ResearchGate

See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/230587434

The Ubiquity of Iron

ARTICLE in ACS CHEMICAL BIOLOGY · JULY 2012

Impact Factor: 5.33 · DOI: 10.1021/cb300323q · Source: PubMed

CITATIONS READS

13 36

2 AUTHORS, INCLUDING:

Perry Allen Frey
University of Wisconsin–Madison
302 PUBLICATIONS 10,253 CITATIONS

SEE PROFILE



Perspective

pubs.acs.org/acschemicalbiology

1 The Ubiquity of Iron

5

6

8 Q

10

11

12

13

14

15

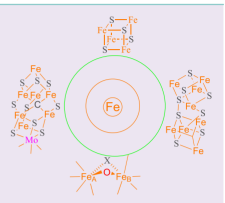
16

17

² Perry A. Frey* and George H. Reed

3 Department of Biochemistry, University of Wisconsin-Madison, 1710 University Avenue, Madison, Wisconsin 53726, United States

ABSTRACT: The importance of iron in living systems can be traced to the many complexes within which it is found, to its chemical mobility in undergoing oxidation reduction reactions, and to the abundance of iron in Earth's crust. Iron is the most abundant element, by mass, in the Earth, constituting about 80% of the inner and outer cores of Earth. The molten outer core is about 8000 km in diameter, and the solid inner core is about 2400 km in diameter. Iron is the fourth most abundant element in Earth's crust. It is the chemically functional component of mononuclear iron complexes, dinuclear iron complexes, [2Fe-2S] and [4Fe-4S] clusters, [Fe-Ni-S] clusters, iron protophorphyrin IX, and many other complexes in protein biochemistry. Metals such as nickel, cobalt, copper, and manganese are present in the crust and could in principle function chemically in place of iron, but they are scarce in Earth's crust. Iron is plentiful because of its nuclear stability in stellar nuclear fusion reactions. It seems likely that other solid planets, formed by the same processes as



Earth, would also foster the evolution of life and that iron would be similarly important to life on those planets as it is on Earth.

recent issue of Current Opinion in Chemical Biology for April 2011, which is devoted to Bioinorganic Chemistry, 20 focuses attention on iron in biochemistry. The opening 21 sentence of the Editorial reads "Iron, the most abundant 22 metal in the earth, dominates the field of inorganic prosthetic 23 groups in proteins". This sentence understates the importance 24 of iron in the Earth, of life on Earth, and of the evolution of life. 25 As used here, understates refers to the fact that iron not only is 26 the most abundant metal in the Earth but is in fact the most 27 abundant element by mass in the Earth. The core of Earth, 28 comprising the inner and outer cores, is 7-8000 km in 29 diameter and contains many elements, but iron dominates the 30 core.

31 IRON AND THE EARTH

32 Iron is prevalent in Earth because of the physics of nuclear 33 fusion. Earth and presumably other solid planets originated 34 from accretion of matter arising from a primordial supernova. 35 The predominance of iron within this matter is a consequence 36 of the physics of nucleosynthesis in dying stars. Fusion of 37 lighter elements to form heavier ones in late-stage stellar 38 evolution produces a net release of energy up to elements in the 39 iron group (Fe, Co, Ni). Nuclear fusion to form elements 40 heavier than iron and nickel absorbs energy and therefore is not 41 sustaining. A major sequence in the complicated fusion chain of 42 nucleosynthesis involves the alpha process whereby the 43 equivalent of a helium nucleus is fused to the precursor 44 element. The masses increase by 4 units up to mass number 56. 45 The ⁵⁶Ni that is produced in this chain decays to ⁵⁶Fe via ⁵⁶Co 46 through sequential electron captures with production of 47 characteristic γ rays. Thus, iron becomes a major component 48 of dying stars that have or acquire sufficient mass to undergo a 49 cataclysmic supernova. Consequently iron is abundant in the 50 debris resulting from supernovae of all types.

Figure 1 depicts the approximate relative diameters of the 51 f1 inner and outer cores, the mantle, and the crust of Earth. The 52 core is very hot, and for this reason the outer core is molten. 53 However, the inner core is solid because of the high pressure, 54 330 GPa or 3.5×10^6 atm, despite the temperature of 5700 ss

The crust, mantle, and core of Earth all contain large 57 amounts of iron, but most is in the core, ~8000 km in diameter 58 with 80% iron.⁵ One theory holds that the inner core, 2400 km 59 in diameter, consists of a "nearly perfectly aligned aggregate of 60 hcp (hexagonal close-packed) crystals of iron.⁶ Physical 61 evidence in support includes seismic sound measurements 62 proving that the core is anisotropic. The only known crystalline 63 form of iron that displays anisotropy at the temperature and 64 pressure of the core is the hexagonal close-packed crystal.⁶⁻⁹ 65 The compositions of the inner and outer cores suggests that 66 iron might be the most abundant element in the Earth, and this 67 is borne out by detailed geologic analysis showing that the iron 68 content of the Earth is 32% by weight. 10,11

Earth's mantle also contains large amounts of iron. A lesser 70 percentage of iron is found on Earth's surface than in the core, 71 but it is still the fourth most abundant element (6.3%) in the 72 crust. The other major elements in the crust are oxygen (47%), 73 silicon (26%), aluminum (8.1%), calcium (5.0%), magnesium 74 (2.9%), sodium (2.3%), and potassium (1.5%).

■ IMPORTANCE OF IRON TO LIFE

The central role of iron in living cells and organisms is widely 77 known but not fully appreciated. Broadly defined, the 78 biochemical reactions in a living cell may be regarded as 79

Received: June 29, 2012 Accepted: July 30, 2012

ACS Chemical Biology

Perspective

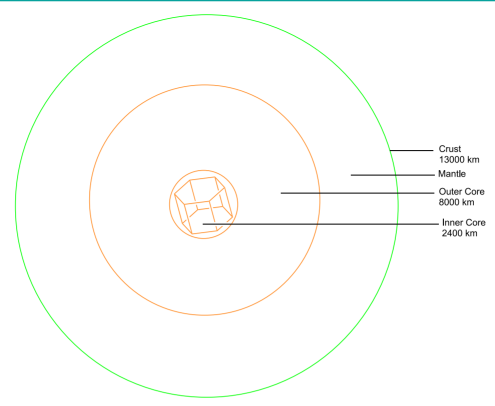


Figure 1. Schematic representation of approximate relative diameters of substructures in Earth. The inner core consists of aligned hcp crystals of iron and is represented here as a single hexagonal solid. The inner and outer cores consist of about 80% iron. The outer core is molten. The mantle is solid rock, and the crust contains about 6.3% iron, the fourth most abundant element in the crust.

80 cellular metabolism. Many branches of metabolism deal with 81 the full range of cellular processes, energy production, 82 biosynthesis, replication, and locomotion. Iron contributes to 83 each of these processes. Iron even participates in the regulation 84 of gene expression. 12 This summation seeks to explain salient 85 aspects of the importance and significance of iron to life on 86 Earth.

Multiplicity of Iron Complexes in Proteins. It is useful 88 to consider the sheer number of different chemical classes of 89 iron complexes in living cells. Figure 2 shows a sampling of the many iron complexes in living cells. New ones are being discovered all the time. For example, the structure of the 92 complex of S-adenosyl-L-methionine (SAM) with [4Fe-4S] 93 clusters in radical SAM enzymes was discovered within the past 94 decade, although it is an ancient coenzyme. 13 This year, the 95 final assignment of carbide as the central interstitial species in 96 the FeMoco cofactor of nitrogenase completes the structure of this molecule, which is central to the fixation of nitrogen. 14,15 It 98 is interesting that the iron carbide in FeMoco appeared in 99 nature millions of years before the carbonization of iron by man 100 in the manufacture of steel. A very recent article reveals that the 101 nucleolus of plant cells is a "hot spot" of iron. 16 The chemical constitution of nucleolar iron is not yet known. There are many 103 more biological iron coenzymes and prosthetic groups than of any other metal.

Iron complexes facilitate electron transfer, from fast intramolecular processes to the long distances (12–20 Å) separating active sites in macromolecular complexes and across membranes. Iron complexes catalyze oxygenation and dehydrogention of metabolites. They catalyze isomerization of metabolites; essential steps in DNA and RNA biosynthesis; certain types of DNA repair; and numerous steps in the biosynthesis of vitamins, coenzymes, cofactors, and antibiotics. All of these 112 processes were important in the genesis and evolution of life. 113 Finally, the iron in heme facilitates transport of molecular 114 oxygen in the bloodstreams of higher animals and humans.

The iron-cysteine and iron sulfide complexes at the top of 116 Figure 2 are likely the earliest biological iron complexes in 117 evolution. They are the most efficient and versatile structures 118 engaged in electron transfer and display reduction potentials 119 ranging from +360 to -650 mV. They are essential for 120 oxidation—reduction processes that provide the energy for ATP 121 production and function in critical roles in many other 122 metabolic processes. These simple iron complexes date to the 123 earliest anaerobic Archeae and bacteria and have persisted in 124 their roles in energy production and metabolism since the 125 appearance of molecular oxygen and the evolution of aerobic 126 metabolism.

Mononuclear iron complexes, such as that at the right of the 128 first line in Figure 2, ¹⁸ and the oxo-Fe₂ complexes in the second 129 line catalyze oxygenation, dehydrogenation, and reduction of 130 metabolites and xenobiotics. Class I ribonucleotide reductases 131 required for DNA biosynththesis and methane monooxyge- 132 nases have di-iron complexes that initiate catalysis by 133 generating free radicals. ^{19–21} The Fe–Ni hydrogenase complex 134 catalyzes the reduction of protons to molecular hydrogen. The 135 C-cluster in bifunctional CO dehydrogenase contains both Fe 136 and Ni, and the A-cluster in this enzyme contains up to three 137 metals, Fe–Zn–Ni, Fe–Cu–Ni, or Fe–Ni–Ni. The latter is 138 thought to be the most biologically active. ²² The [4Fe–4S/ 139 SAM complex catalyzes many radical reactions, including 140 complex metabolic reactions; DNA repair; deoxyribonucleotide 141 production in DNA biosynthesis; maturation of RNA; and 142

ACS Chemical Biology Perspective

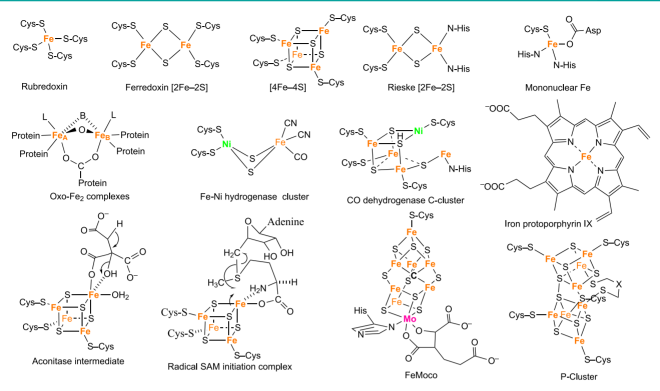


Figure 2. Chemical structures of representative iron complexes in proteins. Rubredoxin, ferredoxin, [4Fe-4S] clusters, and Rieske [2Fe-2S] clusters are found in proteins that function as electron transfer processes. [2Fe-2S] and [4Fe-4S] clusters are also engaged in regulation of the transcription of genes that are under redox control. Mononuclear iron complexes are found in enzymes that catalyze oxygenation reactions. Oxo-Fe₂ complexes catalyze dehydrogenation or reduction of ribonucleotides to deoxyribonucleotides. Fe-Ni hydrogenase and Fe-Fe hydrogenase complexes are found in hydrogenases. The CO dehydrogenase cluster shown is cluster B in carbon monoxide dehydrogenase. The aconitase intermediate occurs as the dehydration intermediate in aconitase. Iron protoporphyin IX is found in heme proteins, cytochromes c and a, and cytochrome P450. The radical SAM initiation complex is found in radical SAM enzymes. FeMoco and P-cluster are essential cofactors in nitrogenase. The interstitial carbide (C^{4-}) has recently been identified. An interstitial carbide (C^{4-})

143 chemically difficult steps in vitamin, coenzyme, and antibiotic 144 biosynthesis. ¹³

Iron protoporphyrin IX in Figure 2 carries out many essential 146 biological functions. It is the coenzyme of cytochrome P450s, ²³ 147 which catalyze the oxygenation of many substrates incorporat-148 ing unactivated and unreactive alkanyl-C-H bonds. It is also 149 the cofactor of peroxidases and catalases. As the cofactor in 150 cytochromes and cytochrome oxidase it facilitates and guides electron transfer processes in ATP biosynthesis. As the 152 prosthetic group in heme, it binds molecular oxygen reversibly 153 and transfers it to cells where it is needed in energy metabolism. Nitrogen and Iron. Unlike iron, nitrogen is not abundant 154 155 in the crust of Earth at only 0.002%. However, nitrogen constitutes 2.6% of the human body. Nitrogen contributes about 16% of the mass in a typical protein and slightly more in 158 DNA. The nitrogen in living cells comes from the atmosphere, which is 78% nitrogen gas. Thus, life depends upon the harvest of nitrogen from the atmosphere by nitrogen fixation, in which N₂ gas is chemically reduced to ammonia (NH₃). Nitrogenase, 162 found in the anaerobic bacteria in root nodules of leguminous 163 plants and in free-living diazotrophs such as Azotobacter 164 vinelandii, catalyzes the fixation of nitrogen. The iron complexes FeMoco and P-cluster shown in Figure 2 drive the action of 166 nitrogenase, and a [4Fe-4S] cluster in nitrogenase relays 167 reducing electrons through the P-cluster to FeMoco, the site at 168 which the hydrogenation of N₂ takes place. An iron-containing 169 siroheme and [4Fe-4S] cluster are essential cofactors in 170 ferredoxin-nitrite reductases that carry out a six-electron 171 reduction of nitrite to ammonia occurring in plants, algae,

and cyanobacteria. Iron is essential to life by making nitrogen $_{172}$ available to all living cells, and this process appeared in nature $_{173}$ millions of years before the invention of the Haber Process for $_{174}$ the nonenzymatic, hydrogenolytic reduction of N_2 to ammonia. $_{175}$

Carbon and Iron. Carbon is not a dominant element in 176 Earth's crust (0.18%) but constitutes 23% of the human body. 177 Most carbon in the biosphere originates with plants that fix 178 CO₂ from the atmosphere. Carbon dioxide fixation requires 179 ATP, which is produced in photosynthesis. In photosynthetic 180 ATP production, the light harvested by chlorophyll produces 181 highly reducing electrons, which are relayed by the [4Fe-4S] 182 clusters of ferredoxin to sites that generate proton gradients 183 across membranes and energize ATP biosynthesis by photo-184 synthetic ATP synthase. Thus, iron plays an essential role in 185 ATP production and CO₂ fixation.

Iron Chemistry and Life. Iron can exist in eight oxidation 187 states from -2 to +6. In biological cells the most common 188 states are +2 and +3, and the +4 and +5 states are frequently 189 brought into play in oxygenation reactions. The standard 190 reduction potential for free iron is in the medium range, -0.44 191 V, so that it is easily oxidized to Fe²⁺, an important state in 192 biochemistry. The standard potential for reduction of Fe³⁺ to 193 Fe²⁺ is +0.77 V, making iron useful for electron transfer 194 reactions in nature. The standard reduction potentials for iron 195 in the complexes shown in Figure 2 are not exactly the same as 196 for free iron and cover a larger range, as pointed out above for 197 the iron–sulfur clusters.

ACS Chemical Biology Perspective

99 LIFE AND OTHER METALS

200 While the midrange reduction potentials for iron complexes 201 and the ability of iron to accept a broad range of ligands enable 202 it to function efficiently in essential biological processes, these 203 properties are not unique to iron. Other transition metals such 204 as nickel ($E^{\circ}=-0.26$ V), cobalt ($E^{\circ}=-0.28$ V), manganese 205 ($E^{\circ}=-1.17$ V), and copper $E^{\circ}=0.34$ V) also bind a broad 206 range of ligands and could in principle function in place of iron. 207 In fact, nickel (Figure 2) and cobalt in vitamin B_{12} do function 208 analogously in a small number of biological reactions. 209 Manganese can function in place of iron in the oxo-Fe₂ 210 complexes, 24,25 and copper proteins function in electron 211 transfer and even reversible binding of molecular oxygen in 212 the case of hemocyanin. Zinc ($E^{\circ}=-0.76$ V) is a special case 213 because Zn^{2+} is the only stable oxidized form, so it does not 214 engage in reversible electron transfer.

Despite redox analogies among iron, cobalt, nickel, 216 manganese, and copper, iron is by far the dominant player in 217 biochemical processes. One reason for this bias is likely the 218 abundance of iron in the Earth's crust. At 6.3% of the crust iron 219 is 700 times more plentiful than nickel, 2100 times more than 220 cobalt, 1000 times more than copper, and 57 times more than 221 manganese. Therefore, although other metals display chemical 222 properties that could be exploited to support life, they are too 223 scarce in Earth's crust to compete with iron. Even the recent 224 discovery of mononuclear carbide (C4-) as the interstitial 225 species in FeMoco does not distinguish iron as uniquely 226 suitable to sustain life. Most metals form stable complexes with 227 mono- or binuclear carbide, and interstitial C⁴⁻ contributes to the hardness and structural rigidity of many alloys, including 229 steel and tungsten carbide, the principal constituent in armor 230 piercing ammunition employed by the military. Interstitial 231 carbide might structurally stabilize the FeMoco cluster and 232 could be expected to do so for other metals as well.

3 WHY NOT SILICON AND ALUMINUM IN LIFE?

234 Among the principal elements in Earth's crust, silicon and 235 aluminum are present as aluminates and silicates in greater 236 amounts than iron; however, they are not important in the 237 chemistry of life. Aluminum binds ligands, especially oxygen, 238 very strongly and does not easily exchange ligands. Aluminum 239 exists in nature in the +3 oxidation state and is not easily 240 reduced (reduction potential of -1.66 V). Because Al $^{3+}$ does 241 not easily engage in either electron transfer or ligand exchange, 242 it has not been adopted as a component of living cells.

Silicon is in the same chemical group as carbon and shares an analogous valence electronic configuration with carbon. For this 245 reason, the question arises why silicon cannot support life as 246 well as carbon. There are at least two reasons for this. Certain chemical properties of silicon vary from carbon and would make it impossible for silicon to stand in for carbon. Carbon binds four ligands in a tetrahedral array that display right and left handedness, or stereochemistry. All sugars are constructed with tetrahedral carbon and are right-handed. All amino acids 252 are constructed with tetrahedral carbon and are left-handed. This property of carbon translates into proteins, all of which are 254 constructed with left-handed amino acids. The handedness of 255 proteins allows them to bind molecules upon which they act 256 with a high degree of specificity. Although silicon forms 257 tetrahedral compounds, the right- and left-handed forms tend 258 to be configurationally mobile. Relatively few chiral silanes are 259 known. Thus, left or right-handedness would be difficult to

translate into a silicon based protein, and such proteins would 260 likely not have stereochemically defined structures and not be 261 specific in binding molecules.

A perhaps more serious problem for silicon in life is the 263 reactivity of silanes. While the silicates in Earth's crust are 264 structurally stable, silanes are unstable and undergo sponta- 265 neous inflammation in the air. This property would make all 266 silicon-based fatty acids and many silicon analogs of amino 267 acids chemically unstable under physiological conditions.

269

287

293

NOT ONLY IRON

Iron is not the only metal required for life. Other common 270 metals in the Earth's crust such as sodium (2.3%), potassium 271 (1.5%), and magnesium (2.9%) are essential cations in all cells. 272 These Group I and II metals do not participate chemically in 273 the complex processes of oxidation-reduction, electron 274 transfer, N2 reduction or O2-processing. Magnesium chelates 275 oxyanions in polyphosphates such as ATP and in polynucleo- 276 tides such as RNA. In plants and photosynthetic bacteria 277 magnesium chelates the heterocyclic bases in chlorophyll. In 278 this function magnesium works very well precisely because it 279 displays chemical properties very different from those of iron. 280 Chlorophyll collects energy from light to generate high 281 potential reducing equivalents and drive biosynthesis in plants 282 and photosynthetic bacteria. Iron would not do as the central 283 chelator because of its propensity to undergo reduction. 284 Magnesium ideally serves the purpose as an excellent chelator 285 but very poor electron acceptor ($\tilde{E}^{\circ} = -2.4 \text{ V}$). 2.86

CONCLUSION

Iron is the most abundant element because of its high nuclear 288 binding energy. Together with other transition metals, iron 289 possesses the ligand binding and electron transfer properties 290 essential to many biological processes. Iron dominates these 291 processes, most likely because of its abundance in the Earth. 292

AUTHOR INFORMATION

Corresponding Author *E-mail: frey@biochem.wisc.edu. Notes The authors declare no competing financial interest. 294 295 296 297

REFERENCES

- (1) Frey, P. A., and Outten, C. E. (2011) Forging ahead: new 299 mechanistic insights into iron biochemistry. *Curr. Opin. Chem. Biol.* 15, 300 257–259.
- (2) Woosley, S. E., Heger, A., and Weaver, T. A. (2002) The 302 evolution and explosion of massive stars. *Rev. Mod. Phys.* 74, 1015–303 1071.
- (3) Alfé, D., Gillan, M., and Price, G. D. (2002) Composition and 305 temperature of the Earth's core constrained by combining *ab initio* 306 calculations and seismic data. *Earth Planet. Sci. Lett.* 195, 91–98.
- (4) Lide, D. R. (2006–2007) CRC Handbook of Chemistry and 308 Physics, 87th ed., pp. 13–14, CRC Press, Boca Raton, FL.
- (5) Jordan, T. H. (1979) Structural Geology of the Earth's Interior. 310 Proc. Natl. Acad. Sci. U.S.A. 76, 4192–4200.
- (6) Stixrude, L., and Cohen, R. E. (1995) High-pressure elasticity of 312 iron and anisotropy of Earth's inner core. *Science 267*, 1972–1975. 313
- (7) Stixrude, L., and Cohen, R. E. (1995) Constraints on the 314 crystalline structure of the inner core: Mechanical instability of BCC 315 iron at high pressure. *Geophys. Res. Lett.* 22, 125–128.
- (8) Vocadlo, L., Alfé, D., Gillan, M. J., Wood, I. G., Brodholt, J. P., 317 and Price, G. D. (2003) Possible thermal and chemical stabilization of 318 body-centered-cubic iron in the Earth's core. *Nature 424*, 536–539.

ACS Chemical Biology Perspective

- 320 (9) Tateno, S., Hirose, K., Ohishi, Y., and Tatsumi, Y. (2010) The 321 structure of iron in the Earth's inner core. *Science* 330, 359–361.
- 322 (10) McDonough, W. F. The composition of the Earth. http://323 quake.mit.edu/histgroup/CoreMantle/EarthCompo.pdf (Table 3).
- 324 (11) McDonough, W. F., and Sun, S.-s. (1995) The composition of 325 the Earth. *Chem. Geol.* 120, 223–253.
- 326 (12) Fleischhacker, A. S., and Kiley, P. J. (2011) Iron-containing 327 transcription factors and their roles as sensors. *Curr. Opin. Chem. Biol.* 328 15, 335–341.
- 329 (13) Frey, P. A., Hegeman, A. D., and Ruzicka, F. J. (2008) The 330 radical SAM superfamily. *Crit. Rev. Biochem. Mol. Biol.* 43, 63–88.
- 331 (14) Spatzal, T., Aksoyoglu, M., Zhang, L., Andrade, S. L., Scheicher,
- 332 E., Weber, S., and Rees, D. C. (2011) Science 334, 940–941.
- 333 (15) Lancaster, K. M., Roemelt, M., Ettenhuber, P., Hu, Y., Ribbe, M.
- 334 W., Neese, F., Bergmann, U., and DeBeer, S. (2011) X-ray emission 335 spectroscopy evidences a central carbon in the nitrogenase iron-
- 335 spectroscopy evidences a central carbon in the nitrogenase iro 336 molybdenum cofactor. *Science* 334, 974–977.
- 337 (16) Roschzttardtz, H., Grillet, L., Isaure, M.-P., Conéjéro, G., 338 Ortega, G. R., Curie, C., and Mari, S. (2011) Plant cell nucleolus as a 339 hot spot for iron. *J. Biol. Chem.* 286, 27863–27866.
- 340 (17) Gray, H. B., and Winkler, J. R. (2005) Long range electron 341 transfer. *Proc. Natl. Acad. Sci. U.S.A. 102*, 3534–3539.
- 342 (18) Koehntop, K. D., Emerson, J. P., and Que, L., Jr. (2005) The 2-
- 343 His-1-carboxylate facial triad: a versatile platform for dioxygen 344 activation by mononuclear non-heme iron(II) enzymes. *J. Biol. Inorg.* 345 *Chem.* 10, 87–93.
- 346 (19) Nordlund, P., and Reichard, P. (2006) Ribonucleotide 347 reductases. *Annu. Rev. Biochem.* 75, 681–706.
- 348 (20) Tinberg, C. E., and Lippard, S. J. (2011) Dioxygen activation in
- 349 soluble methane monooxygenase. Acc. Chem. Res. 44, 280–288.
- 350 (21) Kovaleva, E. G., Neibergall, M. B., Chakrabarty, S., and 351 Lipscomb, J. D. (2007) Finding intermediates in the O₂ activation
- 352 pathways of non-heme iron oxygenases. Acc. Chem. Res. 40, 475–483.
- 353 (22) Drennan, C. L., Doukov, T. I., and Ragsdale, S. W. (2004) The
- 354 metalloclusters of carbon monoxide synthase: a story in pictures. *J.* 355 *Biol. Inorg. Chem. 9*, 511–515.
- 356 (23) Ortiz de Montellano, P. R. (1995) Cytochrome P450: Structure, 357 Mechanism and Biochemistry, Springer, New York.
- 358 (24) Jiang, W., Yun, D., Saleh, L., Bollinger, J. M., and Krebs, C.
- 359 (2008) Biochemistry 47, 13736–13744. 360 (25) Zhang, Y., and Stubbe, J. (2011) Bacillus subtilis class b
- 361 ribonucleotide reductase is a dimanganese(III) radical enzyme.
- 362 Biochemistry 50, 5615-5623.