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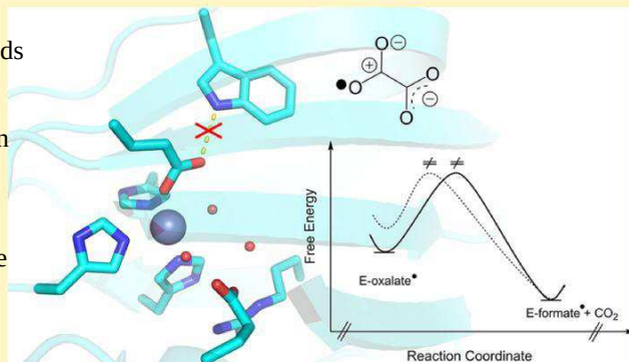
# Second-Shell Hydrogen Bond Impacts Transition-State Structure in *Bacillus subtilis* Oxalate Decarboxylase

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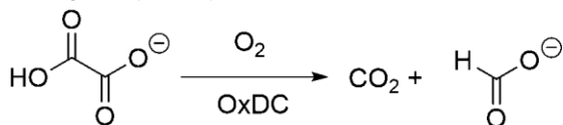
**ABSTRACT:** There is considerable interest in how “second-shell” interactions between protein side chains and metal ligands might modulate Mn(II) ion redox properties and reactivity in metalloenzymes. One such Mn-dependent enzyme is oxalate decarboxylase (OxDC), which catalyzes the disproportionation of oxalate monoanion into formate and CO<sub>2</sub>. Electron paramagnetic resonance (EPR) studies have shown that a mononuclear Mn(III) ion is formed in OxDC during catalytic turnover and that the removal of a hydrogen bond between one of the metal ligands (Glu101) and a conserved, second-shell tryptophan residue (Trp132) gives rise to altered zero-field splitting parameters for the catalytically important Mn(II) ion. We now report heavy-atom kinetic isotope effect measurements on the W132F OxDC variant, which test the hypothesis that the Glu101/Trp132 hydrogen bond modulates the stability of the Mn(III) ion during catalytic turnover. Our results suggest that removing the Glu101/Trp132 hydrogen bond increases the energy of the oxalate radical intermediate from which decarboxylation takes place. This finding is consistent with a model in which the Glu101/Trp132 hydrogen bond in WT OxDC modulates the redox properties of the Mn(II) ion.



## O

oxalate decarboxylase (OxDC),<sup>1</sup> which catalyzes the disproportionation of oxalate monoanion into formate and CO<sub>2</sub> (Scheme 1), is one of only five enzymes that take advantage of redox changes in a mononuclear manganese center to mediate catalysis.<sup>3</sup>

Scheme 1. Reaction Catalyzed by Oxalate Decarboxylase (OxDC)<sup>a</sup>



<sup>a</sup>Although the overall transformation is a disproportionation, catalytic activity requires dioxygen.

OxDC activity is solely Mn-dependent,<sup>4,5</sup> and recent electron paramagnetic resonance (EPR) studies have demonstrated the existence of enzyme-bound Mn(III) during catalytic turnover.<sup>6,7</sup> Current mechanistic models therefore assume that Mn(III) oxidizes oxalate to form a radical anion intermediate in which the barrier to decarboxylation is significantly lowered (Scheme 2).<sup>8</sup> The resulting Mn-bound radical anion intermediate then acquires an electron and a proton to regenerate Mn(III) and produce formate.

Although OxDC possesses two manganese-binding sites located in two cupin domains (Figure 1), substantial evidence

exists to support the hypothesis that catalysis takes place only at the metal center located in the N-terminal domain of the enzyme.<sup>9,10</sup> In addition, the X-ray crystal structure of an Co(II)-substituted OxDC loop variant shows that oxalate coordinates the catalytically important manganese center in a monodentate fashion.<sup>11</sup> Maximal enzyme activity, however, also requires the presence of Mn(II) in the metal-binding site of the

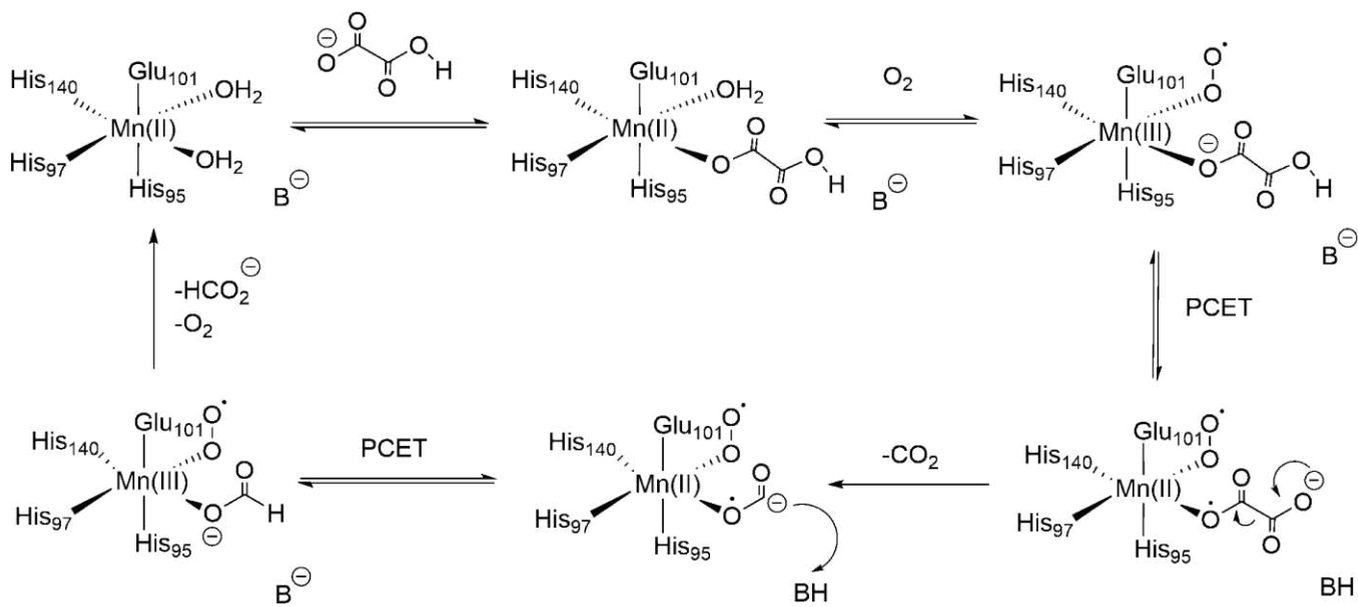
C-terminal cupin domain.<sup>4</sup>

The apparent differential activity of the two Mn(II) centers in the enzyme is not understood.<sup>12</sup> It is possible, however, that “second-shell” interactions between protein side chains and metal ligands might modulate Mn(II) ion reactivity (Figure 1).<sup>13</sup> For example, the putative hydrogen bond between the Trp132 side chain and Glu101, which coordinates the metal ion, in the N-terminal Mn-binding site is replaced by an alternate hydrogen bond between the side chains of Gln232 and Glu280 in the C-terminal site (*Bacillus subtilis* numbering).

Removing the Glu101/Trp132 hydrogen bond by site-specific replacement of Trp132 by a phenylalanine residue leads to altered zero-field splitting parameters for the N-terminal Mn(II) ion as a result of altered charge density on the

Glu101 carboxylate.<sup>15</sup> This observation has led to the proposal

Scheme 2. Current Model for the Catalytic Mechanism Used by OxDC<sup>a</sup>



<sup>a</sup>Although dioxygen is assumed to be the reagent that oxidizes Mn(II) after substrate binding, the exact nature of the oxidizing agent remains to be determined. In addition, the proton-coupled electron transfers (PCETs) may proceed via independent electron-transfer and protonation steps. We assume that decarboxylation is irreversible. The identity of the general base (B) that removes the proton from the substrate monoanion is not yet determined.

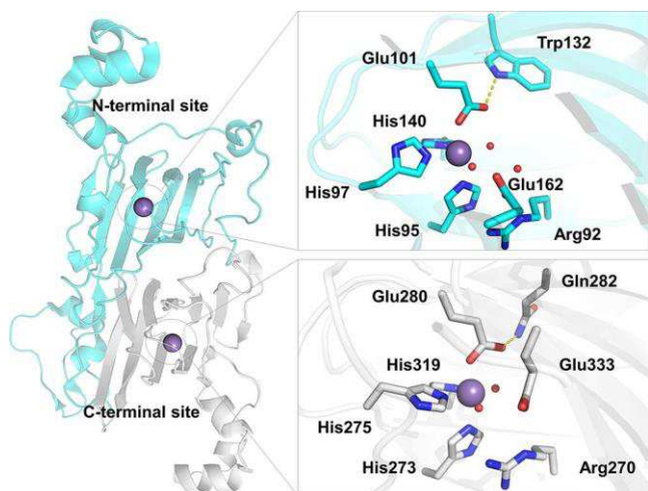


Figure 1. Mn(II)-binding sites present in the monomer of WT OxDC (PDB 1UW8). The enzyme is composed of two cupin domains, which are rendered in cyan (N-terminal) and gray (C-terminal). Active-site residue numbers are those for the OxDC present in *Bacillus subtilis*. Metal ions and water molecules are rendered as purple and red spheres, respectively, and the yellow dashed lines show second-shell hydrogen-bonding interactions.

62 that Mn(III) should be more stable in the W132F OxDC  
 63 variant than in the WT enzyme. We now report heavy-atom  
 64 kinetic isotope effect measurements that seek to test this idea.  
 65 Our experimental measurements are consistent with a model in  
 66 which removing the Glu101/Trp132 hydrogen bond results in  
 67 altered bond polarization in the oxalate radical intermediate.  
 68 This finding, in turn, suggests that the oxalate radical  
 69 intermediate in the W132F OxDC variant is higher in energy  
 70 than the cognate species formed in WT OxDC during turnover,  
 71 consistent with the hypothesis that the Glu101/Trp132

hydrogen bond modulates the redox properties of the Mn(II) 72  
 ion. 73

## EXPERIMENTAL PROCEDURES 74

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106 **Steady-State Enzyme Assays.** OxDC activity was  
107 measured using a standard end-point, coupled assay with  
108 formate dehydrogenase (FDH) in which NADH production  
109 was monitored at 340 nm.<sup>17</sup> Thus OxDC (5.5  $\mu$ M) was  
110 incubated for 1 min with varying amounts of potassium oxalate  
111 (0–200 mM, pH 4.2) in 50 mM acetate buffer, pH 4.2,  
112 containing 0.5 mM o-phenylenediamine and 0.2% TritonX.  
113 After quenching the reaction by the addition of 0.1 M NaOH,  
114 the resulting solution was incubated overnight at 37 °C in the  
115 presence of FDH and NAD<sup>+</sup> (1.5 mM final concentration).  
116 The absorbance at 340 nm was then converted to NADH  
117 concentration using a standard curve. Measurements were  
118 made at specific substrate and enzyme concentrations in  
119 triplicate. All data were processed using GraphPad Prism and  
120 analyzed by standard methods to obtain the values of V and V/  
121 K.<sup>18</sup>

122 **Kinetic Isotope Effect Nomenclature.** In this paper,  
123 <sup>13</sup>(V/K) represents the ratio of V/K for the <sup>12</sup>C-containing  
124 substrate relative to the <sup>13</sup>C-containing substrate.<sup>19</sup> In a similar  
125 manner, <sup>18</sup>(V/K) represents the ratio of V/K for the <sup>16</sup>O-  
126 containing substrate relative to the <sup>18</sup>O-containing substrate.  
127 **<sup>13</sup>C and <sup>18</sup>O Kinetic Isotope Effect Measurements.** As  
128 described in detail elsewhere,<sup>17</sup> the internal competition  
129 method<sup>20</sup> was used to determine the primary <sup>13</sup>C and  
130 secondary <sup>18</sup>O isotope effects on the decarboxylation reaction  
131 catalyzed by the W132F OxDC variant. All experiments  
132 employed oxalate in which the heavy-atom isotopes were at  
133 natural abundance. Partial and total conversion reactions at 22  
134 °C were performed at either pH 4.2 or pH 5.7 using 100 mM  
135 1,4-bis(2-hydroxyethyl)-piperazine (BHEP) or 100 mM  
136 piperazine, respectively. All buffer solutions contained 0.5  
137 mM o-phenylenediamine and were sparged with N<sub>2</sub>(g) for 1 h  
138 before use to remove adventitious CO<sub>2</sub> in solution. Similarly, all  
139 gases were passed over Ascarite to remove CO<sub>2</sub>(g) prior to use.  
140 Solutions of 40 mM potassium oxalate dissolved in the  
141 appropriate buffer, which had been sparged with O<sub>2</sub>(g) for 1  
142 h prior to use, were placed in a sealed flask. Reactions were  
143 initiated by the addition of enzyme in N<sub>2</sub>-saturated buffer  
144 and

145 subsequently quenched by the addition of 500 mM Tris-Cl, pH  
146 7.5. Incubation times were varied from 37 min to 3 h so as to  
147 obtain mixtures in which a different fraction of reaction  
148 had  
149 taken place. Complete oxalate consumption required incubation  
150 with the W132F OxDC variant at 22 °C for 14 h in either  
151 of the two buffers. CO<sub>2</sub> produced during the reaction was  
152 collected and purified through a vacuum line, and the isotopic  
153 composition was determined using an isotope ratio mass  
154 spectrometer (IRMS). After quenching, the solution was passed  
155 through an Amicon ultrafiltration system to remove enzyme,  
156 and an aliquot (50  $\mu$ L) was taken to determine the fraction of  
157 conversion, f, using an oxalate assay kit (Trinity Biotech, NY).  
158 In addition, formate produced in the reaction was measured  
159 using the standard FDH-based assay outlined above.

160 The isotopic composition of residual oxalate and formate  
161 produced in the reaction was also determined in these studies.  
162 Thus formate and oxalate were separated by anion-exchange  
163 chromatography (Bio-Rad AG-1 resin) using dilute H<sub>2</sub>SO<sub>4</sub>, pH  
164 2.7, as eluent. Fractions that contained either oxalate or formate  
165 were pooled, and the pH of these solutions was adjusted to  
166 neutral pH using 0.1 N NaOH before the volume was reduced.  
167 After sparging with N<sub>2</sub>(g) for 30 min, water was removed  
168 completely from the resulting solutions by heating overnight at  
169 70 °C under high vacuum. DMSO (2 mL) containing I<sub>2</sub> (250–  
170 400 mg) was then used to oxidize the dried samples of oxalate

or formate (45 min) with the isotopic composition of the CO<sub>2</sub> 169  
produced in the reaction being measured by IRMS. Control 170  
experiments were performed at pH 4.2 using 2% H<sub>2</sub><sup>18</sup>O to 171  
examine whether <sup>18</sup>O/<sup>16</sup>O exchange took place between solvent  
172 water and either the substrate or products under the reaction 173  
conditions. The observed isotope effects were analyzed using 174  
procedures that our group has detailed elsewhere.<sup>17,21</sup> 175

## RESULTS AND DISCUSSION

176 Standard measurements of formate production at pH 4.2 gave 177  
steady-state kinetic parameters for the W132F OxDC variant 178  
and indicated that the removal of the Glu101/Trp132 179  
hydrogen bond had little impact on the turnover number of 180  
the enzyme (Table 1). On the contrary, the oxalate K<sub>M</sub> was 181 11

Table 1. Steady-State Kinetic Parameters for the Decarboxylation Reaction Catalyzed by Recombinant, WT OxDC, and the C-Terminally His<sub>6</sub>-Tagged W132F OxDC Variant at pH 4.2 and 25 °C

enzyme	K <sub>M</sub> (mM)	k <sub>cat</sub> (s <sup>-1</sup> )	k <sub>cat</sub> /K <sub>M</sub> /Mn (M <sup>-1</sup> s <sup>-1</sup> )	Mn content
WT OxDC	4.0 ± 0.5	60 ± 2	10 000 ± 1000	1.6
W132F OxDC	27 ± 5	60 ± 4	1300 ± 300	1.8

182 increased approximately seven-fold, leading to a decrease in 182  
k<sub>cat</sub>/K<sub>M</sub>/Mn. To obtain more detail about the effects of this 183  
mutation on catalysis, we determined the primary <sup>13</sup>C and 184  
secondary <sup>18</sup>O isotope effects (IEs) for the W132F-catalyzed 185  
decarboxylation by internal competition using oxalate in which 186  
the heavy-atom isotopomers were at natural abundance (Table 187 12  
2). As discussed elsewhere,<sup>20</sup> these measurements report on 188 12  
isotopically sensitive steps in the catalytic mechanism up to, and 189  
including, the first irreversible step, which we assume to be CO<sub>2</sub> 190  
formation. Given that CO<sub>2</sub> hydration and isotope exchange 191  
might have impacted the <sup>18</sup>O IEs at pH 5.7 differently from 192  
those at pH 4.2, control experiments using 2% H<sub>2</sub><sup>18</sup>O in the 193  
solvent were performed. These studies showed that our results 194  
were not severely affected by <sup>18</sup>O/<sup>16</sup>O isotope exchange 195  
between substrate and water, especially when the reaction 196  
was carried at pH 4.2. On the contrary, in analyzing the 197  
observed isotope effects, the <sup>13</sup>C IEs are viewed as providing 198  
more reliable and accurate values. <sup>13</sup>(V/K) on CO<sub>2</sub> is 1.3% at 199  
pH 5.7 in the W132F-catalyzed decarboxylation, which is 200  
substantially larger than that observed for WT OxDC (0.8%), 201  
and <sup>13</sup>(V/K) on formate is 3.6% as compared with 1.9% 202  
measured for WT OxDC. Given that <sup>13</sup>(V/K) is 3–5% for 203  
decarboxylases in which the loss of CO<sub>2</sub> is rate-limiting,<sup>22</sup> our 204  
values indicate that the observed IEs arise from two (or more) 205  
steps that are sensitive to isotopic substitution. As in previous 206  
work from our laboratory,<sup>17,21</sup> we interpreted the observed IEs 207  
using a minimal kinetic model (Scheme 3) in which an initial 208  
proton-coupled electron transfer (PCET) takes place after 209  
oxalate binding to give an intermediate that then undergoes 210  
irreversible decarboxylation. The discrepancy in the two <sup>13</sup>(V/211  
K) values is therefore associated with different sensitivities to 212  
isotopic substitution of the two carbon atoms in the step(s) 213

preceding C–C bond cleavage.<sup>17</sup>

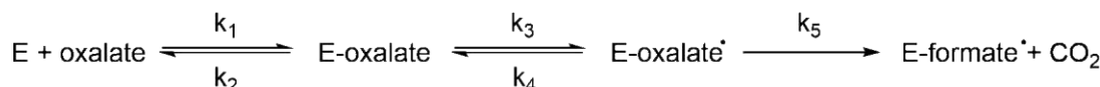
214 Analysis of the Observed <sup>13</sup>(V/K) and <sup>18</sup>(V/K) Isotope 215  
Effects. To understand how the commitments to catalysis 216  
might have been altered by the removal of the Glu101/Trp132 217  
hydrogen bond, we undertook a quantitative analysis of the 218

Table 2.  $^{13}\text{C}$  and  $^{18}\text{O}$  Isotope Effects on the Reactions Catalyzed by Recombinant, C-Terminally His<sub>6</sub>-Tagged WT OxDC, and the W132F and T165V OxDC Variants at 22 °C<sup>a</sup>

OxDC variant	pH	$^{13}\text{C}$ (V/K)		$^{18}\text{O}$ (V/K)		citation
		CO <sub>2</sub>	HCO <sub>2</sub> <sup>-</sup>	CO <sub>2</sub>	HCO <sub>2</sub> <sup>-</sup>	
WT	4.2	1.005 ± 0.001	1.015 ± 0.001	0.998 ± 0.002	1.011 ± 0.002	17
W132F	4.2	1.010 ± 0.001	1.024 ± 0.001	0.993 ± 0.001	1.009 ± 0.001	
T165V	4.2	0.998 ± 0.001	1.008 ± 0.001	0.991 ± 0.001	1.004 ± 0.001	15
WT	5.7	1.008 ± 0.001	1.019 ± 0.001	0.993 ± 0.002	1.010 ± 0.001	17
W132F	5.7	1.013 ± 0.001	1.036 ± 0.001	0.989 ± 0.002	1.014 ± 0.002	
T165V	5.7	0.997 ± 0.001	1.009 ± 0.001	0.984 ± 0.001	1.006 ± 0.001	15

<sup>a</sup>Data for WT OxDC and the T165V OxDC variant have been previously published<sup>15,17</sup> and are included here for ease of comparison.

Scheme 3. Minimal Kinetic Model for the OxDC-Catalyzed Reaction Used in the Quantitative Interpretation of the  $^{13}\text{C}$  and  $^{18}\text{O}$  Isotope Effects<sup>15,17,21</sup>



219 data using our minimal kinetic model and the following  
220 equation

$$x \left( \frac{V}{K} \right) = \frac{x K_{\text{eq}3} \frac{x}{k_5} \frac{x}{k_3} + \left( \frac{k_5}{k_4} \right) + \frac{k_3 k_5}{k_2 k_4}}{1 + \left( \frac{k_5}{k_4} \right) \left( 1 + \frac{k_1}{k_2} \right)} \quad (1)$$

221 where  $x(V/K)$  is the ratio of V/K for the lighter isotopomer in  
222 the enzyme-catalyzed reaction relative to that for the heavy  
223 isotopomer ( $x = 13$  or  $18$ ).  $k_3$ ,  $k_4$ , and  $k_5$  are rate constants in  
224 the minimal model that are assumed to be sensitive to isotopic  
225 substitution, and  ${}^x k_3$  and  ${}^x k_5$  are the isotope effects on the  
226 formation of oxalate radical anion and decarboxylation,  
227 respectively. Finally,  ${}^{13}K_{\text{eq}3}$  and  ${}^{18}K_{\text{eq}3}$  are  $^{13}\text{C}$  and  $^{18}\text{O}$   
228 equilibrium isotope effects (EIEs) on the putative oxalate  
229 radical anion intermediate. The derivation of this equation and  
230 a full discussion of many of the assumptions used below in  
231 analyzing the IE data for the reaction catalyzed by the W132F  
232 OxDC variant have been discussed elsewhere.<sup>17</sup> For example,  
233 we assume that oxalate monoanion remains the substrate when  
234 the Glu101/Trp132 hydrogen bond is absent in the active site  
235 and that the catalytic mechanism is unaffected by the  
236 introduction of a phenylalanine residue. As a result, and in a  
237 similar manner to WT OxDC, the initial rate of the T132F-  
238 catalyzed reaction is slower at pH 5.7 because of the increased  
239 concentration of the oxalate dianion in solution.<sup>17</sup>

240 **CO<sub>2</sub>-Based Analysis.** As discussed in detail elsewhere,<sup>17</sup> in  
241 analyzing the  $^{13}\text{C}$ (V/K) IE value on CO<sub>2</sub> at pH 5.7, we assume  
242 that  $k_3/k_2$  can be ignored, that  ${}^{13}k_5$  on CO<sub>2</sub> during  
243 decarboxylation is 1.04 (which is an average value for this  
244 reaction<sup>17,22</sup>), and that  ${}^{13}K_{\text{eq}3}$  and  ${}^{13}k_3$  are both unity; that is,  
245 proton removal from the carboxylic acid is assumed to proceed  
246 with a negligible  $^{13}\text{C}$  isotope effect.<sup>23</sup> We therefore obtain the  
247 following expression

$${}^{13} \left( \frac{V}{K} \right) = \frac{1.04 + \left( \frac{k_5}{k_4} \right)}{1 + \left( \frac{k_5}{k_4} \right)} = 1.013 \quad (2)$$

249 Solving this equation yields a value of  $k_5/k_4 = 2.08$  (Table 3),  
250 which differs from the value of  $k_5/k_4 = 4.00$  determined for WT  
251 OxDC under the same conditions. The reduction in the  
252 commitment factor for the W132F OxDC variant is associated  
253

Table 3. Commitment Factors and C–O Bond Orders (See Text) in the Decarboxylation Transition States for the Reactions Catalyzed by WT OxDC and the W132F and T165V OxDC Variants<sup>a</sup>

enzyme	$k_3/k_2$	$k_5/k_4$	${}^{13}K_{\text{eq}3}$	${}^{18}k_3$	C–O bond order	citation
WT OxDC	0.75	4.00	1.021	1.016	1.15	17
W132F OxDC	0.44	2.08	1.039	1.012	0.97	
T165V OxDC	3.29	12.33	1.013	1.004	1.26	15

<sup>a</sup>Data for WT OxDC and the T165V OxDC variant have been previously published and are included here for ease of comparison.

with an increase in the  $k_4$  rate constant because, for reasons that  
254 are discussed below, the magnitude of  $k_5$  is increased compared  
255 with the cognate rate constants in the reaction catalyzed by WT  
256 OxDC (Table 3).

Given that the carboxylic acid at the end of the substrate that  
257 becomes CO<sub>2</sub> is protonated in our mechanistic model, the  
258  ${}^{18}\text{O}$ (V/K) value on CO<sub>2</sub> at pH 5.7 must be multiplied by 0.98.<sup>17</sup>  
259 In addition,  ${}^{18}K_{\text{eq}3}$  is assigned a value of 1.02 as a result of  
260 proton removal in the step(s), leading to the formation of the  
261 oxalate radical anion.<sup>17</sup> These assumptions then yield the  
262 following equation

$${}^{18} \left( \frac{V}{K} \right) = \frac{0.98[(1.02)(0.9835) + {}^{18}k_3(2.077)]}{1 + 2.077} = 0.989 \quad (3)$$

263 making the additional assumptions that (i)  $k_3/k_2$  is small  
264 enough to ignore at pH 5.7 and (ii) the IE on decarboxylation  
265 is midway between an estimated  ${}^{18}K_{\text{eq}5}$  (0.967) and unity.<sup>17</sup> As  
266 a result, we obtain a value of 1.0121 for  ${}^{18}k_3$ , which is a  
267 reasonable value for the deprotonation step, albeit smaller than  
268 the value of 1.0159 computed for WT OxDC at pH 5.7 (Table  
271 3).

**Formate-Based Analysis.** In the case of  $^{13}\text{C}$ (V/K) on  
272 formate at 5.7, we can write the following equation

$${}^{13} \left( \frac{V}{K} \right) = \frac{(1.03) {}^{13}K_{\text{eq}3} + (2.08) {}^{13}K_{\text{eq}3+1}}{1 + 2.08} = 1.036 \quad (4)$$

273 where we again ignore  $k_3/k_2$  and assume that (i)  $k_5/k_4$  has a  
274 value of 2.08 as derived in the CO<sub>2</sub> analysis (see above), (ii) 277

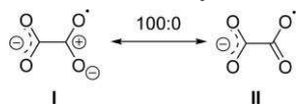
<sup>13</sup>k<sub>3</sub> lies midway between <sup>13</sup>K<sub>eq3</sub> and unity,<sup>17</sup> and (iii) <sup>13</sup>k<sub>5</sub> for decarboxylation is 1.03. The latter value reflects the fact that the change in bond order at this carbon is less than that at the carbon, which becomes CO<sub>2</sub>. Solving the resulting equation gives a value of 1.039 for <sup>13</sup>K<sub>eq3</sub> and a C–O bond order in the transition state of 0.88 based on calculations of the dependence of fractionation factor upon C–O bond order that we have previously reported.<sup>17</sup> Of course, it is possible that assumption (ii), which was used in our original study on WT OxDC to simplify data analysis,<sup>17</sup> may be incorrect for the reaction catalyzed by the W132F OxDC variant. We therefore examined the effect of setting <sup>13</sup>k<sub>3</sub> to <sup>13</sup>K<sub>eq3</sub> in our analysis. Setting <sup>13</sup>k<sub>3</sub> to <sup>13</sup>K<sub>eq3</sub> and solving gives <sup>13</sup>K<sub>eq3</sub> = 1.026 and a corresponding C–O bond order of 1.09, which is still smaller than the bond order computed for WT OxDC (1.16).<sup>17</sup> Moreover, because we have shown the chemical bonding in oxalate and the oxalate radical anion to differ on the basis of high-level ab initio calculations,<sup>8</sup> and due to the fact that a larger IE is observed for the W132F OxDC variant, <sup>13</sup>k<sub>3</sub> is unlikely to be unity. Thus our conclusions are unaffected, and, for the sake of comparison, we compare the IE data for WT OxDC and the W132F OxDC variant using an identical set of assumptions.

An alternate estimate of the C–O bond order could be obtained by determining the value of <sup>18</sup>K<sub>eq3</sub> using the following equation

$$18\left(\frac{V}{K}\right) = \frac{(1.003)^{18}K_{eq3} + (2.08)^{\frac{18K_{eq3}+1}{2}}}{1 + 2.08} = 1.014 \quad (5)$$

where we assume that (i) <sup>18</sup>k<sub>3</sub> lies midway between <sup>18</sup>K<sub>eq3</sub> and unity, (ii) <sup>18</sup>k<sub>5</sub> = 1.003 due to the bond angle change in the formate radical intermediate,<sup>17</sup> and (iii) k<sub>3</sub>/k<sub>2</sub> is small enough to be ignored. Solving this equation gives a value of 1.02 for <sup>18</sup>K<sub>eq3</sub>, which corresponds to a C–O bond order of 1.054. Taking the average of the estimates obtained from the <sup>13</sup>K<sub>eq3</sub> and <sup>18</sup>K<sub>eq3</sub> values then yields an estimate of 0.97 for the C–O bond order, considerably smaller than 1.15 estimated for the transition state in the reaction catalyzed by WT OxDC (Table 3).

We therefore conclude that the oxalate radical anion in the W132F-catalyzed reaction can be represented solely by resonance form I (see below) in which a full positive charge is located on the carbon that is finally converted into formate.



The C–O bonds in this Mn-bound carboxylate are therefore considerably more polarized than the cognate bonds in the radical anion intermediate formed during the reaction catalyzed by WT OxDC (1.15).<sup>17</sup> As a result, the carbon atom is more electron-deficient, which will promote C–C bond cleavage and hence increase the magnitude of the k<sub>5</sub> rate constant. Thus the increase in k<sub>4</sub> must be greater to give a lower k<sub>5</sub>/k<sub>4</sub> ratio in the W132F-catalyzed transformation; therefore, we propose that the removal of the Glu101/Trp132 hydrogen bond raises the energy of the putative oxalate-based radical intermediate (Scheme 2).

At pH 4.2, the smaller observed values of the heavy atom IEs suggest that k<sub>3</sub>/k<sub>2</sub> is no longer negligible. We therefore repeated our calculations to obtain an estimate of this commitment factor. For the end of the substrate that becomes CO<sub>2</sub>, substitution of the observed <sup>13</sup>(V/K) value at pH 4.2 gives the following expression

$$13\left(\frac{V}{K}\right) = \frac{(1.04) + (2.077)\left(1 + \frac{k_5}{k_4}\right)}{1 + (2.077)\left(1 + \frac{k_5}{k_4}\right)} = 1.01 \quad (6)$$

where <sup>13</sup>k<sub>5</sub> on CO<sub>2</sub> for the decarboxylation step is assumed to be 1.04, (ii) k<sub>5</sub>/k<sub>4</sub> has the same value as at pH 5.7, and (iii) <sup>13</sup>K<sub>eq3</sub> and <sup>13</sup>k<sub>3</sub> are both unity.<sup>17</sup> Solving this equation gives k<sub>3</sub>/k<sub>2</sub> = 0.44, which is smaller than that observed for WT OxDC (0.75) (Table 3). Given that K<sub>M</sub> is larger for the W132F OxDC variant, an increase in the k<sub>2</sub> value might be expected, although the k<sub>3</sub> rate constant may also be smaller given that the oxalate-based radical intermediate has a higher energy (see above). With estimates for <sup>13</sup>k<sub>3</sub>, k<sub>3</sub>/k<sub>2</sub>, k<sub>5</sub>/k<sub>4</sub>, and <sup>13</sup>K<sub>eq3</sub> in hand, we calculated the value of <sup>13</sup>(V/K) on formate at pH 4.2 to evaluate the validity of using our minimal kinetic model to interpret the observed IEs. Substitution of the <sup>13</sup>k<sub>3</sub>, k<sub>3</sub>/k<sub>2</sub>, k<sub>5</sub>/k<sub>4</sub>, and <sup>13</sup>K<sub>eq3</sub> values into the master equation gave an estimate of 1.027 for <sup>13</sup>(V/K), which is in good agreement with that observed experimentally (1.026) (Table 2) and confirms that the catalytic mechanism is unchanged by the removal of the Glu101/Trp132 hydrogen bond.

**Second-Shell Interactions in OxDC and Their Impact on the Energy of the Oxalate Radical Anion Intermediate and the Barrier to Decarboxylation.** The role of noncovalent interactions between residue side chains in the local protein environment and the ligands that coordinate metal ions is of considerable general interest, especially for the development of transition-metal complexes with novel catalytic activities. In the case of OxDC, questions remain about how Mn(III) is generated during turnover and how its intrinsic activity as an oxidizing agent is controlled by the metal ligands or oxalate binding. Given that Trp132 (*Bacillus subtilis* numbering) is conserved in all known oxalate decarboxylases and forms a hydrogen bond with the metal ligand Glu101, we speculated that removing this interaction would impact the energy difference between Mn(III)-bound oxalate and the Mn(II)-bound oxalate radical anion intermediate. Importantly, the X-ray crystal structure of the Co-substituted W132F OxDC variant (PDB: 4MET) shows that removing the Glu101/Trp132 hydrogen bond has no impact on the overall fold of the enzyme, and the aromatic rings of Phe132 and Trp132 are positioned identically in the N-terminal Mn(II)-binding site (Figure 2).

On the contrary, there is a slight alteration in the position of the Glu-162 side chain and a rotation of the imidazole ring in the metal ligand His-97. This modification of metal coordination is likely associated with substituting Co(II) for Mn(II),<sup>14</sup> however, and so we assume that the geometry of the Mn/ligand interactions is unchanged by the removal of the Glu-101/Trp-132 hydrogen bond. We also note that removing the Glu101/Trp132 hydrogen bond perturbs the number and locations of active-site water molecules in the W132F OxDC variant compared with WT OxDC (Figure 2). Thus the Mn-bound water oxygens in the W132F variant occupy equivalent positions to the oxygen atoms of oxalate and Mn-bound water seen in the X-ray crystal structure of an OxDC variant in which Glu162 is deleted (PDB 5HI0).<sup>11</sup>

Despite these small alterations in active-site geometry, however, the absence of the second-shell Glu101/Trp132 hydrogen bond does impact the partition ratios k<sub>5</sub>/k<sub>4</sub> and k<sub>3</sub>/k<sub>2</sub> relative to those determined for WT OxDC (Table 3). Our working hypothesis is that removing the hydrogen bond

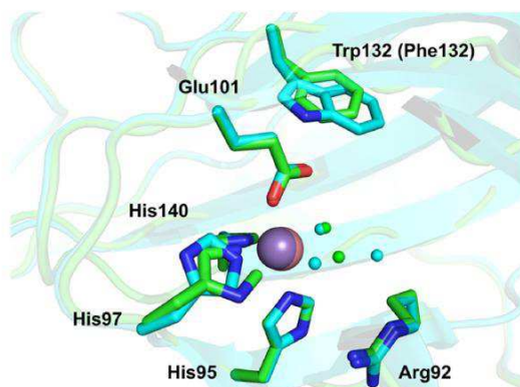


Figure 2. Superimposition of the X-ray crystal structures of WT OxDC (PDB 1UW8) and the Co-substituted W132F OxDC variant (PDB 4MET). The ring “flip” seen for His97 in the W132F OxDC variant likely results from metal replacement, as discussed elsewhere.<sup>11,14</sup> Carbon atoms in WT OxDC and the W132F OxDC variant are rendered in cyan and green, respectively. Metal ions are shown as purple (WT OxDC) and salmon (W132F) spheres, and active-site waters are rendered as cyan (WT OxDC) and green (W132F) spheres.

393 increases the charge density on the Glu101 side chain with a  
 394 concomitant change in the metal midpoint potential so that  
 395 Mn(III) is more stable in the W132F OxDC variant compared  
 396 with the WT enzyme. As a result, we would expect that the rate  
 397 constant,  $k_3$ , for the formation of the oxalate-based radical to be  
 398 decreased and the energy of the Mn(II)/oxalate radical  
 399 intermediate to be increased, thereby increasing the rate of  
 400 reversion ( $k_4$ ) back to the Michaelis complex and Mn(III). The  
 401 higher energy of the Mn-bound oxalate-based radical  
 402 intermediate in the W132F OxDC variant is also consistent  
 403 with the larger magnitude of the decarboxylation rate constant  
 404  $k_5$  in that decarboxylation proceeds through an “earlier”

transition state in which the C–O bonds at the carbon  
 proximal to the metal are more polarized (Figure 3). We note  
 that the oxalate radical anion intermediate formed in the  
 reaction catalyzed by WT OxDC can be considered as a 70:30  
 mixture of the resonance structures I and II (see above),<sup>17</sup> and  
 heavy-atom IE measurements on the reaction catalyzed by the  
 R92K OxDC variant support the view that decarboxylation is  
 slower when resonance structure II makes a larger contribution  
 to the transition-state structure; that is, there is less C–O bond  
 polarization.<sup>21</sup>

These findings for the W132F OxDC variant contrast with  
 those that we have reported for the T165V OxDC variant in  
 which the Arg92/Thr165 hydrogen bond is removed by site  
 specific substitution of Thr165 by a valine residue (Figure 4).<sup>15</sup>  
 Thus both partition ratios  $k_5/k_4$  and  $k_3/k_2$  are decreased in this

variant relative to those determined for WT OxDC (Table 3),  
 suggesting that the oxalate-based radical intermediate becomes  
 more stable (Figure 5). As a result, the removal of Arg92/  
 Thr165 hydrogen bond results in decarboxylation proceeding  
 via a “later” transition state, as evidenced by the increased bond  
 order (1.26) and lower polarization of the C–O bonds at the  
 carbon proximal to the metal in the T165V OxDC variant.

## CONCLUSIONS

The functional roles of second-shell residues in metalloenzyme  
 catalysis is receiving increased attention,<sup>24,25</sup> and may underlie  
 the inability of simple Mn(II)-containing complexes to mediate  
 cleavage of the C–C bond in oxalate.<sup>26</sup> Indeed, the heavy-  
 atom isotope effect measurements reported herein clearly demon-  
 strate the importance of individual hydrogen bonds in the  
 OxDC active site for determining transition-state structure and  
 the free energy of radical intermediates in the catalytic  
 mechanism. In part, these effects may result from these small  
 active-site changes in modulating the midpoint potential of  
 Mn(II) in the N-terminal domain of OxDC, although further

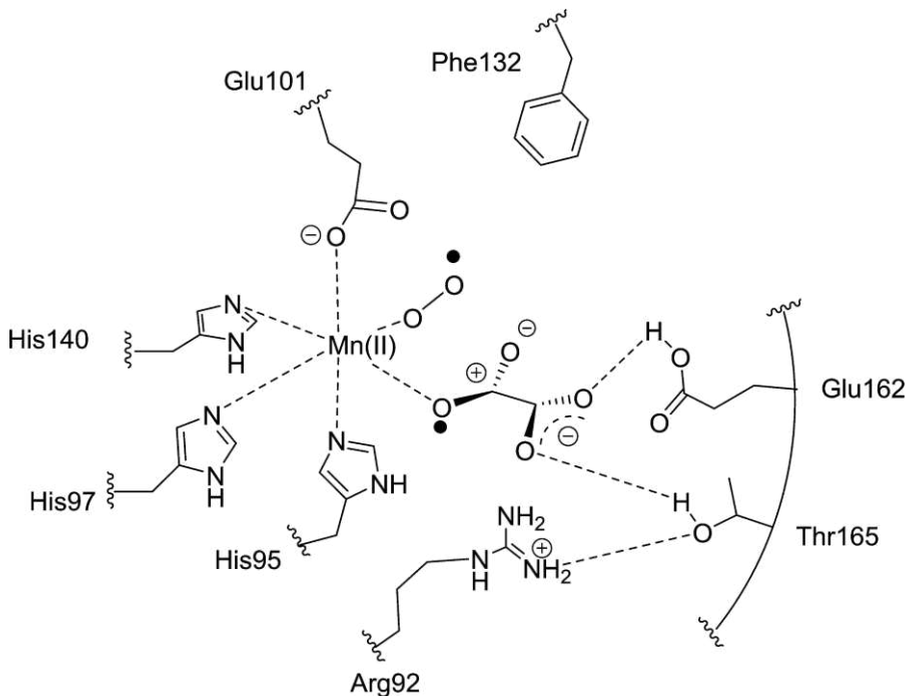


Figure 3. Model for the radical anion intermediate in the W132F OxDC variant.

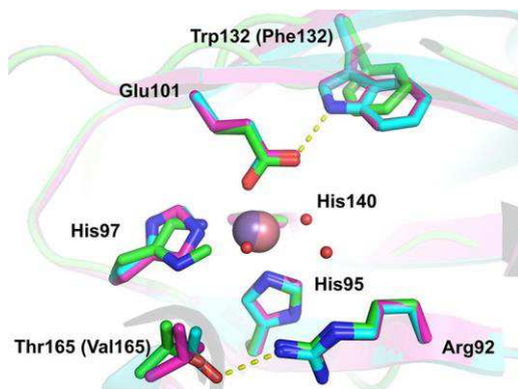


Figure 4. Superimposition of the X-ray crystal structures of WT OxDC (PDB 1UW8) and the T165V (PDB 3SOM) and co-substituted W132F OxDC (PDB 4MET) variants. Note that the removal of the Arg92/Thr165 or Glu101/Trp132 hydrogen bonds does not impact the positions of other active-site residues. Carbon atoms in WT OxDC are rendered in cyan. Carbon atoms in the T165V and the co-substituted W132F OxDC variants are rendered in magenta and green, respectively. Metal ions are shown as purple (WT OxDC and the T165V OxDC variant) and salmon (W132F) spheres, and the three active-site waters in WT OxDC are shown as red spheres.

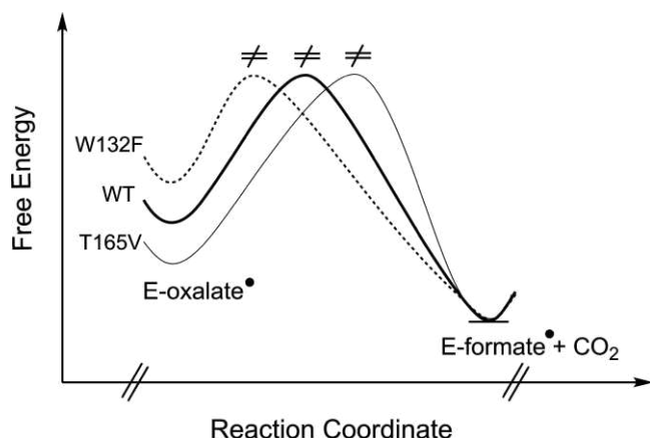


Figure 5. Qualitative representation of the decarboxylation free energy barriers in WT OxDC (thick line) and the W132F (dashed line) and T165V (thin line) OxDC variants.

studies will be needed to determine the validity of this hypothesis and the magnitude of such changes.

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Notes

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#### DEDICATION

This paper is dedicated to the memory of W. W. Cleland (1930–2013).

#### ABBREVIATIONS

OxDC, oxalate decarboxylase; MD, molecular dynamics; FDH, formate dehydrogenase; BHEP, 1,4-bis(2-hydroxyethyl)-piperazine; IRMS, isotope ratio mass spectrometer; IE, isotope effect.

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