## Lawrence Berkeley National Laboratory

**LBL Publications** 

## Title

The Catalytic Manganese-Cluster: Organization of the Metal Ions", in Photosystem II: The Light-Driven Water:Plastoquinone Oxidoreductase

Permalink https://escholarship.org/uc/item/0181c544

Author

Yachandra, VK

## **Publication Date**

2022-12-17

Peer reviewed

eScholarship.org

# Chapter 10

## The Catalytic Manganese-Cluster: Organization of the Metal Ions

Vittal K. Yachandra\*

Melvin Calvin Laboratory, Physical Biosciences Division, Lawrence Berkeley National Laboratory, Berkeley, CA 94720, U.S.A.

Summary	1
I. Introduction	2
II. Oxidation States of the Manganese	3
A. Mn X-ray Absorption Near Edge Spectroscopy (XANES)	3
B. Mn Kβ X-ray Emission Spectroscopy (XES)	3
C. Oxidation State Changes During S-State Transitions	5
1. $S_1$ to $S_2$ Transition	5
2. S <sub>0</sub> to S <sub>1</sub> Transition	6
3. S <sub>2</sub> to S <sub>3</sub> Transition	6
D. Summary of Oxidation State Assignments	7
III. Structure of the Mn Cluster	8
A. Heterogeneity in the Mn-Mn Distances and Structural Implications	.10
B. S-State Transitions and Structural Changes	12
1. $S_1$ to $S_2$ (MLS) Transition	12
2. S <sub>1</sub> to S <sub>2</sub> (g=4.1) Transition	13
3. $S_2$ to $S_3$ Transition	13
4. $S_2$ to $S_3'$ ( $S_2Y_2$ ) Transition	14
5. So to SI Transition	15
IV. Structural Role of the Calcium Cofactor	15
A. Mn Extended X-ray Absorption Fine Structure (EXAFS) and Ca	16
B. Sr Extended X-ray Absorption Fine Structure (EXAFS) and Ca	16
1. Isotropic Samples	16
2. Oriented Samples	16
3. Orientation of the Mn Complex	17
C. Ca Extended X-ray Absorption Fine Structure (EXAFS)	17
V. Structural Role of the Chloride Cofactor	19
VI. Mechanism of Water Oxidation and O <sub>2</sub> Evolution	20
Acknowledgments	22
References	22

## Summary

The light-induced oxidation of water to  $O_2$  is catalyzed by a four-manganese atom cluster associated with Photosystem II (PS II). This chapter summarizes ongoing investigations of the oxidation state, the structure and the associated cofactors calcium and chloride of the catalytic Mn cluster using X-ray and electron paramagnetic resonance (EPR) spectroscopy. Manganese K-edge X-ray spectroscopy, K $\beta$  X-ray emission spectroscopy (XES), and extended X-ray absorption fine structure (EXAFS) studies have not only determined the oxida-

<sup>\*</sup>Email: VKYachandra@LBL.gov

T. Wydrzynski and K. Satoh (eds): Photosystem II: The Water/Plastoquinone Oxido-Reductase in Photosynthesis, pp. 000–000. © 2005 Kluwer Academic Publishers. Printed in The Netherlands.

1

2

3

4

5

6

7

8

9

10

11 12

13

14

15

16

17

#### PSII010

53

54

55

56

57 58

59

60

tion states and structural features, but also changes that occur in oxidation state of the Mn cluster and in its structural organization during the accumulation of oxidizing equivalents leading to  $O_2$  formation. Combining X-ray spectroscopy information with X-ray diffraction studies, and consistent with the available EPR data, we have succeeded in limiting the range of likely structures of the Mn cluster. EXAFS studies at the strontium and calcium K-edges have provided evidence that the catalytic center is a Mn/Ca heteronuclear complex. Based on the X-ray spectroscopy data, models for the structure and a mechanism for  $O_2$  evolution are presented.

## I. Introduction

Most of the  $O_2$  in the atmosphere that supports life on earth is generated by plants, algae, and cyanobacteria by the photo-induced oxidation of water to dioxygen:

$$2\mathrm{H}_{2}\mathrm{O} \rightarrow \mathrm{O}_{2} + 4e^{-} + 4\mathrm{H}^{+} \tag{1}$$

This reaction is catalyzed by a manganese/calcium 18 complex (Mn<sub>4</sub>Ca), which sequentially stores four 19 20 oxidizing equivalents (i.e. the  $S_n$  states, where n =(0-4) that are used to oxidize two molecules of water 21 to molecular oxygen (Kok et al., 1970; for a review, 22 23 see Joliot, 2003). The Mn complex is part of the multiprotein Photosystem II (PS II) assembly (Chapters 24 3-6), which contains the reaction center involved 25 in photosynthetic charge separation (Chapter 7) 26 and an antenna complex of chlorophyll molecules 27 28 (Chapter 2). The assembly also contains cytochrome  $b_{559}$  (Chapter 15) and a Fe-quinone electron acceptor 29 complex (Chapter 8). 30

31 X-ray absorption spectroscopy (XAS) (Sauer et al., 1992; Robblee et al., 2001) and electron para-32 magnetic resonance (EPR) (Dismukes and Siderer, 33 1981; Messinger et al., 1997b; Britt et al., 2000) 34 studies have emerged as the primary methods to 35 provide structural and chemical information about 36 the oxygen evolving complex (OEC) and devise a 37 working model for its Mn cluster. XAS has been used 38 39 to examine the structural environment of metal ions in many proteins (Cramer, 1988; Yachandra, 1995). 40 The energy of the incoming X-ray photons and of 41 42 the outgoing fluorescence is specific to the Mn atom;

43 Abbreviations: Chl – chlorophyll; <E> – 1<sup>st</sup>-moment value; 44 EPR - electron paramagnetic resonance; EXAFS - extended 45 X-ray absorption fine structure; FT-Fourier transform; IPE-in-46 flection point energy; MLS - multiline EPR signal; Mn<sub>4</sub> - catalytic manganese cluster; OEC - oxygen evolving complex; 47 PS II – Photosystem II;  $S_n$  – states of the OEC for n = 0, 1, 2,48 3, or 4; XANES – X-ray absorption near edge spectroscopy; 49 XAS – X-ray absorption spectroscopy; XES – X-ray emission 50 spectroscopy;  $Y_D^{ox}$  – oxidized form of the redox active tyrosine 51 on D2;  $Y_{z}^{*}$  – oxidized form of the redox active tyrosine on D1

hence, other metals or the protein matrix normally 61 co-purified with the OEC in a PS II preparation do 62 not interfere. Element-specificity and applicability to 63 non-crystalline samples have made X-ray spectros-64 copy a useful technique for probing the structure of 65 the Mn complex in the complicated environment of 66 PS II, which has many components such as non-heme 67 iron, cytochrome, chlorophylls that interfere with 68 other techniques. The structural studies can also be 69 performed on crystals, or on frozen solutions and 70 several of the S state intermediates mentioned above 71 72 have been stabilized as frozen solutions and studied by X-ray spectroscopy. The different regions of the 73 X-ray spectrum provide complementary information: 74 X-ray absorption near edge spectroscopy (XANES) 75 76 yield information about the oxidation states and site 77 symmetry of the absorbing atom while extended X-78 ray absorption fine structure (EXAFS) is sensitive 79 to distances, numbers, and atomic number of atoms around the absorbing atom. The more recently used 80 technique of X-ray emission spectroscopy (XES) has 81 also provided information regarding the identity of 82 the oxidation states of Mn complex in the various 83 S-states and the changes accompanying the S-state 84 85 advance.

Critical questions related to the process of pho-86 tosynthetic water oxidation are: (i) What are the 87 oxidation state(s) and structural changes in the Mn 88 complex as the OEC proceeds through the S-state 89 cycle? and (ii) What is the mechanism by which four 90 electrons are removed from two water molecules 91 by the Mn complex to produce an  $O_2$  molecule? 92 93 Electron paramagnetic resonance (EPR) and X-ray spectroscopy studies and the interplay between these 94 95 two methods have provided significant insights into the structure and the mechanism of the OEC. This 96 chapter focuses on the application of X-ray spec-97 troscopic techniques to resolve structural questions 98 regarding the Mn cluster in the OEC, with emphasis 99 on the results from our laboratory. Many excellent 100 reviews are available for comprehensive surveys of the 101 Mn complex and the OEC, EPR, and XAS literature 102 (Debus, 1992; Rutherford et al., 1992; Britt, 1996; 103

52

Yachandra et al., 1996, 2002; Penner-Hahn, 1998; Robblee et al., 2001).

## II. Oxidation States of the Manganese

1

2

3

4

5

6

7

8

9

10

11

12

13

14

15

16

17

18

19

21

Two key questions for the understanding of photosynthetic water oxidation are: (i) do all four oxidizing equivalents necessary to oxidize water to O<sub>2</sub> accumulate on the four Mn ions of the OEC, or do some ligand-centered oxidations take place before the formation and release of  $O_2$  during the  $S_3 \rightarrow [S_4]$  $\rightarrow$  S<sub>0</sub> transition; and (ii) what are the oxidation state assignments for the Mn during S-state advancement. X-ray absorption and emission spectroscopy of Mn, along with EPR have provided answers to these two pivotal questions.

#### A. Mn X-ray Absorption Near Edge Spectros-20 copy (XANES)

22 X-ray absorption near-edge spectroscopy (XANES) 23 is the most well-known of the X-ray techniques and 24 has been extensively used to investigate the oxida-25 tion states of redox-active metals in metalloprotein 26 active sites. XANES results from the excitation of a 27 1s electron (K-shell) to a higher, bound orbital (Fig. 28 1, *Left*). The higher the oxidation state of the metal, 29 the more positive the overall charge of the atom, and 30 higher energy is required to excite an electron out of 31 an orbital (Shulman et al., 1976). The first formally 32 allowed electric dipole transition is the 1s->4p tran-33 sition. Due to the size of the 4p orbital, it overlaps 34 with p-orbitals of the ligands, either through  $\sigma$  or  $\pi$ 35 bonding. Consequently, this transition is sensitive 36 to the oxidation state and the ligand environment of 37 the metal. For certain symmetries around the metal, 38 the formally electric-dipole forbidden 1s→3d tran-39 sition can be observed (arrow points to it in Fig. 1, 40 Middle which is expanded in the inset), occurring at 41 a lower energy than the main edge transitions (Roe 42 et al., 1984). This transition is due to mixing of metal 43 3d and 4p orbitals, and gives information about the 44 ligand as well as about the oxidation state and sym-45 metry of the metal complex (Westre et al., 1997). To increase the sensitivity of XANES, absorption is 46 47 detected as an excitation spectrum by measuring the 48 Ka fluorescence (2p to 1s) of the Mn atoms (Jaklevic 49 et al., 1977).

50 The pioneering application of Mn XANES to PS II 51 was performed by Kirby et al. (1981) on chloroplasts 52

53 and Goodin et al. (1984) on PS II preparations in the early 1980s. These studies, although extremely dif-54 55 ficult because of the low Mn concentration, showed that the Mn K-edge from the S<sub>2</sub> state is shifted to 56 57 higher energy relative to the Mn K-edge from the S<sub>1</sub> state. Dramatic improvements in detector technology 58 and cryostat cooling capabilities have made XANES 59 experiments practical using concentrated PS II prepa-60 rations (McDermott et al., 1988; Guiles et al., 1990a; 61 DeRose et al., 1994), and the collection of XANES 62 spectra from dilute, single-flash saturable PS II sam-63 ples is now achievable. Three different groups have 64 65 investigated the oxidation states of Mn for the  $S_0, S_1$ , S<sub>2</sub>, and S<sub>3</sub> states using XANES on single-flash satu-66 rable PS II samples. Based on shifts in the absorption 67 edge, or lack thereof, Roelofs et al. (1996) proposed 68 that Mn is oxidized during the  $S_0 \rightarrow S_1$  and  $S_1 \rightarrow S_2$ 69 70 transitions, but is not oxidized during the  $S_2 \rightarrow S_3$ 71 transition. In contrast, Ono et al. (1992) and Iuzzolino et al. (1998) interpret their XANES results to indicate 72 73 that Mn is oxidized during each S-state transition, although Ono et al. (1992) reported no independent 74 75 S-state determination for their samples and Iuzzolino et al. (1998) had significant S-state inhomogeneity 76 77 in their samples. However, improved XANES data for S-state transitions, have shown a Mn K-edge shift 78 of only 0.4 eV for the S<sub>2</sub> to S<sub>3</sub> transition (Ono et al., 79 1994). The results of Roelofs et al (1996) indicating 80 81 the absence of Mn centered oxidation during the  $S_2$ to S<sub>3</sub> transition has been reproduced by Messinger 82 et al. (2001) using both XANES (Fig. 1, Middle and 83 *Right*) and Mn XES studies (see below). 84 85

## B. Mn K<sub>β</sub> X-ray Emission Spectroscopy (XES)

87 The recent studies by Messinger et al. (2001) have 88 combined EPR, XANES and XES studies to ad-89 dress the questions of oxidation state assignments 90 and changes during the S state cycle. In contrast to 91 XANES, K $\beta$  XES detects the X-ray emission from 92 the relaxation of a 3p electron to a 1s hole, which is 93 created by excitation of a 1s electron into the con-94 95 tinuum (Fig. 2, Left). In a simplified model, two final states exist due to a constructive  $(K\beta_{1,3})$  or destructive 96 97  $(K\beta')$  spin-exchange interaction between the unpaired 98 electrons in the 3p and 3d orbitals (Tsutsumi et al., 1976). The magnitude of the exchange interaction 99 depends on the number of unpaired electrons in 100 the 3d orbital. In the high spin case, increasing the 101 oxidation state of the metal decreases the number of 102 unpaired 3d electrons; concomitantly, the spin ex-103 104



Fig. 1. Left. A schematic for the excitation and emission processes involved in Mn XANES. The weak 1s to 3d transition is dipole-forbidden and the principal transition is from 1s to a molecular orbital with p character. To enhance sensitivity, the X-ray absorption spectra are collected as excitation spectra using Mn K $\alpha$  (2p to 1s) fluorescence detection. *Middle Top.* Mn K-edge XANES spectra of flash-il-luminated PS II samples. Pure S-state spectra (bottom) were obtained from the flash spectra (top) by deconvolution using multiline EPR spectra. The pre-edge region (principally a 1s  $\rightarrow$  3d transition) for the S<sub>0</sub> – S<sub>3</sub> states is shown in the inset. The inflection point energy (IPE) changes from S<sub>0</sub> to S<sub>1</sub> and S<sub>1</sub> to S<sub>2</sub> transitions are substantial but are much smaller for the S<sub>2</sub> to S<sub>3</sub> transition. *Middle Bottom*. S-state Mn XANES difference spectra show that each transition is unique. The horizontal dashed lines show the zero value for each difference spectrum. Adapted from Messinger et al. (2001) and Visser et al. (2001). Right. Second derivatives of the normalized, pure S-state Mn K-edge spectra of the Mn-cluster of PS II. For clarity a vertical dashed line has been drawn at the inflection point energy (IPE) of the  $S_1$ -state. The change in IPE ( $\Delta$  in eV) for each S-state advance is given at the right shows the significant shifts for the  $S_0$  to  $S_1$  and  $S_1$  to  $S_2$  transitions but is much smaller for the  $S_2$  to  $S_3$  transition indicating that a Mn centered oxidation is unlikely for the  $S_2$  to  $S_3$  transition. For reference, the K-edge spectrum of the S<sub>1</sub>-state is plotted as well (dashed line). Adapted from Messinger et al. (2001). 

change interaction decreases. Accordingly, the  $K\beta_{1,3}$ transition shifts to a higher and the  $K\beta'$  transition shifts to a lower energy (Peng et al., 1994). Compared to the 4p orbitals, the 3p orbitals have less overlap with the ligand orbitals, because they are smaller and more buried within the electronic shells. Therefore, K $\beta$  XES is less sensitive to the ligand environment compared to XANES. The  $K\beta_{13}$  transition is better resolved than the  $K\beta'$  transition due to a difference in relaxation processes. Hence, the  $K\beta_{1,3}$  transition is used as a indicator of the oxidation state of the metal. A more accurate view on KB XES requires the ligand-field multiplet formalism. This indicates that there is some dependence of the ligand environment on K<sub>β</sub> XES spectra.

Shown in Fig. 2, Middle, and inset are the Kβ emis-sion spectra of Mn(IV)O<sub>2</sub>, Mn<sub>2</sub>(III)O<sub>3</sub>, and Mn(II)O, 

which illustrate the features of K $\beta$  emission spectra and their sensitivity to the oxidation states of Mn. As the oxidation state of Mn increases from Mn(II) to Mn(III) to Mn(IV), fewer unpaired 3d valence elec-trons are available to interact with the 3p hole; thus, the magnitude of the 3p - 3d exchange interaction becomes smaller, leading to a decrease in the  $K\beta'$  $-K\beta_{1,3}$  splitting. The consequence is that, if one fo-cuses only on the more intense  $K\beta_{1,3}$  emission peak, it shifts to lower energy as Mn is oxidized. Whereas the K $\beta$  emission spectrum is sensitive to oxidation state through a 3p-3d exchange interaction, the technique of XANES spectroscopy is sensitive to oxidation state through a different mechanism: core-hole shielding effects. This is why, upon Mn oxidation,  $K\beta_{1,3}$  emis-sion spectra shift to lower energy, while XANES spectra shift to higher energy. 

PSII010



*Fig. 2.* Kβ spectra of Mn(II), (III) and (IV) oxides. *Left.* Energy level diagram showing the absorption (1s to continuum) and emission processes (3p to 1s) involved in Kβ Mn XES. In a simplified model, two final spin states exist with either a constructive (K $\beta_{1,3}$ ) or destructive (K $\beta'$ ) spin exchange interaction between the unpaired 3p and 3d electrons. The magnitude of the interaction depends on the number of unpaired 3d electrons, which is related to the oxidation state of the metal. Adapted from Messinger et al. (2001) and Visser et al. (2001). *Middle*. Changes in the Kβ spectrum due to Mn oxidation. The inset shows a pictorial representation of the Kβ fluorescence transition for each of the oxides. Kβ XES arises from the emission of a 3p electron to 1s hole, which is formed following X-ray absorption. *Right*. The S<sub>1</sub>-S<sub>0</sub>, S<sub>2</sub>-S<sub>1</sub>, S<sub>3</sub>-S<sub>2</sub> and S<sub>0</sub>-S<sub>3</sub> difference high resolution Mn X-ray Kβ-emission spectra. The derivative shapes indicate shifts in the energy of the spectra between the S<sub>0</sub> and S<sub>1</sub>, and S<sub>2</sub> spectra, indicating oxidations of Mn. The S<sub>3</sub>-S<sub>2</sub> difference spectra demonstrates the similarity of the spectra, suggesting that this advance does not involve a Mn centered oxidation. Adapted from Messinger et al. (2001).

Using a 1<sup>st</sup>-moment analysis, the position of the main  $K\beta_{1,3}$  peak has been calculated for each S-state. Based on the shifts, or lack thereof, of the 1<sup>st</sup> moments (<E>), it was concluded that Mn is oxidized during the S<sub>0</sub> to S<sub>1</sub> and S<sub>1</sub> to S<sub>2</sub> but not oxidized during the S<sub>2</sub> to S<sub>3</sub> transition. The results (shown as difference spectra in Fig. 2, *Right*) show a shift in energy between the S<sub>0</sub> and S<sub>1</sub> state spectra and between the S<sub>1</sub> and S<sub>2</sub> state spectra. However, there is very little change between the S<sub>2</sub> and S<sub>3</sub> states, as seen in the difference spectrum (Messinger et al., 2001).

# C. Oxidation State Changes During S-State Transitions

Besides determining whether Mn is oxidized or not
during the S-state transitions, both XANES and XES
also allow one to assign the oxidation states of the
Mn atoms. This is often done by comparison to Mn
model compounds.

## 1. $S_1$ to $S_2$ Transition

Mn K-edge XANES spectra (Fig. 1, Middle and Right) and the K $\beta_{1,3}$  emission spectra (Fig. 2, *Right*) indicate the presence of Mn oxidation from  $Mn_4(III_2,IV_2)$  to  $Mn_4(III, IV_3)$  during the S<sub>1</sub> to S<sub>2</sub> transition (Roelofs et al., 1996; Messinger et al., 2001), where the S<sub>2</sub> state is characterized by the multiline (MLS) or the g=4.1EPR signal (Cole et al., 1987). This conclusion is corroborated by several other spectroscopic studies (Saygin and Witt, 1987; Kretschmann et al., 1988; Styring and Rutherford, 1988; Evelo et al., 1989; Dekker, 1992; Sharp, 1992) and is also confirmed by EPR spectroscopic measurements. S<sub>2</sub>-state EPR multiline signal simulations (Hasegawa et al., 1998; Hasegawa et al., 1999) and 55Mn ENDOR spectros-copy on the  $S_2$  state (Peloquin et al., 2000) are most consistent with the oxidation states of  $Mn_4(III, IV_3)$ ; however, other simulations (Zheng and Dismukes, 1996) prefer oxidation states of  $Mn_4(III_3,IV)$  in the  $S_2$  state. 

If the  $S_2$  state is proposed to be either in the 103

1  $Mn_4(III,IV_3)$  or the  $Mn_4(III_3,IV)$  oxidation states, 2 then the oxidation states of Mn in the  $S_1$  state must 3 be either  $Mn_4(III_2, IV_2)$  or  $Mn_4(III_4)$ , which are both 4 one-electron reductions from the oxidation-state pro-5 posals for the  $S_2$  state. Although the oxidation state 6 of  $Mn_4(III_4)$  for the S<sub>1</sub> state has been suggested by a 7 few groups (Zheng and Dismukes, 1996; Kuzek and 8 Pace, 2001; Chapter 12, Ahrling et al.), the XANES 9 spectrum or its  $2^{nd}$  derivative of the S<sub>1</sub> state (Figs. 1 10 Middle and Right) shows that its edge shape is un-11 like the edge shape observed for Mn(III) complexes. 12 When the XANES spectrum of the S<sub>1</sub> state, or its 13 2<sup>nd</sup>-derivative, is fit using Mn(III) and Mn(IV) model 14 compounds, it cannot be fit well using only Mn(III) 15 model compounds (Riggs et al., 1992; Yachandra 16 et al., 1993). Furthermore, inspection of the only 17 available set of tetranuclear Mn complexes with all O ligation, four distorted Mn<sub>4</sub>(III<sub>2</sub>,IV) cubanes, 18 19 shows XANES inflection point energy (IPE) values 20 at ~6551 eV for these complexes (Cinco et al., 1999) 21 consistent with the  $S_0$ -state XANES IPE of 6550.8 22 eV and lower than the  $S_1$ -state XANES IPE of 6552.9 23 eV. These results are consistent with the conclusion 24 that the oxidation states of Mn in the  $S_1$  state are 25  $Mn_4(III_2, IV_2)$ , not  $Mn_4(III_4)$ .

26 An oxidation-state assignment of  $Mn_4(III_2,IV_2)$  is 27 also consistent with recent K $\beta$  XES experiments on 28 the S<sub>1</sub> state. In fitting the K $\beta_{1,3}$  emission spectrum for 29 the  $S_1$  state, Bergmann et al. (1998) found that the 30 best fit to the experimental data was obtained using 31 equal amounts of the  $K\beta_{1,3}$  emission spectra from 32 Mn(III) and Mn(IV) model compounds; the fit was 33 significantly worse if only Mn(III) or only Mn(IV) 34 was used for the fit (Bergmann et al., 1998). It is 35 therefore most likely that the oxidation states for Mn 36 are  $Mn_4(III_2,IV_2)$  in the S<sub>1</sub> state and  $Mn_4(III,IV_3)$  in 37 the  $S_2$  state.

#### 2. So to S, Transition 39

38

40

41 The Mn K-edge XANES spectra for the  $S_0$  to  $S_1$ 42 transition, displayed in Fig. 1, Middle, and the 2<sup>nd</sup> 43 derivatives (Fig. 1, *Right*) show a clear shift of 2.1 44 eV (as determined by the 2<sup>nd</sup>-derivative IPE values) in 45 the XANES edge position. In addition, the  $S_1/S_0$  K $\beta$ 46 XES difference spectrum, shown in Fig. 2, *Right*, is 47 derivative-shaped, which is indicative of Mn oxidation 48 during this transition. Because the  $S_1$  state has Mn 49 oxidation states of  $Mn_4(III_2, IV_2)$ , as detailed above, 50 the changes in the X-ray spectroscopic data could 51 be due to either a Mn(II) to Mn(III) oxidation or a 52

53

72

73

74

75

76 77

78 79

80

81

82 83

84

85

86

87

88

89

90

91

92

93

94 95

96 97

Mn(III) to Mn(IV) oxidation.

The finding that the S<sub>1</sub>-S<sub>0</sub> XANES difference 54 55 spectrum (Fig. 1, Middle) is somewhat different from 56 the  $S_2$ - $S_1$  difference spectrum suggests that the  $S_0$  to 57 S<sub>1</sub> transition most likely reflects a Mn(II) to Mn(III) oxidation. The shape of the XANES edge is different 58 for the  $S_0$ -state and  $S_1$ -state spectra; this is shown in 59 part by the peak at 6548 eV in the 2<sup>nd</sup> derivative of 60 the S<sub>0</sub>-state XANES spectrum (Fig. 1, *Right*). This 61 62 is suggestive of the presence of Mn(II), based on comparison to the 2nd derivatives of Mn(II)-containing 63 model compounds. The significantly larger shift in 64 65 the  $2^{nd}$ -derivative IPE values for the S<sub>0</sub> to S<sub>1</sub> transition compared to the  $S_1$  to  $S_2$  transition is also indicative 66 for a Mn(II) to Mn(III) oxidation. This is based 67 on the finding that, for homologous sets of model 68 compounds, the shifts in the IPE value for Mn(II) to 69 70 Mn(III) oxidations are usually larger than those seen 71 for Mn(III) to Mn(IV) oxidations.

Other spectroscopic studies (Saygin and Witt, 1987; Kretschmann et al., 1988; Styring and Rutherford, 1988; Evelo et al., 1989; Guiles et al., 1990a; Dekker, 1992; Sharp, 1992; Messinger et al., 1997a,b; Åhrling et al., 1998) concur that Mn oxidation occurs during the  $S_0$  to  $S_1$  transition. The greater spectral width of the S<sub>0</sub>-state EPR multiline signal compared to the S<sub>2</sub>-state EPR multiline signal has been attributed to the presence of Mn(II) in the S<sub>0</sub> state (Messinger et al., 1997a,b; Ahrling et al., 1998), which suggests a Mn(II) to Mn(III) oxidation for the  $S_0$  to  $S_1$  transition. The presence of Mn(II) in the S<sub>0</sub> state may also explain why  $Y_D^{ox}$  can oxidize the  $S_0$  state to the  $S_1$ state, but not the  $S_1$  state to the  $S_2$  state.

The K $\beta$  XES and XANES difference spectra, in addition to other spectroscopic data are consistent with the assignment of the  $S_0$  to  $S_1$  transition as a Mn(II) to Mn(III) oxidation. Therefore, the oxidation states of Mn in the S<sub>0</sub> state are proposed to be  $Mn_4(II,III,IV_2)$ , which change upon Mn oxidation to  $Mn_4(III_2, IV_2)$  in the S<sub>1</sub> state, although the oxidation states of  $Mn_4(III_3, IV)$  for the S<sub>0</sub> state cannot be unequivocally excluded.

## 3. $S_2$ to $S_3$ Transition

98 The most controversial S-state transition has been 99 the  $S_2$  to  $S_3$  transition. Most debate has focused on whether this transition is a Mn-centered (Ono et 100 al., 1992; Iuzzolino et al., 1998) or ligand-centered 101 (Guiles et al., 1990b; Roelofs et al., 1996; Messinger 102 et al., 2001) oxidation. In addition, a redox isomerism

103

1 between Mn and ligands has been proposed for the 2  $S_3$  state (Renger, 1997). If Mn were to be oxidized 3 during the  $S_2$  to  $S_3$  transition, the oxidation would 4 have to be a Mn(III) to Mn(IV) transition. A Mn(IV) 5 to Mn(V) transition is unlikely to occur with Mn(III) 6 still present in the complex given the proposed re-7 activity of the Mn(V) ion. Moreover, Mn(V) has a 8 very distinctive Mn XANES spectrum that has not 9 been observed in PS II preparations (V. Yachandra, 10 unpublished). Thus, it would be expected that the 11 XANES and K $\beta$  XES difference spectra, as well as 12 the observed shifts in the XANES IPE values and the 13 K $\beta$  emission spectra 1<sup>st</sup>-moment values, <E>, would 14 be essentially identical to the  $S_1$  to  $S_2$  transition, where 15 a Mn(III) to Mn(IV) oxidation also occurs. However, as described below, the spectroscopic results for the 16 17  $S_2$  to  $S_3$  transition are completely different from those 18 of the  $S_1$  to  $S_2$  transition.

19 The XANES results from Fig. 1 provide strong 20 support for a ligand-based oxidation of the OEC 21 occurring during the  $S_2$  to  $S_3$  transition, based on the 22 small (0.3 eV) shift in the XANES 2<sup>nd</sup>-derivative IPE 23 values. The fact that the  $S_3$ - $S_2$  XANES difference 24 spectrum is significantly different from the S<sub>2</sub>-S<sub>1</sub> dif-25 ference spectrum (Fig. 1, Middle) is inconsistent with 26 a Mn(III) to Mn(IV) oxidation occurring during the 27 S<sub>2</sub> to S<sub>3</sub> transition. Instead, the S<sub>3</sub>-S<sub>2</sub> XANES differ-28 ence spectrum shows how the shape of the XANES 29 edge changes between the  $S_2$  and  $S_3$  states, which is 30 consistent with a ligand-based oxidation.

31 Therefore, based on the XANES data, it is reason-32 able to suggest that, a ligand radical is formed in the 33 S<sub>3</sub> state in lieu of Mn oxidation; this will be denoted 34 by  $Mn_4(III, IV_3)$ . This interpretation is reinforced by 35 the K $\beta$  emission spectra and especially, the K $\beta$  XES 36 difference spectra from Fig. 2, *Right*. The difference 37 spectra show that the derivative-shaped difference 38 spectrum that is expected if Mn is oxidized, as in the 39  $S_0$  to  $S_1$  and  $S_1$  to  $S_2$  transitions, is absent in the  $S_2$  to 40  $S_3$  transition. In addition, the  $\langle E \rangle$  value of 6490.157 41 eV for the S<sub>3</sub> state argues against Mn oxidation dur-42 ing the  $S_2$  to  $S_3$  transition, because it is inconsistent 43 with a  $Mn_4(IV_4)$  oxidation state, which is required 44 for the  $S_3$  state if Mn is oxidized during the  $S_2$  to  $S_3$ 45 transition. Comparison of the  $S_3$ -state  $\langle E \rangle$  value to 46 the  $\leq$ E> values from 18 different monomeric, dimeric, 47 trimeric, and tetrameric Mn(IV) model compounds 48 with different ligands, including Cl-, shows that the 49  $S_3$ -state  $\langle E \rangle$  value is higher than the  $\langle E \rangle$  value for 50 any of the Mn(IV) compounds studied. It is difficult to explain this result unless Mn(III) is still pres-51 52

ent in the S<sub>3</sub> state, which means that, based on the  $Mn_4(III,IV_3)$  redox states derived for the S<sub>2</sub> state, a Mn-based oxidation cannot occur during the  $S_2$  to  $S_3$  transition. The EXAFS studies from the  $S_2$  state described below also support this assignment.

PSII010

## D. Summary of Oxidation State Assignments

The Mn K-edge energy and shape do not depend 61 62 solely on the oxidation state of the Mn atoms, but also on the ligand environment. This is caused by mixing 63 64 of the ligand and the Mn atomic orbitals. It is possible 65 that a change in structure and ligand environment will influence the main-edge energy and shape. Therefore, 66 care must be exercised when correlating the Mn K-67 edge energies to Mn oxidation states without taking 68 into account the nature of the ligand environment. 69 This is especially a problem when sets of compounds 70 71 with different ligand structures, different nuclearities, and different analysis methods to determine edge 72 73 positions are used to infer oxidation-state information about biological systems, such as in Kuzek and Pace 74 75 (2001) and Carrell et al. (2002). However, numerous studies have shown that there is a clear correlation 76 between Mn oxidation state and Mn K-edge energy 77 if the ligand environment stays the same, i.e. the IPE 78 shifts to a higher energy upon oxidation within sets 79 of homologous compounds. 80

A detailed XANES and XES study by Visser et 81 al. (2001) of homologous di-µ-oxo and mono-µ-oxo 82 bridged binuclear compounds in different oxidation 83 states shows that in contrast to Mn K-edge XANES, 84 K $\beta$  XES spectra show much less dependence on li-85 gand environment. The shifts in XES spectra are the 86 same for a Mn(III) to Mn(IV) oxidation, irrespective 87 of whether the Mn is part of a di- or mono-µ-oxo 88 bridged structure (Visser et al., 2001). We have also 89 completed a XES study of homologous sets of tri- and 90 tetranuclear Mn compounds in different oxidation 91 states and we have found that the XES spectra are 92 much less dependent on the ligation or structure and 93 are primarily dependent on the oxidation state of Mn 94 95 (V. Yachandra, unpublished).

Considering both the limitations and strengths of 96 the two methods of XANES and XES, by comparison 97 98 with numerous Mn models and the knowledge from EXAFS data (see below), we have proposed the oxida-99 tion states shown in Fig. 3. Figure 3 summarizes the 100 Kβ XES and XANES flash patterns from our most recent and in our opinion the most comprehensive 102 study that leads to the conclusions about Mn oxidation 103

53

54

55

56

57

58

59

60



*Fig. 3. Left.* (A) Inflection-point energy (IPE) in eV of the Mn K-edge of PS II membranes as a function of the number of applied flashes. (B) The first moments of the Mn K $\beta$  emission spectra (<E>) as a function of flash number. The Mn K-edge shifts to higher energy as Mn is oxidized. The first moment of the Mn K $\beta$  fluorescence shifts to lower energy as Mn is oxidized. The complementary and mutually reinforcing results provide a strong case for Mn oxidation during the S<sub>0</sub> to S<sub>1</sub> and S<sub>1</sub> to S<sub>2</sub> advances but no Mn-centered oxidation at the S<sub>2</sub> to S<sub>3</sub> advance. *Right*. The S-state scheme for O<sub>2</sub> evolution and the oxidation states that are consistent with the EPR signals and the X-ray absorption and emission spectroscopy results. Adapted from Messinger et al. (2001).

states. Both spectroscopies show a clear shift in the spectra to higher energy for XANES spectroscopy and lower energy for K $\beta$  XES between the samples given 0 and 1 flashes indicating the presence of Mn oxidation during the  $S_1$  to  $S_2$  transition. The small change between samples given 1 and 2 flashes in both spectroscopies provides strong support for the  $S_2$  to  $S_3$ transition proceeding without a Mn-based oxidation. On the next transition,  $S_3 \rightarrow [S_4] \rightarrow S_0$ ,  $O_2$  is released, shifting the position of the spectra of samples given 3 flashes to lower energy for XANES spectroscopy and higher energy for K $\beta$  XES. These flash patterns are explained by the proposed oxidation states of the Mn cluster in the  $S_0$ ,  $S_1$ ,  $S_2$ , and  $S_3$  states, as shown. These oxidation states are:  $S_0$ :  $Mn_4(II,III,IV_2)$  or  $Mn_4(III_3,IV), S_1: Mn_4(III_2,IV_2), S_2: Mn_4(III,IV_3), S_3:$  $Mn_4(III, IV_3)$ , where the superscript dot represents an oxidation that is not Mn centered.

## 4748 III. Structure of the Mn Cluster

49
50 Extended X-ray absorption fine structure (EXAFS)
51 spectra at the Mn K-edge results from the modulation
52

of the absorption cross section by the constructive and destructive interference of the 1s photoelectron wave with that backscattered from neighboring at-oms (Savers et al., 1971). The EXAFS modulations can be theoretically modeled to derive exquisitely detailed information about the distance from Mn to neighboring atoms to  $\sim 4$  Å, often to an accuracy as precise as 0.015 Å (Yachandra, 1995). However, determination of the number of such interactions is susceptible to many uncertainties and can often be as large as 25%. The EXAFS approach has been used to determine the distances from Mn and identify the backscattering atoms of the Mn cluster present in the OEC. It is not possible to distinguish between O or N. But it is often facile to distinguish between light atoms such as N, or O, which are potential ligand atoms and backscattering from a heavier atom such as Mn or Ca that would only be present in a multi- or hetero-nuclear Mn complex. To interpret the results in terms of the coordination environment of the Mn, it has been necessary to examine a number of inorganic complexes of known structure, generously provided to us by our collaborators (Guiles et al., 1990a; DeRose et al., 1994; Cinco et al., 1999). The EXAFS 

spectrum is usually presented as a Fourier transform (FT), which is similar to a radial distribution function with each Fourier peak representing shells of neighboring atoms from the absorbing atom. Dichroism 5 resulting from oriented membranes can be used to 6 derive information about the vectorial direction of 7 the particular interaction. A typical Fourier transform 8 from an  $S_1$  state is shown in Fig. 4, *Top* (Yachandra 9 et al., 1993; DeRose et al., 1994) and from oriented PS II membranes in the  $S_1$  state in Fig 4, Bottom 10 (Mukerji et al., 1994). Three distinct Fourier peaks 11 12 are resolved in Fig. 4, Top, and dichroism is evident 13 in all of the peaks in Fig. 4, Bottom.

14 On the basis of the latest findings, we conclude that 15 each Mn is surrounded by a first coordination shell of O or N atoms at  $\sim 1.8$  to  $\sim 2.1$  Å (Fourier peak I), 16 17 a set of Mn atoms at ~2.7 Å (Fourier peak II) and additional Mn and Ca at ~3.3 Å (Fourier peak III). 18 19 Using model-compound data, it was straightforward 20 to assign the two or three 2.7 to 2.8 Å Mn-Mn vectors 21 to di-µ-oxo or hydroxo bridges between pairs of Mn 22 atoms and the 3.3 Å Mn-Mn vector to one or two 23 mono-µ-oxo bridged or similar longer interactions 24 (Kirby et al., 1981; Yachandra et al., 1986, 1987; 25 McDermott et al., 1988; George et al., 1989; Penner-26 Hahn et al., 1990; MacLachlan et al., 1992; DeRose et 27 al., 1994; Mukerji et al., 1994; Kusunoki et al., 1995; 28 Robblee et al., 2001). Furthermore, studies using 29 Sr to replace Ca and, subsequently, direct measure-30 ments using Ca EXAFS, show unambiguously the 31 presence of one or two 3.4 Å Mn-Ca vectors where 32 the bridging motif could also be mono-µ-carboxylato 33 or mono-µ-hydroxo (see below). The use of vector 34 terminology to describe these interatomic interactions 35 is particularly appropriate because of the direction 36 information available from dichroism measurements 37 on oriented samples. Furthermore, analysis of the 38 EXAFS leads to the conclusion that the di- or mono-39 µ-oxo bridged Mn and Ca vectors are not co-linear; 40 that is any three metal atoms in the cluster cannot 41 be linear.

42 On the basis of such data many topological tet-43 ranuclear Mn structures (A through K) compatible 44 with the observed EXAFS data have been presented 45 (DeRose et al., 1994) (Fig. 5). These models have been described in detail (DeRose et al., 1994; Rob-46 47 blee et al., 2001). However, until recently only one 48 of those options, A, was widely used as a working 49 model, although options E, F and G (the nomencla-50 ture was initially introduced in DeRose et al. (1994)) 51 have been shown to be preferred on the basis of EPR

1 2

3

4



PSII010

Fig. 4. Typical Fourier transform of the EXAFS spectrum from a S<sub>1</sub> state sample from an isotropic sample (Top) and oriented sample (Bottom). The three main Fourier peaks labeled I, II and III are assigned to ~ 2 O ligand atoms per Mn at 1.8 Å and 2-4 N/O ligand atoms per Mn at 1.9-2.1 Å (Fourier peak I), 2-3 Mn-Mn interactions at ~2.7 Å (Fourier peak II), and Mn and Ca at 3.3 and 3.4 Å (Fourier peak III), respectively. The dichroism of the Fourier peaks I, II and III is clear and can be used to determine the orientation of the vectors in the PS II membrane. Fourier peak II, that has been assigned to Mn-Mn interactions at ~2.7 Å, is maximal when the e-vector of the X-rays is parallel to the membrane plane (the angle on the figure is between the membrane normal and the X-ray e-vector.). Adapted from Yachandra et al. (1993), DeRose et al. (1994) and Mukerji et al. (1994).

and ENDOR simulations (Hasegawa et al., 1999; 94 95 Peloquin and Britt, 2001). Some of the structures, A through K, are less likely. Structure C and variations 96 97 consisting of two isolated di-µ-oxo Mn-Mn moieties are preferred by Pace and co-workers based on their 98 EPR simulations (Smith and Pace, 1996) but C is not 99 widely accepted on the basis of EPR simulations by 100 other groups (Boussac and Rutherford, 2000; Britt 101 et al., 2000; Peloquin et al., 2000) or EXAFS data 102 (Latimer et al., 1998). It is also physically impossible 103 104

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

76 77

78

79

80

81

82

83

84

85

86

87

88

89

90

91

PSII010



Fig. 5. Topological models for the Mn complex identified as compatible with the EXAFS data by DeRose et al. (1994) and Robblee at al. (2002). The numbers in parenthesis are the number of short 2.7 to 2.8 Å Mn-Mn vectors and long 3.3 to 3.4 Å Mn-Mn vectors. The options G, I, L and M all have 3 short 2.7 to 2.8 Å Mn-Mn vectors and are in better agreement with the X-ray diffraction studies; options E, F and G have been identified as compatible with EPR and ENDOR data. Option G or a topological isomer seems to be the best model by all presently available criteria; X-ray diffraction, EXAFS and EPR.

with four Mn atoms to obtain three Mn-Mn dis-tances (see below) at 2.7 - 2.85 Å with two separated di-µ-oxo Mn-Mn moieties. Structure H, with three 3.3 Å Mn–Mn distances is unlikely, as shown by the EXAFS spectra from a similar set of complexes -aseries of distorted cubanes (Cinco et al., 1999). The more likely candidates were considered to be A, E, **F**, **G** or **K** on the basis of EXAFS and EPR data; only G contained three Mn-Mn distances at ~2.7 Å, while the others had two such interactions.

## A. Heterogeneity in the Mn-Mn Distances andStructural Implications

As mentioned above determining the number of Mn-backscattering atoms can be difficult. The num-ber of the Mn-Mn 2.7 and 3.3 Å vectors (shown in parenthesis in Fig. 5) are especially important in this endeavor as these data directly lead or preclude certain arrangements of the Mn atoms that comprise the Mn cluster. The distance of the Mn-Mn vectors (at ~2.7 and 3.3 Å) is particularly important because it is easily the most reliably determinable parameter from the Mn EXAFS spectra of PS II. It is informative to focus on the ~2.7 Å Mn-Mn distance. When only one Mn-Mn 2.7 Å distance can be derived from the data as in the case of the S1 and S2 states, then one is constrained by 

## Chapter 10 Mn-Cluster and X-ray Spectroscopy

1 2

3

4

5

6

7

8

9

10

11 12

13

14

15

16

17

18

19

20

52

Table 1. Curve-fitting results for Fourier peak at ~2.7 Å that is characteristic of oxo-bridged Mn moieties from the S-states

S-state	Ref.	$R_{I}(\text{\AA})$	$R_2$ (Å)	$N_I$	$N_2$	$N_1:N_2$
$S_0^*$ state	Table III from Guiles et al. (1990a)	2.69	2.87	1.0	0.5	2:1
$S_2 (g = 4.1)$ state	Table 4B from Liang et al. (1994)	2.72	2.85	0.76	0.44	1.7:1
$NH_3$ -inhibited $S_2$ state	Table 1 from Dau et al. (1995)	2.71	2.86	0.75	0.5	1.5:1
F <sup>-</sup> inhibited S <sub>2</sub> state	Table 2 from DeRose et al. (1995)	2.71	2.85	0.8	0.4	2:1
S <sub>3</sub> state	Table 1B from Liang et al. (2000)	2.82	2.95	0.7	0.4	1.8:1
S <sub>0</sub> state	Table 3 from Robblee et al. (2002)	2.72	2.86	1.05	0.46	2.3:1
$S_0$ state	Table 4 from Robblee et al. (2002)	2.72	2.85	1.02	0.51	2:1

 $N_i$  and  $N_j$  are the number of Mn-Mn interactions, normalized to one Mn in the Mn<sub>4</sub> complex, at the two distances  $R_i$  and  $R_j$  respectively. An N of 1 translates to two Mn-Mn distances and a N of 0.5 translates to 1 Mn-Mn distance at the corresponding distance. The  $N_1 N_2$ ratio of 2:1 in the ~2.7 and ~2.8 Å Mn-Mn distances shows there are a total of 3 such short Mn-Mn distances in the OEC.

21 the uncertainty in the number of such interactions that 22 is inherent to the EXAFS methodology; it is at pres-23 ent two or three such Mn-Mn interactions. However, 24 when the distance degeneracy is lifted and there are two resolvable 2.7 or 2.8 Å Mn-Mn distances then 25 26 one can determine the number of such vectors to a 27 much higher accuracy because one can only have an 28 integral number of each of such interactions.

29 Interestingly, when we examined states that exhibited such heterogeneity in the 2.7 Å Mn-Mn distances, 30 it became quite clear that there was distinct prefer-31 32 ence for the existence of three such Mn-Mn vectors 33 as described below (summarized in Table 1). Figure 34 6, Left shows the Fourier transforms (FT) of the  $k^3$ weighted EXAFS data from the S<sub>0</sub> and S<sub>1</sub> states. The 35 Fourier peak II was isolated and fit to the EXAFS 36 37 equation with one or two distances for the S<sub>0</sub> and 38  $S_1$  states. Figure 6, *Right* shows the best fits to two Mn-Mn distances for the S<sub>0</sub> state as a function of 39 40 the number of each of the Mn-Mn interactions. The 41 contour plot graphically illustrates that the global minimum at two Mn-Mn interactions at ~2.7 Å and 42 43 one Mn-Mn interaction at ~2.85 Å is very well defined 44 (Robblee et al., 2002).

45 We have re-examined the data from the  $S_0^*$  ( $S_0$ 46 state induced by  $NH_2OH$  incubation), the g=4.1 S<sub>2</sub>, 47 the NH<sub>3</sub>- or F-inhibited  $S_2$ , and the  $S_3$  states, where 48 we have shown that there is distance heterogeneity in 49 the Mn-Mn vectors. The results of this re-examination are given in Table 1, which show that none of the 50 51 S-states that exhibit distance heterogeneity are best

fit by an equal N value for both Mn–Mn shells; the fit results from these S-states are in fact more consistent with a 2:1  $N_1$ :  $N_2$  ratio, where  $N_1$  corresponds to the shorter distance. In each case, the ratio of the number of the two Mn-Mn interactions was ~2:1. This leads us to think that there may be three Mn interactions at ~2.7 Å in the Mn cluster in its native state, one of which is perturbed in the S<sub>0</sub>, F<sup>-</sup>- and NH<sub>3</sub>-inhibited  $S_2$  and the g=4.1  $S_2$  states, and all three are perturbed in the S<sub>3</sub> state.

83 On the basis of these re-evaluations of the results from the S<sub>0</sub> state and earlier data, it is important 84 85 to consider the options that include three di-µ-oxo bridged moieties in the Mn cluster. Figure 5 shows 86 several such options, G, I, and J, that were proposed 87 earlier, and two newer options, L and M, among 88 several others that can be conceived. Options J and 89 I are less likely structures because J lacks a Mn-Mn 90 interaction at 3.3 Å, and I has two such interactions. 91 The EXAFS data from an inorganic compound (Au-92 ger et al., 1990) with the motif in option J is very 93 different from that obtained from a PS II sample (V. 94 95 Yachandra, unpublished). Options G, L and M all have three di-µ-oxo bridges and one mono-µ-oxo 96 bridge; that is three 2.7 Å and one 3.3 Å Mn-Mn 97 interactions. Options L and M are also similar to the 98 structure proposed on the basis of density functional 99 theory calculations although Ca is not included in L 100 and M and is an integral part of the Siegbahn model 101 (Siegbahn, 2000). 102

In addition to the constraints imposed by consider-103

104

Note: when referring to unpublished data, the word "unpublished" stands alone; "data" or "results" are not added as this is not useful information.

53

68

69

70

71

72

73

74

75

76

77

78

79

80

81

PSII010



Fig. 6. Left. Fourier Transforms of the  $S_0$  (dashed line) and  $S_1$  state (solid line) of the Mn-EXAFS spectra. The main Fourier peaks are labeled I, II and III. There are clear differences in amplitude and position of these peaks between the  $S_0$  and  $S_1$  states. The peak labeled II is from Mn-Mn interactions at ~2.7 Å, and in the  $S_0$  state this vector is best fit to two distances at 2.72 and 2.85 Å. *Right*. The contour plot shows the fit-quality-parameter  $\epsilon^2$  for the  $S_0$  state plotted versus the number of the two Mn-Mn interactions. The dashed lines shows the distinct minimum at one Mn-Mn vector at 2.85 Å and two Mn-Mn vectors at 2.7 Å. Adapted from (Robblee et al., 2002).

ing the EXAFS and EPR data, a preliminary structure of the OEC has been recently reported based on X-ray crystallographic data from PS II (Zouni et al., 2001; Kamiya and Shen, 2003; Ferreira et al., 2004; Chapters 19-21) and a further criterion for narrowing the choice of favored cluster geometries comes from these X-ray crystallography data of PS II. These data are most consistent with an OEC that is asymmetric and shaped somewhat like a 'Y'. The arrangement of the Mn atoms has been described as pear-shaped (an elongated ellipsoid) with all four Mn lying ap-proximately in the same plane. Presently, G, L and M are qualitatively in agreement with the asymmetry seen in the electron density of the Mn cluster (Zouni et al., 2001) and should be considered as possibili-ties for a topological model of the OEC based on the insights developed from the EXAFS spectroscopy results, with option G being the best in terms of being compatible with the most criteria; electron density, EPR, and EXAFS. However, as confirmed by Mn and Sr EXAFS studies (Latimer et al., 1995; Cinco et al., 1998) the OEC is most accurately described as a Mn/Ca heteronuclear cluster; therefore, Ca should be incorporated into each of the proposed structures so that 1-2 Mn-Ca vectors exist which are oriented close to the membrane normal (Ferreira et al., 2004; Chapter 21).

## 

#### *B. S-State Transitions and Structural Changes*

Although EXAFS studies may not be able to uniquelyidentify the structure of the Mn complex, they are

invaluable in understanding the changes in structure, if any, that might be occurring during the S-state transitions. As noted above, EXAFS is exquisitely sensitive to distances of neighboring atoms from Mn and hence can provide an unique window into how the Mn cluster changes as it proceeds through the catalytic cycle, as summarized in Fig. 7. These changes in structure provide a rationale for testing various mechanisms that have been proposed as described below.

### 1. $S_1$ to $S_2$ (MLS) Transition

EXAFS studies of the S<sub>1</sub> state and the S<sub>2</sub> state characterized by the multiline EPR signal revealed that the Mn–Mn distances are essentially the same: ~2.7 Å and 3.3 Å in both states. This distance is consistent with those found in numerous studies of di- and mono- $\mu$ -oxo-bridged and Mn<sub>2</sub>(III,IV) and Mn<sub>2</sub>(IV,IV) complexes. Oxidation of a mono- or di- $\mu$ -oxo bridged Mn motif from (III,III) to (III,IV) or from (III,IV) to (IV,IV) has minimal effect on the Mn-Mn separation in many inorganic models (Wieghardt, 1989; Pecoraro, 1992; Pecoraro and Hsieh, 2000).

Interestingly, EXAFS studies on oriented PS II in97the  $S_1$  and  $S_2$  states show that there maybe heterogene-98ity in the 2.7 Å vector, a contention that is supported99by related studies on modified  $S_2$  states. This Mn-Mn100vector has been determined to be at an average angle101of ~60° to the membrane normal (Mukerji et al., 1994; Dau et al., 1995).103

Note: the "new" Fig. 6 sent on the disk with the corrected 1st proof does not contain the dashed line indicated by the author. No version of this figure that I have received contains this dashed line.



*Fig.* 7. Summary of the changes in oxidation states and Mn–Mn distances at 2.7 to 3.0 Å in the native  $S_1$ , native  $S_2$ , modified  $S_2$ ,  $S_3'$  ( $S_2Y_2'$ ), native  $S_3$  and  $S_0$  states of PS II as determined by Mn EXAFS.

## 2. $S_1$ to $S_2$ (g=4.1) Transition

The inequivalence of the di-µ-oxo bridged Mn units becomes more evident in the S<sub>2</sub> state that is prepared by illumination at 130 K and characterized by the g=4.1 EPR signal (Liang et al., 1994). Similar results were obtained with NH<sub>3</sub>-treated and annealed S<sub>2</sub> state samples (Dau et al., 1995) and in F--treated samples (DeRose et al., 1995). The scheme in Fig. 7 shows that there are paths from the MLS S<sub>2</sub> state that lead to two states: the  $S_2$ -g=4.1 state inhibited by  $NH_3/F^-$  or the  $S_2 Y_z$  states; neither of these states can proceed to the physiologically relevant S<sub>3</sub> state. These states are depicted in Fig. 7 as branching away from the normal pathway leading to the S<sub>3</sub> state by photon absorption. However, the  $S_2$ -g=4.1 state generated by 820 nm illumination at 130 K (Boussac et al., 1996) can proceed to an S<sub>3</sub> state (Zimmermann and Rutherford, 1986). In the modified  $S_2$  states, one of the Mn-Mn distances increases to  $\sim 2.85$  Å, whereas there is very little change in the other one or two 2.7 Å or the 3.3 Å Mn-Mn distance. With the degeneracy of the 2.7 Å lifted, it has been possible by studying the dichroism of the Fourier peak, to assign the relative orientation of the 2.7 and 2.85 Å vectors (Dau et al., 1995), which are oriented at 55° and 67° respectively to the membrane normal.

46 The  $NH_3$ -treated and F-treated samples cannot 47 advance beyond the modified  $S_2$  state ( $S_2Y_2$ , state). 48 It is likely that  $F^-$  or  $NH_3$  prevent the oxidation of 49 a ligand atom, thereby blocking structural changes 50 from occurring that are necessary for the forma-51 tion of the  $S_3$  state, and thus inhibiting  $O_2$  evolution activity. In the case of  $NH_3$ , it is probably due to an amido-group  $(NH_2^{-})$  replacement of the oxo-bridge involved in oxidation. The modification of the MLS spectra upon addition of  $NH_3$  (Beck et al., 1986) and ESEEM studies using <sup>14</sup>NH<sub>3</sub> and <sup>15</sup>NH<sub>3</sub> demonstrated that NH<sub>3</sub> becomes a ligand of Mn (Britt et al., 1989). The asymmetry parameter derived from ESEEM results suggested that the amido group is likely to be a bridging ligand (Britt et al., 1989). It is probable that the oxo-bridge that is displaced by NH<sub>3</sub> is oxidized during the S<sub>2</sub> to S<sub>3</sub> transition (see next section).

PSII010

### 3. $S_2$ to $S_3$ Transition

Earlier XAS data from the S<sub>3</sub> state samples produced by a cryogenic double-turnover method indicated in-creased disorder in the peak at 2.7 Å; that a structural change that involved the lengthening of the Mn-Mn distance was occurring between the S<sub>2</sub> and S<sub>3</sub> states and was absent during the  $S_1$  to  $S_2$  state transition (Guiles et al., 1990b). More recent detailed analysis of data from native S<sub>3</sub> state samples created under physiological conditions with saturating actinic flash illumination show there is a significant change in the 2.7 Å Mn-Mn distances that characterize the di-µoxo bridged Mn-Mn; with an increase in the short Mn-Mn distances from 2.7 Å to ~2.8 and ~3.0 Å (Liang et al., 2000).

The perturbation in the Mn-Mn distances is different from that observed in the modified  $S_2$  states. Each of the altered  $S_2$  states gives a fit that is consistent with the alteration of one of the Mn-Mn distances from ~2.72 to 2.85 Å. Unlike the modified  $S_2$  states, 



*Fig. 8. Left.* Fourier-filtered *k*-space EXAFS data from  $S_2$  (dashed line) and  $S_3$  (solid line) state of PS II samples. The differences in phase, frequency and amplitudes between the raw  $S_2$  and  $S_3$  state EXAFS spectra are very obvious in these spectra. *Right.* Fourier transform power spectra of  $S_2$  (dashed line) and  $S_3$  (solid line) states of PS II. The major Fourier peaks are labeled I, II and III. The spectra are clearly different between the  $S_2$  and  $S_3$  states. There is a reduction in amplitude in all three peaks in the  $S_3$  state compared to the  $S_2$  state. More importantly, peaks II and III are at a greater apparent distance R' for the  $S_3$  state compared to the  $S_2$  state as shown.

the position of Fourier peak II from the native S<sub>3</sub> state clearly shifts to a longer distance in the FTs (Fig. 8) with one or two Mn-Mn distances of 2.82 Å and one Mn-Mn distance of 2.95 Å (Liang et al., 2000). These results underscore the point that during the  $S_2$  to  $S_3$ transition under physiological conditions, all of the di-µ-oxo-bridged Mn-Mn units undergo structural changes that lead to an increase in the Mn-Mn dis-tances. A change in the Mn-Mn distance from 2.7 to 3.0 Å was also observed in samples that had been depleted of Ca by NaCl treatment (MacLachlan et al., 1994). The best fits for the Mn-O/N shell shorter Mn-O distance is characteristic of Mn-O bridging distances and it increases from 1.80 Å in the S<sub>2</sub> state to 1.86 Å in the  $S_3$  state. This is a significant change, and it provides additional evidence for the involve-ment of the bridging oxygen atoms during the  $S_2$  to S<sub>2</sub> transition.

The results from isotropic S<sub>3</sub> samples are supported by polarized EXAFS studies on oriented PS II in the native S<sub>3</sub> state. The data confirm that the two Mn-Mn vectors units are not equivalent. The polarized EX-AFS data are different from those observed in the S<sub>2</sub> state (Mukerji et al., 1994; Dau et al., 1995). Fourier peak II of S<sub>3</sub> is dichroic and is readily resolved to Mn-Mn distances of ~2.8 Å and ~3.0 Å, each with its own distinct projection on the membrane normal (Fernandez et al., 1998). Schiller et al. (1998) have reported results on the oriented S<sub>3</sub> state, showing an increase in the amplitude of Fourier peak II that is 

interpreted as an increase in the number of 2.7 Å Mn-Mn interactions in the  $S_3$  state. However, a closer examination of their FTs indicates that Fourier peak II occurs at a longer distance and is broader compared to that in the  $S_2$  state.

In the  $S_3$  state it is seen that Fourier peak III at ~3.3 Å occurs at a greater apparent distance than that of the  $S_2$  state indicating a lengthening of the two Mn-Mn distances or the Mn-Mn and Mn-Ca distances compared to the  $S_2$  state by 0.04-0.2 Å.

Significant changes are observed in the Mn-Mn distances in the  $S_3$  state compared to the  $S_1$  and the  $S_2$  states (Fig. 8). These changes in Mn-Mn distances are interpreted as consequences of the onset of substrate water oxidation in the  $S_3$  state. Mn-centered oxidation is evident during the  $S_0$  to  $S_1$  and  $S_1$  to  $S_2$  transitions. During the  $S_2$  to  $S_3$  transition, we propose that the changes in Mn-Mn distances are the result of ligand or water oxidation, leading to the formation of an oxyl radical intermediate formed at a bridging or terminal position. The reaction of the oxyl radical with OH<sup>-</sup>, H<sub>2</sub>O, or an oxo group during the subsequent S-state conversion is proposed to lead to the formation of the O-O bond (Chapter 25).

## 4. $S_2$ to $S_3'(S_2Y_z)$ Transition

The other unproductive state generated from the  $S_2$ -MLS state is denoted in Fig. 7 as the  $S_3'$  state or  $S_2Y_2$ ' state generated in Ca-depleted samples. The

69

70

71

72

73

74

75

91

92

93

94

104

Ca-depleted samples are inactive in  $O_2$  evolution, while a broad g = 2 EPR signal has been observed in such samples. The g = 2 broadened EPR signal has been confirmed to arise from the tyrosine  $Y_z$ radical (Gilchrist et al., 1995; Peloquin et al., 1998) and it is proposed that the signal is broadened by interaction with the spin on the Mn cluster (Lakshmi et al., 1998).

1

2

3

4

5

6

7

8

52

9 EXAFS analysis from Ca-depleted S<sub>3</sub>' state 10 samples  $(S_2Y_z)$  (Latimer et al., 1998) shows that 11 Fourier peak II (Mn-Mn distance of ~2.7 Å) in the FT of  $S_3$ ' state samples ( $S_2Y_z$ ) was invariant relative 12 13 to that of the native  $S_2$  state sample. The EXAFS fits 14 showed that the ~2.7 Å Mn-Mn distances did not 15 lengthen as observed in the native  $S_3$  state samples 16 and are essentially unchanged from those of the na-17 tive S<sub>2</sub> state. This finding is surprising because the 18 Mn K-edges from these Ca-depleted samples showed 19 a behavior similar to the native PS II in that little 20 or no shift was observed in the  $S_2'$  to  $S_3'$  (or  $S_2Y_z$ ) 21 transition. This difference between the native S<sub>3</sub> and 22 Ca-depleted S<sub>3</sub>' states indicate that the core di-µ-oxo-23 bridged structure is probably dissimilar in the native 24  $S_3$  and the  $S_3'(S_2Y_z)$  states, with the structure in the 25  $S_2Y_z$  state resembling the native  $S_2$  state structure. 26 It is intriguing how the transfer of one electron from 27 the Mn cluster onto Y<sub>z</sub> result in major changes in 28 the Mn-Mn distances as is observed in the native S<sub>3</sub> 29 state. The roles of the  $Y_z$  and the Mn cluster in the 30 process of water oxidation are clearly delineated by 31 the comparison of the  $S_3$  and the  $S_3'$  (or  $S_2Y_7$ ) states. 32 Instead of the Mn cluster just providing a scaffolding 33 for water oxidation, this comparison shows that the 34 structural change in the Mn cluster initiated by the 35 transfer of an electron from the Mn cluster to Y<sub>z</sub> dur-36 ing the  $S_2$  to  $S_3$  transition might provide the trigger 37 to the chemistry of the formation of the O-O bond, 38 via the formation of a ligand radical. The results 39 show that the Mn cluster is involved in a much more 40 intimate manner in the catalysis than just providing 41 the framework. The implications to the mechanism 42 are significant (Chapter 25).

43 In Ca-depleted systems the tyrosine  $Y_z$  radical 44 is stabilized and the oxidation of the Mn-OEC and 45 the concomitant changes in Mn-Mn distances are 46 prevented. In the S<sub>3</sub> state, with Ca present, the oxida-47 tion equivalent resides on the Mn cluster leading to 48 profound changes in the Mn-Mn distances. Thus, Ca 49 is proposed to play a crucial role in controlling the 50 redox potential and thus the course of the mechanism 51 of water oxidation.

## 5. $S_0$ to $S_1$ Transition

Early EXAFS experiments by Guiles et al. (1990a) 55 with the S<sub>0</sub> state used chemical treatments to get 56 around the problem of low concentrations. However, 57 the  $S_0$  state generated in this manner was designated 58 as  $S_0^*$  to emphasize that it is generated through 59 chemical treatment and is thus not a native S-state. 60 Although hampered by a low signal-to-noise ratio and 61 the uncertainty about the relationship between the 62 chemically generated  $S_0^*$  state and the native  $S_0$  state, 63 those experiments provided the first evidence from 64 EXAFS that heterogeneity may exist in the 2.7 Å 65 Mn–Mn distances in the S<sub>0</sub> state in the form of a 66 reduced amplitude of Fourier peak II in the  $S_0^*$  state 67 relative to that in the  $S_1$  state. 68

Riggs-Gelasco et al. (1996a) examined reduced S states of the OEC generated by treatment with  $NH_2OH$  or hydroquinone and observed a decrease in the amplitude of the 2.7 Å Mn–Mn Fourier peak. This was interpreted as a reduction in the number of Mn–Mn vectors instead of the appearance of distance heterogeneity.

Recent EXAFS experiments by Robblee et al. 76 (2002) show that, in the  $S_0$  state, heterogeneity most 77 likely exists in the 2.7 Å Mn-Mn distances with one 78 or two Mn-Mn distances at 2.72 Å and one Mn-Mn 79 distance at 2.85Å (as described above, see Fig. 6), 80 which can be explained through the protonation of a 81 di-µ-oxo-bridged Mn–Mn moiety and/or the presence 82 of Mn(II) (Chapter 25). The presence of distance 83 heterogeneity in the  $S_0$  state has been exploited in 84 the curve-fitting procedure, whose results are sug-85 gestive of the possibility that three di-u-oxo-bridged 86 Mn-Mn moieties may exist in the OEC instead of 87 the two di-µ-oxo-bridged Mn-Mn moieties that are 88 widely used in proposed structural models for the 89 OEC (see above). 90

## IV. Structural Role of the Calcium Cofactor

Along with Mn and Cl-, Ca is an essential cofactor in 95 O<sub>2</sub> evolution (Debus, 1992; Chapter 13, van Gorkom 96 and Yocum). Depleting this cofactor suppresses 97 OEC activity, which can be restored (up to 90%) by 98 replenishing with Ca<sup>2+</sup>. Partial reactivation (up to 99 40%) results from addition of Sr<sup>2+</sup> to Ca-depleted 100 PS II membranes (Ghanotakis et al., 1984; Bous-101 sac and Rutherford, 1988b) and no other metal ions 102 can restore activity, making this requirement highly 103 specific for Ca or Sr (Ghanotakis et al., 1985; Ono and Inoue, 1989).

3 Although Sr<sup>2+</sup> replenishes the Ca-depleted centers 4 to a similar extent as added Ca<sup>2+</sup>, the slower kinetics 5 of the OEC turnover yields an overall lower steadystate rate (Boussac and Rutherford, 1988a) (40%) at 6 7 saturating light intensities. Substitution of Ca with 8 Sr also alters the EPR multiline signal (MLS) from 9 the S<sub>2</sub> state, giving narrower hyperfine splitting and different intensity patterns (Boussac and Rutherford, 10 11 1988b; Sivaraja et al., 1989). Most researchers ad-12 dressing the stoichiometry of the Ca cofactor in PS II 13 now conclude that functional water oxidase activity requires one essential Ca<sup>2+</sup>, which can be removed 14 15 by low-pH/citrate or 1.2 M NaCl wash (Cammarata and Cheniae, 1987; Katoh et al., 1987; Ono and In-16 oue, 1988; Shen et al., 1988; Adelroth et al., 1995). 17 In higher plants, another more tightly bound Ca is 18 19 associated with the light-harvesting complex (LHC 20 II) and requires harsher treatments for its removal 21 (Han and Katoh, 1993; Chen et al., 1995). 22

#### 23 A. Mn Extended X-ray Absorption Fine Struc-24 ture (EXAFS) and Ca

26 Several investigations have involved removal of Ca and substitution of various metals into this binding 27 28 site, followed by EXAFS studies on the Mn cluster. One set of experiments using Mn EXAFS on Sr-reac-29 30 tivated PS II membranes was interpreted to indicate 31 a 3.4–3.5 Å distance between the Ca (Sr) and the Mn 32 cluster (Latimer et al., 1995). This conclusion was 33 based on the observation of increased amplitude in Fourier peak III at 3.3 Å upon replacement of Ca with 34 Sr, a heavier atom and better X-ray scatterer. Analysis 35 of EXAFS spectra from purified PS II membrane 36 37 preparations indicated a Mn-Ca interaction at slightly longer distance (MacLachlan et al., 1992) (~3.6-3.7 38 39 Å). Ca depletion by NaCl-washing of PS II membranes removed the 16 and 23 kDa extrinsic proteins 40 and led to a reduced amplitude for this 3.6 Å feature 41 42 and because of the lower X-ray scattering ability of 43 Na, this result was interpreted as possible Na<sup>+</sup> sub-44 stitution for Ca<sup>2+</sup> at this distance (MacLachlan et al., 45 1994). Another Mn EXAFS study (Riggs-Gelasco et 46 al., 1996b) did not detect any changes in the Fourier 47 peak at 3.3 Å when Ca was replaced with Sr<sup>2+</sup> or 48 Dy<sup>3+</sup> in PS II reaction center complexes lacking the 16 and 23 kDa extrinsic polypeptides; however, it 49 50 was proposed that Ca might be linked via a hydrogen 51 bond to the oxo ligand of the Mn cluster. 52

## B. Sr Extended X-ray Absorption Fine Structure (EXAFS) and Ca

## 1. Isotropic Samples

The most common approach, as described above (Latimer et al., 1995; Riggs-Gelasco et al., 1996b), was to substitute other metals (such as Sr) for Ca and then use Mn EXAFS to detect changes in the cluster. Isolating the Mn-Ca or Sr component of the EXAFS spectrum from the combined EXAFS from all Mn-ligand and Mn-Mn interactions can be difficult. The reverse experiment where one probes for backscattering from Mn using Ca or Sr EXAFS (Ca/Sr cofactor point-of-view for nearby Mn) is an elegant alternative and is more definitive than Mn EXAFS results. Such studies on both isotropic and oriented PS II membranes have yielded unequivocal evidence for the proximity of Ca to the Mn cluster (see below).

Several factors favor Sr as the better cofactor for XAS study. First, the X-ray energies involved (16 keV for the K-edge) are more penetrating and not attenuated by air. The higher X-ray absorption cross-section and fluorescence yield of Sr also make the experiment practicable. The Sr EXAFS-based experiment requires PS II samples with Sr substituted for Ca while maintaining O<sub>2</sub>-evolving activity and a stoichiometry of 1 Sr per PS II, to focus on the functional cofactor binding site. Along with reactivated Sr-PS II, an inactivated sample can be prepared by treating with hydroxylamine (NH<sub>2</sub>OH) to disrupt the Mn cluster and suppress water oxidase activity (Tamura and Cheniae, 1985).

By using Sr EXAFS on isotropic Sr-reactivated PS II membranes, the proximity of Sr (and implicitly Ca) to within 3.5 Å of the Mn cluster (Cinco et al., 1998) has been confirmed. The finding was based on the presence of a second Fourier peak (peak II, Fig. 9, Top) in the Sr EXAFS from functional samples, a peak that is absent from inactive, hydroxylaminetreated PS II. This Fourier peak was found to fit best to two Mn at ~3.5 Å rather than lighter atoms (C, O, P, S, Cl). Both types of samples share similar first coordination shells of oxygen (Peak I, Fig. 9, Top).

## 2. Oriented Samples

The technique of using Sr EXAFS has been extended to using polarized Sr EXAFS on layered Sr-substituted samples, to provide important angle 103

16

1

2

25

53 54 55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

87

88

89

90

91

92

93

94

95

96

97

98

99

100

101

102



*Fig. 9. Top.* Fourier transforms of Sr EXAFS for intact and inactive Sr-substituted PS II samples (Chelex-treated). The dominant Fourier Peak I is due to ligating oxygens in the first coordination sphere and is common to both samples. The two Sr-PS II samples differ mainly at the 'R' = 3.0 Å region, where the intact samples exhibit Fourier Peak II. Disruption of the Mn cluster in NH<sub>2</sub>OH-treated sample leads to the absence of Peak II. *Bottom.* Fourier Transform of Ca EXAFS from Chelex-treated, layered samples with 2 Ca/PS II with O<sub>2</sub>-evolving activity and an S<sub>2</sub> EPR multiline signal. The FTs show the presence of a second Fourier peak in the 2Ca/PS II sample that fits to Ca-Mn that is absent in the control sample where the Mn complex has been disrupted with NH<sub>2</sub>OH.

42 information. Polarized EXAFS involves collecting 43 spectra for different incident angles ( $\theta$ ) between 44 the membrane normal of the layered sample and 45 the X-ray electric field vector. Dichroism in the 46 EXAFS can occur, depending on how the particular 47 absorber–backscatterer (A–B) vector is aligned with 48 the electric field. Through analysis of the dichroism, 49 the average orientation ( $\phi$ ) of this A–B vector relative 50 to the membrane normal, and the average number of 51 scatterers per absorbing atom ( $N_{iso}$ ) can be extracted.  Constraints on the structural model are then imposed by these parameters.

PSII010

Sr-substituted PS II samples made by a process of Ca depletion, Sr<sup>2+</sup> reactivation, and Chelex treatment to remove excess Sr were layered onto flat Mylar films to produce oriented samples (Mukerji et al., 1994; Dau et al., 1995). The Fourier transforms from the polarized Sr EXAFS showed extreme dichroism in Fourier peak II (Fig. 10, Left). Nonlinear least-squares regression analysis produced the solid curve shown in Fig. 10, Right as the best fit of the 15 data points (angles from six separate samples) and the result translates to 1-2 Sr-Mn vectors with an average angle of 0° or 23° for the average relative angle between the Sr-Mn vectors and the membrane normal depending on the method of analysis. 

#### 3. Orientation of the Mn Complex

The orientation data from the Sr EXAFS experiments can be combined with the dichroism data from Mn EXAFS data to calculate the orientation of the 3.3 Å Mn-Mn vector. The Fourier peak in the Mn EXAFS which contains the Mn–Mn (3.3 Å) and Mn–Ca (3.4 Å) contributions, is dichroic, with an average angle of  $43 \pm 10^{\circ}$  with respect to the membrane normal (Mukerji et al., 1994). By including the Mn-Ca vec-tor at 23°, an angle of  $\sim$ 62° for the 3.3 Å Mn–Mn vector can be calculated. Previous polarized Mn EXAFS experiments on PS II have shown angles of 55° and 67° for the 2.7 Å Mn–Mn vectors (Mukerji et al., 1994; Dau et al., 1995). Thus it follows that all Mn-Mn vectors lie at roughly the same angle  $(\sim 61^{\circ})$  with respect to the membrane normal, but are not restricted to being collinear, because the PS II membranes are ordered in one dimension only. The electron density from the X-ray diffraction studies (Zouni et al., 2001) are in agreement with such an assignment; the plane containing the Mn electron density is at  $\sim 67^{\circ}$  to the membrane normal. 

## C. Ca Extended X-ray Absorption Fine Structure (EXAFS)

In a complementary and definitive experiment, Ca97K-edge EXAFS studies have been used to probe the98binding site of the native cofactor for any nearby Mn,99within ~ 4 Å. The use of Ca EXAFS spectroscopy100has produced results essentially congruent with those101found by other independent methods: Sr EXAFS102on Sr-reactivated PS II (Cinco et al., 1998) and Mn103104



Fig. 10. Left. Fourier transforms of Sr-EXAFS from oriented Sr-PS II samples at two angles (θ). The dichroism is most readily apparent in Fourier peak II (R' = 3.0 Å) which is assigned to backscattering from Mn. *Right*. The polar plot shows N<sub>exp</sub> (filled circles) plotted vs. the angle of detection,  $\theta$ . The solid line is the best fit from which we obtain N<sub>iso</sub> (number of Mn neighbors to Sr is ~2) and  $\phi$  (~23°), the angle the Mn-Sr vector makes with the membrane normal.

EXAFS on similar samples (Latimer et al., 1995), but it focuses on the native cofactor and avoids the treatments involving Ca depletion and Sr substitution. Like the earlier Sr EXAFS, the Ca EXAFS study has focused on the Ca cofactor of PS II (poised in the S<sub>1</sub> state). This technique is a more sensitive and direct probe of the Ca binding site in PS II and the Ca EXAFS experiment directly probed the Ca cofactor in as close to a native system as possible.

The FT of the Ca EXAFS is shown in Fig. 9, Bottom and the spectra are remarkably similar to the Fourier transforms of the earlier Sr EXAFS study with Sr substituted for Ca. The first (largest) Fourier peak corresponds to the coordinating oxygen atoms closest to Ca. In contrast to the control (NH<sub>2</sub>OHtreated) sample, the Chelex-treated PS II shows a second Fourier peak. When Fourier peak II is isolated and simulated with possible scattering atoms, it corresponded best to Mn at 3.4 Å, rather than to light atom (C, O or Cl) neighbors. These results were consistent with the earlier Sr EXAFS studies.

The results are summarized in a motif shown in Fig. 11 and this motif depicts the Ca linked to two Mn by single-O bridges, which can be supplied by protein residues or water. Only single-oxygen bridges (not bidentate bridges) can provide the required 3.4 Å distance indicated by the Ca EXAFS fitting. This bridge may be derived from carboxylate ligands (aspartate or glutamate protein residues), protein 

backbone carboxyl, water or hydroxide.

Because significant information about Mn-Mn and Mn-Ca vector angles is now available, topo-logical models previously discussed can be refined to include the presence of Ca and account for the dichroism data (Fig. 11). We have chosen to modify two options: option  $\mathbf{F}$  (from Fig. 5) where the two di-µ-oxo motifs are formed using a common Mn atom and the mono- $\mu$ -oxo motif is placed at the end of the trinuclear unit, and option G can be thought of as at the corners of the base of a trigonal pyramid, with an O atom at the apex and three additional O atoms forming bridges pairwise among the Mn atoms be-low the base with the fourth Mn linked to one of the corner Mn atoms by a single O-atom bridge. Option F has two 2.7 to 2.8 Å Mn-Mn vectors while G has three such vectors. These structures are consistent with simulations of EPR and ENDOR data and also qualitatively in agreement with the reported electron density of the Mn cluster, especially the first option for **G** with Ca. 

It has been speculated that Ca controls substrate water binding to the catalytic Mn site (Chen et al., 1995) and recent mechanisms have suggested the crucial involvement of the cofactor (Ananyev and Dismukes, 1997; Renger, 1997; Limburg et al., 1999; Siegbahn, 2000). The results Mn EXAFS, Sr EXAFS and Ca EXAFS are mutually consistent and converge toward the conclusion that the Ca co-



*Fig. 11. Top.* Model of Ca-binding site of the oxygen-evolving complex in PS II. From the results of the Ca EXAFS studies on PS II, the Ca cofactor is linked by two single-O bridges to two Mn. The oxygens can be provided by water, hydroxyl or protein residues (carboxylate, phenolate). Other ligands to Ca are not shown. The arrangement shown here is not unique as other placements of the two Mn around the Ca are conceivable (Cinco et al., 2002). *Bottom.* Refined models for the active site of the OEC in PS II. These models combine the finding from oriented Sr-substituted PS II samples with previous results from Mn EXAFS on oriented PS II samples. These are derived from core structures that have been described in earlier studies and the models presented here are based on options F and G from Fig. 5 (F with Ca is a variation on a model originally proposed by Siegbahn (2000).

45 factor is intimately structurally linked with the Mn 46 cluster in PS II. Taken together, the three methods 47 offer compelling evidence that the catalytic center 48 of photosynthetic  $O_2$  evolution is a heteronuclear 49 Mn<sub>4</sub>Ca cluster (for further details, see Chapter 21, 50 Barber and Iwata).

V. Structural Role of the Chloride Cofactor

Despite a multitude of spectroscopic studies, defini-tive structural evidence for Mn-Cl ligation has not yet been reported. There is one functional Cl- per PS II unit (Lindberg and Andréasson, 1996; Olesen and Andréasson, 2003), but it is unclear whether chloride is a ligand to one of the Mn atoms in any of the S-states (Lindberg et al., 1990). Steady-state kinetic studies indicate the presence of a halide binding site on the Mn complex (Sandusky and Yocum, 1986; Yocum, 1992; Lindberg and Andréasson, 1996) and activity is inhibited by some compounds which compete with the Cl- binding site, such as fluoride (Sandusky and Yo-cum, 1986), primary amines (Sandusky and Yocum, 1986) and acetate (Clemens et al., 2002). Ono et al. (1986) have shown that Cl<sup>-</sup> depleted PS II particles in the  $S_2$  state do not have the usual multiline EPR signal but rather have a signal at g=4.1, signifying that an alternative S<sub>2</sub> state is formed. A similar EPR signal is observed on treatment with F-. It was found that when the OEC is in this alternative state, it can no longer advance to higher S-states. However, the EPR multiline signal is restored following illumina-tion and subsequent addition of Cl-. Recent studies indicate that the presence of the Cl- is necessary only for the  $S_2$  to  $S_3$  and  $S_3$  to  $S_0$  transitions of the OEC, while the earlier steps of the cycle can proceed in its absence (Wincencjusz et al., 1997, 1998). These studies indicate that Cl- is closely associated with the structure of the OEC and the mechanism of  $O_2$ evolution, but its detailed role is as yet unclear. 

F- perturbation of the Mn–Mn distances by treat-ment with F<sup>-</sup> is one of the most direct structural data available that implies halide as a ligand of Mn. XAS studies of F- inhibited samples show that one of the two 2.7 Å Mn–Mn distances is increased to ~2.8 Å, which is suggestive of F<sup>-</sup> binding to the Mn cluster (DeRose et al., 1995). The recent ESEEM results from acetate treated PS II samples (Clemens et al., 2002) also provide the most direct spectroscopic evidence to date for the presence of a chloride ligand to Mn. 

There is evidence from EXAFS studies with oriented S<sub>3</sub> state samples indicating the presence of Mn-Cl ligation in the S<sub>3</sub> state (Fernandez et al., 1998). EXAFS results from isotropic samples in the S<sub>3</sub> state have shown that there is considerable change in the Mn-Mn distances as the system advances from the  $S_2$  to the  $S_3$  state (Liang et al., 2000). The two 2.7 Å Mn–Mn distances in the S<sub>2</sub> state increase to ~2.8 and ~3.0 Å in the  $S_3$  state. With the aim of determining the relative orientation of the 2.8 and 



*Fig. 12.* The Fourier transform of an oriented  $S_3$  sample. The Mn-Mn vectors represented by Fourier peaks labeled II and III are dichroic. Interestingly, a Fourier peak labeled 1.5 is discernible that is not observed in other S-states and it fits best to Cl, indicating the possible presence of a Cl ligand to Mn at a distance of 2.2 Å (Fernandez et al., 1998) in the  $S_3$  state.

2.9-3.0 Å Mn-Mn vectors, we initiated XAS stud-ies of oriented  $S_3$  state samples. The FT in Fig. 12 shows that the two Mn-Mn vectors are dichroic. Interestingly, maybe because of the lengthening of the Mn–Mn vectors, a new Fourier peak is observed between the first and second Fourier peaks, labeled 1.5 in Fig. 12. Comparing this FT with that of a Mn binuclear complex with one Cl- as a terminal ligand to a Mn atom shows that the peak corresponding to Cl backscattering in the model complex is found at approximately the same apparent distance as the new Fourier peak in the PS II samples (Fernandez et al., 1998). In other model compounds, the amplitude of this peak increases as the ratio of Cl/Mn increases and the peak is absent when Cl<sup>-</sup> is not present as a ligand. In the  $S_3$  state sample, this peak does not fit to lower Z atoms such as C, N, or O. The fit is significantly better for Cl backscattering at ~2.2 Å. The fits for the model compound are very similar. Dichroism stud-ies on  $S_3$  state samples show that the Fourier peak is larger at the 10° orientation compared to the 80° orientation, which suggests that the Mn–Cl vector is more parallel to the membrane normal.

## 

# VI. Mechanism of Water Oxidation and $\rm O_2$ Evolution

The mechanisms for water oxidation and oxygen evolution that have been proposed can be broadly divided into four categories in which the character of the oxygen atoms that ultimately form dioxygen is different. In these four groups the oxygen atoms come from: two terminal oxygens bound to two separate Mn atoms, one terminal oxygen and one oxygen not bound to the Mn cluster, one terminal and one bridging oxygen, or two  $\mu$ -oxo bridges. Each of these mechanisms involves distinctly different Mn–O bonds formed and broken during the catalytic cycle (see Chapter 25 for details).

The contribution of X-ray studies to the mechanism of O<sub>2</sub> evolution is in resolving whether an oxygen atom derived from a bridging position in the Mn cluster is involved in the ultimate formation of dioxygen and in the determination of whether Mn centered oxidation occurs during each stage of the S-state transitions. Our X-ray spectra are strongly supportive of Mn cluster oxidation during the  $S_0 \rightarrow S_1$  and  $S_1 \rightarrow S_2$ transitions, they do not support a similar Mn-centered oxidation during  $S_2 \rightarrow S_3$ . Furthermore, EXAFS analyses show clear evidence of non-degeneracy in the three 2.7 Å vector lengths in  $S_0$  and  $S_3$ , but not in S<sub>1</sub> and S<sub>2</sub>, and shows significant structural change 



Fig. 13. Summary of changes in Mn oxidation states and Mn-Mn distances during photosynthetic water oxidation using option G in Fig. 5. 35 In the  $S_1 \rightarrow S_2$  transition, one Mn(III) is oxidized to Mn(IV), and the Mn–Mn distances do not change. During the  $S_2 \rightarrow S_3$  transition, the 36 oxidizing equivalent is localized on the  $\mu$ -oxo bridge, which triggers the increase in Mn–Mn distances. In the  $S_3 \rightarrow [S_4] \rightarrow S_0$  transition, 37 a short-lived peroxo intermediate might be formed in the  $S_4$  state. (a), (b), (c) and (d) are 4 mechanisms for O-O bond formation in the 38  $S_4$  state. Mechanisms (a), (b) and (c) are where one or two bridging O atoms are directly involved in O-O bond formation; a motif that is 39 consistent with XAS data. Mechanism (d) would require localization of an oxidizing equivalent on an oxo bridge that later exchanges or rearranges to a terminal O ligand before O-O bond formation. During the  $S_0 \rightarrow S_1$  transition, a Mn(II)  $\rightarrow$  Mn(III) oxidation or a Mn(III) 40 to Mn(IV) concurrent with a deprotonation of a hydroxy to an oxo causing the decrease of the Mn–Mn distance is indicated. For clarity, 41  $Y_{z}$ , the cofactors Ca<sup>2+</sup> and Cl<sup>-</sup>, and terminal Mn ligands are not shown. Mn–Mn distances were determined by EXAFS spectroscopy. As 42 mentioned in DeRose et al. (1994), Cinco et al. (2002) and Robblee et al. (2002) other possible topological models exist for the OEC; 43 similar mechanisms that can be proposed for each of the alternative topological models should be considered equally viable. 44

46 during the  $S_2$  to  $S_3$  transition that is not accompanied 47 by a Mn-centered oxidation. The X-ray spectroscopy 48 evidence supports the involvement of a bridging O 49 atom in the mechanism of the formation of the O-O 50 bond. These observations must be considered in any 51 postulation of mechanistic aspects of the successive 52

45

steps in water oxidation.

Our proposed mechanism (illustrated in Fig. 13 99 using the proposed model **G**) where the O-O bond 100 is formed between one critical bridging oxygen and 101 another oxygen atom derived from a bridging/terminal oxygen ligand or an exogenous oxygen also 103

87

88

89

90

91

92

93

94

95

96

97 98

1 2

3

4

5

6

7

8

9

11

16

17

18

### PSII010

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

87

88

89

90

91

92

93

94

95

96

97

98

99

100

101

avoids the formation of the O-O bond until the most oxidized state  $(S_4)$  is reached (Fig. 13, (a), (b), (c)). All four options for the  $S_4$  state generate  $O_2$  which is derived from inequivalent oxygen atoms in the  $S_{2}$ state, as shown by <sup>16</sup>O/<sup>18</sup>O water-exchange measurements on the S<sub>2</sub> state (Messinger et al., 1995; Hillier et al., 1998). In addition, a recent  $H_2^{16}O/H_2^{18}O$  FTIR experiment has shown the presence of an exchangeable di- $\mu$ -oxo-bridge in the S<sub>1</sub> and S<sub>2</sub> states (Chu et 10 al., 2000). This mechanism precludes the formation and release of peroxide or other oxidation products 12 of water in the earlier S-states, thus preventing the 13 system from 'short circuiting' and avoiding the risk 14 of damaging the polypeptides of PS II. 15

## Acknowledgments

19 This work was supported by the National Institutes 20 of Health grant (GM 55302) and by the Director, 21 Office of Science, Office of Basic Energy Sciences, 22 Division of Energy Biosciences, U.S. Department 23 of Energy under contract DE-AC03-76SF00098. I 24 am grateful to every one of the very talented and 25 motivated graduate students and postdocs, and many 26 sabbatical visitors to the Calvin Lab that I have had 27 the privilege to work with on the oxygen evolving 28 complex over the last two decades and more; each 29 and every one of them can recognize some aspect of 30 the work presented here that they were responsible for 31 and that has contributed to the present understand-32 ing of the Mn cluster in the OEC. Specifically, the 33 contributions to the research presented in this article 34 by Roehl Cinco, John Robblee, Johannes Messinger, 35 Henk Visser, Uwe Bergmann, Pieter Glatzel, Carmen 36 Fernandez, Elodie Anxolabéhère-Mallart, Shelly 37 Pizarro, Karen McFarlane, Junko Yano, Steve Cra-38 mer and Ken Sauer is gratefully acknowledged. Mel 39 Klein's profound influence and contributions to the 40 EPR and X-ray spectroscopy research pervades the 41 work presented in this chapter. I extend my thanks 42 to our collaborators Profs. W. H. Armstrong, G. W. 43 Brudvig, G. Christou, T. Collins, J.-J. Girerd, R. N. 44 Mukherjee, V. L. Pecoraro and K. Wieghardt who 45 have generously provided all the inorganic Mn compounds that has made all the difference in analyzing 46 47 the data from PS II. Synchrotron radiation facilities 48 were provided by the Stanford Synchrotron Radiation 49 Laboratory (SSRL) and the Advanced Photon Source 50 (APS), both supported by the U.S. Department of 51 Energy and the National Institutes of Health.

## References

- Adelroth P, Lindberg K and Andréasson LE (1995) Studies of Ca2+ binding in spinach Photosystem II using 45Ca2+. Biochemistry 34:9021-9027
- Åhrling KA, Peterson S and Styring S (1998) The S<sub>0</sub> state EPR signal from the Mn cluster in Photosystem II arises from an isolated S=1/2 ground state. Biochemistry 37: 8115-8120
- Ananyev GM and Dismukes GC (1997) Calcium induces binding and formation of a spin-coupled dimanganese (II, II) center in the apo-water oxidation complex of Photosystem II as precursor to the functional tetra-Mn/Ca cluster. Biochemistry 36: 11342-11350
- Auger N, Girerd J-J, Corbella M, Gleizes A and Zimmermann J-L (1990) Synthesis, structure, and magnetic properties of the stable triangular [Mn(IV)<sub>3</sub>O<sub>4</sub>]<sup>4+</sup> core. J Am Chem Soc 112:448-450
- Beck WF, Depaula JC and Brudvig GW (1986) Ammonia binds to the manganese site of the O2-evolving complex of Photosystem-II in the S<sub>2</sub> state. J Am Chem Soc 108: 4018–4022
- Bergmann U, Grush MM, Horne CR, DeMarois P, Penner-Hahn JE, Yocum CF, Wright DW, Dubé CE, Armstrong WH, Christou G, Eppley HJ and Cramer SP (1998) Characterization of the Mn oxidation states in Photosystem II by KB X-ray fluorescence spectroscopy. J Phys Chem B 102: 8350-8352
- Boussac A and Rutherford AW (1988a) S-state formation after Ca<sup>2+</sup> depletion in the Photosystem II oxygen-evolving complex. Chem Script 28A: 123–126
- Boussac A and Rutherford AW (1988b) Nature of the inhibition of the oxygen-evolving enzyme of Photosystem II induced by NaCl washing and reversed by the addition of Ca<sup>2+</sup> or Sr<sup>2+</sup>. Biochemistry 27: 3476-3483
- Boussac A and Rutherford AW (2000) Comparative study of the g=4.1 EPR signals in the S<sub>2</sub> state of Photosystem II. Biochim Biophys Acta 1457: 145-156
- Boussac A, Girerd J-J and Rutherford AW (1996) Conversion of the spin state of the manganese complex in Photosystem II induced by near-infrared light. Biochemistry 35: 6984-6989
- Britt RD (1996) Oxygen evolution. In: Ort DR and Yocum CF (eds) Oxygenic Photosynthesis: The Light Reactions, pp 137–164. Kluwer Academic Publishers, Dordrecht
- Britt RD, Zimmermann J-L, Sauer K and Klein MP (1989) Ammonia binds to the catalytic manganese of the oxygenevolving complex of Photosystem II. Evidence by electron spin-echo envelope modulation spectroscopy. J Am Chem Soc 111: 3522-3532
- Britt RD, Peloquin JM and Campbell KA (2000) Pulsed and parallel-polarization EPR characterization of the Photosystem II oxygen-evolving complex. Ann Rev Biophys Biomol Struct 29:463-495
- Cammarata KV and Cheniae GM (1987) Studies on 17, 24 kDa depleted Photosystem II membranes. Plant Physiol 84: 587-595
- Carrell TG, Tyryshkin AM and Dismukes GC (2002) An evaluation of structural models for the photosynthetic water-oxidizing complex derived from spectroscopic and X-ray diffraction signatures. J Biol Inorg Chem 7: 2-22
- Chen C, Kazimir J and Cheniae GM (1995) Calcium modulates the photoassembly of Photosystem II (Mn)<sub>4</sub>-clusters by preventing ligation of nonfunctional high-valency states of manganese.

52

Biochemistry 34: 13511-13526

- Chu H-A, Sackett H and Babcock GT (2000) Identification of a Mn-O-Mn cluster vibrational mode of the oxygen-evolving complex in Photosystem II by low-frequency FTIR spectroscopy. Biochemistry 39: 14371–14376
- Cinco RM, Robblee JH, Rompel A, Fernandez C, Yachandra VK, Sauer K and Klein MP (1998) Strontium EXAFS reveals the proximity of calcium to the manganese cluster of oxygen-evolving Photosystem II. J Phys Chem B 102: 8248–8256
- Cinco RM, Rompel A, Visser H, Aromí G, Christou G, Sauer K, Klein MP and Yachandra VK (1999) Comparison of the manganese cluster in oxygen-evolving Photosystem II with distorted cubane manganese compounds through X-ray absorption spectroscopy. Inorg Chem 38: 5988–5998
- Cinco RM, Holman KLM, Robblee JH, Yano J, Pizarro SA, Bellacchio E, Sauer K and Yachandra VK (2002) Calcium EXAFS establishes the Mn-Ca cluster in the oxygen-evolving complex of Photosystem II. Biochemistry 41: 12928–12933
- Clemens KL, Force DA and Britt RD (2002) Acetate binding at the Photosystem II oxygen evolving complex: An S<sub>2</sub>-state multiline signal ESEEM study. J Am Chem Soc 124: 10921–10933
- Cole J, Yachandra VK, Guiles RD, McDermott AE, Britt RD, Dexheimer SL, Sauer K and Klein MP (1987) Assignment of the g=4.1 EPR signal to manganese in the S<sub>2</sub> state of the photosynthetic oxygen-evolving complex: An X-ray absorption edge spectroscopy study. Biochim Biophys Acta 890: 395–398
- Cramer SP (1988) Biochemical application of X-ray absorption spectroscopy. In: Koningsberger DC and Prins R (eds) X-Ray Absorption: Principles, Applications, Techniques of EXAFS, SEXAFS and XANES, pp 327–320. John Wiley & Sons, New York
  - Dau H, Andrews JC, Roelofs TA, Latimer MJ, Liang W, Yachandra
     VK, Sauer K and Klein MP (1995) Structural consequences of ammonia binding to the manganese cluster of the photosynthetic oxygen-evolving complex: An X-ray absorption study of isotropic and oriented Photosystem II particles. Biochemistry 34: 5274–5287
- 32 Debus RJ (1992) The manganese and calcium ions of photosynthetic oxygen evolution. Biochim Biophys Acta 1102: 269–352
- Dekker JP (1992) Optical studies on the oxygen-evolving complex
   of Photosystem II. In: Pecoraro VL (ed) Manganese Redox
   Enzymes, pp 85–103. VCH Publishers, New York
- 37 DeRose VJ, Mukerji I, Latimer MJ, Yachandra VK, Sauer K and Klein MP (1994) Comparison of the manganese oxygen-evolving complex in Photosystem II of spinach and *Synechococcus* sp. with multinuclear manganese model compounds by X-ray absorption spectroscopy. J Am Chem Soc 116: 5239–5249
- 41 DeRose VJ, Latimer MJ, Zimmermann J-L, Mukerji I, Yachandra
   42 VK, Sauer K and Klein MP (1995) Fluoride substitution in the
   43 Mn cluster from Photosystem II: EPR and X-ray absorption
   43 spectroscopy studies. Chem Phys 194: 443–459
- Dismukes GC and Siderer Y (1981) Intermediates of a polynuclear
   manganese cluster involved in photosynthetic oxidation of
   water. Proc Natl Acad Sci USA 78: 274–278
- Evelo RG, Styring S, Rutherford AW and Hoff AJ (1989) EPR
  relaxation measurements of Photosystem-II reaction centers — influence of S-state oxidation and temperature. Biochim Biophys Acta 973: 428–442
- Fernandez C, Cinco RM, Robblee JH, Messinger J, Pizarro SA,
   Sauer K, Klein MP and Yachandra VK (1998) Calcium and

chloride cofactors of the oxygen evolving complex — X-ray<br/>absorption spectroscopy evidence for a Mn/Ca/Cl heteronuclear<br/>cluster. In: Garab G (ed) Photosynthesis: Mechanisms and<br/>Effects, Vol II, pp 1399–1402. Kluwer Academic Publishers,<br/>Dordrecht5355565657

- Ferreira K, Iverson TM, Mughlouni K, Barber J and Iwata S (2004) Architecture of the photosynthetic oxygen-evolving center. Science 303: 1831–1838
- George GN, Prince RC and Cramer SP (1989) The manganese site of the photosynthetic water-splitting enzyme. Science 243: 789–791
- Ghanotakis DF, Babcock GT and Yocum CF (1984) Calcium reconstitutes high rates of oxygen evolution in polypeptide depleted Photosystem II preparations. FEBS Lett 167: 127–130
- Ghanotakis DF, Babcock GT and Yocum CF (1985) Structure of the oxygen-evolving complex of Photosystem II: Calcium and lanthanum compete for sites on the oxidizing side of Photosystem II which control the binding of water-soluble polypeptides and regulate the activity of the manganese complex. Biochim Biophys Acta 809: 173–180
- Gilchrist ML, Jr., Ball JA, Randall DW and Britt RD (1995) Proximity of the manganese cluster of Photosystem II to the redoxactive tyrosine Y<sub>2</sub>. Proc Natl Acad Sci USA 92: 9545–9549
- Goodin DB, Yachandra VK, Britt RD, Sauer K and Klein MP (1984) The state of manganese in the photosynthetic apparatus.
  3. Light-induced changes in X-ray absorption (K-edge) energies of manganese in photosynthetic membranes. Biochim Biophys Acta 767: 209–216
- Guiles RD, Yachandra VK, McDermott AE, Cole JL, Dexheimer SL, Britt RD, Sauer K and Klein MP (1990a) The  $S_0$  state of Photosystem II induced by hydroxylamine: Differences between the structure of the manganese complex in the  $S_0$  and  $S_1$  states determined by X-ray absorption spectroscopy. Biochemistry 29: 486–496
- Guiles RD, Zimmermann JL, McDermott AE, Yachandra VK, Cole JL, Dexheimer SL, Britt RD, Wieghardt K, Bossek U, Sauer K and Klein MP (1990b) The S<sub>3</sub> state of Photosystem II: Differences between the structure of the manganese complex in the S<sub>2</sub> and S<sub>3</sub> states determined by X-ray absorption-spectroscopy. Biochemistry 29: 471–485
- Han K-C and Katoh S (1993) Different localization of two Ca<sup>2+</sup> in spinach oxygen-evolving Photosystem II membranes. Evidence for involvement of only one Ca<sup>2+</sup> in oxygen evolution. Plant Cell Physiol 34: 585–593
- Hasegawa K, Kusunoki M, Inoue Y and Ono T-A (1998) Simulation of S<sub>2</sub>-state multiline EPR signal in oriented Photosystem II membranes: Structural implications for the manganese cluster in an oxygen-evolving complex. Biochemistry 37: 9457–9465
- Hasegawa K, Ono T-A, Inoue Y and Kusunoki M (1999) Spinexchange interactions in the S<sub>2</sub>-state manganese tetramer in photosynthetic oxygen-evolving complex deduced from g=2 multiline EPR signal. Chem Phys Lett 300: 9–19
- Hillier W, Messinger J and Wydrzynski T (1998) Kinetic determination of the fast exchanging substrate water molecule in the  $S_3$ state of Photosystem II. Biochemistry 37: 16908–16914
- state of Photosystem II. Biochemistry 37: 16908–16914 Iuzzolino L, Dittmer J, Dörner W, Meyer-Klaucke W and Dau H (1998) X-ray absorption spectroscopy on layered Photosystem II membrane particles suggests manganese-centered oxidation of the oxygen-evolving complex for the S<sub>0</sub>-S<sub>1</sub>, S<sub>1</sub>-S<sub>2</sub>, and S<sub>2</sub>-S<sub>3</sub> transitions of the water oxidation cycle. Biochemistry 103

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

87

88

89

90

91

92

93

94

95

96

97

98

Jaklevic J, Kirby JA, Klein MP, Robertson AS, Brown GS and

Eisenberger P (1977) Fluorescence detection of EXAFS:

Sensitivity enhancement for dilute species and thin films. Solid

Joliot P (2003) Period-four oscillations of the flash-induced oxygen

Kamiya N and Shen JR (2003) Crystal structure of oxygen-evolv-

Katoh S, Satoh K, Ohno T, Chen J-R and Kashino Y (1987)

Numbers of calcium ions associated with oxygen evolving

Photosystem II preparations with different affinities. In: Biggins

J (ed) Progress in Photosynthesis Research, Vol I, pp 625-628.

Kirby JA, Robertson AS, Smith JP, Thompson AC, Cooper SR

and Klein MP (1981) State of manganese in the photosynthetic

apparatus. 1. Extended X-ray absorption fine structure stud-

ies on chloroplasts and di-µ-oxo-bridged dimanganese model

Kok B, Forbush B and McGloin M (1970) Cooperation of charges

Kretschmann H, Dekker JP, Saygin Ö and Witt HT (1988) An

agreement on the quaternary oscillation of ultraviolet absorption

changes accompanying the water splitting in isolated Photo-

system II complexes from the cyanobacterium Synechococcus

Kusunoki M, Takano T, Ono T, Noguchi T, Yamaguchi Y,

Oyanagi H and Inoue Y (1995) Advanced EXAFS studies of

the  $S_1$  state manganese cluster in plant Photosystem II. In:

Mathis P (ed) Photosynthesis: From Light to Biosphere, Vol II,

Kuzek D and Pace RJ (2001) Probing the Mn oxidation states

in the OEC Insights from spectroscopic, computational and

pp 251-254. Kluwer Academic Publishers, Dordrecht

kinetic data. Biochim Biophys Acta 1503: 123-137

in photosynthetic oxygen evolution. I. A linear four step mecha-

ing Photosystem II from Thermosynechococcus vulcanus at 3.7

formation in photosynthesis. Photosynth Res 76: 65-72

Å resolution. Proc Natl Acad Sci USA 100: 98-103

Martinus Nijhoff Publishers, Dordrecht

compounds. J Am Chem Soc 103: 5529-5537

nism. Photochem Photobiol 11: 457-475

sp. Biochim Biophys Acta 932: 358-361

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

87

88

89

90

91

92

93

94

95

96

97

98

99

100

101

102

103

24

25

26

27

28

29

30

31

32

33 34

35

43

47

52

45 46

X-ray absorption spectroscopy. Biochemistry 33: 4923-4932 48 of the oxygen-evolving complex of Photosystem II. Does it

Liang W, Roelofs TA, Cinco RM, Rompel A, Latimer MJ, Yu WO, Sauer K, Klein MP and Yachandra VK (2000) Structural change of the Mn cluster during the  $S_2 \rightarrow S_3$  state transition

49 reflect the onset of water/substrate oxidation? Determination 50 by Mn X-ray absorption spectroscopy. J Am Chem Soc 122: 51 3399-3412

Limburg J, Szalai VA and Brudvig GW (1999) A mechanistic and 53 structural model for the formation and reactivity of a Mnv=O 54 species in photosynthetic water oxidation. J Chem Soc, Dalton 55 Trans 1353-1361 56

- Lindberg K and Andréasson LE (1996) A one-site, two-state model for the binding of anions in Photosystem II. Biochemistry 35: 14259-14267
- Lindberg K, Wydrzynski T, Vänngård T and Andréasson L-E (1990) Slow release of chloride from 36Cl-labeled Photosystem II membranes. FEBS Lett 264: 153-155
- MacLachlan DJ, Hallahan BJ, Ruffle SV, Nugent JHA, Evans MCW, Strange RW and Hasnain SS (1992) An EXAFS study of the manganese oxygen-evolving complex in purified Photosystem II membrane fractions. The S<sub>1</sub> and S<sub>2</sub> states. Biochem J 285: 569-576
- MacLachlan DJ, Nugent JHA, Bratt PJ and Evans MCW (1994) The effects of calcium depletion on the O2-evolving complex in spinach PS II: The  $S_1^*$ ,  $S_2^*$  and  $S_3^*$  states and the role of the 17 kDa and 23 kDa extrinsic polypeptides. Biochim Biophys Acta 1186: 186-200
- McDermott AE, Yachandra VK, Guiles RD, Cole JL, Dexheimer SL, Britt RD, Sauer K and Klein MP (1988) Characterization of the manganese O2-evolving complex and the iron-quinone acceptor complex in Photosystem II from a thermophilic cyanobacterium by electron paramagnetic resonance and X-ray absorption spectroscopy. Biochemistry 27: 4021-4031
- Messinger J, Badger M and Wydrzynski T (1995) Detection of one slowly exchanging substrate water molecule in the S<sub>3</sub> state of Photosystem II. Proc Natl Acad Sci USA 92: 3209-3213
- Messinger J, Nugent JHA and Evans MCW (1997a) Detection of an EPR multiline signal for the S<sub>0</sub><sup>\*</sup> state in Photosystem II. Biochemistry 36: 11055-11060
- Messinger J, Robblee J, Yu WO, Sauer K, Yachandra VK and Klein MP (1997b) The  $S_0$  state of the oxygen-evolving complex in Photosystem II is paramagnetic: Detection of an EPR multiline signal. J Am Chem Soc 119: 11349-11350
- Messinger J, Robblee JH, Bergmann U, Fernandez C, Glatzel P, Visser H, Cinco RM, McFarlane KL, Bellacchio E, Pizarro SA, Cramer SP, Sauer K, Klein MP and Yachandra VK (2001) Absence of Mn-centered oxidation in the  $S_2 \rightarrow S_3$  transition: Implications for the mechanism of photosynthetic water oxidation. J Am Chem Soc 123: 7804-7820
- Mukerji I, Andrews JC, DeRose VJ, Latimer MJ, Yachandra VK, Sauer K and Klein MP (1994) Orientation of the oxygen-evolving manganese complex in a Photosystem II membrane preparation: An X-ray absorption spectroscopy study. Biochemistry 33: 9712-9721
- Olesen K and Andréasson LE (2003) The function of the chloride ion in photosynthetic oxygen evolution. Biochemistry 42: 2025-2035
- Ono T-A and Inoue Y (1988) Discrete extraction of the Ca atom functional for O2 evolution in higher plant Photosystem II by a simple low pH treatment. FEBS Lett 227: 147-152

Ono T-A and Inoue Y (1989) Roles of Ca2+ in O2 evolution in higher plant Photosystem II: Effects of replacement of Ca2+ site by other cations. Arch Biochem Biophys 275: 440-448

Ono T-A, Zimmermann JL, Inoue Y and Rutherford AW (1986) EPR evidence for a modified S-state transition in chloridedepleted Photosystem II. Biochim. Biophys. Acta 851: 193-201

Ono T-A, Noguchi T, Inoue Y, Kusunoki M, Matsushita T and

104

24

1

37: 17112-17119

State Commun 23: 679-682

Lakshmi KV, Eaton SS, Eaton GR, Frank HA and Brudvig GW (1998) Analysis of dipolar and exchange interactions between manganese and tyrosine Z in the  $S_2Y_z$  state of acetate-inhibited

Photosystem II via EPR spectral simulations at X- and Q-bands. J Phys Chem B 102: 8327-8335 Latimer MJ, DeRose VJ, Mukerji I, Yachandra VK, Sauer K and

- Klein MP (1995) Evidence for the proximity of calcium to the 36 manganese cluster of Photosystem II: Determination by X-ray 37 absorption spectroscopy. Biochemistry 34: 10898-10909 38
- Latimer MJ, DeRose VJ, Yachandra VK, Sauer K and Klein MP 39 (1998) Structural effects of calcium depletion on the manganese 40 cluster of Photosystem II: Determination by X-ray absorption 41 spectroscopy. J Phys Chem B 102: 8257-8265
- Liang W, Latimer MJ, Dau H, Roelofs TA, Yachandra VK, Sauer 42 K and Klein MP (1994) Correlation between structure and magnetic spin state of the manganese cluster in the oxygen-evolving 44 complex of Photosystem II in the S2 state: Determination by

Oyanagi H (1992) X-ray detection of the period-four cycling

of the manganese cluster in photosynthetic water oxidizing

Ono T-A, Noguchi T, Inoue Y, Kusunoki M, Yamaguchi H and

Oyanagi H (1994) Study of the intermediate S-states for wa-

ter oxidation in the normal and Ca-depleted photosynthetic

oxygen-evolving enzyme by means of flash-induced X-ray

absorption near edge structure spectroscopy. Biochem Soc

Pecoraro VL (ed) (1992) Manganese Redox Enzymes. VCH

Pecoraro VL and Hsieh W-Y (2000) The use of model complexes

to elucidate the structure and function of manganese redox

enzymes. In: Sigel A and Sigel H (eds) Manganese and Its

Role in Biological Processes, pp 429-504. Marcel Dekker

Peloquin JM and Britt RD (2001) EPR/ENDOR characterization

Peloquin JM, Campbell KA and Britt RD (1998) 55Mn pulsed

ENDOR demonstrates that the Photosystem II 'split' EPR

signal arises from a magnetically-coupled mangano-tyrosyl

Peloquin JM, Campbell KA, Randall DW, Evanchik MA, Pecoraro

VL, Armstrong WH and Britt RD (2000) 55Mn ENDOR of the

S2-state multiline EPR signal of Photosystem II: Implications

on the structure of the tetranuclear cluster. J Am Chem Soc

Peng G, de Groot FMF, Hämäläinen K, Moore JA, Wang X,

Grush MM, Hastings JB, Siddons DP, Armstrong WH, Mullins

OC and Cramer SP (1994) High-resolution manganese X-ray

fluorescence spectroscopy. Oxidation-state and spin-state

Penner-Hahn JE (1998) Structural characterization of the Mn

Penner-Hahn JE, Fronko RM, Pecoraro VL, Yocum CF, Betts

SD and Bowlby NR (1990) Structural characterization of

the manganese sites in the photosynthetic oxygen-evolving

complex using X-ray absorption- spectroscopy. J Am Chem

Renger G (1997) Mechanistic and structural aspects of photosyn-

Riggs PJ, Yocum CF, Penner-Hahn JE and Mei R (1992) Reduced

gen-evolving complex. J Am Chem Soc 114: 10650-10651

Riggs-Gelasco PJ, Mei R, Yocum CF and Penner-Hahn JE (1996a)

Reduced derivatives of the Mn cluster in the oxygen-evolving

complex of Photosystem II: An EXAFS study. J Am Chem

Riggs-Gelasco PJ, Mei R, Ghanotakis DF, Yocum CF and Penner-

Hahn JE (1996b) X-ray absorption spectroscopy of calcium-

derivatives of the manganese cluster in the photosynthetic oxy-

thetic water oxidation. Physiol Plant 100: 828-841

site in the photosynthetic oxygen-evolving complex. Struct

of the physical and electronic structure of the OEC Mn cluster.

enzyme. Science 258: 1335–1337

Biochim Biophys Acta 1503: 96–111

complex. J Am Chem Soc 120: 6840-6841

sensitivity. J Am Chem Soc 116: 2914-2920

Trans 22: 331-335

Inc., New York

122: 10926-10942

Bond 90: 1-36

Soc 112: 2549-2557

Soc 118: 2387–2399

Publishers, New York

1

2

3

20 21

29

30 31

32 33

34 35

36

37 38

39 40

> 41 42

43 44

52

substituted derivatives of the oxygen-evolving complex of 45 Photosystem II. J Am Chem Soc 118: 2400-2410

Robblee JH, Cinco RM and Yachandra VK (2001) X-ray spec-46 troscopy-based structure of the Mn cluster and mechanism 47 of photosynthetic oxygen evolution. Biochim Biophys Acta 48 1503: 7-23

49 Robblee JH, Messinger J, Cinco RM, McFarlane KL, Fernandez 50 C, Pizarro SA, Sauer K and Yachandra VK (2002) The Mn 51 cluster in the S<sub>0</sub> state of the oxygen-evolving complex of

Photosystem II studied by EXAFS spectroscopy: Are there three di-µ-oxo-bridged Mn<sub>2</sub> moieties in the tetranuclear Mn complex? J Am Chem Soc 124: 7459-7471

- Roe AL, Schneider DJ, Mayer RJ, Pyrz JW, Widom J and Que L, Jr (1984) X-ray absorption spectroscopy of iron-tyrosinate proteins. J Am Chem Soc 106: 1676-1681
- Roelofs TA, Liang W, Latimer MJ, Cinco RM, Rompel A, Andrews JC, Sauer K, Yachandra VK and Klein M (1996) Oxidation states of the manganese cluster during the flash-induced S-state cycle of the photosynthetic oxygen-evolving complex. Proc Natl Acad Sci USA 93: 3335-3340
- Rutherford AW, Zimmermann J-L and Boussac A (1992) Oxygen evolution. In: Barber J (ed) The Photosystems: Structure, Function, and Molecular Biology, pp 179-229. Elsevier B. V., Amsterdam
- Sandusky PO and Yocum CF (1986) The chloride requirement for photosynthetic oxygen evolution: Factors affecting nucleophilic displacement of chloride from the oxygen-evolving complex. Biochim Biophys Acta 849: 85–93
- Sauer K, Yachandra VK, Britt RD and Klein MP (1992) The photosynthetic water oxidation complex studied by EPR and X-ray absorption spectroscopy. In: Pecoraro VL (ed) Manganese Redox Enzymes, pp 141–175. VCH Publishers, New York
- Sayers DE, Stern EA and Lytle F (1971) New technique for investigating noncrystalline structures. Fourier analysis of the extended X-ray-absorption fine structure. Phys Rev Lett 27: 1204-1207
- Saygin Ö and Witt HT (1987) Optical characterization of intermediates in the water-splitting enzyme system of photosynthesis-possible states and configurations of manganese and water. Biochim Biophys Acta 893: 452-469
- Schiller H, Dittmer J, Iuzzolino L, Dörner W, Meyer-Klaucke W, Solé VA, Nolting H-F and Dau H (1998) Structure and orientation of the oxygen-evolving manganese complex of green algae and higher plants investigated by X-ray absorption linear dichroism spectroscopy on oriented Photosystem II membrane particles. Biochemistry: 37: 7340-7350
- Sharp RR (1992) Proton NMR relaxation due to the photosynthetic oxygen-evolving center. In: Pecoraro VL (ed) Manganese Redox Enzymes, pp 177–196. VCH Publishers, New York
- Shen J-R, Satoh K and Katoh S (1988) Calcium content of oxygen-evolving Photosystem II preparations from higher plants. Effects of NaCl treatment. Biochim Biophys Acta 933: 358-364
- Shulman RG, Yafet Y, Eisenberger P and Blumberg WE (1976) Observation and interpretation of X-ray absorption edges in iron compounds and proteins. Proc Natl Acad Sci USA 73: 1384–1388
- Siegbahn PEM (2000) Theoretical models for the oxygen radical mechanism of water oxidation and of the water oxidizing complex of Photosystem II. Inorg Chem 39: 2923-2935
- Sivaraja M, Tso J and Dismukes GC (1989) A calcium-specific site influences the structure and activity of the manganese cluster responsible for photosynthetic water oxidation. Biochemistry 28: 9459-9464
- Smith PJ and Pace RJ (1996) Evidence for two forms of the 99 g=4.1 signal in the S<sub>2</sub> state of Photosystem II. Two magneti-100 cally isolated manganese dimers. Biochim Biophys Acta 1275: 101 213-220
- 102 Styring SA and Rutherford AW (1988) The microwave power saturation of  $S_{IIslow}$  varies with the redox state of the oxygen-evolving 103

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

87

88

89

90

91

92

93

94

95

96

97

98

104

PSII010

complex in Photosystem II. Biochemistry 27: 4915-4923

Tamura N and Cheniae G (1985) Effects of Photosystem-II extrinsic proteins on microstructure of the oxygen-evolving complex and its reactivity to water analogs. Biochim Biophys Acta 809: 245–259

Tsutsumi K, Nakamori H and Ichikawa K (1976) X-ray manganese Kβ emission spectra of manganese oxides and manganates. Phys Rev B 13: 929–933

- Visser H, Anxolabéhère-Mallart E, Bergman U, Glatzel P, Robblee JH, Cramer SP, Girerd J-J, Sauer K, Klein MP and Yachandra VK (2001) Mn K-edge XANES and Kβ XES studies of two Mn-oxo binuclear complexes. Investigation of three different oxidation states relevant to the oxygen-evolving complex of Photosystem II. J Am Chem Soc 123: 7031–7039
- Westre TE, Kennepohl P, DeWitt JG, Hedman B, Hodgson KO and Solomon EI (1997) A multiplet analysis of Fe K-edge 1s to 3d pre-edge features of iron complexes. J Am Chem Soc 119: 6297–6314
- Wieghardt K (1989) The active centers in manganese-containing metalloproteins and inorganic model complexes. Angew Chem Int Ed Engl 28: 1153–1172
- Wincencjusz H, van Gorkom HJ and Yocum CF (1997) The photosynthetic oxygen evolving complex requires chloride for its redox state S<sub>2</sub> to S<sub>3</sub> and S<sub>3</sub> to S<sub>0</sub> transitions but not for S<sub>0</sub> to S<sub>1</sub>
   or S<sub>1</sub> to S<sub>2</sub> transitions. Biochemistry 36: 3663–3670
- Wincencjusz H, Yocum CF and van Gorkom HJ (1998) S-state
   dependence of chloride binding affinities and exchange dynamics in the intact and polypeptide-depleted O<sub>2</sub> evolving complex of Photosystem II. Biochemistry 37: 8595–8604

 Yachandra VK (1995) X-ray absorption spectroscopy and applications in structural biology. In: Sauer K (ed) Methods of Enzymology, Vol 246, pp 638–675. Academic Press, San Diego

Yachandra VK (2002) Structure of the manganese complex in
Photosystem II: Insights from X-ray spectroscopy. Phil Trans
R Soc London B 357: 1347–1357

- Yachandra VK, Guiles RD, McDermott A, Britt RD, Dexheimer SL, Sauer K and Klein MP (1986) The state of manganese in the photosynthetic apparatus. 4. Structure of the manganese complex in Photosystem II studied using EXAFS spectroscopy. The S<sub>1</sub> state of the oxygen-evolving Photosystem II complex from spinach. Biochim Biophys Acta 850: 324–332
  Yachandra VK, Guiles RD, McDermott AE, Cole JL, Britt RD, 58
- Yachandra VK, Guiles RD, McDermott AE, Cole JL, Britt RD, Dexheimer SL, Sauer K and Klein MP (1987) Comparison of the structure of the manganese complex in the  $S_1$  and  $S_2$  states of the photosynthetic  $O_2$ -evolving complex: An X-ray absorption spectroscopy study. Biochemistry 26: 5974–5981
- Yachandra VK, DeRose VJ, Latimer MJ, Mukerji I, Sauer K and Klein MP (1993) Where plants make oxygen: A structural model for the photosynthetic oxygen-evolving manganese cluster. Science 260: 675–679
- Yachandra VK, Sauer K and Klein MP (1996) Manganese cluster in photosynthesis: Where plants oxidize water to dioxygen. Chem Rev 96: 2927–2950
- Yocum CF (1992) The calcium and chloride requirements for photosynthetic water oxidation. In: Pecoraro VL (ed) Manganese Redox Enzymes, pp 71–84. VCH Publishers, New York
- Zheng M and Dismukes GC (1996) Orbital configuration of the valence electrons, ligand field symmetry, and manganese oxidation states of the photosynthetic water oxidizing complex: Analysis of the  $S_2$  state multiline EPR signals. Inorg Chem 35: 3307–3319
- Zimmermann JL and Rutherford AW (1986) Electron paramagnetic resonance properties of the  $S_2$  state of the oxygen-evolving complex of Photosystem II. Biochemistry 25: 4609–4615
- Zouni A, Witt HT, Kern J, Fromme P, Krauss N, Saenger W and Orth P (2001) Crystal structure of Photosystem II from *Synechococcus elongatus* at 3.8 Å resolution. Nature 409: 739–743