

# Effective intermediate-spin iron in O<sub>2</sub>-transporting heme proteins

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Proteins carrying an iron-porphyrin (heme) cofactor are essential for biological O2 management. The nature of Fe-O2 bonding in hemoproteins is debated for decades. We used energy-sampling and rapid-scan X-ray Kβ emission and K-edge absorption spectroscopy as well as quantum chemistry to determine molecular and electronic structures of unligated (deoxy), CO-inhibited (carboxy), and O2-bound (oxy) hemes in myoglobin (MB) and hemoglobin (HB) solutions and in porphyrin compounds at 20-260 K. Similar metrical and spectral features revealed analogous heme sites in MB and HB and the absence of lowspin (LS) to high-spin (HS) conversion. Amplitudes of KB main-line emission spectra were directly related to the formal unpaired Fe(d) spin count, indicating HS Fe(II) in deoxy and LS Fe(II) in carboxy. For oxy, two unpaired Fe(d) spins and, thus by definition, an intermediatespin iron center, were revealed by our static and kinetic X-ray data, as supported by (time-dependent) density functional theory and complete-active-space self-consistent-field calculations. The emerging Fe-O<sub>2</sub> bonding situation includes in essence a ferrous iron center, minor superoxide character of the noninnocent ligand, significant double-bond properties of the interaction, and three-center electron delocalization as in ozone. It resolves the apparently contradictory classical models of Pauling, Weiss, and McClure/Goddard into a unifying view of O<sub>2</sub> bonding, tuned toward reversible oxygen transport.

heme cofactor  $\mid$  X-ray spectroscopy  $\mid$  quantum chemistry  $\mid$  O<sub>2</sub> binding  $\mid$  spin state

Proteins binding an iron-porphyrin (heme) cofactor are nature's versatile dioxygen (O<sub>2</sub>) managers, facilitating O<sub>2</sub> sensing, transport, activation chemistry, and catalysis (1). Scrutiny over decades has been devoted to the heme chemistry in myoglobin (MB) and hemoglobin (HB) proteins of vertebrates (Fig. 1), which is the basis for their vital O<sub>2</sub>-transporting properties (1–5). Renewed interest in the topic stems from ongoing discovery of globins in plants and microorganisms, which further extends the spectrum of biological functions (6–8). A controversy remains on the nature of the iron-oxygen bonding in HB/MB, dating back to pioneering work of Pauling, Monod, and Perutz starting in the 1930s (3, 9).

Three physiologically relevant heme species are distinguished in MB/HB for which crystal structures at ~1 Å resolution are available (10, 11): the ligand-free (deoxy), carbon-monoxide inhibited (carboxy), and O<sub>2</sub>-bound (oxy) forms (Fig. 1), besides the ferric (met) species resulting from autoxidation. Paramagnetic behavior has established high-spin (HS) iron in deoxy [d<sup>6</sup> Fe(II); spin (S) multiplicity, M = 2S + 1 = 5] and met  $[d^5]$  Fe(III); M = 6], but carboxy is diamagnetic because of low-spin (LS) Fe(II) (M = 1) (12). For oxy, a diamagnetic (M = 1) cofactor was proven (3, 12, 12)13). However, the O<sub>2</sub> triplet ground state provides several options for spin flipping or charge transfer (Fig. 1): Pauling suggested a neutral singlet O<sub>2</sub> bound to a formal LS Fe(II) (13, 14), McClure suggested a triplet O<sub>2</sub> and an intermediate-spin (IS) Fe(II) with antiferromagnetic (afc) coupling (15), an IS Fe(II) with d-level population inversion in an ozone-like model was also suggested by Goddard (16), and Weiss suggested an Fe(III) afc coupled to a superoxo (O<sub>2</sub><sup>-</sup>) ligand (17). The possible resonance character of these Lewis structures was emphasized already in early theory work (18). Density functional theory (DFT), providing access to

experimental observables (19, 20), and complete-active-space self-consistent-field (CASSCF) quantum chemical calculations facilitated description of many structural and spectroscopic features of the  $O_2$  bonding (1, 18). A McClure-like oxy (21), and Pauling/Weiss formulations have emerged, to some extent depending on model terms used to interpret multiconfigurational wavefunctions (21–23). Modification of porphyrin groups and heme/ligand geometries by the protein matrix and hydrogen bonding may bias the Fe- $O_2$  interaction to the diverse functions (1, 24–26).

Core level spectroscopy is powerful for electronic structure analysis of transition metal complexes (27, 28). X-ray absorption spectroscopy (XAS) probes unoccupied valence levels by resonant 1s electron excitation in the pre-K-edge region (core-to-valence transitions, ctv), ligation symmetries in the X-ray absorption near edge structure (XANES), and bond lengths in the extended X-ray absorption fine structure (EXAFS). X-ray emission spectroscopy (XES) of the Kβ line probes occupied levels via electronic decay from metal-3p (main lines) or metal-valence (satellite lines; valence-to-core transitions, vtc) levels to the core hole (29). The core/valence-to-valence/core electronic transitions (ctv/vtc) spectra can be calculated by DFT methods for theory benchmarking (30, 31). 3p-3d spin coupling splits the Kβ main-line emission into the Kβ' and Kβ<sub>1,3</sub> features, and the Kβ' intensity is a measure of the metal spin state and bonding covalency (32-34). XAS has been used to address structural metrics, solution vs. crystalline materials, and redox states in MB/HB (35-39). ctv data were interpreted in favor of the Weiss model of O<sub>2</sub> binding (36, 40). Systematic Kβ XES on hemoproteins has not been done yet and is challenging on protein solutions because of radiation-induced modifications (41). Temperature effects further may contribute, because in met MB/HB partial spin crossover (i.e., HS population) was found (42).

We used advanced X-ray spectroscopy to study heme species in MB and HB proteins in solution. Structural and electronic features

# **Significance**

Tuning of the iron-oxygen bonding in hemoproteins is crucial for their versatile  $O_2$  chemistry. The nature of the Fe- $O_2$  bond in myoglobin (MB) and hemoglobin (HB) has been debated for a long time. High-resolution X-ray K $\beta$  emission and K-edge absorption spectroscopy on MB and HB solutions and quantum chemical (time-dependent) density functional theory and complete-active-space self-consistent-field calculations revealed an essentially ferrous, intermediate-spin iron in an ozone-like configuration in  $O_2$ -bound oxy heme. The emerging bonding situation resolves the three classical models into a consistent view of the metal-ligand interaction, biased toward reversible  $O_2$  binding in the oxygen transporters.

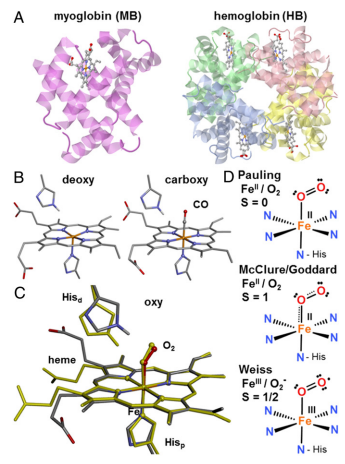
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**Fig. 1.** Heme protein structures and O<sub>2</sub> binding. (A) Crystal structures of human MB/HB (*Left/Right*, PDB ID codes 2DN2/3RGK). (*B*) Structures of deoxy/carboxy hemes in MB (PDB ID codes 3QM6/3QM7, 0.91/0.96 Å resolution). (C) Overlay of oxy hemes in MB/HB (balls-and-sticks/yellow-sticks, PDB ID codes 1A6M/2DN1, 1.00/1.25 Å resolution). (*D*) Schematic Fe-O<sub>2</sub> bonding in oxy in the three literature models (His<sub>p/d</sub> in C, proximal/distal histidine; color code: gray, C; blue, N; red, O; orange, Fe). Hydrogen bonding of His<sub>d</sub> to bound O<sub>2</sub> has been established by neutron diffraction (68).

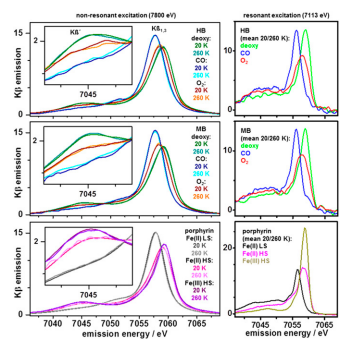
were similar for both proteins and constant at 20–260 K. Static and kinetic X-ray absorption/emission spectroscopy (XAS/XES) and (time-dependent) density functional theory (TDDFT) and CASSCF analyses revealed HS Fe(II) in deoxy, LS Fe(II) in carboxy, and IS iron in a delocalized Fe-O $_2$  bonding situation in oxy, which resolves the classical models in a unifying view of the metal–ligand interaction.

# Results

Heme Species Quantification. MB and HB proteins in solution were prepared with ligand-free (deoxy), carbon monoxide inhibited (carboxy), or oxygenated (oxy) hemes. Similar optical absorption spectra of MB and HB (SI Appendix, Fig. S2) with typical Soret band shifts (MB/HB: met, 407/405 nm; deoxy, 433/429 nm; carboxy, 423/419 nm; oxy, 419/414 nm) and differences in the Q-band region (500-600 nm) agreed with previous reports (36, 43). Fe K-edge XANES spectra of three sets of MB/HB samples were collected at 20 K and 260 K at two monochromator resolutions and using broad-band Kα or narrow-band Kβ emission detection (SI Appendix, Figs. S3–S5). XANES shapes of MB/HB differed only slightly, and K-edge energies of MB/HB for a given heme species at 20/260 K were similar within  $\pm 0.1$  eV (deoxy, 7,121.0 eV; carboxy, 7,122.3 eV; oxy, 7,123.9 eV; SI Appendix, Table S1), in agreement with previous reports (35, 36, 40). The ~1.3 eV or ~2.9 eV energy differences for deoxy/carboxy or deoxy/oxy reflected iron ligation and spin-state changes as outlined below. Precise iron-ligand bond

lengths were determined from EXAFS analysis. Coordination numbers close to unity were found for the Fe-CO and Fe-O<sub>2</sub> bonds in MB/HB. The Fe-O<sub>2</sub> (1.90 Å), Fe-CO (1.79 Å), Fe-N<sub>porphyrin</sub> (deoxy, 2.07 Å; carboxy, 2.01 Å; oxy, 2.04 Å), and Fe-H<sub>His</sub> (deoxy, 2.18 Å; carboxy, 2.07 Å; oxy, 2.16 Å) distances for MB/HB in the three states and at 20/260 K deviated by <0.02 Å, and the distances in the first and second iron coordination spheres mostly agreed within <0.05 Å with crystal structures (*SI Appendix*, Table S2). These results verified near-quantitative (>90%) presence of deoxy, carboxy, and oxy hemes and showed that the iron site configurations in MB and HB are similar and independent of the temperature.

Spin States from K $\beta$  XES. Nonresonantly excited K $\beta$  main-line emission spectra of HB and MB with deoxy, carboxy, and oxy hemes were collected at 20 K and 260 K by using an energysampling approach (Fig. 2) (44). As references for Kβ spectral features, three synthetic iron-porphyrin compounds [1, LS Fe(II); 2, HS Fe(II); 3, HS Fe(III)] and several nonporphyrin iron complexes with known spin and redox states were studied (SI Appendix, Figs. S7 and S14). At both temperatures, the spectral shapes for MB and HB were similar and the  $K\beta_{1,3}$  line energies (deoxy, 7,058.6 eV; carboxy, 7,058.0 eV; oxy, 7,058.4 eV; SI Appendix, Table S1) agreed within ±0.1 eV. Significant line-shape differences between the MB/ HB heme species and the porphyrins, i.e., variations in the relative  $K\beta_{1,3}$  amplitudes at increasing energies for carboxy, deoxy, and oxy and of the Kβ' feature at ~7,045 eV, reflected iron-spin and redoxstate differences. Both proteins in all states and the porphyrins showed temperature-independent Kβ spectra (Fig. 2 and SI Appendix, Fig. S6). The Kβ' amplitudes for MB/HB, porphyrins, and nonporphyrin compounds were determined and plotted versus the formal number of unpaired Fe(d) spins (Fig. 3). For porphyrin and nonporphyrin compounds, the doubling of the Kβ' amplitude for a Fe(d) spin count increase from 0 in LS Fe(II) to 5 in HS Fe(III) was well described by a linear relation. The Kβ' amplitudes of deoxy MB/HB agreed well with a Fe(d) count of 4 as in compound 2, indicating HS Fe(II). The Kβ' amplitudes of carboxy MB/HB were centered close to 0 unpaired Fe(d) spins in LS Fe(II) as in



**Fig. 2.** Kβ emission spectra of MB, HB, and porphyrin references. (*Left*) Nonresonantly excited, normalized Kβ main-line spectra at 20 K or 260 K and magnified Kβ′ feature in *Inset (Top.* MB solution; *Middle*, HB solution; *Bottom*, porphyrin compounds **1–3**). (*Right*) Resonantly excited, normalized Kβ main-line spectra (mean of data at 20 K and 260 K).

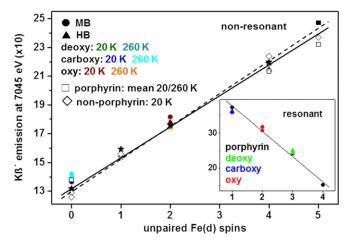


Fig. 3. Correlation of Kβ' intensity and formal unpaired Fe(d) spin count. Data points refer to Kβ spectra in Fig. 2 (Left) and SI Appendix, Fig. S7 (nonporphyrin compounds). Kβ data for dehydrated deoxy, carboxy, and oxy HB and met (at Fe(d) = 1) MB/HB (solid asterisks) and solid hemin (solid square) at room temperature derived from ref. 45 were scaled to our data at Fe(d) = 0. Fe(d) spins correspond to: 0, LS Fe(II); 1, LS Fe(III); 4, HS Fe(II), and 5, HS Fe(III). Kβ' intensities for oxy MB/HB were placed at Fe(d) = 2 for a formal IS Fe(II) (solid or dashed lines, linear fits to porphyrin/nonporphyrin or porphyrin data, R =0.99). (Inset) Data from Kβ spectra in Fig. 2, Right (for Fe(d) count calculation see the text; line, linear fit to porphyrin data, R = 0.98).

compound 1. For oxy MB/HB, the Kβ' amplitudes varied within only  $\sim$ 5%, and the mean value was in best agreement with an Fe(d) spin count of 2. We note that the hypothetical conversion of the  $K\beta$ ' amplitude of oxy HB/MB to the smaller Kβ' amplitude for 1 spin [as for LS Fe(III)] required subtraction of a large (~40%) theoretical contribution of deoxy from the oxy spectra, which resulted in distorted line shapes (SI Appendix, Fig. S8). Such a large deoxy contribution further was unlikely regarding near-quantitative oxy formation as deduced from the optical absorption and XAS data. A LS Fe(III) contribution from met heme would even lead to underestimation of the Kβ' amplitude of oxy. An in-depth fit analysis of the KB spectra revealed detailed relations between peak energies/intensities and converged toward the same spin assignments as above (SI Appendix, Figs. \$9-\$11). These results strongly suggested that the HS Fe(II) in deoxy MB/HB is converted to an effective IS iron species in oxy upon O2 binding.

A single earlier Kβ study on dried HB/MB at room temperature is available (45). Respective K<sub>B</sub>' amplitudes showed that HS Fe(III) in hemin and deoxy and carboxy HB were close to our data for Fe(d) counts of 5, 4, or 0, and the Fe(d) count of ~1 in met HB/ MB agreed with LS Fe(III) dominance (42). The Kβ' amplitude of oxy HB suggested an Fe(d) count of 2, as did our data. The "IS" Fe(II) population in oxy thus seems not to depend on the protein hydration level and persists at ambient temperatures. Resonantly excited Kβ spectra were collected at an energy in the K-preedge region (Fig. 2). The spectra for MB/HB and the porphyrins revealed pronounced shape differences and larger Kβ' amplitudes for decreasing Fe(d) counts. A plot of the Kβ' amplitudes versus the by one unit diminished spin numbers due to a surplus Fe(d) electron showed an approximately linear relation for the porphyrins and positioned deoxy and carboxy MB/HB at values of 3 or 1 for HS or LS Fe(II) (Fig. 3). For oxy HB/MB, the Kβ' amplitudes close to 2 Fe(d) spins indicated a higher spin state than in carboxy. A mean spin count of 2 was calculated for 1s excitation, resulting in a surplus spin in the initially unoccupied d-levels or in spin pairing [Fe(d) = 3 or 1, see SI Appendix for further discussion], which supported an effective IS iron ion in oxy.

X-Ray-Induced Ligation Changes. X-ray fluorescence time traces were recorded at a fixed excitation energy in the K-edge rise to monitor radiation-induced changes at the iron centers (Fig. 4). Similar kinetic behavior was observed for MB and HB. At 20 K, the deoxy K-edge energy remained almost unchanged during ~3.5 min of X-ray exposure because of the absence of photoreduction of the ferrous iron. For carboxy and oxy, a biphasic increase ( $t_{1/2} \sim 5/75$  s or ~10/150 s) of the fluorescence level due to an edge energy decrease to the deoxy level was observed and Kβ main-line emission, preedge absorption, and XANES spectra after ~150 s of X-ray exposure revealed that the resulting states showed similar spectral features as deoxy (SI Appendix, Fig. S12). At 260 K, the deoxy fluorescence decreased to the carboxy level within  $\sim 3.5$  min ( $t_{1/2}$  $\sim$ 25 s) (Fig. 4) and final carboxy formation was revealed by K $\beta$ main-line emission, preedge absorption, and XANES spectra (SI Appendix, Fig. S12). For carboxy, however, the X-ray fluorescence and, thus, the K-edge energy, as well as the Kβ and XANES features remained unchanged for ~3.5 min of X-ray exposure. Notably, porphyrin compound 3 showed no X-ray photoreduction at 20 K and monophasic Fe(III) to Fe(II) reduction at 260 K within ~5 min (SI Appendix, Fig. S12). For oxy, the K-edge energy shifts according to the fluorescence traces and the Kβ/XANES spectra after ~20 s and ~150 s of X-ray exposure suggested initial formation of deoxy ( $t_{1/2} \sim 8$  s) followed by carboxy formation ( $t_{1/2} \sim 18$  s). In particular, the oxy emission revealed an initial increase of the Kβ' amplitude, which was expected for deoxy formation, but not deoxy to carboxy conversion. The 260 K data for oxy were explainable by initial reduction and detachment of the O<sub>2</sub> ligand, followed by binding of a mobile CO species to the vacancy, whereas similarly fast CO binding than detachment explained the invariant data for carboxy. Mobile CO species may be created for example by X-ray induced decarboxylation of amino acid side chains or CO photolysis (46, 47). At 20 K, carbon oxide species were immobile and rebinding was impaired. The biphasic kinetics in carboxy and oxy suggested a similar two-step ligand-centered reduction, which seems incompatible with an initial true superoxide ligand in oxy.

Electronic Structure and Bonding. High-resolution preedge absorption (ctv) and nonresonantly excited Kβ satellite emission (vtc) spectra were collected on MB/HB (SI Appendix, Fig. S13) and the porphyrins. Spectral features were almost identical at 20/260 K and similar for MB/HB, which justified averaging to obtain mean ctv/ vtc spectra of the porphyrins (SI Appendix, Fig. S14) and the heme species (Fig. 5). Subtle ctv differences of MB/HB were emphasized by narrow-band  $K\beta_{1,3}$  emission detection (SI Appendix, Fig. S13). The ctv spectra of the hemes showed two to three peaks varying ~1 eV in energy and ~twofold in intensity. The vtc spectra showed ~1 eV energy differences and  $K\beta_{2.5}$  intensity variations (Fig. 5). ctv

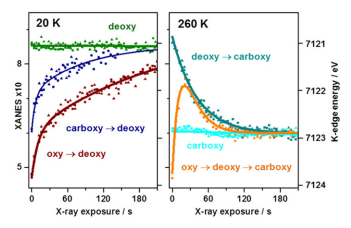
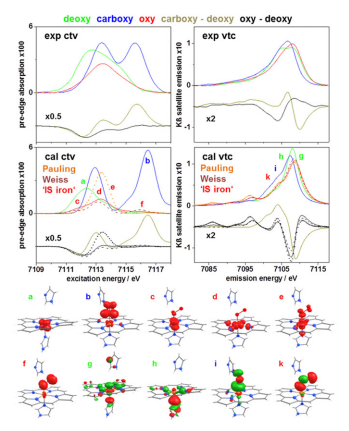


Fig. 4. Heme state conversion during X-ray exposure. Fluorescence changes monitored at 7,123 eV in the XANES (SI Appendix, Fig. S13) for indicated starting species in MB/HB (triangles/squares) at 20/260 K (Left/Right), K-edge amplitudes/energies shown on left/right y axes. Time traces correspond to ctv, XANES, and  $K\beta$  main-line changes (SI Appendix, Fig. S12) due to transitions between heme species. Lines: biexponential (carboxy/oxy, 20 K; oxy, 260 K), monoexponential (deoxy, 260 K), or linear (deoxy, 20 K; carboxy, 260 K) fits.



**Fig. 5.** Electronic excitation and decay transitions. (*Top*) Experimental ctv/vtc spectra (background-corrected mean MB/HB spectra/differences at 20/260K, individual spectra in *SI Appendix*, Fig. S13). (*Middle*) ctv/vtc spectra from (TD)DFT (vtc area-normalized; mean spectra of relaxed MB/HB heme structures correspond to deoxy, HS Fe(II); carboxy, LS Fe(II); and oxy, LS Fe(II) (Pauling), LS Fe(III) (Weiss), or to a mean of 2 Fe(d) spins ('IS iron'; *SI Appendix*, Figs. S17 and S18). (*Bottom*) Representative target/source MOs for transitions in spectral regions a–k.

spectra were collected by using narrow-band (7,045 eV)  $K\beta$ ' emission detection. For the HS porphyrins, in contrast to the  $K\alpha$ -or  $K\beta_{1,3}$ -detected spectra, small ctv amplitudes reflected impaired 1s  $\alpha$ -electron excitation due to the occupied  $\alpha$ -spin valence levels, whereas for the LS porphyrin,  $K\beta_{1,3}$ - and  $K\beta$ '-detected spectra were similar because of similar 1s excitation into unoccupied levels (*SI Appendix*, Fig. S14). MB/HB showed similar  $K\beta$ '-detected ctv spectra (*SI Appendix*, Fig. S15). Negligible or similar ctv amplitudes vs. the  $K\alpha$ -detected spectra agreed with HS Fe(II) in deoxy and LS Fe(II) in carboxy. Smaller amplitudes vs. the  $K\alpha$ -detected spectra indicated a higher or lower spin state in oxy vs. carboxy or deoxy, further supporting two unpaired spins in oxy.

(TD)DFT was used to simulate ctv/vtc spectra. Calculated spectra for the porphyrins matched the experimental data well, which confirmed the spin states, revealed spin density distributions, and attributed target/source molecular orbitals (MOs) to excitation/decay transitions (SI Appendix, Fig. S16). For MB/HB, ctv spectra for HS Fe(II) in deoxy and LS Fe(II) in carboxy for calculations on crystal structures agreed reasonably with the experiment (SI Appendix, Fig. S17) and improvement resulted from geometry optimization (Fig. 5), yielding bond lengths/angles in excellent agreement with EXAFS (SI Appendix, Table S3). For deoxy, the ctv features reflected transitions mostly into Fe(d)centered MOs, and the vtc features for decreasing energies were dominated by decay from MOs centered at iron, histidine, or heme; the spin density was located mostly on iron (SI Appendix, Fig. \$18). For carboxy, two ctv peaks reflected transitions into Fe(d) or CO dominated MOs and increased lower-energy vtc

intensity was due to decay from CO-centered MOs (SI Appendix, Fig. S18). For oxy, depending on whether a nonafc or afc solution (within the "broken-symmetry" DFT formalism) was searched, Pauling or Weiss models emerged and a canonical IS Fe(II) was not produced even for extensive Fe-O2 geometry variation (SI Appendix, Fig. S19). The structures showed similar vtc spectra with increased lower-energy intensity because of decay from O<sub>2</sub>-centered MOs. A single large ctv peak for the Pauling model disagreed with the experiment. Three ctv features for the Weiss model were in better agreement and because of a lowest-energy transition into the Fe(d<sub>yz</sub>) LUMO, intermediate transitions into  $Fe(d_{x-y}^{2})$  and  $Fe(d_{z}^{2})$  dominated MOs, and a highest-energy transition into an O<sub>2</sub>-centered MO (Fig. 5). The ctv/vtc spectra for a mean of 2 Fe(d) spins, as estimated from (TD)DFT calculations for increasing spin states (SI Appendix, Fig. S17), were expected to be compatible with the experimental data (Fig. 5). A method for calculation of ctv/vtc spectra from a CASSCF wavefunction (below) is not available, but the respective MO occupancies suggested that the expected ctv transitions would yield three main ctv features, similar to the TDDFT and experimental spectra of oxy (SI Appendix, Fig. S18).

CASSCF calculations were carried out on porphyrin and heme structures (SI Appendix, Tables S4-S6 and Fig. S20), and spin densities (SD) in the natural molecular orbitals and effective unpaired electron densities (EUED = SD  $\times$  (2 – SD)) (48) on iron and ligands were determined from the wavefunctions. Three doubleoccupied Fe(d) MOs for the porphyrin (1) and heme (carboxy) LS Fe(II) species and four or five single-occupied MOs for the Fe(II) (deoxy, 2) or Fe(III) (3) HS species, and almost integer EUEDs on iron (0, 4, or 5) were in full agreement with the formal redox/spin states (Table 1). The oxy wavefunction was dominated (~90%) by two contributions, both with unoccupied (SD < 0.05) Fe( $d_z^2$ ) and  $Fe(d_{x-y}^{2/2})$  MOs and a double-occupied MO representing a polarized dative  $\sigma(\text{Fe-O})$  bond with major O(p) and weak  $Fe(d_z^2)$ contributions (Fig. 6). MOs representing  $\pi$  and  $\pi^*$  Fe-O (back-)bonds showed dominant Fe(d<sub>vz</sub>) and weaker O(p) contributions and double/zero or zero/double occupancy in the two configurations; a double-occupied MO with dominant Fe(d<sub>xz</sub>) and minor O(p) contributions indicates further delocalized  $\pi(\text{Fe-O})$  backbonding (SI) Appendix, Table S5). The SD in the two  $Fe(d_{vz})$ -dominated MOs was  $\sim 1.3$  and  $\sim 0.7$  (EUED  $\sim 0.89$  and  $\sim 0.91$ ), revealing essentially two spatially separated unpaired spins in oxy, independent of, e.g., moderate Fe-O<sub>2</sub> bond length variations (*SI Appendix*, Fig. S20). The latter values together with the small EUEDs in the other Fe-assigned MOs yielded a total EUED on iron of ~2.0, which summed up with the EUED on  $O_2$  (~0.3) to a value of ~2.3, close to the EUED ( $\sim$ 2.4) for the ozone radical ( $O_3^+$ ) (48).

### Discussion

We used Fe  $K\beta$ -XES and K-edge XAS as well as quantum chemistry to study molecular and electronic structures of heme

Table 1. Effective unpaired electron density from CASSCF

Species	Redox and spin state	EUED	
		Fe	Ligand*
Heme			
deoxy	Fe(II) HS	4.000 (4)	_
carboxy	Fe(II) LS	0.001 (0)	0.352
оху	Fe(II) IS <sup>†</sup>	2.014 (2)	0.302
Porphyrin			
1	Fe(II) LS	0.000 (0)	0.000
2	Fe(II) HS	4.000 (4)	0.092
3	Fe(III) HS	5.000 (5)	0.001

\*Axial CO (carboxy),  $O_2$  (oxy),  $N_{Py}$  (1),  $O_{THF}$  (2), or  $Cl^-$  (3) ligand on top of the porphyrin in Fig. 1 and *SI Appendix*, Fig. S14.

<sup>†</sup>Simplified annotation (see text); formal unpaired Fe(d) spins in parentheses. The EUED on the ring systems in all heme and porphyrin species was zero.

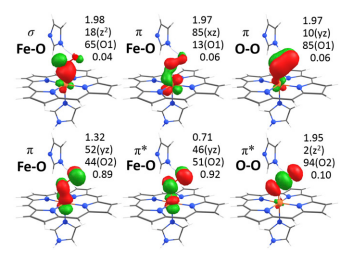


Fig. 6. Iron-oxygen bonding in oxy heme. Indicated spin densities, Fe(d) and O(p) characters (main contribution in parenthesis, O1 is Fe-bound), and EUEDs refer to (clockwise) MOs 2, 4, 3, 5, 7, 6 from CASSCF (14, 11) calculations (Table 1 and SI Appendix, Fig. S20).  $\sigma$ -donation and delocalized  $\pi/\pi^*$  backbonding stabilize the O<sub>2</sub> ligand. For further analysis, see text and SI Appendix, Table S5.

species in MB and HB solutions. Fast spectral changes for all species even at cryogenic temperatures mandated rapid-scan XAS and energy-sampling XES approaches for unperturbed data collection. Apparent bond lengths and angle variations particularly in crystal structures of oxy globins likely relate to such radiationinduced modifications (41). We interpret our data as sequential ligand-centered two-electron reduction (at 20 K) of CO or O<sub>2</sub> and product detachment to form apparent deoxy heme (46, 47, 49). The similar kinetic behavior of carboxy and oxy implies similar electron uptake capability of both ligands, which is in agreement with the absence of a superoxide ligand, thereby supporting our computational results and the redox/spin state assignment for oxy.

Similar metrical, electronic, and kinetic parameters indicated similar metal site configurations in MB and HB. A significant influence of amino acid environment or hydrogen-bonding variations in the two heme proteins on these parameters of the iron centers and on the ligand bonding is unlikely, which agrees with earlier spectroscopic studies (50-53). Heme side-chain orientation differences in MB/HB presumably were reflected in the XANES by slight virtual level, i.e., Fe(4p), energy changes (54). All relevant heme features in MB/HB were invariant within 20-260 K, excluding population of higher spin states even far above the protein glass-transition point (~200 K) (55). This finding and, e.g., gradual spin crossover in iron porphyrins (56–58) suggest that also at room temperature, the spin state in the three heme species is the same and, therefore, physiologically relevant.

Kβ spectra of MB/HB and porphyrin compounds revealed a direct correlation of the formal unpaired Fe(d) spin count to the relative Kβ' intensity, in agreement with results for other iron species (32, 34), establishing further that the iron-centered radiative 3p→1s decay is a reliable probe of the valence level configuration. The steep correlation locates the overall iron bonding in the MB/HB hemes and the porphyrins closer to the noncovalent than covalent limit (32, 34, 59, 60). Agreement of our and earlier Kβ data of MB/HB (45) supports the spin state changes. For the hemes and porphyrins, our Kβ main-line, ctv/vtc, and (TD)DFT/ CASSCF data revealed ferrous HS (deoxy, 2) or LS (carboxy, 1) and ferric HS (3) species (1, 40), emphasizing the method's sensitivity for valence and spin-state discrimination. Further discussion of relations between the XAS/XES data and iron spin states is given in SI Appendix.

For oxy, the Kβ spectra indicated two unpaired Fe(d) spins on iron, which was compatible with the ctv/vtc data, by definition implying an IS metal center, as suggested by McClure/Goddard (15, 16). No significantly decreased ligand character of Fe(d) levels in oxy vs. deoxy/carboxy in the calculations rendered diminished iron-porphyrin bonding covalency as an explanation for the Kβ data of oxy unlikely. The Pauling model with a LS Fe(II) and an innocent O<sub>2</sub> ligand (13) was essentially excluded (and higher spin states, M > 1) by our data, strengthening previous conclusions (36, 40). DFT converged to the Weiss model with a ferric iron and a superoxide ligand (17) and yielded ctv/vtc spectra more close to our and earlier data (36, 40), but this preference reflected overestimation of metal-to-ligand charge transfer (22, 23). CASSCF on oxy vielded a two-component wavefunction as found earlier (21– 23). It comprised in effect two unpaired spins on iron, in perfect agreement with our  $K\beta$  data, thereby benchmarking the quantum chemical prediction and providing a quantitative link between theory and spectroscopy. The emerging bonding situation includes an essentially ferrous iron center, little charge transfer to O<sub>2</sub> and minor superoxide character of the noninnocent ligand, significant double-bond properties of the Fe-O<sub>2</sub> interaction, and three-center ozone-like electron delocalization. It combines the classical models into a more realistic view of O<sub>2</sub> bonding in MB/HB, featuring limited radical character and delocalized backbonding stabilization of the ligand as prerequisites for reversible O<sub>2</sub> transport.

### **Materials and Methods**

Sample Preparation. Porphyrin compounds (Fe(TPP)(Py)2, LS Fe(II), 1; Fe(TPP)(THF)2, HS Fe(II), 2; and Fe(TPP)Cl, HS Fe(III), 3; TPP, tetraphenylporphyrin; Py, pyridine; THF, tetrahydrofurane) were synthesized as described (40, 61) and grinded with solid BN (wt/wt 1:10). Bovine HB and equine-heart MB (met) were purchased from Sigma-Aldrich, at 12 mM heme dissolved in buffer A (100 mM Tris-HCl, 150 mM NaCl, 2.5 mM EDTA, pH 8), and reduced with sodiumdithionite (50 mM) to yield deoxy. MB/HB deoxy solutions (350  $\mu$ L) were gently stirred under pure CO or O2 gas for 10 min to yield carboxy or oxy (62). Aliquots (50 µL) were loaded in Kapton-covered acrylic-glass holders and frozen in liquid nitrogen. Samples were handled in a glove box (99/1% dry N<sub>2</sub>/H<sub>2</sub> atmosphere). Optical absorption spectra of HB/MB (12 µM in buffer A under each atmosphere in a sealed 1-cm cuvette) were collected on a Shimadzu UV-2450 spectrometer.

X-Ray Spectroscopy. High-resolution Fe K-edge XANES/Kβ-XES spectra were collected at beamline ID26 at European Synchrotron Radiation Facility (ESRF) (Grenoble, France) by using established methods (see SI Appendix and ref. 34 for details). Rapid-scan (~1 s) XANES spectra were obtained in transmission mode (porphyrins) and by broad-band total (Kα) (scintillation detector) or narrow-band Kβ fluorescence detection (HB/MB). XES data were collected on a vertical-plane Rowland-circle spectrometer (5 spherical Ge[620] analyzers at R = 1,000 mm) with an avalanche photodiode detector (34). The X-ray beam was attenuated by Al foils. Kß spectra were collected by using an energysampling approach (~1 s data acquisition per 0.35-eV step; SI Appendix, Fig. S1) (44). Monochromator and emission spectrometer energies were calibrated with ±0.1-eV precision (44). Lower-resolution XANES/EXAFS spectra were collected at beamline KMC-3 at Berlin Electron Storage Ring Society for Synchrotron Radiation (BESSY) (Helmholtz-Center Berlin; SI Appendix) (63). Signal-to-noise ratios were improved by averaging of spectra (XAS, ≤150 scans; XES, ≤10 point-by-point datasets) from separate sample spots. XAS/XES spectra were processed as described in ref. 34. Kβ' amplitudes were derived from data within 7,044-7,046 eV. EXAFS spectra were simulated with in-house software and phase functions from FEFF9 ( $S_0^2 = 0.85$ ) (64).

Quantum Chemistry. Calculations at (TD)DFT and Møller-Plesset second-order perturbation theory/complete-active-space self-consistent-field calculation theory levels on porphyrin and heme structures were performed with ORCA (65) and Gaussian09 (66) for geometry-optimization and spectral calculations [for (TD)DFT: B3LYP functional, def2-TZVP basis set; COSMO solvation model] (67) (SI Appendix, Datasets S1 and S2). For comparison with experimental data, ctv/vtc stick spectra from (TD)DFT were broadened and scaled (c2v, Gaussian FWHM = 0.5 eV, x0.025; v2c, Lorentzian FWHM = 3.5 eV, x0.5) and 152 eV shifted. For further computational details see SI Appendix.

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