFS of Sr-reactivated spinach PSII membranes showed the proximity of Ca to Mn at 3.4 Å, and Sr to Mn at 3.5 Å in the dark-stable S1 state (36,42,43). A more recent study on Sr-substituted PSII core complexes from *Synechococcus elongatus* shows Sr to be proximate to all four of the Mn in the Mn4 cluster in all S-states (37). The study shows that there are two to three Sr-Mn interactions in which Sr and Mn are likely to be bridged by µ-oxo groups around 3.5 Å, while there are two or one longer interactions around 3.9 Å. The fitting results of the published Sr-substituted PSII core complexes from *Synechococcus elongatus* is shown in Table S2 in the Suppl. Info.

FT peak I in Fig. 4(b) is best simulated by one shell of 7 to 8 oxygen atoms at R= 2.56 Å. For convenience, the N number of the Sr-O 2.5 Å interaction was fixed to 8 in Table 3. The Debye-Waller factor of this peak is much smaller (0.005) in the mutant than in the WT (Table S2). Assuming that the coordination number of Sr is the same in the WT and mutant samples, the result suggests that the Sr-O distance is more uniform in the mutant compared to the WT. On the other hand, FT peak II, which is best fit with two to three shorter Sr-Mn interactions and two to one longer Sr-Mn interaction (37) in the Sr-WT PSII core complexes, appeared with the same intensity in the two samples, suggesting that the Mn-Sr binding modes are similar between the WT and the mutant. Fit#1 and 2 are shown in Figure 6. The R² value of Fit#1 is slightly better than others and preferable, suggesting that there are three short (~3.5 Å) and one long (~3.9 Å) Sr-Mn interactions similar to that seen in the WT.

**Discussion**

**Structural changes of the OEC in the H332E mutant.** The D1-H332E mutant of *Synechocystis* sp. PCC 6803 has previously been characterized with chlorophyll fluorescence (24,34) and thermoluminescence (24) measurements in intact cells, and with optical absorption (23), EPR (20,23) and ESEEM (20) measurements in PSII core complexes. These data show that, in the mutant, the temperature threshold for forming the S2 state is approximately 100 K higher than in WT, that the quantum yield for forming the S2 state is very low, corresponding to a dramatic slowing of electron transfer from the Mn cluster to YZ⁺, that the S2/S1 midpoint potential is substantially decreased, that Mn oxidation beyond the S2YZ⁺ state is blocked, and that the S1 state multiline EPR signal is altered, exhibiting more hyperfine lines and narrower splittings than in WT. In addition, D1-H332E PSII core complexes exhibit no S1 state multiline EPR signal (23). These data were interpreted as showing that although the Mn cluster is assembled in the altered
ligand environment, the mutation perturbs the structure of the Mn$_4$Ca cluster and the network of hydrogen bonds that facilitates the $S_2$ to $S_1$ transition (20,23,24). Compared to the energy scale of EPR or IR spectroscopy, the energy scale of XAS is much larger (1 eV shift in the edge corresponds to 8066 cm$^{-1}$). Thus changes in XANES would indicate much larger changes in the electronic/geometric structure that would also be reflected in the substantial changes in the EXAFS. In the current study, the XAS data show that the D1-H332E mutation substantially alters the structure of the Mn$_4$Ca cluster in both the $S_1$ and $S_2$ states, resulting in an elongation of Mn-ligand and di-$\mu$-oxo bridged Mn-Mn interactions in the mutant. These structural alterations are far greater than those caused by any mutation (unpublished data) or biochemical treatment examined previously by X-ray absorption spectroscopy, including Ca-depletion (44), Sr/Ca exchange (45), and NH$_3$-inhibition (46).

The lower rising edge energy, and the changes in the shape in the XANES spectrum of the D1-H332E PSII core complexes in its dark-stable ($S_1$) state could originate from a change in Mn oxidation state or from a change in charge density on the Mn cluster caused by a change in cluster structure, ligand symmetry, or ligand exchange. Because dark-adapted D1-H332E PSII core complexes (EPR silent) produces a multiline EPR signal upon illumination, which is evidence for a paramagnetically coupled 4Mn cluster, we conclude that the formal oxidation state of dark-adapted D1-H332E PSII centers is the same as that of the WT, i.e. Mn(III)$_2$Mn(IV)$_2$. Therefore, the XANES shift to lower energy compared to WT probably results from a combination of (1) increased negative charge density on Mn caused by, for example, replacing a neutral histidyl nitrogen with a negative carboxylate oxygen and (2) accompanying structural changes of the Mn$_4$Ca cluster.

The presence of a Ca (Sr) ion in the Mn cluster of D1-H332E is confirmed by the similar intensity of the Mn-Sr EXAFS interactions observed in the Sr EXAFS of the mutant.

The Mn EXAFS curve fitting result suggests that the Mn$_4$Ca cluster in D1-H332E contains two to three di-$\mu$-oxo bridged Mn-Mn units, one to two mono-$\mu$-oxo bridged Mn-Mn unit, and two to three ~ 3.5 Å Mn-Ca interactions. In the one shell fit, the averaged distance of di-$\mu$-oxo-bridged Mn-Mn interaction in the mutant is changed from ~2.72 Å (WT) to ~2.77 Å (Tables 1 and 2). A possible reason for this elongation is a protonation of the oxo-bridge (Figure 7a), presumably to restore the charge balance because of the replacement of a neutral histidine ligand by a negatively charged glutamate residue. Another is a ligand structural change, where the replacement of the histidine residue by a glutamate triggers a previously bridged bidendate carboxylate changing its coordination mode from bidentate to monodentate (Figure 7b). In the case of protonation of an oxo-bridge, we expect about a 0.1 Å elongation of the Mn-Mn distance (47). In the case of a carboxylate changing its coordination mode, removal of a bidentate-type bridge could elongate the di-$\mu$-oxo bridged Mn-Mn distance also by 0.1 Å, such as is observed between the di-$\mu$-oxo/$\mu$-carboxylato and the di-$\mu$-oxo species (48). The proposed scenarios for the distance elongation suggested above could also occur by replacing His332 by another proximal carboxylate residue or water molecule.

Regarding the structural motif of the altered Mn$_4$Ca cluster in the mutant, the following two possibilities are considered. If there are three di-$\mu$-oxo-bridged Mn-Mn interactions, as in models having (3:1:x) or (3:2:x) configurations (shown in Fig. S3, Suppl. Info.), the distance heterogeneity would be greater in D1-H332E than in WT in addition to the Mn-Mn distance being elongated. In recent polarized XAS and range extended XAS studies, the presence of three di-$\mu$-oxo-bridged Mn-Mn interactions having 2.7Å:2.8Å distances in a 2:1 ratio has been suggested (39,49). In the D1-H332E mutant, we would expect an elongation (to >2.8 Å) of the longer component and/or an increased fraction of the longer component: in either case, peak II will be shifted to a longer distance and the intensity will be decreased. If there are two di-$\mu$-oxo-bridged Mn-Mn interactions as in the model having a (2:1:x) or (2:2:x) configurations in Fig. S3, Suppl. Info., one or both of the distance(s) would be elongated in the mutant.

Accompanied by the elongation and weak intensity of the ~2.7 Å Mn-Mn interaction (peak II), there is a strong Mn-ligand FT peak I in the D1-H332E dark state. This suggests a more
symmetric Mn-O ligand environment, which could happen if an oxo-bridge is broken, ligands are ligated as monodentate instead of bidentate, or water is ligated instead of carboxylate in the altered OEC (Figure 7b).

In the Sr XAS experiments, a stronger Sr-O FT peak was observed in D1-H332E than in the WT, just as a stronger Mn-ligand peak was observed in the Mn XAS of the mutant. This is likely due to a more uniform Sr-O environment and elongation of the Sr-O distances in the mutant. Although typical Sr-O(carboxylate) and Sr-O(water) distances are both within the range of 2.5 – 2.7 Å and therefore cannot be discriminated on the basis of the distance, a more symmetric Sr-O environment may occur, for example, when a carboxylate residue ligated to Sr is replaced with a water molecule. One possibility is that the mutation-induced alteration of the Mn₄ cluster structure causes the rearrangement of the ligands around the Sr(Ca), and that a water molecule ligates to the Mn₄ cluster in place of a carboxylate group. On the other hand, the Sr-Mn interactions are similar in the mutant and WT. On the basis of these observations, we consider the following two cases for the Mn-Sr(Ca) interactions: (i) the Mn-Sr(Ca) binding modes are the same in the WT and the mutant, but the hydrogen bonding network is changed; for example, more water molecules ligate to Sr(Ca) instead of carboxylate oxygen atoms, or (ii) the Mn-Sr(Ca) binding mode is altered, but the number of the Mn-Sr(Ca) interactions remains the same.

**D1-His332 is a Ligand of the Mn₄Ca cluster.**

As noted above, the structural perturbations caused by the D1-H332E mutation are substantially larger than those produced by any biochemical treatment or mutation examined previously with XAS, including the extraction of Ca from the Mn₄Ca cluster. Whereas Ca-depletion, Sr/Ca exchange, and NH₃-inhibition all alter the S₁ state multiline EPR signal in the same manner as the D1-H332E mutation, Mn-EXAFS studies show that these treatments alter the structure of the Mn₄Ca cluster either negligibly (in the case of Sr substitution) or to much lesser extents than the D1-H332E mutation (in the case of Ca extraction). Only in the case of NH₃-inhibition, where NH₃ is proposed to enter the coordination sphere of the Mn₄Ca cluster as a amido bridge between two Mn ions (50), was elongation of a Mn-Mn distance observed (from 2.72 Å to 2.87 Å) (46).

One significant difference of the mutation compared to the above treatments is that the Mn₄Ca cluster must necessarily be assembled in a different ligand environment in the mutant. In the WT, the D1 and CP43 protein subunits of PSII provide ligands to the OEC: the current view of the assembly process is that the first two Mn²⁺ ions are bound and photooxidized into Mn³⁺ during their assembly into the apoprotein and then the binding of the subsequent Mn²⁺ ions occurs (51,52). Whether the ligands that are required for the assembly are the same as the proposed direct ligands to the Mn₄Ca cluster remains a question. Nevertheless, the fact that the Mn₄Ca cluster can assemble in the His332Glu mutant but in an altered structure, suggests that a different ligand environment has modified the Mn₄Ca cluster. On the basis of the earlier Mn-EXAFS studies of Cd-depleted, Sr/Ca-exchanged, and NH₃-inhibited PSII preparations, it is difficult to imagine how the D1-H332E mutation could cause the substantial structural perturbations to the Mn₄Ca cluster observed in our Mn-EXAFS data if D1-His332 is not directly coordinated to a Mn ion. Therefore, we conclude that our data are in support of the X-ray crystallographic models depicting D1-His332 as a ligand to a Mn ion. Upon mutation of D1-His332 to glutamate, the missing imidazole ligand must be replaced by another ligand/residue, such as (i) a glutamate carboxylate, (ii) another carboxylate residue, (iii) a water molecule, or (iv) another histidine ligand. Another possibility is that, (v) no new residue is ligated to the Mn and the coordination number therefore changes from six to five. Alternatively a combination of those possibilities could occur if the reorganization of several ligands is triggered by the D1-H332E mutation. The Mn XANES spectrum of D1-H332E helps to distinguish between these possibilities: as discussed above, its shift to lower energy in the S₁ dark state, indicative of an increased negative charge density on the Mn₄Ca cluster compared to WT, implies that the imidazole group of D1-His332 is likely replaced by D1-Glu332 or by another nearby carboxylate group.

**Comparison of D1-H332E with Other Mutants.**

As described in the Introduction, there are two ESEEM studies in the literature on
D1-H332 mutants from *Synechocystis* sp. PCC 6803 (D1-H332E, same as in this study) (19,21) and *Thermostrehlococcus elongatus* (D1-H332S/Q) (22). Although both studies agree that D1-H332 is a ligand of Mn, there are differences in the assignment of the spin-echo signals attributed to Mn-histidine ligand N atoms; one providing support for D1-His332 and the other preferring D1-His337 which is not a direct ligand of Mn in the current crystal structures (9,10). It is suggested by the authors of ref. (22), that (i) all of the published ESEEM studies have detected D1-His337 instead of D1-His332, and (ii) the structural perturbations caused by the D1-H332E mutation eliminate the nitrogen couplings of D1-His337 (for a more detailed description see Suppl. Info.). Clarification of the source of the histidyl nitrogen coupling in PSII will require additional ESEEM studies of the S₂ state Mn₄Ca. The current XAS study on the D1-H332E mutant from *Synechocystis* sp. PCC 6803 study cannot directly resolve all the questions posed by the ESEEM studies. However, the XAS study points to the conclusion that D1-His332 directly coordinates to a Mn ion in the Mn₄Ca cluster and the altered EPR signal is a consequence of the altered OEC structure. The substantial changes seen in the structure of the Mn cluster in the mutant points to the consequences of the different ligand environment that influences the assembly, and the importance of His332 in *Synechocystis* sp. PCC 6803. The several differences of the D1-H332S and D1-H332Q mutants compared to D1-H332E, such as (1) D1-H332S and D1-H332Q evolve O₂, whereas D1-H332E does not, (2) D1-H332S and D1-H332Q exhibit a normal S₂ EPR multiline signal, whereas D1-H332E does not, and (3) D1-H332S and D1-H332Q shows small changes in the S₁ state, whereas D1-H332E cannot advance beyond the S₂Y₉ state, make the D1-H332S and D1-H332Q mutants attractive candidates for future XAS studies. Such XAS studies may also shed light on the differences between the mutants in their assembly/stability and coordination compensation/mutant rescue properties.

**Concluding Remarks**

On the basis of XANES and EXAFS data, we show that the D1-H332E mutation substantially alters the structure of the Mn₄Ca cluster in both the S₁ and S₂ states, resulting in an elongation of Mn-ligand and Mn-Mn interactions in the mutant. These structural perturbations are larger than those produced by any biochemical treatment or mutation examined previously with X-ray absorption spectroscopy. Because it is difficult to imagine how the D1-H332E mutation could cause these substantial structural perturbations if D1-His332 is not directly coordinated to a Mn ion, we conclude that D1-His332 ligates a Mn ion, in support of the current crystallographic structural models. In the D1-H332E mutant, Mn₄Ca cluster can be assembled but in an altered manner. This cluster advances one oxidation state (dark to illuminated state) by absorbing light, but the altered OEC structure prevents the complete catalytic cycle of the water splitting. This suggests that the first oxidation step (S₁ to S₂) takes place relatively easily, while the second step (S₂ to S₁) is critical and structurally more demanding.

**References**


Footnotes

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The abbreviations used are: PSII, photosystem II; OEC, oxygen-evolving complex; WT, wild type; XAS, X-ray absorption spectroscopy; XANES, X-ray absorption near edge structure, EXAFS; extended X-ray absorption fine structure, EPR; electron paramagnetic resonance, ESEEM; electron spin echo envelope Modulation.
Figure Legends

Fig. 1. Ligand environment of the OEC obtained from X-ray crystallography (a,b),(9,10) and putative ligand environment from the single crystal EXAFS study (c)(38). In (c), Model II and III from the ref.(38) are shown.

Fig. 2. Mn XANES and 2nd derivative spectra of (a) Synechocystis WT S1 (black) and S2 (gray) states, (b) a comparison of Synechocystis WT (black) and D1-H332E mutant (gray) S1 state spectra, and (c) D1-H332E mutant dark (gray) and illuminated (black) samples. The bottom part shows the 2nd derivative spectra of XANES, and zero-crossing energy gives the rising edge inflection point, which is indicated.

Fig. 3. Fourier transforms of the Mn EXAFS spectra of (a) Synechocystis WT S1 (black) and S2 (gray) states and (b) D1-H332E mutant dark (black) and illuminated (gray) states.

Fig. 4. Fourier transforms of the Sr XANES and EXAFS spectra of Synechocystis D1-H332E mutant dark state (gray). For comparison, Sr XANES and EXAFS spectra of Synechococcus elongatus WT S1 state (black) are shown (37).

Fig. 5. Mn EXAFS curve fitting result of (a) Synechocystis WT S1 state and (b) D1-H332E dark state. Black is the experimental data. Fit# 2 and 4 in Table 1 for the WT and Fit# 2 and 8 for the D1-H332E in Table 2 are shown in the figure.

Fig. 6. Sr EXAFS curve fitting result of Synechocystis D1-H332E S1 state. Black is the experimental data. Fit# 1 and 2 in Table 3 are shown in the figure.

Fig. 7. Possible structural reasons for the Mn•••Mn elongation in the mutant: (a) protonation of the bridging oxygen and (b) bidentate to monodentate change.

Table 1. EXAFS curve fitting parameters for the WT S1 state.

Table 2. EXAFS curve fitting parameters for Synechocystis D1-H332E mutant dark state.

Table 3. EXAFS curve fitting parameters for the Sr-substituted D1-H332E mutant dark state.
Table 1: Curve Fitting Results of Synechocystis WT S₁ state EXAFS spectrum.

<table>
<thead>
<tr>
<th>Fit No.</th>
<th>Mn-O</th>
<th>Mn-Mn</th>
<th>Mn-Mn</th>
<th>Mn-Ca</th>
<th>Mn-C</th>
<th>$R_f$ (%)</th>
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<td></td>
<td>R</td>
<td>N*a</td>
<td>$\sigma^2$</td>
<td>R</td>
<td>N*b</td>
<td>$\sigma^2$</td>
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<td>0.012</td>
<td>2.72</td>
<td>1.5</td>
<td>0.004</td>
</tr>
<tr>
<td>3</td>
<td>1.87</td>
<td>6.0</td>
<td>0.012</td>
<td>2.72</td>
<td>1.5</td>
<td>0.004</td>
</tr>
<tr>
<td>4</td>
<td>1.86</td>
<td>6.0</td>
<td>0.012</td>
<td>2.72</td>
<td>1.5</td>
<td>0.004</td>
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</tbody>
</table>

*a k-range of 2.5-11.4 Å⁻¹ was used for the fitting.

*b Bold letters indicate fixed parameters.

*c The MnO (~1.8 Å) coordination number was fixed to N=6.0 for simplicity, assuming hexa-coordination for all four Mn. Note that if one Mn out of four is penta-coordinate as suggested by the literature (53), the N value becomes 5.75. However, this does not cause a noticeable effect on the fit quality.

*d N values are defined as the total number of absorber-backscatterer vectors divided by the number of absorber atoms per OEC. For example, N=1.5 implies that there are three Mn-Mn ~ 2.7 Å interactions.

*e The MnC (~3.2 Å) coordination number was fixed to N=2.75, based on the ligand model of the 2.9 Å crystal structure (11). Approximately, each Mn sees two to four carbons of carboxylate groups and an imidazole ring, although these numbers remain somewhat ambiguous at the 2.9 Å structure due to its moderate resolution. However, the contribution of this path to the fitting quality is minor due to the high Debye-Waller factor.
Table 2: Curve Fitting Results of His332Glu dark state EXAFS spectrum.

<table>
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<tr>
<th>Model</th>
<th>Fit No.</th>
<th>Mn-O</th>
<th>Mn-Mn</th>
<th>Mn-Mn</th>
<th>Mn-Ca</th>
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<td></td>
<td></td>
<td>R (Å)</td>
<td>N^b</td>
<td>σ² (Å²)</td>
<td>R (Å)</td>
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<tr>
<td>3:1:x^a</td>
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<td>6.0</td>
<td>0.009</td>
<td>2.77</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>1.89</td>
<td>6.0</td>
<td>0.009</td>
<td>2.77</td>
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<tr>
<td>3:2:x</td>
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<td>6.0</td>
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<td>1.89</td>
<td>6.0</td>
<td>0.009</td>
<td>2.77</td>
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* k-range of 2.5-11.4 Å⁻¹ was used for the fitting.
* Bold letters indicate fixed parameters.
* E₀=6561.3 eV, S₀²=0.85.

^a This indicates the number of interactions in the model, di-μ-oxo bridged Mn-Mn(2.7 Å) : mono-μ-oxo bridged Mn-Mn(3.3 Å) : Mn-Ca(3.4 Å). The MnCa fragment (x) was changed either to x=2 (N=0.5) or x=3 (N=0.75) based on our previous study using Sr-substituted PSII (37).

^b Hexa-coordination for all four Mn was assumed for simplicity.
Table 3: Curve Fitting Results of Sr-substituted His332Glu dark state Sr EXAFS spectrum.

<table>
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<th>Fit No.</th>
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<th>N</th>
<th>σ² (Å²)</th>
<th>Sr-Mn R (Å)</th>
<th>N</th>
<th>σ² (Å²)</th>
<th>Sr-Mn R (Å)</th>
<th>N</th>
<th>σ² (Å²)</th>
<th>Rt (%)</th>
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<tr>
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<td>0.005</td>
<td>3.48</td>
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<tr>
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<td>0.012</td>
<td>14.4</td>
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* k-range of 2.7-12.8 Å⁻¹ was used for the fitting.
* Bold letters indicate fixed parameters.
* $E_0=16120.0$ eV, $S_0^2=0.99$.
* The coordination number N was fixed to 8.
Figure 1
Figure 2
Figure 3

(a) WT

(b) H332E
Figure 4

(a) Sr XANES

(b) Sr EXAFS
Figure 5
Figure 7
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