

## Supporting Information

For

### Closed-shell Phenalenyl-based Dinuclear Iron(III) Complex as a Robust Cathode for One-Compartment H<sub>2</sub>O<sub>2</sub> Fuel Cell

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## 1. Materials and general methods

All the chemicals were obtained from commercial sources, and used without further purification. Carbon black, and N-methyl pyrrolidine (NMP) were procured from Alfa Aesar, and Nafion solution (5 wt. %) was procured from Sigma Aldrich. The commercially available H<sub>2</sub>O<sub>2</sub>, 30% w/v, procured from Alfa Aesar, was used to perform the experiments. The solvents required for synthesis, and conducting CV measurements were dried and distilled using standard techniques. *9-hydroxy-1H-phenalen-1-one* was prepared using the literature method.<sup>1</sup>

Crystals suitable for single crystal X-ray diffractions (SCXRD) were loaded on a Bruker Smart Apex CCD diffractometer at 100 K. All the structures were solved by direct methods using SHELXS-97 and refined by full-matrix least squares on F<sup>2</sup> using SHELXL-97 softwares.<sup>2</sup> The details of crystallographic refinement parameters is given in Table S1. Anisotropic displacement parameters are used to refine the non-hydrogen atoms. All the hydrogen atoms were included in idealized positions, and were refined isotropically by a riding model.

The cyclic voltammetric (CV) tests were performed using a three-electrode assembly; glassy carbon (GC) working electrode, Pt counter electrode, and Ag/AgCl (3.0 M KCl) reference electrode. The electrochemical studies of complex **1** with CV were conducted in 10 ml DMF solution of complex **1** (1 mM) and the supporting electrolyte, tetrabutylammonium hexafluorophosphate (0.1 M). The CV measurements were carried out under an oxygen-free atmosphere by purging nitrogen gas, and the scan rate was fixed at 50 mV s<sup>-1</sup>. To test H<sub>2</sub>O<sub>2</sub> fuel cell activity, the modified GC electrode and Ni foam anode were immersed in the supporting electrolyte for at least 5 min before adding H<sub>2</sub>O<sub>2</sub> to the electrolytic solution. The performance tests are repeated at least 5 times to ensure the reproducibility.

## 2. Crystallographic data for the hmbh-PLYH<sub>2</sub> ligand and complex **1**.

**Table S1.** Crystal data and structure refinement parameters for the ligand, hmbh-PLYH<sub>2</sub> and complex **1**.

<b>Identification code</b>	hmbh-PLYH <sub>2</sub>	Complex <b>1</b>
<b>Empirical formula</b>	C <sub>21</sub> H <sub>16</sub> N <sub>2</sub> O <sub>3</sub>	C <sub>65</sub> H <sub>44</sub> N <sub>6</sub> O <sub>9</sub> Cl <sub>6</sub> Fe <sub>2</sub>
<b>Formula weight</b>	344.36	1377.46
<b>Temperature/K</b>	100	100
<b>Crystal system</b>	orthorhombic	monoclinic
<b>Space group</b>	<i>Aba2</i>	<i>P2<sub>1</sub>/c</i>
<b>a/Å</b>	16.38(2)	23.2460(12)
<b>b/Å</b>	29.87(4)	13.2054(7)
<b>c/Å</b>	6.843(9)	19.6226(10)
<b>α/°</b>	90	90
<b>β/°</b>	90	101.358(2)
<b>γ/°</b>	90	90
<b>Volume/Å<sup>3</sup></b>	3348(7)	5905.6(5)
<b>Z</b>	8	4
<b>ρ<sub>calc</sub>/g/cm<sup>3</sup></b>	1.366	1.549
<b>μ/mm<sup>-1</sup></b>	0.093	0.829
<b>F(000)</b>	1440.0	2808.0
<b>Crystal size/mm<sup>3</sup></b>	0.21 × 0.19 × 0.16	0.2 × 0.19 × 0.16
<b>Radiation</b>	MoKα (λ = 0.71073)	MoKα (λ = 0.71073)
<b>2θ range for data collection/°</b>	5.996 to 49.998	4.234 to 50.698
<b>Index ranges</b>	-14 ≤ h ≤ 19, -34 ≤ k ≤ 35, -8 ≤ l ≤ 8	-28 ≤ h ≤ 28, -15 ≤ k ≤ 15, -23 ≤ l ≤ 16
<b>Reflections collected</b>	10271	52543
<b>Independent reflections</b>	2837 [R <sub>int</sub> = 0.0935, R <sub>sigma</sub> = 0.1068]	10810 [R <sub>int</sub> = 0.0501, R <sub>sigma</sub> = 0.0430]
<b>Data/restraints/parameters</b>	2837/7/231	10810/0/805
<b>Goodness-of-fit on F<sup>2</sup></b>	1.133	1.080
<b>Final R indexes [I ≥ 2σ (I)]</b>	R <sub>1</sub> = 0.0973, wR <sub>2</sub> = 0.1617	R <sub>1</sub> = 0.0797, wR <sub>2</sub> = 0.1928
<b>Final R indexes [all data]</b>	R <sub>1</sub> = 0.1336, wR <sub>2</sub> = 0.1759	R <sub>1</sub> = 0.0990, wR <sub>2</sub> = 0.2043
<b>Largest diff. peak/hole / e Å<sup>-3</sup></b>	0.25/-0.28	1.01/-1.18

**3. Bond parameters of complex **1**, and comparison with the iso-structural [Fe<sub>2</sub><sup>III</sup>(salhn)<sub>3</sub>] complex.**

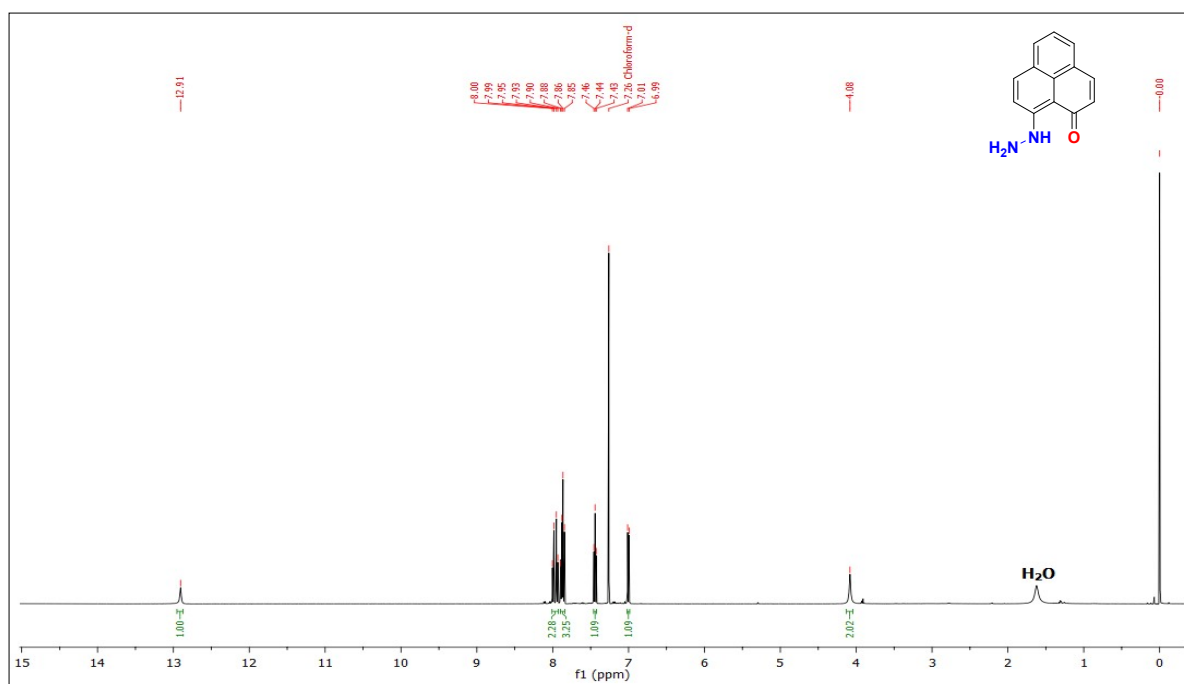
**Table S2.** Comparison of the bond distances of the reported  $[\text{Fe}_2^{\text{III}}(\text{salhn})_3]$  complex<sup>3</sup> with complex **1**.

$[\text{Fe}_2^{\text{III}}(\text{salhn})_3]$		$[\text{Fe}^{\text{III}}_2(\text{hmbh-PLY})_3]$ , complex <b>1</b>	
Bond	Bond Distance (Å)	Bond	Bond Distance (Å)
Fe1-O1	1.930(5)	Fe1-O1	1.935(4)
Fe1-O3	1.925(3)	Fe1-O8	1.923(4)
Fe1-O5	1.912(4)	Fe1-O5	1.906 (4)
Fe1-N1	2.164(5)	F1-N1	2.123(4)
Fe1-N3	2.197(6)	Fe1-N4	2.194(4)
Fe1-N5	2.178(4)	Fe1-N6	2.222(4)
Fe2-O2	1.910(5)	Fe2-O2	1.940(4)
Fe2-O4	1.927(3)	Fe2-O4	1.916(4)
Fe2-O6	1.919(5)	Fe2-O7	1.933(4)
Fe2-N2	2.167(4)	Fe2-N5	2.087(4)
Fe2-N4	2.183(5)	Fe2-N3	2.115(4)
Fe2-N6	2.182(5)	Fe2-N2	2.188(4)

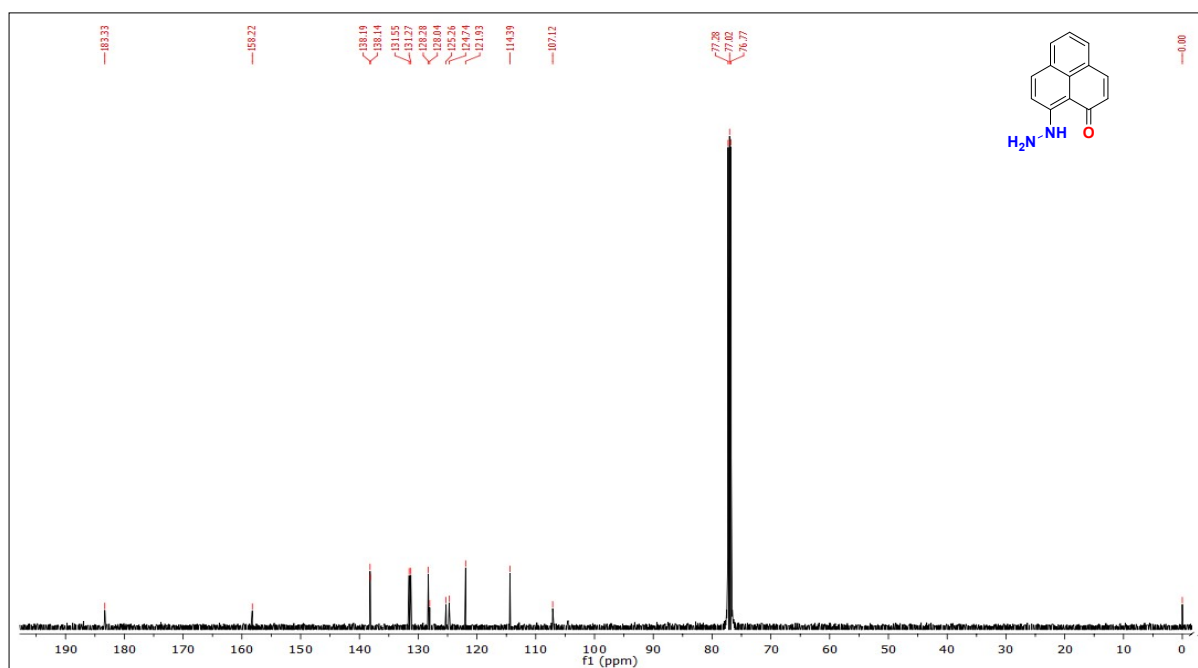
**Table S3.** Comparison of the bond angles of the reported  $[\text{Fe}_2^{\text{III}}(\text{salhn})_3]$  complex with complex **1**.

$[\text{Fe}_2^{\text{III}}(\text{salhn})_3]$		$[\text{Fe}^{\text{III}}_2(\text{hmbh-PLY})_3]$ , complex <b>1</b>	
Atoms	Bond Angle (°)	Atoms	Bond Angle (°)
O1-Fe1-N3	169.6(2)	O8-Fe1-N4	168.74(17)
O5-Fe1-N1	169.9(1)	O5-Fe1-N1	170.38(16)
O3-Fe1-N3	84.8(2)	O1-Fe1-N6	82.71(15)
O1-Fe1-N5	92.6(2)	O5-Fe1-O1	95.11(17)
O3-Fe1-O5	97.8(2)	O1-Fe1-N1	94.49(17)
O1-Fe1-N1	84.4(2)	O5-Fe1-N6	87.97(15)
N1-Fe1-N3	85.3(2)	N1-Fe1-N6	87.16(15)
O2-Fe2-N6	171.0(1)	O7-Fe2-N2	171.76(17)
O4-Fe2-N2	167.6(2)	O4-Fe2-N5	167.45(16)
O2-Fe2-N4	92.1(2)	O2-Fe2-N3	90.90(16)
O2-Fe2-O4	99.1(2)	O4-Fe2-O2	89.83(16)
N4-Fe2-N6	86.9(2)	N5-Fe2-N3	96.14(17)
O2-Fe2-O6	84.4(2)	O2-Fe2-N5	84.09(16)
O6-Fe2-N6	170.7(2)	O4-Fe2-N3	165.94(16)

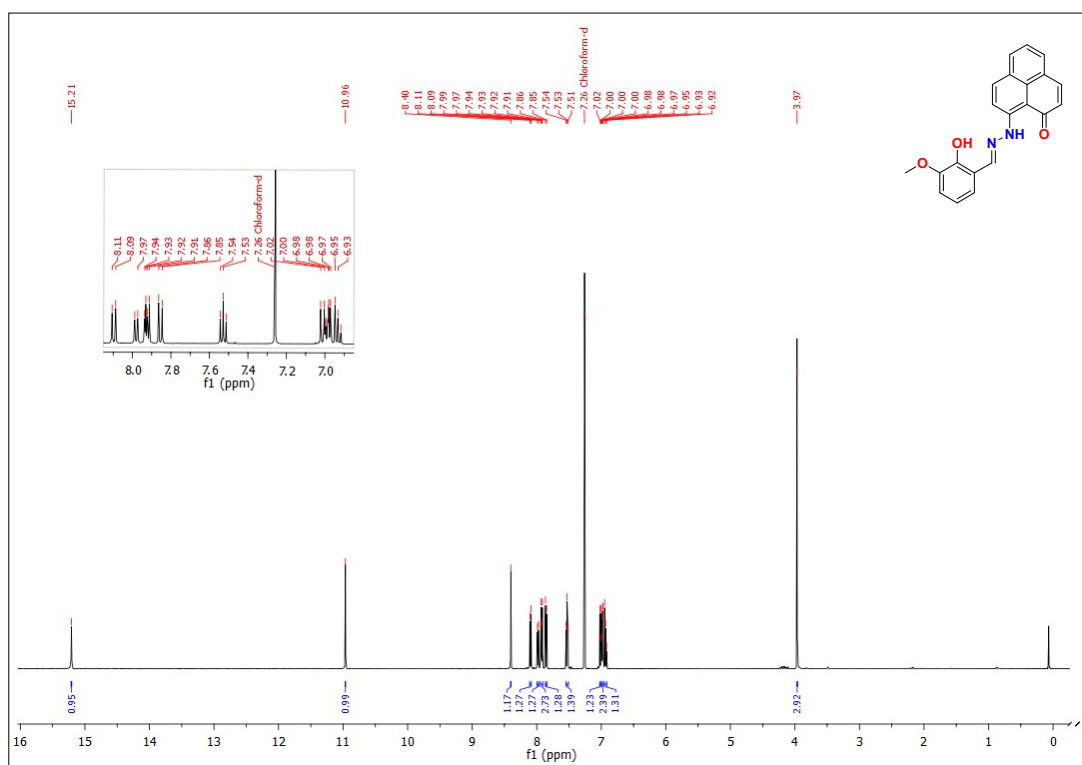
#### 4. $^1\text{H}$ NMR and $^{13}\text{C}$ NMR Spectra



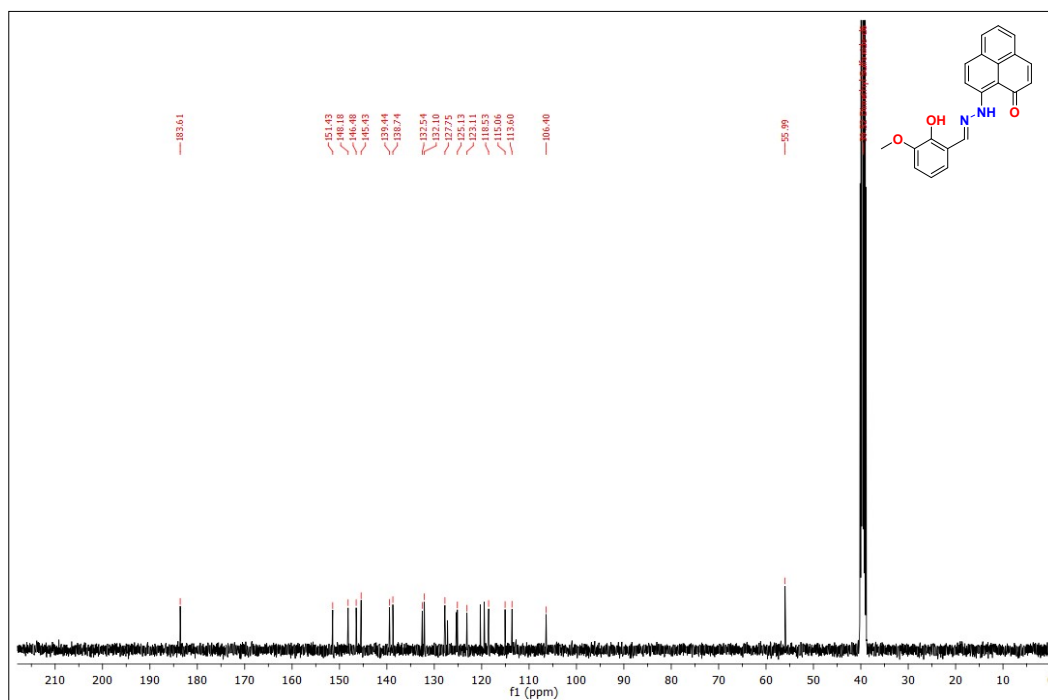
**Figure S1.**  $^1\text{H}$  NMR spectrum of Hz-PLY in  $\text{CDCl}_3$  solvent.



**Figure S2.**  $^{13}\text{C}$  NMR spectrum of Hz-PLY in  $\text{CDCl}_3$  solvent.

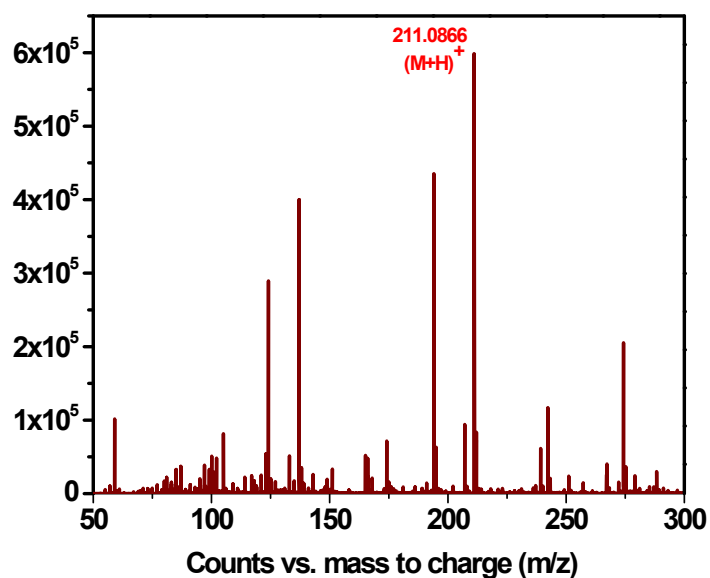


**Figure S3.**  $^1\text{H}$  NMR spectrum of hmbh-PLYH<sub>2</sub> in CDCl<sub>3</sub> solvent.

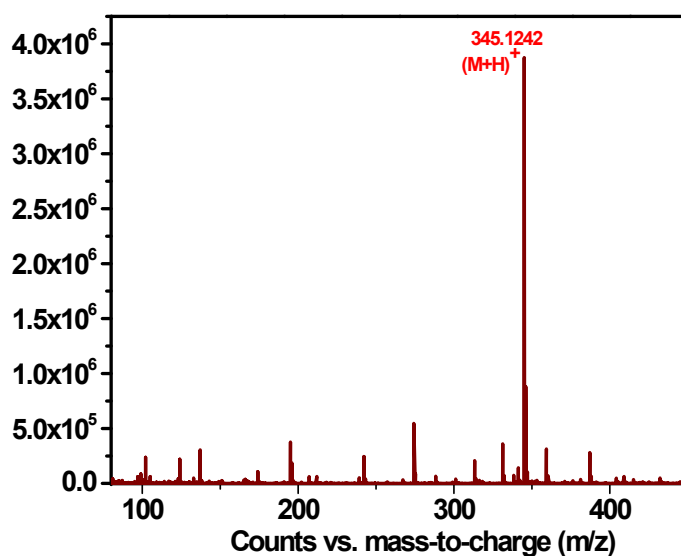


**Figure S4.**  $^{13}\text{C}$  NMR spectrum of hmbh-PLYH<sub>2</sub> in DMSO-d<sub>6</sub> solvent.

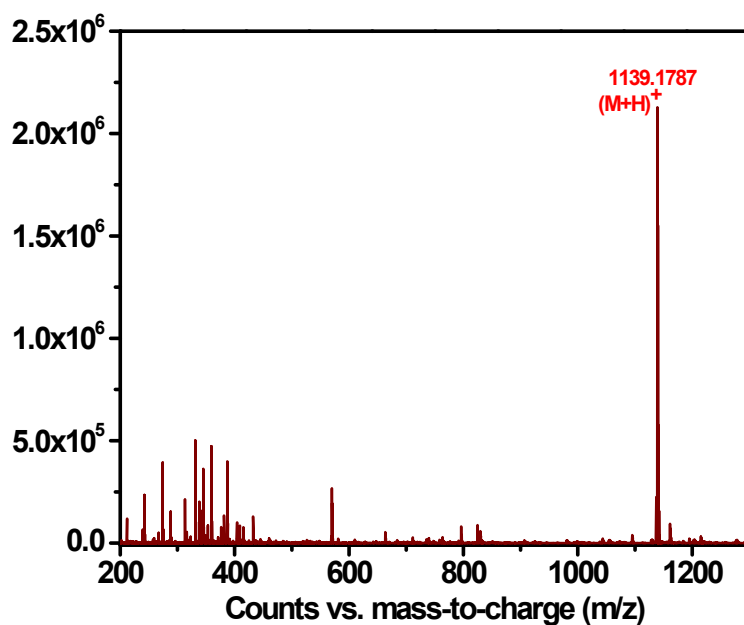
## 5. HRMS Spectra of Hz-PLY, hmbh-PLYH<sub>2</sub> and complex 1.



**Figure S5.** HRMS spectrum of Hz-PLY. HRMS (CH<sub>3</sub>CN, positive ionization): calcd. for C<sub>13</sub>H<sub>10</sub>N<sub>2</sub>O m/z = 211.0867 [M + H]<sup>+</sup>, found 211.0866 [M + H]<sup>+</sup>.

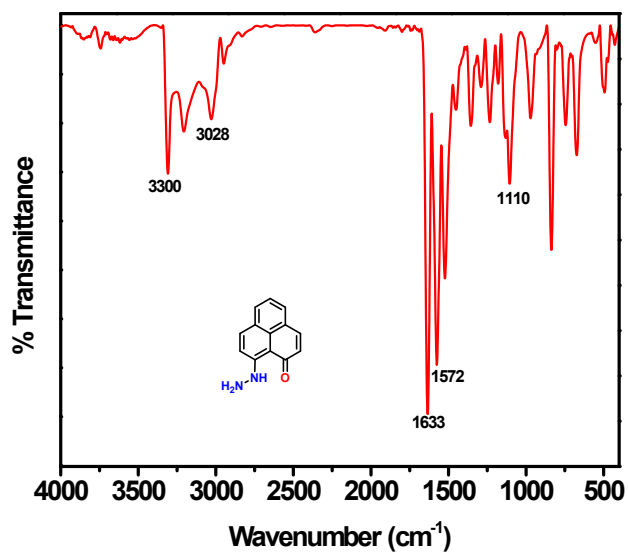


**Figure S6.** HRMS spectrum of hmbh-PLYH<sub>2</sub>. HRMS (CH<sub>3</sub>CN, positive ionization): calcd. for C<sub>21</sub>H<sub>16</sub>N<sub>2</sub>O<sub>3</sub> m/z = 345.1234 [M + H]<sup>+</sup>, found 345.1242 [M + H]<sup>+</sup>.

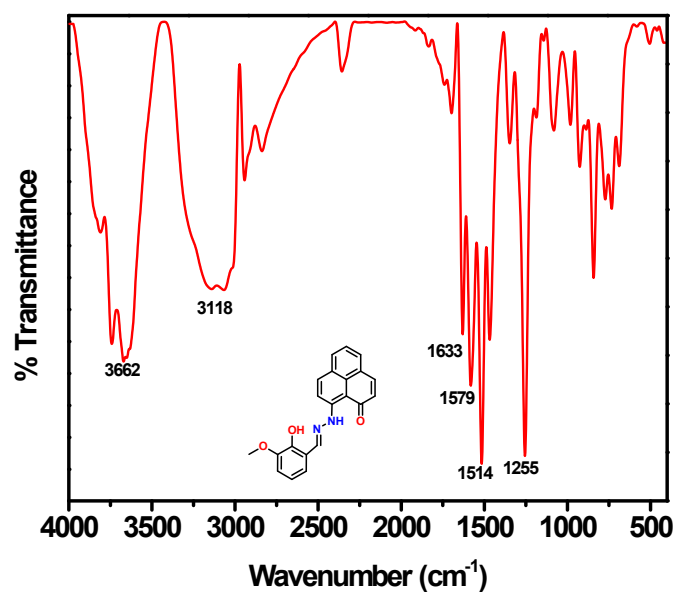


**Figure S7.** HRMS spectrum of complex **1**. HRMS (CH<sub>3</sub>CN, positive ionization): calcd. for C<sub>63</sub>H<sub>42</sub>Fe<sub>2</sub>N<sub>6</sub>O<sub>9</sub> m/z = 1139.1785 (M+H)<sup>+</sup>, found 1139.1787 [M + H]<sup>+</sup>.

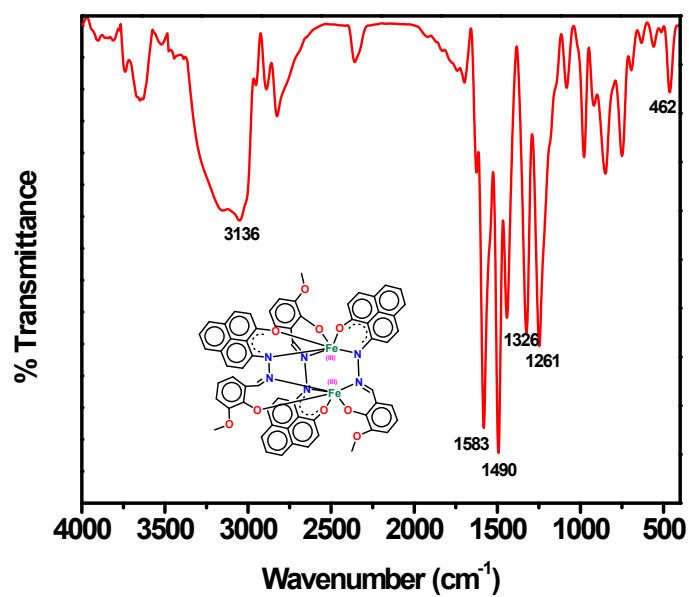
**6. FT-IR Spectra of 9-hydrazineyl-1H-phenalen-1-one, hmbh-PLYH<sub>2</sub> and complex **1**.**



**Figure S8.** FT-IR spectrum of Hz-PLY.

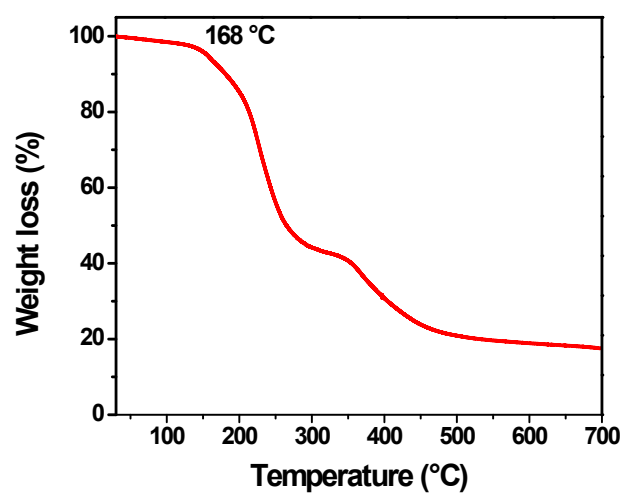


**Figure S9.** FT-IR spectrum of hmbh-PLYH<sub>2</sub>.

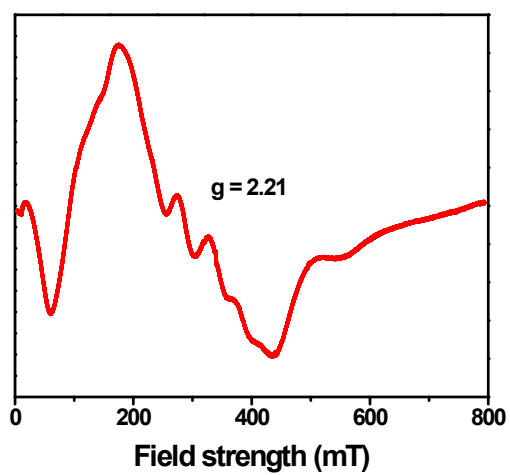


**Figure S10.** FT-IR spectrum of complex 1.

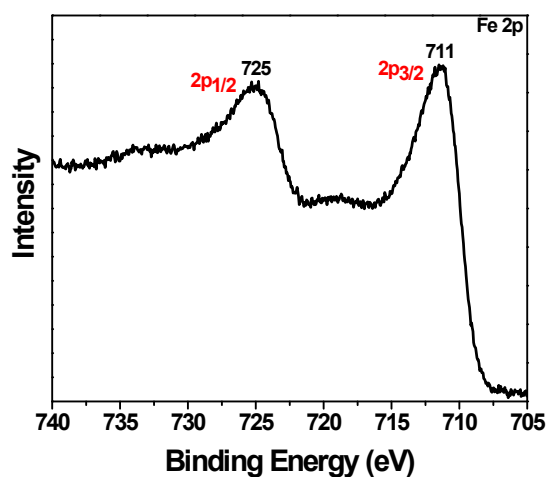
## 7. Thermal stability, EPR, XPS and elemental analysis of the complex 1.



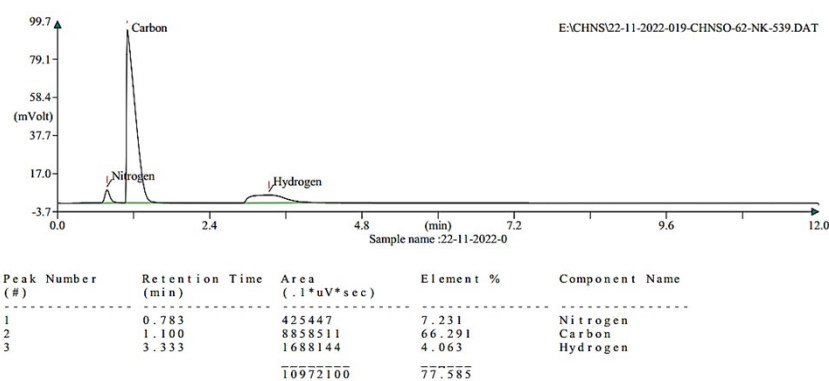
**Figure S11.** Thermogravimetric analysis (TGA) plot of the complex **1**.



**Figure S12.** Solid state X-band EPR of complex **1** recorded at room temperature.

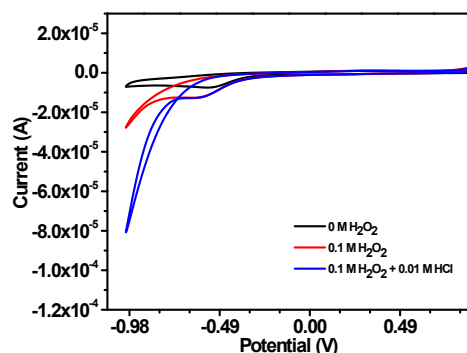


**Figure S13.** XPS analysis of the complex 1.

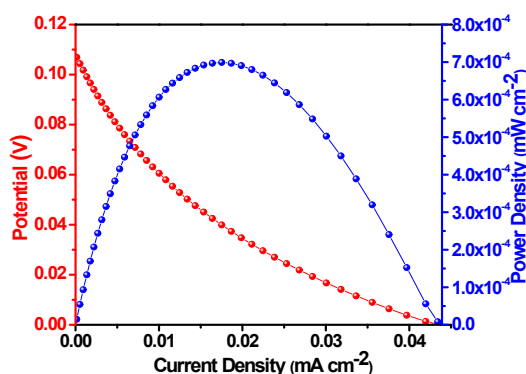


**Figure S14.** Elemental analysis of complex 1: calcd. for  $C_{63}H_{42}Fe_2N_6O_9$ : C, 66.45; H, 3.72; N, 7.38. Found: C, 66.29; H, 4.06; N, 7.23.

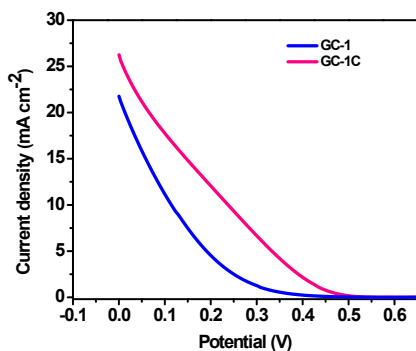
## 8. Electrocatalytic reduction of $\text{H}_2\text{O}_2$ .



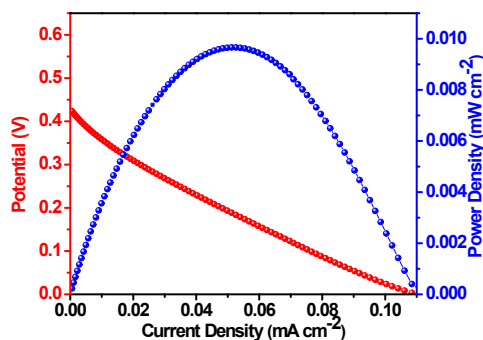
**Figure S15.** Electrocatalytic reduction of  $\text{H}_2\text{O}_2$  with bare glassy carbon (GC) electrode in acetate buffer (pH 3), and acetate buffer (pH 3) containing 0.01 M HCl solution in the presence of  $\text{H}_2\text{O}_2$  (0 M and 0.1 M, as mentioned in the graph). Ag/AgCl(3.0 M KCl) electrode is used as the reference, Pt-wire as the counter electrode, and a scan rate of  $100 \text{ mV s}^{-1}$  was maintained during the experiment.



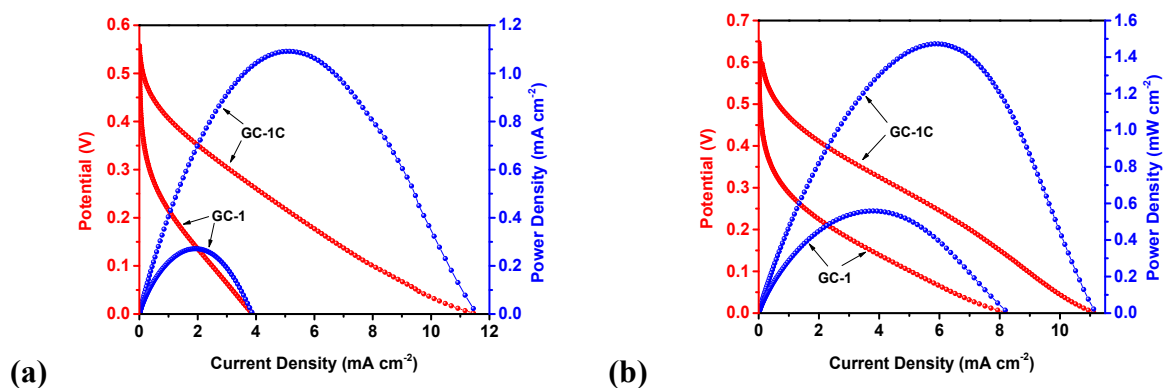
**Figure S16.**  $I$ - $V$  (red) and  $I$ - $P$  (blue) curves of one-compartment  $\text{H}_2\text{O}_2$  fuel cell with GC-1 cathode, and Ni foam anode at a scan rate  $10 \text{ mV s}^{-1}$ . The performance tests were conducted in an acetate buffer (pH 3) containing 0.3 M  $\text{H}_2\text{O}_2$ . Current density is normalized by the geometrical area of glassy carbon electrode.



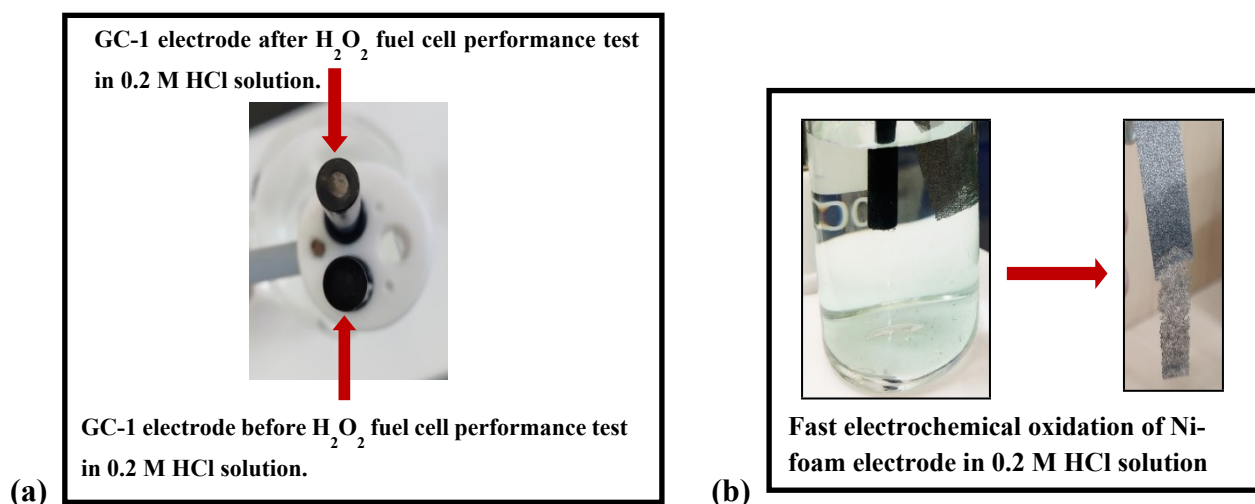
**Figure S17.** Polarization curves for **GC-1** (blue curve)/**GC-1C** (pink curve) as cathode in 0.1 M HCl containing 0.3 M H<sub>2</sub>O<sub>2</sub>, and Ni foam as the anode in one compartment H<sub>2</sub>O<sub>2</sub> fuel cell at a scan rate 10 mV s<sup>-1</sup>. Current density is normalized by the geometrical area of glassy carbon electrode.



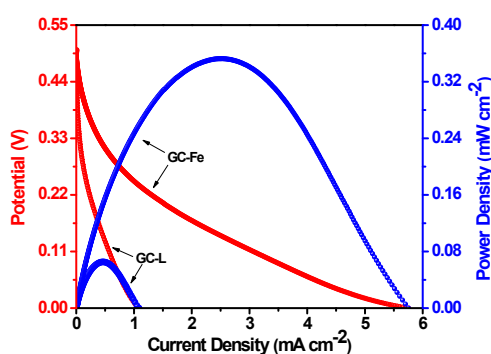
**Figure S18.** *I-V* (red) and *I-P* (blue) curves of a one-compartment H<sub>2</sub>O<sub>2</sub> fuel cell with bare glassy carbon (GC) cathode and Ni foam as anode, at a scan rate 10 mV s<sup>-1</sup>. The performance tests were conducted in 0.1 M HCl containing 0.3 M H<sub>2</sub>O<sub>2</sub>. Current density is normalized by the geometrical area of glassy carbon electrode.



**Figure S19.**  $I$ - $V$  (red) and  $I$ - $P$  (blue) curves of a one-compartment  $\text{H}_2\text{O}_2$  fuel cell with GC-1/GC-1C cathode and Ni foam anode, at a scan rate  $10 \text{ mV s}^{-1}$ . The performance tests were conducted in (a) 0.05 M and (b) 0.2 M HCl containing 0.3 M  $\text{H}_2\text{O}_2$ . Current density is normalized by the geometrical area of glassy carbon electrode.

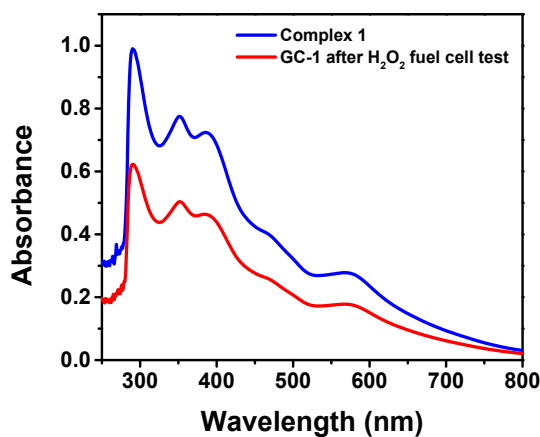


**Figure S20.** Image showing (a) GC-1 electrode, before and after the fuel cell performance test in 0.2 M HCl as the electrolyte, and (b) the corrosion of Ni foam anode while running the performance tests in 0.2 M HCl as the electrolyte.



**Figure S21.**  $I$ - $V$  (red) and  $I$ - $P$  (blue) curves of a one-compartment  $\text{H}_2\text{O}_2$  fuel cell with Ni-foam anode and  $\text{FeCl}_3$  (GC-Fe) modified cathode; and Ni-foam anode and ligand, hmbh-PLYH<sub>2</sub> (GC-L) modified cathode at a scan rate  $10 \text{ mV s}^{-1}$ . The performance tests were

conducted in 0.1 M HCl containing 0.3 M H<sub>2</sub>O<sub>2</sub>. Current density is normalized by the geometrical area of glassy carbon electrode.



**Figure S22.** UV-vis spectra of complex **1** (blue line) and the recovered catalyst from **GC-1** electrode (red line) after running the H<sub>2</sub>O<sub>2</sub> fuel cell test.

**Anode:**  $\text{H}_2\text{O}_2 \rightarrow \text{O}_2 + 2\text{H}^+ + 2\text{e}^-$  ( $E^\circ_{\text{ox}} = 0.68 \text{ V vs. NHE}$ ) equation.....S1

**Cathode:**  $\text{H}_2\text{O}_2 + 2\text{H}^+ + 2\text{e}^- \rightarrow 2\text{H}_2\text{O}$  ( $E^\circ_{\text{red}} = 1.78 \text{ V vs. NHE}$ ) equation.....S2

**Overall reaction:**  $2\text{H}_2\text{O}_2 \rightarrow \text{O}_2 + 2\text{H}_2\text{O}$  ( $E^\circ_{\text{cell}} = 1.09 \text{ V vs. NHE}$ ) equation.....S3

**Scheme S1.** Electrochemical equations for the one-compartment H<sub>2</sub>O<sub>2</sub> fuel cell, and the potentials vs. Normal Hydrogen Electrode (NHE).

## 9. DFT calculations

The optimized geometry parameters around the metal centre of the distorted octahedral complex **1** in its high spin ( $S = 5$ ) and broken symmetry spin states are presented in Table S4 along with respective experimental X-ray values for comparison. The DFT calculated bond distance and bond angle values obtained at method 1 (B3LYP/def2-SVP), method 2 (B3LYP+D3BJ/def2-SVP+def2-TZVP), method 3 (BP86+D3BJ/def2-SVP+def2-TZVP) are in good accord with the X-ray values with mean absolute deviation for bond distance values are estimated as 0.031, 0.028, 0.031 Å. The method 2 is selected for the present study. Note that, the metal ions are treated with def2-TZVP basis function during optimization.

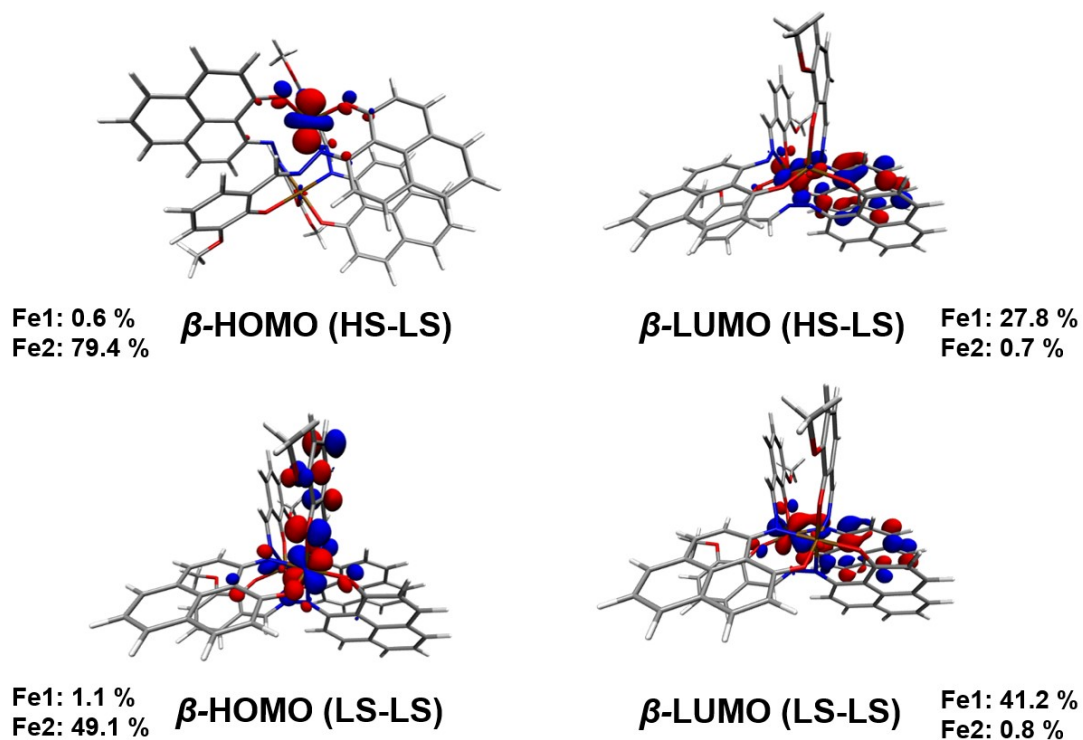
Broken symmetry analysis is performed for **1** to **1**<sup>-4</sup> complexes. The magnetic coupling constant ( $J$ ) is obtained using eq.

$$J = - \frac{(E_{HS} - E_{BS})}{\langle S_{max}^2 \rangle - \langle S_{min}^2 \rangle}$$

Where,  $(E_{HS} - E_{BS})$  is the energy difference between high spin (HS) and broken symmetry (BS) spin states and  $S_{max}^2$ ,  $S_{min}^2$  are maximum and minimum magnetic spin moment values.

**Table S4.** Experimental (X-ray) and DFT optimized bond lengths around the transition metal (Fe) centers of the diiron(III) complex in its HS-HS ( S = 5) and HS-BS (S = 0). M1 = B3LYP/def2-SVP, M2 = B3LYP+D3BJ/def2-SVP+def2-TZVP, and M3 = BP86+D3BJ/def2-SVP+def2-TZVP). Bond lengths are given in Å.

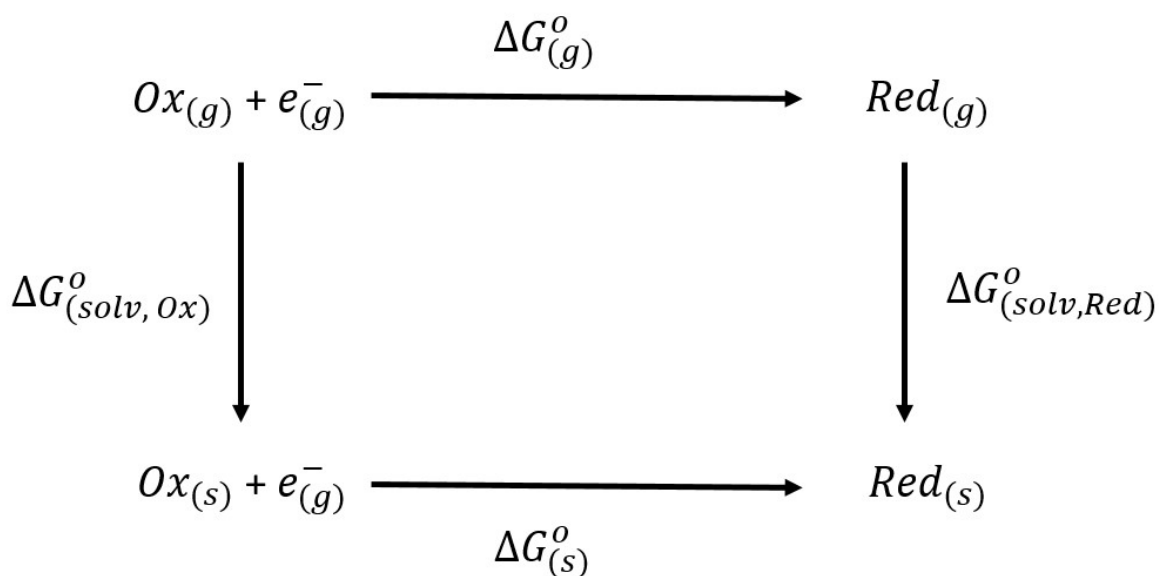
	<i>X-ray</i>	<i>Theoretical values</i>			
			<b>HS-HS</b>		<b>HS-BS</b>
		Method 1	Method 2	Method 3	Method 1
<b><i>Fe1-O8</i></b>	1.92284	1.93388	1.92668	1.93293	1.93476
<b><i>Fe1-O1</i></b>	1.93482	1.96268	1.95195	1.97649	1.96321
<b><i>Fe1-O5</i></b>	1.90552	1.93444	1.92497	1.93918	1.93568
<b><i>Fe1-N1</i></b>	2.12272	2.17895	2.17002	2.14833	2.17701
<b><i>Fe1-N6</i></b>	2.22154	2.21055	2.22931	2.21829	2.20973
<b><i>Fe1-N4</i></b>	2.19359	2.24161	2.23546	2.24145	2.24051
<b><i>Fe2-O4</i></b>	1.91546	1.96415	1.95795	1.95753	1.96518
<b><i>Fe2-O2</i></b>	1.94024	1.931	1.92471	1.93036	1.93279
<b><i>Fe2-O7</i></b>	1.93334	1.96135	1.94929	1.97399	1.96222
<b><i>Fe2-N2</i></b>	2.18752	2.24085	2.24258	2.24662	2.23982
<b><i>Fe2-N5</i></b>	2.08708	2.17166	2.16348	2.13945	2.1701
<b><i>Fe2-N3</i></b>	2.11589	2.18755	2.17127	2.1794	2.18635
<b><i>N1-N2</i></b>	1.4278	1.39802	1.39377	1.38572	1.39643
<b><i>N6-N5</i></b>	1.41773	1.3971	1.39418	1.38655	1.39531
<b><i>N4-N3</i></b>	1.43062	1.39811	1.39435	1.38692	1.39653
<b><i>O1-Fe1-N6</i></b>	170.3849	170.8865	169.0994	170.3281	170.9741
<b><i>O5-Fe1-N1</i></b>	168.7467	166.7828	166.6057	165.4511	166.8412
<b><i>O8-Fe1-N4</i></b>	165.9458	168.2884	165.2623	166.2056	168.3589
<b><i>O4-Fe2-N5</i></b>	171.7548	169.9623	169.1666	169.4211	169.9599
<b><i>O2-Fe2-N3</i></b>	167.4431	166.1594	165.4062	165.6744	166.1898
<b><i>O7-Fe2-N2</i></b>	172.2738	170.758	169.3859	169.9418	170.8186
<b><i>Fe1-N4-N3-Fe2</i></b>	64.9497	63.1119	69.0727	64.8317	62.9905
<b><i>Fe1-N1-N2-Fe2</i></b>	65.5179	64.2503	68.34	69.7711	64.1186
<b><i>Fe1-N6-N5-Fe2</i></b>	63.2717	67.0845	69.3545	72.003	66.9174
<i>Average</i>		-0.11038	0.0699	0.1511	-0.1154
<i>Average(distance)</i>		0.0237	0.0182	0.0191	0.0233
<b><i>MAD (distance)</i></b>		<b>0.031</b>	<b>0.028</b>	<b>0.031</b>	<b>0.031</b>



**Figure S23.** Molecular orbital plots of  $\beta$ -HOMO and  $\beta$ -LUMO orbitals of  $1^{-1}$  complex in its LS-HS ( $S = \frac{5}{2}$ ) and LS-LS ( $S = \frac{1}{2}$ ) spin states. Molecular orbitals are generated at an isosurface value of 0.04 au. Compositions of metal ions are also given for each MO.

## Theoretical computation of redox potential using DFT

The most popular method to compute the redox potential uses the Born-Haber cycle presented in Figure S24.



**Figure S24.** The Born-Haber cycle for redox potential calculation.

$\Delta G_{(g)}^o$  is the gas-phase Gibbs energy of the addition of one electron to the oxidant. It is a sum of the negative calculated electron affinity, and the thermal and entropic corrections.

$\Delta G_{(solv, Ox)}^o$  and  $\Delta G_{(solv, Red)}^o$  are solvation energy values of the oxidant and reductant molecule.

This is achieved with the continuum solvation model CPCM. The entity  $\Delta G_{(s)}^o$  is then calculated as:

$$\Delta G_{(s)}^o = \Delta G_{(g)}^o + \Delta G_{(solv, Red)}^o - \Delta G_{(solv, Ox)}^o$$

The absolute reduction potential  $E_{abs}^o$  of the Ox/Red couple in Solvent is defined as

$$E_{abs}^{o(Solv)} = - \frac{\Delta G_{(s)}^o}{nF},$$

where  $n$  is number of electrons involved in the half-reaction, present case  $n = 1$ .  $F$  is the Faraday constant ( $F = 96485 \text{ J. mol}^{-1}.\text{V}^{-1}$  or  $23.061 \text{ kcal/mol}^{-1}.\text{V}^{-1}$ ).

#### DFT optimized atomic Cartesian coordinates of metal complex 1

Fe	14.196272	6.363878	11.622023
Fe	15.348543	6.730234	15.274753
O	15.196000	5.265561	10.355397
O	15.529230	8.523003	15.951398
O	15.018141	10.490514	17.613761
O	17.100960	6.062971	15.838105
O	12.457178	5.588630	11.339114
O	10.046197	5.403265	10.329585
O	14.347902	5.922643	16.739755
O	14.058826	7.982809	10.586517
O	14.389046	9.562321	8.518990
N	16.200643	6.867802	12.283498
N	16.129361	7.691839	13.405315
N	15.398153	4.923480	14.071588
N	14.266346	4.845253	13.260970
N	13.358147	7.089710	14.506797
N	13.432479	7.684123	13.247877
C	16.418350	5.008527	10.063210
C	16.646157	4.105615	8.964571

H	15.756819	3.758128	8.435934
C	17.900758	3.691421	8.623778
H	18.045939	2.994411	7.793157
C	19.047174	4.146701	9.344265
C	20.344882	3.695270	9.023162
H	20.461990	2.990610	8.195068
C	21.455008	4.124674	9.739479
H	22.452624	3.760859	9.484167
C	21.282560	5.023577	10.797952
H	22.145114	5.360365	11.378710
C	20.013162	5.502398	11.142294
C	18.856861	5.070451	10.419650
C	17.546655	5.524004	10.779114
C	17.394921	6.446150	11.894351
C	18.599053	6.865389	12.592702
H	18.494010	7.527799	13.447694
C	19.827306	6.412465	12.236595
H	20.704782	6.725269	12.808295
C	16.285218	8.961232	13.195085
H	16.546642	9.276764	12.174662
C	16.077761	10.009234	14.156116
C	15.679022	9.722256	15.495355
C	15.410832	10.836763	16.366336
C	15.547591	12.140152	15.903147
H	15.338846	12.980042	16.566672

C	15.948572	12.400228	14.572953
H	16.038483	13.433164	14.230798
C	16.207525	11.351883	13.715528
H	16.488549	11.538808	12.676501
C	14.715084	11.508320	18.533585
H	15.588563	12.156789	18.734224
H	14.417895	11.011058	19.466744
H	13.880286	12.144857	18.183942
C	17.985422	5.261744	15.372273
C	19.314187	5.365080	15.920163
H	19.446885	6.086085	16.728655
C	20.351698	4.632126	15.423946
H	21.358073	4.748750	15.836729
C	20.154592	3.708633	14.350948
C	21.231210	2.987200	13.795751
H	22.231790	3.141749	14.209033
C	21.039794	2.105975	12.737799
H	21.886296	1.568091	12.306290
C	19.753135	1.919840	12.224783
H	19.587962	1.237051	11.387840
C	18.656382	2.612636	12.753442
C	18.834849	3.537920	13.827903
C	17.736476	4.306452	14.333047
C	16.425529	4.153783	13.726015
C	16.279285	3.157087	12.681018

H	15.303013	3.031599	12.221674
C	17.336123	2.439432	12.222361
H	17.194077	1.732353	11.401062
C	13.303955	4.080718	13.664034
H	13.474858	3.482325	14.570909
C	12.006948	3.971499	13.051294
C	11.656106	4.747164	11.907254
C	10.317457	4.619175	11.399961
C	9.406808	3.769863	12.017561
H	8.390214	3.685026	11.632054
C	9.772690	3.019393	13.156493
H	9.034776	2.373243	13.635669
C	11.051739	3.120201	13.663117
H	11.340801	2.562027	14.556919
C	8.763479	5.348638	9.760365
H	8.521021	4.336158	9.386352
H	8.762975	6.051012	8.915871
H	7.978378	5.651418	10.479310
C	13.140074	5.569002	16.979777
C	12.915137	4.794270	18.175513
H	13.795676	4.589061	18.786679
C	11.675595	4.337350	18.512251
H	11.532974	3.743435	19.419684
C	10.538850	4.617756	17.691201
C	9.255590	4.137817	18.025209

H	9.141275	3.542009	18.935016
C	8.152750	4.409004	17.222412
H	7.165266	4.029568	17.494111
C	8.320768	5.171602	16.062443
H	7.464853	5.387539	15.417404
C	9.579432	5.665428	15.695768
C	10.725119	5.398440	16.507075
C	12.023297	5.883406	16.137330
C	12.171146	6.655450	14.913227
C	10.979247	6.896143	14.119170
H	11.083369	7.426839	13.176788
C	9.762423	6.428450	14.495253
H	8.894145	6.607521	13.856246
C	13.208633	8.957577	13.183600
H	12.903828	9.467215	14.109217
C	13.365872	9.780331	12.014900
C	13.799218	9.235093	10.769170
C	13.971737	10.145428	9.666486
C	13.722148	11.502852	9.827815
H	13.856882	12.186029	8.988563
C	13.296997	12.020464	11.072338
H	13.115531	13.092290	11.174728
C	13.123586	11.172576	12.145731
H	12.815341	11.561916	13.118795
C	14.563871	10.363455	7.378510

H	13.623300	10.861917	7.077369
H	14.887606	9.694066	6.570033
H	15.337246	11.139757	7.532438

**DFT optimized atomic Cartesian coordinates of metal complex 1<sup>-1</sup>**

Fe	14.228579	6.397862	11.685280
Fe	15.351556	6.740384	15.195331
O	15.195809	5.324699	10.343851
O	15.516799	8.609136	15.950178
O	14.919246	10.608209	17.547328
O	17.175785	6.050899	15.824916
O	12.483232	5.607626	11.343566
O	10.066207	5.426137	10.339519
O	14.346857	5.842825	16.740147
O	14.052589	8.014823	10.618250
O	14.364396	9.604344	8.551319
N	16.242459	6.913791	12.274152
N	16.192595	7.740065	13.396725
N	15.452201	4.883860	14.052483
N	14.318881	4.818375	13.246547
N	13.327516	7.093414	14.531388
N	13.403098	7.699204	13.280134
C	16.411352	5.039454	10.058417
C	16.624943	4.132202	8.957769

H	15.727943	3.806335	8.427600
C	17.869468	3.686143	8.620445
H	18.000672	2.984432	7.790630
C	19.024048	4.111076	9.345357
C	20.312097	3.626377	9.030504
H	20.413727	2.917188	8.203527
C	21.427402	4.024832	9.754207
H	22.416648	3.630570	9.509319
C	21.271798	4.927632	10.814186
H	22.137965	5.235060	11.405945
C	20.015743	5.440525	11.151366
C	18.850653	5.038933	10.421641
C	17.551678	5.524301	10.776473
C	17.419842	6.453101	11.892218
C	18.632889	6.838998	12.598807
H	18.530042	7.498699	13.456579
C	19.848818	6.354685	12.248767
H	20.731339	6.634663	12.829746
C	16.304778	9.013370	13.169983
H	16.526045	9.329489	12.139135
C	16.090519	10.061929	14.125868
C	15.662784	9.780903	15.469114
C	15.347616	10.931175	16.297283
C	15.479500	12.224221	15.817868
H	15.239299	13.074112	16.458953

C	15.917401	12.466658	14.490889
H	16.000777	13.493756	14.128209
C	16.209832	11.402289	13.668137
H	16.509921	11.571521	12.630386
C	14.555256	11.640442	18.416234
H	15.401421	12.321304	18.637209
H	14.226971	11.168061	19.353310
H	13.722841	12.251679	18.014080
C	18.033924	5.242309	15.363138
C	19.380920	5.335388	15.896968
H	19.519604	6.070422	16.692666
C	20.408862	4.583621	15.418920
H	21.417652	4.693567	15.830211
C	20.203418	3.644367	14.357931
C	21.273451	2.905057	13.819085
H	22.272800	3.054843	14.239054
C	21.082074	2.015303	12.766499
H	21.926598	1.464862	12.345227
C	19.798343	1.845955	12.242249
H	19.630880	1.166059	11.402730
C	18.705077	2.556236	12.755353
C	18.879681	3.486130	13.831422
C	17.784684	4.264662	14.327184
C	16.478788	4.120953	13.703799
C	16.340371	3.135386	12.643514

H	15.367407	3.029712	12.170210
C	17.391415	2.402169	12.200496
H	17.255146	1.700135	11.373265
C	13.341379	4.082041	13.663311
H	13.505721	3.491304	14.576308
C	12.036337	3.989678	13.057014
C	11.685541	4.768517	11.914122
C	10.342619	4.639765	11.414368
C	9.428946	3.795472	12.032413
H	8.409800	3.716786	11.651360
C	9.794953	3.048152	13.173164
H	9.055051	2.412018	13.663205
C	11.077539	3.148083	13.673188
H	11.366371	2.597146	14.571763
C	8.776345	5.390465	9.799002
H	8.508162	4.382978	9.424401
H	8.764650	6.096827	8.956723
H	8.006541	5.698181	10.533861
C	13.146802	5.509485	16.968846
C	12.905830	4.689776	18.144709
H	13.792117	4.450761	18.736276
C	11.666122	4.247781	18.488219
H	11.522560	3.631601	19.381775
C	10.521659	4.568199	17.688746
C	9.238190	4.096867	18.025708

H	9.128803	3.483991	18.925681
C	8.128183	4.391603	17.238630
H	7.139147	4.015721	17.512720
C	8.296037	5.168065	16.088420
H	7.439323	5.397186	15.447941
C	9.556063	5.656840	15.718519
C	10.710879	5.374391	16.518033
C	12.008196	5.858906	16.148204
C	12.147152	6.644785	14.930724
C	10.951120	6.883610	14.138226
H	11.057702	7.416929	13.196767
C	9.734475	6.422601	14.518343
H	8.862433	6.604044	13.884205
C	13.209328	8.976974	13.228108
H	12.924690	9.484964	14.160800
C	13.366274	9.805768	12.058588
C	13.789558	9.263308	10.808027
C	13.953994	10.182469	9.710877
C	13.708443	11.540152	9.875720
H	13.839284	12.225915	9.037389
C	13.297583	12.052869	11.126339
H	13.126051	13.125629	11.239967
C	13.134289	11.196689	12.196695
H	12.846925	11.582130	13.177829
C	14.553471	10.416150	7.427688

H	13.620507	10.929244	7.121863
H	14.879388	9.756854	6.610592
H	15.330716	11.187600	7.594806

**DFT optimized atomic Cartesian coordinates of metal complex 1<sup>-2</sup>**

Fe	14.240533	6.266575	11.640320
Fe	15.311822	6.628729	15.170871
O	15.307183	5.219310	10.240663
O	15.489401	8.513336	15.956332
O	14.939356	10.511179	17.577677
O	17.143322	5.936322	15.821461
O	12.418317	5.401452	11.263778
O	9.950078	5.252349	10.375266
O	14.273746	5.764197	16.722467
O	14.003855	7.970253	10.491199
O	14.433868	9.641957	8.510249
N	16.263177	6.848521	12.229124
N	16.175308	7.656315	13.361702
N	15.435858	4.754877	14.017729
N	14.307877	4.659172	13.208618
N	13.274565	7.007073	14.473634
N	13.348813	7.597772	13.215146
C	16.528171	5.001580	9.982747
C	16.817872	4.112686	8.872125
H	15.943531	3.745793	8.329354

C	18.085173	3.742788	8.537489
H	18.264994	3.062913	7.697098
C	19.209847	4.219068	9.281309
C	20.522205	3.809875	8.969725
H	20.665770	3.125371	8.126975
C	21.612532	4.245000	9.714824
H	22.623389	3.906091	9.470959
C	21.397888	5.106679	10.796938
H	22.239060	5.438493	11.412609
C	20.113219	5.547533	11.133628
C	18.971274	5.115496	10.377740
C	17.648108	5.533899	10.724436
C	17.461099	6.430305	11.861589
C	18.649326	6.829388	12.605026
H	18.506152	7.452369	13.484709
C	19.891017	6.411340	12.260059
H	20.751824	6.704153	12.867939
C	16.259418	8.931519	13.154843
H	16.470595	9.266739	12.127451
C	16.038527	9.972924	14.126717
C	15.632310	9.683743	15.475716
C	15.343514	10.835719	16.312806
C	15.468982	12.132798	15.841015
H	15.243238	12.979417	16.492892
C	15.876877	12.380447	14.506992

H	15.948903	13.407990	14.141551
C	16.146579	11.313261	13.675617
H	16.417497	11.484832	12.630048
C	14.569708	11.539826	18.439857
H	15.407588	12.234931	18.657821
H	14.248744	11.070747	19.382411
H	13.729627	12.144649	18.040487
C	18.020003	5.161996	15.335132
C	19.364623	5.265692	15.874626
H	19.484792	5.981222	16.691522
C	20.412451	4.552968	15.376904
H	21.418871	4.674780	15.792834
C	20.231434	3.645314	14.286711
C	21.320326	2.950424	13.723259
H	22.316158	3.113506	14.148413
C	21.151669	2.097297	12.638922
H	22.010528	1.589712	12.192067
C	19.870534	1.916618	12.106748
H	19.720939	1.269474	11.238101
C	18.760215	2.578185	12.643281
C	18.910228	3.475971	13.752519
C	17.796630	4.211136	14.270522
C	16.489018	4.048842	13.643844
C	16.379734	3.095189	12.548795
H	15.407616	2.990000	12.073052

C	17.449453	2.409755	12.080145
H	17.335445	1.740081	11.222893
C	13.304993	3.998756	13.688608
H	13.445486	3.485625	14.653117
C	11.991113	3.906380	13.100744
C	11.632638	4.637971	11.915258
C	10.251829	4.514022	11.486716
C	9.336579	3.738512	12.179423
H	8.300253	3.675178	11.841221
C	9.720067	3.042203	13.350941
H	8.979103	2.463342	13.907225
C	11.023347	3.135059	13.793286
H	11.328483	2.632378	14.715744
C	8.635970	5.249815	9.916343
H	8.295077	4.239098	9.608257
H	8.598739	5.915288	9.040372
H	7.920863	5.624723	10.677556
C	13.059632	5.515216	16.981953
C	12.788467	4.778017	18.205605
H	13.670684	4.520827	18.796547
C	11.530966	4.431402	18.595065
H	11.367145	3.878367	19.526694
C	10.395193	4.770375	17.794153
C	9.090268	4.398049	18.177430
H	8.961745	3.851238	19.117517

C	7.988359	4.704472	17.386282
H	6.982402	4.405018	17.694518
C	8.185324	5.391197	16.181417
H	7.334516	5.621804	15.532777
C	9.463017	5.781230	15.765205
C	10.614965	5.490584	16.571169
C	11.929488	5.882348	16.159140
C	12.091670	6.602300	14.897372
C	10.898017	6.823625	14.090246
H	11.028611	7.272015	13.108217
C	9.667377	6.443841	14.506378
H	8.800806	6.603431	13.858522
C	13.162468	8.877304	13.161015
H	12.893176	9.392080	14.096922
C	13.325993	9.716442	12.001107
C	13.770873	9.194718	10.734362
C	13.985946	10.179475	9.684494
C	13.752737	11.529485	9.887464
H	13.920502	12.244502	9.079115
C	13.304433	12.005360	11.145368
H	13.143040	13.075913	11.295777
C	13.106398	11.107504	12.173331
H	12.801717	11.461550	13.162333
C	14.729920	10.503706	7.458057
H	13.840696	11.068945	7.107989

H	15.100592	9.883732	6.627537
H	15.512189	11.243835	7.725467

**DFT optimized atomic Cartesian coordinates of transition state TS1**

Fe	-0.48461655555535	-0.32077258237994	-1.43211093331095
Fe	0.48491928246371	0.37310699142011	2.02030346559354
O	0.50138817515903	-1.21835370334173	-2.95401906029831
O	0.65230495136602	2.33892670758948	2.57776479242973
O	0.24789330111341	4.46887294338122	4.06189074664703
O	2.33607258825462	-0.23376365491164	2.64593990421755
O	-2.39152057574865	-0.70906226085325	-1.69254652122479
O	-4.80596109601028	-0.88243109759236	-2.69257166447108
O	-0.47708391057206	-0.49193808260250	3.62696625250010
O	-1.26943936792481	1.94264706521954	-2.45566709206751
O	-1.15433149129926	3.77014335725025	-4.34249405546330
N	1.40339032521194	0.39443705284281	-0.97821648933310
N	1.29529187452809	1.28027146083343	0.09140757884903
N	0.59658303338371	-1.57784090199624	1.00216463757802
N	-0.52061767028610	-1.69508255307667	0.17694134735790
N	-1.58388109008104	0.71639043305056	1.45793014385310
N	-1.57345481307875	1.42893529357734	0.27997685484090
C	1.72484889707269	-1.59187303210914	-3.08206078174515
C	2.01600927985967	-2.61331152756961	-4.06114019775686
H	1.16300596290815	-2.99161674868474	-4.62918107859735
C	3.27621110821376	-3.09908124237933	-4.25332513898292

H	3.46376984817522	-3.88313995458575	-4.99441738727400
C	4.38254298224219	-2.60860703555150	-3.49331194254581
C	5.69118827774175	-3.10644561277773	-3.67920655793028
H	5.85422854523022	-3.88182084390639	-4.43459462383660
C	6.75688010220944	-2.63580653676642	-2.92271573181887
H	7.76356566357078	-3.03707946154855	-3.06899545630237
C	6.52736220292003	-1.65145464048285	-1.95421153969987
H	7.35127707403334	-1.28322222713427	-1.33662310408185
C	5.24778074190865	-1.12651705733004	-1.74449472907558
C	4.13419323010858	-1.59491397536069	-2.51373926295796
C	2.81365149191057	-1.08134926128747	-2.29660945652363
C	2.60989643893972	-0.07901613226786	-1.26163161105829
C	3.76938908557422	0.36459095150841	-0.50555753559316
H	3.61177532695222	1.07689605217862	0.29840031381432
C	5.00870294744214	-0.13143911725595	-0.73769185644266
H	5.84863785657642	0.20128766596765	-0.12264670170520
C	1.49237250162310	2.52485782764632	-0.20698802223192
H	1.76609857458245	2.75922657356189	-1.24825357532304
C	1.35586137279268	3.63825308256222	0.68273963937600
C	0.90908940901460	3.46230623985389	2.03807077924558
C	0.70885528092880	4.68282840854304	2.80043436998560
C	0.97535859112782	5.92978069433184	2.25755651405457
H	0.82827871965982	6.83095833122432	2.85481967070805
C	1.43914051896901	6.06130376893737	0.92365362500738
H	1.63950096042328	7.05354948155871	0.51315958820367

C	1.61718337463384	4.93123865736270	0.15810841316710
H	1.94489253552956	5.01106097302611	-0.88271149754083
C	0.01692993115111	5.57636030652130	4.88154056106582
H	0.93989788351740	6.16246531410701	5.06070207952538
H	-0.35184006731219	5.19056094670459	5.84243298019560
H	-0.74368238679036	6.26035047189392	4.45465146619473
C	3.18131442291311	-1.10314234850219	2.28221893767566
C	4.51944481179232	-0.98727772285296	2.83376548089691
H	4.65816319156723	-0.18004211602734	3.55630052126794
C	5.53800035693140	-1.80692938171158	2.45880623863148
H	6.53909206616402	-1.68168855169081	2.88440146051928
C	5.33320944674755	-2.83770045100124	1.48665474936262
C	6.39561615388298	-3.65219569028695	1.05162505257547
H	7.38632800426677	-3.49165129323958	1.48824826173251
C	6.20913855400957	-4.62899568648230	0.07828696061917
H	7.04851896326192	-5.24012594167949	-0.26221889868487
C	4.93812100665833	-4.80520440985635	-0.47405215645799
H	4.77567753718650	-5.55305995043097	-1.25520992781654
C	3.85163333512009	-4.02245148623361	-0.06148952553867
C	4.01940605945854	-3.00921667291627	0.93829210828600
C	2.93024845346179	-2.17028212058299	1.33890047635280
C	1.63226251226999	-2.35084710990503	0.70780293034276
C	1.50703641609775	-3.40073594524241	-0.29146929722331
H	0.54136811751064	-3.52802516393103	-0.77516617039598
C	2.55307023934211	-4.18594368591899	-0.64822921685196

H	2.42409397726545	-4.94777809988111	-1.42211590119659
C	-1.54137598377283	-2.34819093368325	0.61889877215795
H	-1.43929661757593	-2.88278694741214	1.57549779888377
C	-2.83230743522171	-2.41124245274116	-0.02734671822103
C	-3.17977386202751	-1.58272054335544	-1.14356182107619
C	-4.52175018739347	-1.70727591160985	-1.64448162813551
C	-5.43474567163788	-2.58316151708278	-1.06876387119222
H	-6.44851065096608	-2.65666006151204	-1.46251605487834
C	-5.07362637599718	-3.37020383544978	0.04515049716300
H	-5.81309606012984	-4.02602821281692	0.51114741044648
C	-3.79713296229874	-3.27327778918179	0.55442411801227
H	-3.51292918737407	-3.84929770222754	1.43933863713853
C	-6.08186108489654	-0.95256836412241	-3.26064801865575
H	-6.31309023848717	-1.96549427884290	-3.64489802774771
H	-6.09707617464434	-0.24387811968649	-4.10056069740909
H	-6.87595513565856	-0.67314621620236	-2.53973328636658
C	-1.63964693989873	-0.95076203571029	3.83397195582053
C	-1.80389029402275	-1.87258069588116	4.94495204872466
H	-0.89575059262874	-2.08750678751877	5.51214668879503
C	-3.00153734924938	-2.44074741212475	5.25467378016235
H	-3.08519146218125	-3.13977983856802	6.09309249635515
C	-4.17877454605457	-2.14383908092426	4.49480970439322
C	-5.42102058196827	-2.73234585319918	4.80203826277184
H	-5.47216798664147	-3.42878120632709	5.64448488009469
C	-6.56247312001262	-2.44579039543696	4.05827525749101

H	-7.51804433084676	-2.91370780414655	4.30871120650891
C	-6.47108843691782	-1.55594272934001	2.98344784407872
H	-7.35473708555627	-1.32949668283193	2.37972549373878
C	-5.25364481850936	-0.95476902574766	2.63968088122371
C	-4.06643241037749	-1.23325636677323	3.39278908555646
C	-2.80929020232088	-0.64469018218036	3.04304815382960
C	-2.73492280607370	0.21762808134288	1.87462450313282
C	-3.96573622731593	0.47603110748819	1.14263004239455
H	-3.90728029514959	1.08940535575474	0.24661009269323
C	-5.14807070870924	-0.07628906708550	1.50929064035813
H	-6.04526755254888	0.11028156787782	0.91245955333683
C	-1.68655767056190	2.70125595839977	0.38814421238189
H	-1.82889127629369	3.14722976577069	1.38466104715152
C	-1.60912369765009	3.63872576992142	-0.72982575438025
C	-1.43555534535271	3.24619681207662	-2.07155149278357
C	-1.33391108313170	4.23533698293727	-3.07878450765494
C	-1.43898683534631	5.59452965913976	-2.74958543973395
H	-1.37201084423541	6.35837721158557	-3.52340738673233
C	-1.62567139763724	5.97393617013124	-1.41564983228247
H	-1.69166965838724	7.03501642980521	-1.16069309672421
C	-1.71026931892073	5.01193207221956	-0.42117832533464
H	-1.81201990398420	5.30767784200776	0.62444202063810
C	-1.04897122938978	4.69792590971489	-5.39535062014704
H	-1.96447742196874	5.31035674866972	-5.49360849579463
H	-0.90664559576507	4.11380475762844	-6.31367081558909

H	-0.18634271082079	5.37608063388774	-5.25884174126472
H	-2.11132801387276	1.44251020564094	-2.50898437792624
H	-1.01661681631552	1.46879150249066	-4.56664278953885
H	-0.56102087064938	0.02889530636574	-4.36545899245076
O	-1.15396978492420	0.57191821833708	-4.91259749672777

**DFT optimized atomic Cartesian coordinates of transition state TS2**

Fe	-0.27017844274118	-0.28806700320260	-2.11615248868706
Fe	0.64624255226620	0.53828718073248	1.46752918829677
O	0.75013424355554	-1.44969448253170	-3.28569403707432
O	0.87447958701110	2.47953648327632	2.07156983397440
O	0.71734588670623	4.59993291182106	3.60822867096351
O	2.55298244412094	-0.03092904624140	1.99483046113882
O	-2.23178084832349	-0.48663656015053	-2.06283184789559
O	-4.71955228262820	-0.25871903712748	-2.77958333814257
O	-0.04813466114250	-0.38499986326340	3.15480899517698
O	-2.82179868762273	2.32375879277770	-2.19322467867817
O	-3.50724352369451	4.25806518699558	-3.97649031631556
N	1.53272329803460	0.46306131100034	-1.47284156978840
N	1.32896416754146	1.40279084788414	-0.45954347447774
N	0.74206403788503	-1.43619189455579	0.48722485978159
N	-0.37064035818431	-1.54499881815397	-0.34570333565227
N	-1.53855719564163	0.84401966979403	1.20262337576659
N	-1.86462637722133	1.71472092091558	0.17495902827754
C	1.96712163412629	-1.86486694727260	-3.21061584364670

C	2.29886595423604	-3.02537200640477	-3.99538333267755
H	1.49894257275094	-3.41896195487272	-4.62563217403841
C	3.52032116248868	-3.62652289531616	-3.91361710915137
H	3.73491913277063	-4.52941126682614	-4.49353991608104
C	4.53470353027887	-3.10666950320896	-3.05507223411870
C	5.77436938435955	-3.76347737381770	-2.90460631496502
H	5.94624281905580	-4.68636862553169	-3.46599394951816
C	6.74923993892912	-3.27236264329905	-2.04770613013360
H	7.69297522062335	-3.80687329724701	-1.91904754154416
C	6.50551914914825	-2.09188648705718	-1.33703193877604
H	7.25965544122916	-1.69749677232621	-0.65118829023708
C	5.29359719724900	-1.40478292244636	-1.46543475476972
C	4.26200577413951	-1.91013923790162	-2.31854548290844
C	2.98110380774154	-1.26824452497352	-2.39204085420433
C	2.73408663762965	-0.09181196793156	-1.56056789866756
C	3.83887061114319	0.43030336760351	-0.77760317475144
H	3.66196215561737	1.30877579964619	-0.16418573338160
C	5.03810381149430	-0.19912164103488	-0.72849103194786
H	5.83003366980973	0.19176832756543	-0.08471056966979
C	1.47467475415455	2.65196120545792	-0.78593739976436
H	1.69584441198915	2.87862104762445	-1.83625301130264
C	1.37641339739866	3.76963132766725	0.10840620754953
C	1.09817542495353	3.60009138263668	1.50857092553084
C	1.02255242101706	4.81605454816838	2.29998533674792
C	1.23630740941222	6.06265672228956	1.73450334201862

H	1.18046474351205	6.96196786947976	2.35024641565931
C	1.51813077617184	6.19328191826319	0.35077912171441
H	1.67006169354054	7.18613544383223	-0.07849030931048
C	1.57824917556239	5.06621201387578	-0.43794890092321
H	1.76729519865400	5.15053500167859	-1.51172287323554
C	0.60082773638863	5.70428262340794	4.45771481904244
H	1.54586412424291	6.27824305366972	4.53001102703896
H	0.34564937204406	5.31540910066822	5.45392465109435
H	-0.19694688330626	6.40047925107194	4.13053782886491
C	3.26163737717480	-1.07460929195633	1.91692510492872
C	4.53211167687258	-1.06110919655035	2.62190856207311
H	4.72976481569557	-0.16196306696731	3.20937382978829
C	5.42191137942776	-2.08729478659595	2.54767496629981
H	6.37174424016263	-2.04200344920046	3.09056055521726
C	5.14916659187444	-3.24083849097662	1.74508826378522
C	6.08616040901997	-4.28345373366100	1.62217265995658
H	7.02632240708486	-4.20539123282904	2.17677160582172
C	5.84336247005555	-5.38488824618387	0.80634756791068
H	6.58637424752819	-6.18086477890853	0.71316769259062
C	4.64426771422373	-5.45019064619840	0.09226622733883
H	4.44477398670553	-6.29540147878873	-0.57295781187318
C	3.67882032779737	-4.43935590058328	0.19617364295451
C	3.90142998721137	-3.30207666819022	1.03960031209996
C	2.93365590541403	-2.25183122740974	1.14353063799528
C	1.68688446768946	-2.35963510468502	0.39811889867866

C	1.52283087086712	-3.51401742986425	-0.47207540864811
H	0.62380105118982	-3.57027245545157	-1.08138875512009
C	2.46122735539635	-4.48958942542540	-0.56072385105683
H	2.30531837166457	-5.33187542004991	-1.24076321849637
C	-1.37894939515476	-2.22856061870338	0.09465453706618
H	-1.24464709393272	-2.79646239117502	1.02640604574846
C	-2.69691094469748	-2.24283718617943	-0.48915965654132
C	-3.05866473999499	-1.30738568720281	-1.50195411083268
C	-4.44426892839895	-1.23575277656938	-1.87641815402405
C	-5.37931153166624	-2.09794367353988	-1.32012629848473
H	-6.42681340933082	-2.03944414349805	-1.62069235834164
C	-4.99150128778185	-3.04144052270722	-0.34285299496310
H	-5.74238765564767	-3.69374516493279	0.10781641912207
C	-3.67729059821012	-3.09664056650018	0.07552559802909
H	-3.38154295820814	-3.77728112257194	0.87766439849613
C	-6.05604829521412	0.06443053667562	-3.04125472290292
H	-6.59066559111472	-0.74955383491500	-3.56996477206801
H	-6.04330753838132	0.95934453157688	-3.67927841116878
H	-6.61273819681869	0.29481457141369	-2.11221418905362
C	-1.10959051135103	-1.01547306940437	3.43268041934611
C	-1.02718705321966	-2.03433491006848	4.46498298275067
H	-0.03724419232959	-2.17670829911119	4.90352723652525
C	-2.10380087517447	-2.77092526295018	4.85164334213833
H	-2.00476641581842	-3.54014953689861	5.62432537749457
C	-3.39228169890038	-2.55466227530836	4.26435163812293

C	-4.51586952059977	-3.29996039222090	4.67094515642095
H	-4.38226869805637	-4.06208692297340	5.44478622441742
C	-5.77235664539458	-3.08002250185081	4.11249932551615
H	-6.63387177278366	-3.66813280371793	4.43891031875063
C	-5.91884212837974	-2.09509968942548	3.13139310719425
H	-6.89764461639405	-1.90756839221537	2.68043741333956
C	-4.82434284352024	-1.33865943927369	2.69209310910431
C	-3.52048946524030	-1.55366141839734	3.24442738326588
C	-2.38670356694343	-0.80184879805660	2.79254618335819
C	-2.55811674719094	0.16635740506999	1.71837070151006
C	-3.90823065351648	0.37016765146194	1.21376301419747
H	-4.04869791637055	1.09639189921179	0.41686125990512
C	-4.96716240965014	-0.34199766585873	1.67033031233671
H	-5.95753663200259	-0.18104436218384	1.23614901497980
C	-1.73633438126826	2.97406373533771	0.40983366897138
H	-1.41677478533931	3.31850890254336	1.40186500109410
C	-2.04631432708486	3.97419020594297	-0.60279322689961
C	-2.59088653257024	3.60462280600122	-1.85843446282984
C	-2.93107397404594	4.61353409457563	-2.78489942742944
C	-2.70447904406485	5.95390142007785	-2.47333255842425
H	-2.98180566297634	6.70523968985400	-3.21718373135000
C	-2.14860261976785	6.32101199100786	-1.24051579138645
H	-1.96340952222818	7.37262109189045	-1.01033282885632
C	-1.82520389570424	5.33433586752850	-0.31657115330771
H	-1.37243063237418	5.59649430873865	0.64172970953638

C	-2.59459935935451	4.07927883591636	-5.05054570656314
H	-2.21039823415810	5.05376820010306	-5.41228607006233
H	-3.14720008301955	3.59180077263396	-5.86768811625071
H	-1.73396017888053	3.45829722969734	-4.75452566983451
H	-2.56237880567163	1.73718149051250	-1.42265608119549
H	-1.62248193996655	1.42645208373203	-3.28919641510301
O	-0.71859428437063	1.07756487800009	-3.47946737506620
O	0.02219996190359	2.61610762785153	-3.54626294325252
H	0.65874126479812	2.32497198442248	-4.21961326699144

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