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Introduction 1

Nitrogenases uniquely reduce atmospheric inert dinitrogen (N₂) to bioavailable ammonia (NH₃) under ambient conditions, providing nearly two-thirds of the worldwide nitrogen resource.1 A catalytic reaction occurs at the dinitrogenase component, while a dinitrogenase reductase provides the required electrons and is also the site of adenosine triphosphate (ATP) hydrolysis to drive the reaction.² Three homologous nitrogenase isoforms, Mo-, V- and Fe-nitrogenases, are known to date, which primarily differ in the architecture of their catalytically active sites, named M-clusters or FeMo-/FeV-/FeFe-cofactors (FeMo-/FeV-/FeFe-cos), respectively.³⁻⁹ Their N₂-reducing activities decrease from MoFe, VFe and FeFe proteins sequentially, and the latter two are thus generally regarded as backup systems.10,11 Three nitrogenases catalyze the reactions as follows:12,13

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Mo-nitrogenase:

 $N_2 + 8e^- + 8H^+ + 16MgATP \rightarrow 2NH_3 + H_2 + 16MgADP + 16P_i$

V-nitrogenase:

$$\begin{array}{l} N_2+18e^-+18H^++36MgATP\rightarrow 2NH_3+6H_2+36MgDP\\ \qquad +36P_i \end{array}$$

Fe-nitrogenase:

 $N_2 + 20e^- + 20H^+ + 40MgATP \rightarrow 2NH_3 + 7H_2 + 40MgADP$ $+40P_{i}$

There are three recognized reversible states of the P-cluster in nitrogenase: the reduced P^N state, the one electron-oxidized P^{1+} state, and the two electron-oxidized P^{2+} state.⁸ It has been proposed that electrons for N2 reduction are transferred from the P-cluster to FeMo/FeV/FeFe-cos.¹⁴⁻¹⁶ The formal electronic configuration of FeMo-co in the resting state has been assigned to Mo³⁺-3Fe²⁺-4Fe³⁺ by spectroscopic studies and theoretical calculations,^{17–19} while a reduced oxidation state of FeV-co is suggested as V^{3+} -3Fe²⁺-4Fe³⁺.²⁰ Although Fe-only nitrogenase has been studied extensively,²¹⁻²⁵ the first crystallographic structure was isolated recently from Azotobacter vine*landii* (PDB code *8BOQ*) and determined by X-ray diffraction. The cryogenic electron microscopy of Fe-only nitrogenase from Rhodobacter capsulatus was also presented (PDB code 80IE).²⁶

Analyses of the electronic structures of FeFe-cofactors compared with those of FeMoand FeV-cofactors and their P-clusters[†]

Zhen-Lang Xie, 🕩 a Wan-Ting Jin^b and Zhao-Hui Zhou 🕩 a

The electronic structures of FeFe-cofactors (FeFe-cos) in resting and turnover states, together with their P^N clusters from iron-only nitrogenases, have been calculated using the bond valence method, and their crystallographic data were reported recently and deposited in the Protein Data Bank (PDB codes: 8BOQ and 8OIE). The calculated results have also been compared with those of their homologous Mo- and V-nitrogenases. For FeFe-cos in the resting state, Fe1/2/4/5/6/7/8 atoms are prone to Fe³⁺, while the Fe3 atom shows different degrees of mixed valences. The results support that the Fe8 atom at the terminal positions of FeFe-cos possesses the same oxidation states as the Mo³⁺/V³⁺ atoms of FeMo-/FeV-cos. In the turnover state, the overall oxidation state of FeFe-co is slightly reduced than those in the resting species, and its electronic configuration is rearranged after the substitution of S2B with OH, compatible with those found in CO-bound FeV-co. Moreover, the calculations give the formal oxidation states of 6Fe²⁺-2Fe³⁺ for the electronic structures of P^N clusters in Fe-nitrogenases. By the comparison of Mo-, V- and Fe-nitrogenases, the overall oxidation levels of 7Fe atoms (Fe1-Fe7) for both FeFe- and FeMo-cos in resting states are found to be higher than that of FeV-co. For the P^N clusters in MoFe-, VFe- and FeFeproteins, they all exhibit a strong reductive character.



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[†]Electronic supplementary information (ESI) available: Details of bond distances and valence calculations of PDB codes 8BOQ and 8OIE. See DOI: https:// doi.org/10.1039/d3dt04126c

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FeFe-co was therefore identified as $Fe_8S_9C[R-(H)$ homocitrate],^{7,26} which carries an organic homocitrate ligand that is strikingly similar to those observed in FeMo- and FeVcos.^{4,27,28} Intriguingly, Fe-only nitrogenase can retain its reactivity towards N₂ without the exchange of the apical iron of the M-cluster with heterometals Mo or V.^{7,26} In particular, recent work has shown that Fe-only nitrogenase has the unique ability to produce significant quantities of methane (CH₄) by CO₂ reduction, not shared by the other two homologous nitrogenases.²⁹

However, the electronic structures of FeFe-co and the P-cluster in Fe-nitrogenases are still uncertain, and their oxidation states are crucial for understanding the potential electron transfer sites and pathways. This prompts us to apply a classical bond-valence method for analysing the oxidation states of FeFe-cos and their P-clusters, where the data are taken from the crystal structures of Fe-nitrogenases in the Protein Data Bank (PDB codes *8BOQ* and *8OIE*). Moreover, the resolution of *8BOQ* is high at 1.55 Å, which indicates good reliability of the bond distances with a high weight scheme in calculations.

The bond valence method is a simple but powerful approach for determining the oxidation states of metal ions based on the bond distances of the metal-ligand from their crystal structures.^{30,31} It was initially used to analyse inorganic crystal structures,³²⁻³⁴ and was gradually applied to other fields, especially in the field of high-resolution protein structures.19,35-39 In contrast to density functional theory (DFT) calculations, whose reliability in nitrogenase models has been questioned by different results,40 the BVS method is much more convenient for studying metalloenzyme systems without considering the spin state of metal ions. Herein, we have employed the BVS method for comparing the resting and turnover states of FeFe-cos in Fe-nitrogenases, trying to figure out the electron distributions of different metal atoms in iron-only nitrogenases and assess the rationality of their structures with the P-cluster. Only P^N states of P-clusters are found in FeFe protein in the PDB codes 8BOQ and 8OIE. Therefore, all the following calculations involve P^N clusters alone. To further investigate the nitrogen fixation mechanism with different nitrogenases, we have also performed comprehensive comparisons among FeMo-, FeV- and FeFe-cos, together with their P^N clusters in nitrogenases.

2. Calculation method

2.1 Criteria for valence assignment

Bond valence sums (BVS) were calculated according to eqn (1)-(3):

$$S_{i} = \sum_{j} \exp[(R_{0} - r_{ij})/B]$$
(1)

$$d = S_{i} - n \tag{2}$$

$$S_{\rm t} = \sum_i S_{\rm i} \tag{3}$$

In the FeFe-co and P^{N} cluster of Fe-nitrogenases, r_{ij} is the bond length between metal *i* (Fe) and coordinated atom *j* (S/O/

N/C), measured by Pymol from the crystal structures of FeFe proteins deposited in the Protein Data Bank (PDB). B is considered a constant and is equal to 0.37 Å.³² Here, R₀ values used for the calculations of FeFe-co and the P^N cluster in Fenitrogenases are listed in Table 1. They were determined empirically so that the BVS is generally quite close to the oxidation states of the metal ions.35 Least-squares fitting procedures were used to determine the initial set of R_0 values by minimizing the differences between the BVS and the known oxidation states for crystallographically characterized materials. Most R_0 values can be found on the web and relevant studies.⁴² S_i refers to the calculated bond valence sum of each Fe ion. The d value allows us to assess the deviation between calculated valence S_i and assumed valence n. The absolute deviation |d| can serve as an evaluation index of the calculated result. Some valences calculated from either R_0 (+2) and R_0 (+3) might give a similar |d| value due to the electron delocalization in FeFe-cos and the P^N cluster.¹¹ In this case, we assume the existence of mixed valences instead of integral valences. If the |d| values calculated from R_0 (+2) and R_0 (+3) differ significantly, the assignment of the oxidation state of the Fe atom can be associated with the smallest |d| value. Although the Thorp group has stated that BVS values calculated from Brown's lengths were trustworthy within ±0.25 units,43 electron delocalizations in FeFe-cos and the P^N cluster require a wider error-tolerance range than those in inorganic crystals. Therefore, we established an acceptable D value of 0.30 as a valence assignment criterion to distinguish integral or mixed valences.^{39,44} If the absolute deviation |d| is less than the *D* value, then iron is assumed to be Fe²⁺ or Fe³⁺; otherwise the valence is designated as indeterminate mixed valence Fe^{n+} (2 < *n* < 3).

 $S_{\rm t}$ in eqn (3) represents the calculated valence sum of all eight Fe atoms Fe1–Fe8 in FeFe-co and the P^N cluster (abbreviated as 8Fe). The valences of 8Fe are compared with the values of 16 and 24, which correspond to the valences of eight all-ferrous and all-ferric iron atoms, respectively. The average value of eight Fe atoms Fe1–Fe8 is denoted as 8Fe_{av}. The detailed bond valence calculations of FeFe-co and the P^N cluster (PDB codes: *8BOQ* and *8OIE*) are given in Tables S1–S10.†

2.2 Weight schemes for the different resolutions of FeFe proteins

The bond valence method requires highly accurate bond distance data.³⁶ To account for variations in the resolutions of crystallographic data, we have used a weighting scheme based on the *d*-spacing resolutions of structures in the PDB to calcu-

Table 1 The R_0 values correspond to different types of bonds with Fe^{*n*+} ions

M-L bonds	R_0 (Å)	M–L bonds	R_0 (Å)
$Fe^{2+}-C$ $Fe^{2+}-N$ $Fe^{2+}-S$ $Fe^{2+}-O$	$1.650^{38} \\ 1.769^{46} \\ 2.120^{42} \\ 1.734^{32}$	Fe ³⁺ -C Fe ³⁺ -N Fe ³⁺ -S Fe ³⁺ -O	$1.689^{45} \\ 1.815^{46} \\ 2.149^{45} \\ 1.759^{32}$

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Fig. 1 The resting states of (a) FeMo-co, (b) FeV-co and (c) FeFe-co (PDB codes: *3U7Q*,⁴ *5N6Y*⁶ and *8BOQ*⁷), respectively. Color code: Fe, brown; Mo, teal; V, violet; S yellow; O red; N blue; C grey; H, turquoise. Hydrogen atoms on homocitrates are added based on the vibrational circular dichroism (VCD) experiment for the FeMo-cofactor.⁴¹

late the bond valence of each iron S_i in all calculated FeFe-co or the P^N cluster. The smaller the value of the *d*-spacing resolution limit A_i , the larger the weight w_i . As a result, we have revised the inverse distance weighted (IDW) interpolation method for assigning multiple weights:

$$w_{i} = A_{i}^{-p} / \sum_{i}^{N} (A_{i}^{-p})$$
 (4)

$$\overline{S_{w}} = \sum_{i}^{N} S_{i} w_{i} / \sum_{i}^{N} w_{i} = \sum_{i}^{N} S_{i} w_{i}$$
(5)

Eqn (4) is of the same general form as the IDW principle.⁴⁷ The bond valence S_i calculated from a higher resolution A_i (*e.g. 8BOQ* at 1.55 Å) has greater effects on weighted average valences than that from a lower resolution (*e.g. 80IE* at 2.35 Å). $\overline{S_w}$ in eqn (5) is obtained from the weighted average of calculated valences S_i of different Fe atoms from all analysed FeFeco or the P^N cluster. N is the number of samples containing FeFe-co or the P^N cluster of two FeFe proteins. $\sum w_i$ is actually equal to 1 in this equation. *p* is an index parameter set by the user.^{48,49} We have previously set p = 1,^{39,44} which would generate concordant w_i with balanced distribution in the P-clusters of MoFe proteins. Here, we also use p = 1 to calculate FeFe proteins for further comparisons with MoFe and VFe proteins.

3. Results and discussion

3.1 Valence analyses of iron in FeFe-co

The $F_o - F_c$ difference in the electron density map of FeFe proteins (PDB code: *8BOQ*) revealed dual conformations, which are in an equilibrium of resting and turnover states.⁷ In the resting state conformation, Fe2 and Fe6 are bridged by S2B, and Gln176^D points away from S2B, with its amide N ε_2 weakly hydrogen-bonded to homocitrate. In the turnover state conformation, residue Gln176^D rotates its side chain to accept a hydrogen bond from the N ε_2 atom of the imidazole side chain of His180D to the O $_{e1}$ atom, forming the turnover state. That is, in the turnover state, S2B in FeFe-co is replaced by OH, which is suggested to represent a reaction intermediate, similar to the *6FEA* structure with the OH group in the FeV- cofactor.⁵⁰ Thus, in the PDB code *8BOQ*, two FeFe-cos are in mixed states (resting and turnover states). In contrast, two FeFe-cos in the PDB code *80IE* are in resting states only.

Fig. 2a and b show the absolute deviations |d| between the calculated and the expected valences of the weighted average value of eight Fe atoms (8Fe_{av}) and individual Fe atoms (Fe1–Fe8) in the resting and turnover states of FeFe-cos, respectively. Fig. 2c and d exhibit the valence distributions of Fe atoms in the resting and turnover states of FeFe-cos. Four FeFe-cos in the resting states from the PDB codes *8BOQ* and *8OIE* were calculated by weighted average values, and two FeFe-cos in the turnover state from the PDB code *8BOQ* were calculated by an average value. Two groups of errors are generated from the parameters R_0 (+2) (green) and R_0 (+3) (orange), respectively. Table 2 lists the weighted average valences and absolute deviations of iron atoms calculated from R_0 (+2) and R_0 (+3) in FeFe-cos. In conjunction with Fig. 2a and Table 2, the two sets of |d| values calculated using R_0 (+2) and R_0 (+3) can be clearly



Fig. 2 (a) The weighted average |d| values of 8Fe_{av} and Fe1–Fe8 atoms calculated from R_0 (+2) and R_0 (+3) for resting states in the FeFe-co (PDB codes: *8BOQ* and *8OIE*). (b) The average |d| values of 8Fe_{av} and Fe1–Fe8 atoms calculated from R_0 (+2) and R_0 (+3) for the turnover state in FeFe-co (PDB code: *8BOQ*). The red dashed line represents D = 0.3. The valence distributions of Fe atoms in resting (c) and turnover (d) states in FeFe-co. Red, orange and green balls represent n values of 2.7 < $n \le 3.0$, 2.3 < $n \le 2.7$, and 2.0 < $n \le 2.3$ for Feⁿ⁺¹, respectively.

Table 2	The valences of Fe1–Fe8 atoms and	8Fe calculated from R_0	(+2) and R ₀ (+3) fo	or resting states (PD	DB codes: 8BOQ and 8OI	E) and turnover
states (PI)B code: 8BOQ) in FeFe-cos					

				$R_0(+2)$									
PDB codes	States	Res (Å)	w_{i}	Fe1	Fe2	Fe3	Fe4	Fe5	Fe6	Fe7	Fe8	8Fe	8Fe _{av}
FeFe-co													
$8BOQ^7$	Resting	1.55	0.301	2.627	2.532	2.390	2.534	2.594	2.763	2.466	2.721	20.627	2.578
$8BOQ^7$	Resting	1.55	0.301	2.598	2.525	2.425	2.478	2.521	2.680	2.508	2.415	20.150	2.519
$8OIE^{26}$	Resting	2.35	0.199	2.625	2.616	2.548	2.418	2.498	2.807	2.712	2.536	20.760	2.595
80IE ²⁶	Resting	2.35	0.199	3.358	2.469	2.693	2.772	2.646	2.430	2.569	2.547	21.484	2.686
$\overline{S_w}$				2.763	2.534	2.492	2.541	2.563	2.681	2.548	2.557	20.679	2.585
Weighted $ d $				0.763	0.534	0.492	0.541	0.563	0.681	0.548	0.557	4.632	0.585
8BOQ ⁷	Turnover	1.55		2.627	2.298	2.390	2.534	2.594	2.441	2.466	2.721	19.909	2.509
$8BOQ^7$	Turnover	1.55		2.598	2.546	2.425	2.478	2.521	2.222	2.508	2.415	19.713	2.464
Average				2.613	2.422	2.408	2.506	2.558	2.332	2.487	2.568	19.894	2.487
d				0.613	0.422	0.408	0.506	0.558	0.332	0.487	0.568	3.894	0.737
				$R_0(+3)$									
PDB codes	States	Res (Å)	w _i	Fe1	Fe2	Fe3	Fe4	Fe5	Fe6	Fe7	Fe8	8Fe	8Fe _{av}
FeFe-co													
$8BOQ^7$	Resting	1.55	0.301	2.841	2.751	2.596	2.753	2.817	2.999	2.678	2.949	22.384	2.798
$8BOQ^7$	Resting	1.55	0.301	2.810	2.742	2.634	2.692	2.737	2.910	2.724	2.620	21.869	2.734
$8OIE^{26}$	Resting	2.35	0.199	2.839	2.767	2.626	2.840	3.047	2.945	2.713	2.740	22.519	2.815
80IE ²⁶	Resting	2.35	0.199	2.799	2.682	2.925	3.010	2.873	2.640	2.791	2.752	22.471	2.809
$\overline{S_w}$	-			2.823	2.738	2.679	2.803	2.850	2.890	2.721	2.769	22.273	2.784
Weighted $ d $				0.177	0.262	0.321	0.197	0.150	0.110	0.279	0.231	1.779	0.216
8BOQ ⁷	Turnover	1.55		2.841	2.491	2.596	2.753	2.817	2.645	2.678	2.949	21.595	2.721
$8BOQ^7$	Turnover	1.55		2.810	2.756	2.634	2.692	2.737	2.410	2.724	2.620	21.383	2.673
Average				2.826	2.624	2.615	2.722	2.777	2.528	2.701	2.785	21.578	2.697
d				0.174	0.376	0.385	0.278	0.223	0.472	0.299	0.215	2.422	0.303

distinguished for Fe1, Fe4, Fe5, Fe6 and Fe8 atoms in the resting state. The assignments of Fe³⁺ for these atoms are reliable with small errors (0.110 < |d| < 0.249), indicating a good fit to the parameters and credible calculated results. Similarly, Fe2 and Fe7 atoms may be favourable for Fe³⁺, due to their |d| values calculated from R_0 (+3) below the valence assignment criteria D value of 0.30. The Fe8 atom is coordinated by R-(H)homocitrate and residue His423D, and the weighted average bond lengths of Fe8-O5 (2.217av Å), Fe8-O7 (2.160_{av} Å) and Fe8-N (2.416_{av} Å) are close than those of heterometals (Mo³⁺ and V³⁺) in Mo- and V-nitrogenases.^{4,6,7} The designation of Fe8 as Fe³⁺ with a small error (0.231) in the R_0 (+3) group in the resting states is consistent with the oxidation states of Mo³⁺/V³⁺ in FeMo-/FeV-cos.¹⁹ This means that the types of metallic elements have no obvious effect on the oxidation states of the metal (Mo/V/Fe) at the end positions of the cofactors. Thus, the octahedral Fe8 atom probably serves a similar purpose as Mo³⁺ or V³⁺ in the other isoforms.⁷ Importantly, Fe6 is identified as the most oxidized site in the ground state with a calculated value of 2.955_{av} from R_0 (+3) in the PDB code 8BOQ, which is in accordance with spatially resolved anomalous dispersion analysis and BVS calculations of FeMo-co,^{18,19} and thus Fe6 is likely a site of reduction from the P-cluster.^{18,19} In contrast, the dots calculated from R_0 (+2)

calization. With respect to the total eight iron atoms in FeFeco, the 8Fe_{av} values are 0.585 and 0.216 calculated from R_0 (+2) and R_0 (+3), respectively. The parameter R_0 (+3) gives a more reliable result with a small error, suggesting more oxidative iron segments in the resting states of FeFe-co. The metal centers in the resting states could be formally equivalent to Fe^{2.5+}–7Fe³⁺. Additionally, Fe1, Fe2, Fe4, Fe5, Fe6, Fe7 and Fe8 are more likely to have a higher oxidation state of +3, whereas Fe3 is more reduced and exhibits some degree of mixed valence. The calculations for the crystal structures of FeFe-co in the resting state seems not to be the same as the traditional view that FeFe-co is "4Fe²⁺–4Fe³⁺", which is based on the analyses conducted in reductive solutions with excess dithionitereduced agents.⁵¹

In the turnover state, the sulfide S2B was replaced by an OH group. As shown in Fig. 1b, calculations for Fe2 and Fe6 atoms in the turnover state show that the |d| values are 0.422 and 0.332 calculated from R_0 (+2), and 0.376 and 0.472 calculated from R_0 (+3), respectively. The two sets of errors for Fe2 and Fe6 atoms are larger than 0.30. Thus, neither the assumed valences +2 nor +3 can be reliably assigned for the oxidation states of these Fe atoms, indicating that some degree of mixed valences exists among these metal centres (Fig. 1d). For the total of eight iron atoms, the average values $8Fe_{av}$ of |d| are 0.487 and 0.303 calculated from R_0 (+2) and R_0 (+3), implying the presence of electron delocalization in the iron segments of FeFe-co in the turnover state. The total formal oxidation states of 8Fe in the turnover state of FeFe-co could be equal to $2Fe^{2+}$ -

and R_0 (+3) for Fe3 are 0.492 and 0.321, respectively, which are

above the valence assignment criteria D value of 0.30 (red

dashed line in Fig. 2a). Thus, as depicted in Fig. 2c, the Fe3

atom may perform as mixed valence with strong electron delo-

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 $6Fe^{3+}$ or Fe^{2+} – $7Fe^{3+}$, where $S_t = 22$ calculated from $2Fe^{2+}$ – $6Fe^{3+}$ is close to 21.578 calculated from R_0 (+3).

Overall, we have found that FeFe-co in the turnover state is more reduced than that in the resting state in he PDB code *8BOQ*. The electron distributions in Fe2 and Fe6 atoms have been rearranged with the change of the structure. OH-bound sites Fe2 and Fe6 undergo reduction and exhibit mixed valence in the turnover state. Electron transfer from the Fe protein promotes FeFe-co from the resting state E_0 to the E_1 state, which leads to the reduction of Fe6, weakening its interaction with S2B, and then receives the proton from His180.^{7,52} Based on the above analyses, the electron configurations of FeFe-co in the turnover state seem to be more complicated and disturbed by the replacement of the bridging sulfur S2B.

3.2 Valence analyses of iron in the P^N clusters in FeFe proteins

Fig. 3a presents the weighted average absolute deviations for $8Fe_{av}$ and individual iron atoms between calculated and expected valences in the P^N cluster. Fig. 3b shows the valence



Fig. 3 (a) The weighted average |d| values of 8Fe_{av} and Fe1–Fe8 atoms in P^N clusters (PDB codes: *8BOQ* and *8OIE*) calculated from R_0 (+2) and R_0 (+3). Red dashed lines represent D = 0.3. (b) The valence distributions of Fe atoms in P^N clusters of FeFe proteins.

distributions of Fe atoms in P^N clusters. Table 3 shows the weighted average calculated valences of iron atoms using R_0 (+2) and R_0 (+3). As shown in Fig. 3a, it can be seen that Fe1, Fe2, Fe5 and Fe6 atoms can be clearly distinguished between the two groups of errors calculated from either R_0 (+2) and R_0 (+3), and the electrons are well localized in these iron atoms. The one-side match means that +2 valences are more suitable for Fe1/2/5/6. In contrast, the |d| values of the two sets of R_0 (+2) and R_0 (+3) for Fe3, Fe4, Fe7 and Fe8 are almost similar, which suggests that Fe3/4/7/8 atoms have higher oxidation states than Fe(π) with mixed valences.

Regarding the total eight iron atoms in the P^N cluster, the 8Fe_{av} value of |d| in the R_0 (+2) group is 0.315, if we assume that the P^N clusters are all-ferrous, while the corresponding |d|in the R_0 (+3) group is 0.505 when the P^N clusters are supposed to be all-ferric. The smaller error calculated from R_0 (+2) suggests that the P^N clusters in FeFe proteins may exhibit strong reductive character, similar to our previous calculations for the P^N clusters in MoFe and VFe proteins.³⁹ The weighted average valence sum of 8Fe calculated by R_0 (+2) is 18.519 instead of 16, resembling the calculated value for the synthetic model compound of the P^N cluster (CSD code DUGNEZ, S_t = 18.373).⁵³ Therefore, we deduce that the oxidation states of the eight irons in the P^N clusters of FeFe proteins are not allferrous, but contain four Fe²⁺ atoms and four mixed-valence irons. The formal oxidation states for 8Fe could be approximately equal to 4Fe^{2.5+}-4Fe³⁺ or 6Fe²⁺-2Fe³⁺ with delocalized electrons.

As mentioned above, the Fe6 atom undergoes a reduction in the turnover state of FeFe-co, which may be due to the electron transfer from the P^N cluster to FeFe-co. Judging from the spatial positions of the FeFe-co and P^N cluster in the PDB code *8BOQ* shown in Fig. 4a, terminal Fe3 is the closest atom among the eight iron atoms in the P^N cluster to Fe6 of the

Table 3 The valences of Fe1–Fe8 atoms and 8Fe of the P^N cluster (PDB codes: 8BOQ and 8OIE) calculated from R_0 (+2) and R_0 (+3)

			$R_0(+2)$									
PDB codes	Res (Å)	w _i	Fe1	Fe2	Fe3	Fe4	Fe5	Fe6	Fe7	Fe8	8Fe	8Fe _{av}
P-cluster												
$8BOQ^7$	1.55	0.301	2.279	2.088	2.634	2.452	2.146	1.881	2.414	2.460	18.345	2.294
8BOQ ⁷	1.55	0.301	2.067	2.208	2.530	2.344	2.274	1.873	2.430	2.532	18.259	2.282
$80IE^{26}$	2.35	0.199	2.195	2.299	2.186	2.353	2.386	2.394	2.458	2.540	18.812	2.351
80IE ²⁶	2.35	0.199	2.189	2.228	2.291	2.616	2.450	2.468	2.347	2.276	18.864	2.358
$\overline{S_w}$			2.181	2.194	2.445	2.433	2.293	2.097	2.415	2.461	18.519	2.315
Weighted d			0.181	0.194	0.445	0.433	0.293	0.097	0.415	0.461	2.519	0.315
			$R_0(+3)$									
PDB codes	Res (Å)	w_{i}	Fe1	Fe2	Fe3	Fe4	Fe5	Fe6	Fe7	Fe8	8Fe	8Fe _{av}
P-cluster												
$8BOO^7$	1.55	0.301	2.465	2.258	2.848	2.652	2.321	2.035	2.611	2.660	19.850	2.481
8BOO ⁷	1.55	0.301	2.236	2.388	2.737	2.535	2.459	2.025	2.629	2.739	19.748	2.469
80IE ²⁶	2.35	0.199	2.374	2.486	2.364	2.545	2.581	2.593	2.659	2.747	20.349	2.544
80IE ²⁶	2.35	0.199	2.368	2.409	2.478	2.829	2.650	2.320	2.538	2.461	20.053	2.507
S_{w}			2.359	2.373	2.645	2.631	2.480	2.200	2.611	2.662	19.961	2.495
Weighted $ d $			0.641	0.627	0.355	0.369	0.520	0.800	0.389	0.338	4.039	0.505



Fig. 4 The spatial positions of FeFe-co and P^N cluster in the PDB codes (a) *8BOQ* and (b) *8OIE*, respectively. The latter contains Fe and FeFe proteins simultaneously.

FeFe-co, with a distance of 14.4 Å. Thus, we deduce that Fe3 might be responsible for electron transfer to FeFe-co. Similarly, Fe3 has the shortest distance (14.5 Å) in the eight iron atoms in the P^N cluster to Fe6 in the M-cluster in the PDB

code *80IE*. Besides, Fe6 is the nearest iron to the electron donor $[Fe_4S_4]$, with a distance of 15.0 Å as shown in Fig. 4b.

3.3 Comparison of the metal oxidation states in FeMo-, FeVand FeFe-cos and their P^N clusters in nitrogenases

As depicted in Fig. 5a–c, FeMo-, FeV- and FeFe-cos in resting states show some similarities that Fe1, Fe2, Fe6 and Fe7 are more likely to be Fe³⁺. Fe3 atoms in FeMo-co and FeFe-co have a tendency to be mixed valence and Fe³⁺, respectively, while those in FeV-co prefers to be the Fe²⁺ state. Fe4 and Fe5 atoms (bound to a CO_3^{2-} ligand) in FeV-co are more inclined toward Fe²⁺, which is different from those in FeMo-co and FeFe-co with higher oxidation states. The overall oxidation level of 7Fe atoms (Fe1–Fe7, S_t = 19.504) calculated from R_0 (+3) for FeFe-co in the resting state is close to the average value of FeMo-co (S_t = 19.950), but obviously higher than that of FeV-co (S_t = 18.649).¹⁹ In the resting state, the



Fig. 5 The weighted average |d| values of 7Fe and Fe1–Fe7 atoms in resting states (a–c) and turnover states (d–f) in the FeMo-, FeV- and FeFe-cos calculated from R_0 (+2) and R_0 (+3). (g–i) The |d| values of 8Fe and Fe1–Fe7 atoms in P^N clusters of FeMo, FeV and FeFe proteins. The data for FeMo, FeV-cos and P^N clusters come from our previous calculated results.^{19,38,39}

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designation of Fe8 as Fe³⁺ at the terminal positions of FeFe-co has the same oxidation states as the Mo^{3+}/V^{3+} atoms of FeMo-/ FeV-co, indicating that the types of metallic elements do not affect the oxidation of metal (Mo/V/Fe) states at the terminal positions of the cofactors. Therefore, we infer that the octahedral Fe8 atom in FeFe-co probably serves a similar purpose to Mo^{3+} or V^{3+} in the other isoforms.

Fig. 5d–f show the |d| values of CO-bound FeMo-co (PDB code: 4TKV), CO-bound FeV-co (PDB code 7ADR) and FeFe-co in turnover states (PDB code 8BOQ), respectively. The 7Fe values of |d| calculated from R_0 (+2) and R_0 (+3) are 3.940 and 1.494, respectively, for CO-bound FeMo-co, suggesting strong oxidative iron segments. The overall oxidation level of 7Fe atoms (S_t = 19.506) calculated from R_0 (+3) for CO-bound FeMo-co is close to that of FeMo-co in the resting state (S_t = 19.950). This indicates that the dissociated S2B ligand (replaced by CO) takes two protons with it.54 For FeV-co, the 7Fe values of |d| produced from R_0 (+2) and R_0 (+3) are 0.973 and 3.451 for CO-bound FeV-co, and 2.292 and 2.391 for FeVco in the resting state, respectively, which implies that FeV-co in the CO-bound state are more reduced than that in the resting state. Fe2 and Fe6 atoms undergo reduction and are assigned accurately as iron(II) after S2B substituted by the CO ligand. This may be helpful for the comprehension that V-nitrogenase holds a unique property for the conversion of CO to hydrocarbons with yields over 93% of C₂H₄.^{52,55} Similarly, the 7Fe values of |d| calculated from R_0 (+2) and R_0 (+3) are 3.326 and 2.207 for FeFe-co in the turnover state, and 4.122 and 1.548 for FeFe-co in the resting state, respectively. As mentioned above, FeFe-co in the turnover state is slightly reduced than that in the resting state, and Fe2 and Fe6 atoms are rearranged as mixed valence with the substitution of S2B by OH. The slightly reduced valences of Fe2 and Fe6 in FeFe-co in the turnover state are compatible with those found in CO-bound FeV-co. These observations in FeV- and FeFe-cos indicate the presence of a reduced and rearranged oxidation state in the substrate-binding state, especially at the bound sites of Fe2 and Fe6.

For the P^N clusters of FeMo, FeV and FeFe proteins shown in Fig. 5g–i, the 8Fe_{av} value of |d| in the R_0 (+2) group is below 0.30, indicating that these P^N clusters exhibit strong reductive character. In particular, Fe1, Fe2, Fe5 and Fe6 tend to be Fe²⁺, while Fe8 exhibit some mixed valences. The electron distributions of Fe3, Fe4 and Fe7 atoms in the P^N clusters of FeMo, FeV and FeFe proteins are somewhat different. From the point of view of the valence sum S_t of 8Fe by R_0 (+2), the P^N clusters in FeMo (S_t = 18.353), FeV (S_t = 17.960) and FeFe proteins (S_t = 18.515) may have a similar formal oxidation state with 6Fe²⁺–2Fe³⁺.

In general, the electron distributions of FeMo- and FeFe-cos are somewhat different from that of VFe-co. These distinctions could be attributed to the different structures of MoFe, VFe and FeFe proteins, which result in different electron transfer channels that induce the formation of the relevant iron oxidation states. Intriguingly, the oxidation states of 7Fe in FeVand FeFe-cos in the turnover state are much reduced than those in the resting species, and Fe2 and Fe6 atoms undergo reduction in the substrate-binding state. MoFe, VFe and FeFe

4. Conclusions

In summary, we have calculated the valences of metal centres in the FeFe-cos and P^N clusters of iron-only nitrogenases (PDB codes: *8BOQ* and *8OIE*) using the bond valence method with weight schemes. The electronic structures of FeMo-, FeV- and FeFe-cos and their P-clusters have been further compared. For FeFe-co in the resting state, Fe1/2/4/5/6/7/8 atoms ought to be designated as Fe³⁺, while Fe3 is more reduced and exhibits some degree of mixed valences. Moreover, the overall oxidation state of FeFe-co in the turnover state is slightly reduced than those in resting states. Interestingly, the electron distributions in Fe2 and Fe6 atoms are rearranged as mixed-valence states with the substitution of S2B by OH. In the P^N clusters of FeFe proteins, the formal oxidation states of 8Fe could be approximately equal to $6Fe^{2+}-2Fe^{3+}$ with delocalized electrons.

A comparative study of the metal oxidation states for the FeMo-, FeV- and FeFe-cos suggests that the overall oxidation levels of 7Fe atoms (Fe1–Fe7) for both FeFe- and FeMo-cos in resting states are higher than that of FeV-co. Fe8 in the resting state of FeFe-co is calculated as the Fe³⁺ state, suggesting that the types of metallic elements have no obvious effect on the oxidation states of the apical Mo/V/Fe of the M-clusters. Moreover, Fe2 and Fe6 atoms undergo reduction in FeV- and FeFe-cos in the turnover state. Despite sharing the same core structure [Fe₈S₇] of the P^N cluster in MoFe, VFe and FeFe proteins, it can be found that their electron distributions of the P^N cluster exhibit some differences. These differences can be attributed to the unique structures of MoFe, VFe and FeFe proteins, which may lead to diverse channels of electron transfer and induce the formation of relevant iron oxidation states.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 L. M. Rubio and P. W. Ludden, *Annu. Rev. Microbiol.*, 2008, 62, 93–111.
- 2 F. A. Tezcan, J. T. Kaiser, D. Mustafi, M. Y. Walton, J. B. Howard and D. C. Rees, *Science*, 2005, **309**, 1377–1380.
- 3 R. R. Eady, Chem. Rev., 1996, 96, 3013-3030.

- 4 T. Spatzal, M. Aksoyoglu, L. Zhang, S. L. A. Andrade, E. Schleicher, S. Weber, D. C. Rees and O. Einsle, *Science*, 2011, **334**, 940–940.
- 5 O. Einsle, J. Biol. Inorg. Chem., 2014, 19, 737-745.
- 6 D. Sippel and O. Einsle, Nat. Chem. Biol., 2017, 13, 956–960.
- 7 C. Trncik, F. Detemple and O. Einsle, *Nat. Catal.*, 2023, 6, 415–424.
- 8 A. J. Jasniewski, C. C. Lee, M. W. Ribbe and Y. L. Hu, *Chem. Rev.*, 2020, **120**, 5107–5157.
- 9 Y. L. Hu, A. W. Fay, C. C. Lee, J. A. Wiig and M. W. Ribbe, *Dalton Trans.*, 2010, **39**, 2964–2971.
- 10 Y. L. Hu, C. C. Lee and M. W. Ribbe, *Dalton Trans.*, 2012, 41, 1118–1127.
- 11 O. Einsle and D. C. Rees, Chem. Rev., 2020, 120, 4969-5004.
- 12 D. F. Harris, D. A. Lukoyanov, S. Shaw, P. Compton, M. Tokmina-Lukaszewska, B. Bothner, N. Kelleher, D. R. Dean, B. M. Hoffman and L. C. Seefeldt, *Biochemistry*, 2018, 57, 701–710.
- 13 D. F. Harris, D. A. Lukoyanov, H. Kallas, C. Trncik, Z. Y. Yang, P. Compton, N. Kelleher, O. Einsle, D. R. Dean, B. M. Hoffman and L. C. Seefeldt, *Biochemistry*, 2019, 58, 3293–3301.
- 14 J. W. Peters, K. Fisher, W. E. Newton and D. R. Dean, J. Biol. Chem., 1995, 270, 27007–27013.
- 15 K. Danyal, D. R. Dean, B. M. Hoffman and L. C. Seefeldt, *Biochemistry*, 2011, **50**, 9255–9263.
- 16 H. L. Rutledge and F. A. Tezcan, *Chem. Rev.*, 2020, **120**, 5158–5193.
- 17 R. Bjornsson, F. Neese and S. DeBeer, *Inorg. Chem.*, 2017, 56, 1470–1477.
- 18 T. Spatzal, J. Schlesier, E. M. Burger, D. Sippel, L. Zhang, S. L. A. Andrade, D. C. Rees and O. Einsle, *Nat. Commun.*, 2016, 7, 10902.
- 19 W. T. Jin, M. Yang, S. S. Zhu and Z. H. Zhou, Acta Crystallogr., Sect. D: Biol. Crystallogr., 2020, 76, 428-437.
- 20 Z. Y. Yang, E. Jimenez-Vicente, H. Kallas, D. A. Lukoyanov, H. Yang, J. S. Martin del Campo, D. R. Dean, B. M. Hoffman and L. C. Seefeldt, *Chem. Sci.*, 2021, **12**, 6913–6922.
- 21 J. R. Chisnell, R. Premakumar and P. E. Bishop, *J. Bacteriol.*, 1988, **170**, 27–33.
- 22 E. Krahn, B. Weiss, M. Kröckel, J. Groppe, G. Henkel, S. Cramer, A. Trautwein, K. Schneider and A. Müller, *J. Biol. Inorg. Chem.*, 2002, 7, 37–45.
- 23 K. Schneider and A. Müller, in *Catalysts for Nitrogen Fixation: Nitrogenases, Relevant Chemical Models and Commercial Processes*, ed. B. E. Smith, R. L. Richards and W. E. Newton, Springer Netherlands, Dordrecht, 2004, pp. 281–307.
- 24 D. A. Lukoyanov, D. F. Harris, Z. Y. Yang, A. Pérez-González, D. R. Dean, L. C. Seefeldt and B. M. Hoffman, *Inorg. Chem.*, 2022, 61, 5459–5464.
- 25 S. Greed, Nat. Rev. Chem., 2023, 7, 379-379.
- 26 F. V. Schmidt, L. Schulz, J. Zarzycki, S. Prinz, N. N. Oehlmann, T. J. Erb and J. G. Rebelein, *Nat. Struct. Mol. Biol.*, 2024, **31**, 150–158.
- 27 K. M. Lancaster, M. Roemelt, P. Ettenhuber, Y. Hu, M. W. Ribbe, F. Neese, U. Bergmann and S. DeBeer, *Science*, 2011, 334, 974–977.

- 28 D. Sippel and O. Einsle, Nat. Chem. Biol., 2017, 13, 956-960.
- 29 Y. Zheng, D. F. Harris, Z. Yu, Y. Fu, S. Poudel, R. N. Ledbetter, K. R. Fixen, Z. Y. Yang, E. S. Boyd, M. E. Lidstrom, L. C. Seefeldt and C. S. Harwood, *Nat. Microbiol.*, 2018, 3, 281–286.
- 30 I. D. Brown and R. D. Shannon, Acta Crystallogr., Sect. A: Cryst. Phys., Diffr., Theor. Gen. Crystallogr., 1973, 29, 266.
- 31 I. D. Brown, Acta Crystallogr., Sect. B: Struct. Crystallogr. Cryst. Chem., 1977, 33, 1305–1310.
- 32 I. D. Brown and D. Altermatt, Acta Crystallogr., Sect. B: Struct. Sci., 1985, 41, 244-247.
- 33 M. H. Chiang, M. R. Antonio, C. W. Williams and L. Soderholm, *Dalton Trans.*, 2004, 801–806.
- 34 S. Matsia, O. Tsave, A. Hatzidimitriou, C. Gabriel and A. Salifoglou, *J. Inorg. Biochem.*, 2021, **222**, 111469.
- 35 W. Liu and H. H. Thorp, Inorg. Chem., 1993, 32, 4102-4105.
- 36 P. Muller, S. Kopke and G. M. Sheldrick, Acta Crystallogr., Sect. D: Biol. Crystallogr., 2003, 59, 32–37.
- 37 I. D. Brown, Chem. Rev., 2009, 109, 6858-6919.
- 38 C. Yuan, W. T. Jin and Z. H. Zhou, New J. Chem., 2022, 46, 9519–9525.
- 39 C. Yuan, W. T. Jin and Z. H. Zhou, *RSC Adv.*, 2022, 12, 5214–5224.
- 40 L. Cao and U. Ryde, Phys. Chem. Chem. Phys., 2019, 21, 2480-2488.
- 41 L. Deng, H. Wang, C. H. Dapper, W. E. Newton, S. Shilov, S. Wang, S. P. Cramer and Z. H. Zhou, *Commun. Chem.*, 2020, 3, 145.
- 42 I. Brown, Accumulated table of bond-valence parameters. https://www.iucr.org/__data/assets/file/0011/150779/ bvparm2020.cif, 2020.
- 43 H. H. Thorp, Inorg. Chem., 1992, 31, 1585-1588.
- 44 Z. L. Xie, C. Yuan and Z. H. Zhou, *Acta Crystallogr., Sect. D: Biol. Crystallogr.*, 2023, **79**, 401–408.
- 45 I. D. Brown and D. Altermatt, Acta Crystallogr., Sect. B: Struct. Sci., 1985, 41, 244-247.
- 46 W. Liu and H. H. Thorp, Inorg. Chem., 1993, 32, 4102-4105.
- 47 Y. Shi, W. He, J. Zhao, A. Hu, J. Pan, H. Wang and H. Zhu, *J. Cleaner Prod.*, 2020, 253, 119965.
- 48 D. Shepard, presented in part at the *Proceedings of the* 1968 23rd ACM national conference, 1968.
- 49 P. M. Bartier and C. P. Keller, *Comput. Geosci.*, 1996, 22, 795–799.
- 50 D. Sippel, M. Rohde, J. Netzer, C. Trncik, J. Gies, K. Grunau, I. Djurdjevic, L. Decamps, S. L. A. Andrade and O. Einsle, *Science*, 2018, **359**, 1484–1489.
- 51 E. Krahn, B. J. R. Weiss, M. Kröckel, J. Groppe, G. Henkel, S. P. Cramer, A. X. Trautwein, K. Schneider and A. Müller, *J. Biol. Inorg. Chem.*, 2002, 7, 37–45.
- 52 M. Rohde, K. Laun, I. Zebger, S. T. Stripp and O. Einsle, *Sci. Adv.*, 2021, 7, eabg4474.
- 53 Y. Ohki, A. Murata, M. Imada and K. Tatsumi, *Inorg. Chem.*, 2009, **48**, 4271–4273.
- 54 J. Bergmann, E. Oksanen and U. Ryde, *J. Inorg. Biochem.*, 2021, **219**, 111426.
- 55 M. Rohde, K. Grunau and O. Einsle, *Angew. Chem., Int. Ed.*, 2020, **59**, 23626–23630.