Direct synthesis of phenol by novel [FeFe]-hydrogenase model complexes as catalysts of benzene hydroxylation with H$_2$O$_2$

Xia Zhang, Tianyong Zhang,* Bin Li,* Guanhui Zhang, Li Hai, Xiaoyuan Ma and Wubin Wu

Three new [FeFe]-hydrogenase model complexes, $\mu$-(SCH$_2$CH$_2$)CH$_2$S)Fe$_2$(CO)$_6$ (complex 1), $\mu$-(SCH$_2$CH$_2$)CH$_2$S)Fe$_2$(CO)$_5$PCy$_3$ (complex 2) and $\mu$-(SCH$_2$CH$_2$)CH$_2$S)Fe$_2$(CO)$_5$PPN$_3$ (complex 3) were prepared. The structures of complexes 1–3 were characterized by FT-IR, UV-vis, $^1$H, $^{13}$C, $^{31}$P NMR spectra and single-crystal analyses. The electron density of these model complexes was studied by IR spectra, UV spectra and electrochemical analysis and evaluated against their respective catalytic performances. The CV (cyclic voltammetry) study of complex 2 showed a less positive oxidation event at 0.6 V and a more negative reduction event at $-1.94$ V, which is in accordance with the enlargement of electron density at diiron centers when CO were substituted by better electron donor ligands. Of all these three complexes, complex 2 exhibited the best catalytic activity, with a yield of phenol of up to 24.6% and selectivity up to 92%, which is consistent with its higher electron density of the Fe–Fe bond. This study revealed the correlations between the electron density of the catalytic site of catalysts and their performance in catalytic hydroxylation of benzene. Based on these experimental results, a catalytic oxidation mechanism via an Fe$^{2+}$–$\mu$–O–Fe$^{2+}$ intermediate as oxygen transfer reagent has been proposed.

Introduction

Being an important industrial feedstock used in production of fibres, resins, antioxidants, plastics, agrochemicals and medicines, phenol is in high demand. The current method for industrial scale production of phenol is mostly based on the cumene process, which produces a large amount of acetone as the by-product. Since the market demand for phenol is much higher than for acetone, a method of direct oxidation of the by-product. Since the market demand for phenol is much higher than for acetone, a method of direct oxidation of benzene to phenol is expected to bring better economic benefits. The direct catalytic hydroxylation of benzene to phenol under mild conditions has attracted great interest and has been extensively investigated due to the reduced expense of the base materials. Some new catalysts have already been tried in this reaction, like iron complexes, TS-1 and Py$_3$PMo$_12$V. However, the yield of phenol by the direct hydroxylation of benzene has been found to be lower than produced through the traditional method. The hydroxylation process can not avoid the excessive oxidation and subsequent formation of by-products derived from phenol thereby leading to an overall decline in desirable yield. Therefore, the seeking for novel catalysts with high selectivity and efficiency for the direct catalytic hydroxylation of benzene to phenols under mild circumstances has been an ongoing process and a challenge.

Among diversified catalysts, iron-based catalysts attracted great research interest for many years due to the convenience, low cost, environmental amity and low toxicity of iron. As reported in the previous studies, iron-based catalysts usually have high selectivity and efficiency in a series of chemically challenging oxidative processes. Especially, iron-based biomimetic catalysts with low barriers to suitable redox states for selective oxidations, have been studied. These catalysts mimic the active site of natural oxygenase enzymes and serve as high-performance catalysts for selective hydrocarbon oxidation.

[FeFe]-hydrogenases are also iron-based bio-catalysts which are much more efficient in the evolution and oxidation of H$_2$ under mild conditions. However, oxygenation is found to occur at the dithiolato-sulfur or at the Fe–Fe bond site of the [FeFe]-hydrogenase active site model complexes. This oxygenation may lead to S-oxygenated products which having relatively stable O atom. The other route is the formation of Fe-based oxidative addition products of Fe$^{2+}$–$\mu$–O–Fe$^{2+}$. Although this species are demonstrated thermodynamically favourable,
they are not isolated to date. According to the reaction kinetics research on [FeFe]-hydrogenase active site model complexes, the diiron Fe−Fe subsite is thermodynamically favored to be oxidized to the active Fe$^{2+}$−μ-O−Fe$^{2+}$ species. The iron-based highly active Fe$^{2+}$−μ-O−Fe$^{2+}$ species as the crucial intermediates during oxygenation could be used for transfer of oxygen in the biomimetic hydrocarbon hydroxylation. This is due to the fact that the [FeFe]-hydrogenases model complexes have an active site that remains unconstrained towards a suitable redox state. This also implies high activity towards the oxygenation reactions. Our group’s previous research showed the Fe−Fe bond based oxidogenated μ-O complex has the ability of transferring oxygen atom to the aromatic substrates. Supporting DFT calculations and the experimental catalytic activity towards hydroxylation tend to confirm the mechanism. Although the selectivity for phenol is high, the yield is not ideal yet. Chen’s group showed the benzene conversions by the catalytic system selectivity for phenol is high, the yield is not ideal yet. Thus, it is essential to prepare some new [FeFe]-hydrogenase model compound with a PPh$_3$ analogue ligand aimed at enhancing the yield of catalytic hydroxylation of benzene.

Herein, a new dinuclear [2FeS] mimic complex μ-(SCH(CH$_2$CH$_3$)$_2$CH$_2$S)−Fe$_2$(CO)$_6$ (1) was obtained by reaction of Fe(CO)$_3$ with excess 1,2-butanedithiol. For studying the impact of the ligand at the active site of [FeFe]-hydrogenase model compound, the complexes namely μ-(SCH(CH$_2$CH$_3$)$_2$CH$_2$S)−Fe$_2$(CO)$_6$L (2: L = PCy$_3$, tricyclohexyl phosphine; 3: L = PPh$_3$, triphenyl phosphine) have been prepared by controllable CO substitution of 1 with PCy$_3$ and PPh$_3$, as shown in Scheme 1. The structural elucidation of these model compounds featured the use of UV-vis, FT-IR, $^1$H, $^{13}$C, $^{31}$P NMR spectra and single-crystal analyses. For investigating the correlations between the catalytic reactivity and the electronic features of the catalysts, their electrochemical properties, FT-IR, UV-vis and their catalytic performance have been discussed. The results indicate that complex 2 has the greatest electron density and also the best catalytic reactivity. This finding suggests that the catalytic reactivity of the catalysts could be improved by enhancing the electron density of active sites in the catalysts.

**Results and discussion**

**Synthesis and spectroscopic characterization of complexes 1–3**

The reaction of Fe(CO)$_3$ with 1,2-butanedithiol in refluxing toluene, yielded the anticipated μ-(SCH(CH$_2$CH$_3$)$_2$CH$_2$S)−Fe$_2$(CO)$_6$ (1). Mono-substituted complexes μ-(SCH(CH$_2$CH$_3$)$_2$CH$_2$S)−Fe$_2$(CO)$_6$L (2: L = PCy$_3$, tricyclohexyl phosphine; 3: L = PPh$_3$, triphenyl phosphine) were obtained by treating 1 with PCy$_3$ and PPh$_3$, respectively, along with Me$_3$NO in THF at ambient temperature with good yields through sufficient stirring. The $^1$H NMR spectra of complexes 1–3 consist of the expected signals for the 1,2-butanedithiol bridges and the phosphine ligands. In the $^{13}$C NMR spectrum, complexes 1–3 show peaks at $\delta = 208.82$, 210.86 and 210.82 ppm, respectively. These peaks can be safely assigned to the Fe=CO carbon atom. The $^{31}$P NMR spectrum of 2 and 3 was display a singlet at $\delta = 70.89$ and 63.95 ppm, respectively. The observations from the elemental analysis in all products agree well with the assumed compositions.

The IR spectra for the compounds 1–3 obtained in the hexane solution show distinct peaks between 2100 and 1900 cm$^{-1}$, characteristic of stretching vibrational modes of terminal metal carbonyls. As a result of the electron-donating phosphine ligands, the IR spectra of complexes 2–3 are shifted by $\nu_{CO} = 30$ cm$^{-1}$ and $\nu_{CO} = 27$ cm$^{-1}$ towards lower frequency compared to the all-carbonyl compound 1 (Fig. 1). The application of $\nu(CO)$ bands is regarded to be a great tool for identifying the alteration of electron density on Fe atoms in the model compounds. The CO stretches of the mononuclear complexes 2–3 are lower in wavenumber in comparison to the all-carbonyl compound 1 which is indicative of the strong donating...
character of PCy3 and PPh3 group. The red-shi

value order for the ν(CO) bands of 2–3 is in accordance with electron-donation capabilities of diverse phosphine ligands, PCy3 ligand reveals its strong electron-donation character compared to the PPh3 ligand. These showing a distinct ranking of 2 > 3 > 1 in electron density on active sites.

To further clarify the electronic effect of the terminal ligands (L) on the activity of the complexes, UV-vis spectra of the compounds 1–3 were recorded in dichloromethane. As shown in the Fig. 2, prominent absorption peaks of complexes 1–3 are observed at 326 nm, 360 nm and 354 nm, respectively. These peaks in the visible region can be attributed to π–π* transition. The electronic transitions of compound 2 (360 nm) and 3 (354 nm) were found to be red-shifted compared to compound 1 (326 nm). These indicated more electron-donation ligand will improve the electron densities at the diiron core. The electronic transition signal compound 2 is about 6 nm red-shifted compared to compound 3, which indicated the PCy3 ligand has the strong electron-donation than the PPh3 ligand. These are in accordance with the result derived from the IR spectra analysis and showed a distinct ranking of 2 > 3 > 1.

Molecular structures of complexes 1–3

X-ray quality single crystals of complexes 1–3 were obtained by slow evaporation of the respective hexane–dichloromethane solutions at low temperature under normal atmospheric pressure. The structures of the compounds 1–3 are illustrated in Fig. 3. As the coordination locations for phosphine ligands on the diiron compounds are not identifiable only via the spectroscopic data, the X-ray diffraction study was conducted for 1–3. The [Fe2S2] skeleton of compounds 1–3 have the famous butterfly structure where every iron center takes on the twisted square-pyramidal coordination shape. Some of the important bond lengths and angles are displayed in Table S1,† while crystallographic data are shown in Table S2.† The Fe–Fe distances (2.5090 (4) Å in 2, and 2.5126(11) Å in 3) were found to be slightly longer compared to the Fe–Fe distances in complex 1 (2.5054 Å in 1). The PCy3 or PPh3 substituent group has a minor impact on the Fe–Fe distances. PCy3 or PPh3 ligand was located in the apical site of Fe atom and was placed trans to the Fe–Fe bond. The Fe–P bond distances of complex 2 (2.2734(5) Å) are longer than that of complex 3 (2.2447(15) Å), showing the larger steric effects on the coordination geometry following the order PCy3 > PPh3. X-ray crystallographic analysis for complexes 2 and 3 implied that the replacement of one CO by tertiary phosphine ligands on the μ-(SCH(CH2CH3)CH2S)-Fe2(CO)6 offers solely an apical isomer as illustrated in Fig. 3. The displacement of CO by the tertiary phosphine ligands (PCy3 or PPh3), which are better electron-donating, increases the electron density on the iron centers of a diiron dithiolate model compound, thereby further strengthening the strong back-bonding from the Fe atoms to each other.
the CO ligands and simultaneous weakening of CO triple bonds.\textsuperscript{11,40,41}

**Electrochemistry**

The electrochemical properties of complexes 1–3 were investigated via cyclic voltammogram (Fig. 4). CV data are shown in Table 1. The cyclic voltammogram of complex 1 displays an irreversible oxidation process at $E_{pa} = 0.74$ V and a quasi-reversible process at $E_{pc} = -1.78$ V. The first oxidation process is ascribed to the process from Fe\textsuperscript{II}Fe\textsuperscript{I} to Fe\textsuperscript{III}Fe\textsuperscript{I}, and the initial process is able to be allocated to the Fe \textsuperscript{II}Fe\textsuperscript{I}/Fe\textsuperscript{II}Fe\textsuperscript{0} reduction.\textsuperscript{11} The CV of the PCy\textsubscript{3}-substituted complex 2 and PPh\textsubscript{3}-substituted complex 3 in CH\textsubscript{3}CN solution display one irreversible reduction process (Fe\textsuperscript{II}Fe\textsuperscript{I}/Fe\textsuperscript{II}Fe\textsuperscript{0}) at $E_{pc} = -1.94$ V and $-1.84$ V, and a quasi-reversible oxidation process at $E_{pa} = 0.6$ V and 0.75 V, respectively. Compared with the first reduction and oxidation event of complex 1, the PCy\textsubscript{3}-substituted complex 2 and PPh\textsubscript{3}-substituted complex 3 reveal cathodic shifts in the reduction potential, all in accordance with the enlargement of electron density in diiron centers when CO were substituted by better electron donating ligands.\textsuperscript{11,12}

**Table 1 Redox potentials of the complexes**

<table>
<thead>
<tr>
<th>Complex</th>
<th>$E_{pc}$ (V) vs. Fe/Fe\textsuperscript{+}</th>
<th>$E_{pa}$ (V) vs. Fe/Fe\textsuperscript{+}</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$-1.78$</td>
<td>0.74</td>
</tr>
<tr>
<td>2</td>
<td>$-1.94$</td>
<td>0.60</td>
</tr>
<tr>
<td>3</td>
<td>$-1.84$</td>
<td>0.75</td>
</tr>
</tbody>
</table>

**Hydroxylation reactions of aromatic compounds**

Although many iron-based catalysts have been studied, but the [FeFe]-hydrogenase model complexes have rarely been used for the direct production of phenol from benzene.\textsuperscript{11} In the present work, the [FeFe]-hydrogenase model compounds 1–3 were used as catalysts in the single-step hydroxylation from benzene into phenol with oxidant. We choose O\textsubscript{2}, H\textsubscript{2}O\textsubscript{2}, PhIO, m-CPBA as oxidants in the single-step hydroxylation. Among the oxidants, H\textsubscript{2}O\textsubscript{2} has a higher phenol yield than m-CPBA and O\textsubscript{2}, PhIO were not have the ability to catalyze the single-step hydroxylation from benzene into phenol with these complexes under the given experimental conditions, as summarized in Table S3.\textsuperscript{†} Thus, we used H\textsubscript{2}O\textsubscript{2} as the oxidant in the single-step hydroxylation from benzene into phenol with [FeFe]-hydrogenase model compounds. The catalytic performance of complex 1 was assessed by following the procedures well established in literature\textsuperscript{42–45} in acetonitrile using H\textsubscript{2}O\textsubscript{2} as oxidant. Some parameters which can potentially impact the catalytic properties, like the amount of H\textsubscript{2}O\textsubscript{2}, amount of catalyst, the reaction time, as well as the temperature, were also examined.

The phenol yield and selectivity is significantly affected by the amount of H\textsubscript{2}O\textsubscript{2} (Fig. 5). The yield increased with increase in amount of H\textsubscript{2}O\textsubscript{2} from 0–5.0 mmol. The maximum yield of phenol was 10.5\% when the amount of H\textsubscript{2}O\textsubscript{2} was 5.0 mmol. But with the further raise in the amount of H\textsubscript{2}O\textsubscript{2}, the yield and selectivity of phenol has an obvious decreased, because too much H\textsubscript{2}O\textsubscript{2} may be resulted in excessive oxidation of phenol. Therefore, the best amount of H\textsubscript{2}O\textsubscript{2} applicable for this reaction is 5.0 mmol.

The phenol yield as well as selectivity is displayed with the variation temperature in Fig. 6. The result showed none of phenol is detected below 40 °C, since induction for a longer period of time at a low-temperature blocks the creation of phenol. The yield of phenol increased with temperatures growing from 40 to 60 °C. But with further rise in temperature, a clear
The yield went to a maximum (12.6%) after 3 h. Declining selectivity due to phenol overoxidation was also discovered with extension in the reaction time.\textsuperscript{31} Under the optimized circumstances (1, 30 μmol; benzene, 1.12 mmol; CH₃CN, 2.0 mL; H₂O₂, 5.0 mmol; reaction time, 3 h), the selectivity and yield of one-step conversion of benzene to phenol with catalyst complex 1 is 12.6% and 83%, respectively.

According to previous results,\textsuperscript{12} the catalytic hydroxylation activity of such diiron complexes is based on their special structural and electronic properties, which was noticed firstly in some precious fundamental studies of the oxygen sensitivity of the [FeFe]-hydrogenase active site models, which was carried out by Daresbourg’ group.\textsuperscript{25} They found the iron based oxidized species was thermodynamically favored in the diiron model complexes, which was further proven by the experimental and theoretical studies. Just based on this idea, we designed the catalytic system of these diiron complexes as an
oxygen transfer species. Actually, the Fe–(μ-O)–Fe species as the iron–oxygen reactive species in the catalytic oxidation was also studied by Can Li’s group.\(^{46}\) They identified the Fe–(μ-O)–Fe sites in Fe/ZSM-35 by in situ resonance Raman spectroscopy. Fe–(μ-O)–Fe species worked as an oxygen transfer species.

Under the optimized reaction conditions, performances of complexes 1–3 as catalysts were investigated and the results are summarized in Table 2, their active properties decline in the ranking of 2 > 3 > 1 under identical conditions. The complexes 1–3 showed better phenol yield and comparable selectivity to that we reported recently,\(^{11}\) namely [(μ-dmedt){Fe(CO)}\(_3\)] \(_2\) [dmedt = SCH(CH\(_3\))\(_2\)CH(CH\(_3\))] \(_2\) (yield 7.5%; selectivity 92.5%), [(μ-dmest){Fe(CO)}\(_3\)] \(_2\) [dmest = SCH(CH\(_3\))CH(CH\(_3\))S(O)] (yield 3.7%; selectivity 95.6%). These observations can probably be attributed to the different activities of 1, 2 and 3 correlations with the diverse Fe–Fe electron densities. The Fe–Fe bond of the complex 2 or complex 3 with the electron-donating PCy\(_3\) ligand or PPh\(_3\) ligand could be oxidized more easily by \(\text{H}_2\text{O}_2\) to form the iron–oxygen reactive species, whereas the less electron-donating complexes were found to be less reactive with this oxidant. These proved the electron density of the Fe–Fe bond of the complexes would affect the catalytic performances.\(^{43,45}\)

According to the inference of the experimental results and based on previously published theoretical results,\(^{20,27,46}\) a probable mechanism has been proposed for hydroxylation of benzene to phenol. The [FeFe]-hydrogenase model complex has the ability to catalyze the hydroxylation of different substrates to phenol, which revealed that hydroxylation reaction involves an electrophilic addition process and the proposed reaction mechanism for the hydroxylation of benzene to phenol is shown in Fig. 9. In the process of hydroxylation, the nucleophilic diiron center of [Fe\(^{2+}\)Fe\(^{2+}\)] complex tended to react with the oxidant of \(\text{H}_2\text{O}_2\) due to the high electron density of the diiron center.

In order to investigate the catalysts stability of complexes 1–3, further investigations were carried out through solution IR spectrum before and after the hydroxylation reaction in CH\(_3\)CN. As shown in Fig. 10, the CO-stretching band positions of complexes 1–3 didn’t change, only the intensities of IR spectrum of complexes 1–3 has slightly changes before and after the hydroxylation reaction in CH\(_3\)CN. This indicates the preservation of molecular structure of complexes 1–3 in the hydroxylation reaction in CH\(_3\)CN. Further research on the catalysts stability of complexes 1–3 is concluded by the TG analysis.

The TG analysis patterns of complexes 1–3 are shown in Fig. 11, these showed great heat resistance of complexes 1–3 below 100 °C. Among them complexes 2 and 3 have better heat resistance than complex 1. A rapid weight loss for complexes 2 and 3 originated from the decomposition ranging from 200 to

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**Table 2** Effects of various catalysts for the direct hydroxylation of benzene to phenol with \(\text{H}_2\text{O}_2\)^\(a\)

<table>
<thead>
<tr>
<th>Catalysts</th>
<th>Yield (%)</th>
<th>Selectivity (%)</th>
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<tbody>
<tr>
<td>1</td>
<td>12.6</td>
<td>85</td>
</tr>
<tr>
<td>2</td>
<td>24.6</td>
<td>92</td>
</tr>
<tr>
<td>3</td>
<td>18.2</td>
<td>87</td>
</tr>
</tbody>
</table>

* a Catalysts, 30 μmol; 60 °C; benzene, 1.12 mmol; CH\(_3\)CN, 2.0 mL; oxidants, 5.0 mmol; reaction time, 3 h. \(^{b}\) Selectivity: yield of phenol/benzene conversion.
were tested as catalysts in the study of the one-step oxidation of CH2S–CH2–CH3, CH2Cl2, toluene, and acetonitrile were deoxygenated and dried and distilled by standard methods. THF, toluene, and acetonitrile were purchased from Jiangtian Chemical and used without further purification. Methylamine oxide) (2) and methylamine hydroxide were directly applied a first time. The Fe(CO)5 was acquired from Jiangsu Tianyi (Shanghai) Chemical and used without further purification. The [FeFe]-hydrogenase model compounds were performed in an inert (N2) atmosphere. Examinations of GC (9890B) equipped by a FID sensor as well as a capillary pillar (OV-1701; 30 m × 0.25 mm × 0.25 μm) were conducted in a Heraeus CHN-Rapid, a fully automatic factor analyzer along with TCD inspection, type: TMT CHN, BESTELL-NR 2215001. By using a Rigaku MM-007 diffractometer along with a Saturn 724 CCD, the single-crystal X-ray diffraction data were gathered at 293 K or 113 K with a confocal monochromator in Mo-Kα radiation (λ = 0.71073 Å). Data gathering, decrease as well as absorption rectification was conducted in the CRYSTALClear program. Structural issues were addressed by direct approaches in SHELXL-97 program and also refined by SHELXL-97 on F2. The UV/visible spectra of the compounds were determined using the UV-vis diffuse reflectance spectra (UV-3600) at room temperature. TG was performed using a TA SDT-Q600 apparatus and conducted in dynamic temperature at 10 °C min−1 from room temperature to a temperature of 400 °C in nitrogen circumstance. The conductivities of the complexes 1–3 were on a Delsa™ Nano C Particle Analyzer (Beckman Coulter Commercial Enterprise Co., Ltd, USA).

Electrochemical examinations were performed with a CHI760B electrochemical workstation. The working electrode (diameter 3 mm) was polished in an α-alumina suspension, washed with CH3CN, and then sonicated with ion-free water for 20 min before usage. The counter electrode referred to a Pt wire. And the reference electrode referred to a non-aqueous Ag/AgCl (0.01 M AgNO3 and 0.1 M n-Bu4NPF6 in CH3CN) reference electrode. A liquid of 0.1 M n-Bu4NPF6 with CH3CN was applied as auxiliary electrolyte that was degassed with dry N2 for 10 min prior to examination. Ferrocene was applied as an internal criterion in the identical examining situation and each potential was referred to the Cp2Fe+/0 couple at 0 V.

Hydroxylation of benzene with H2O2 was conducted in a 25 mL flask that is equipped by a condenser as well as a magnetic agitator. In a characteristic reaction, the composite 1 (4 mg, 0.01 mmol) was mixed in CH3CN (2.0 mL). After heating to the required temperature, benzene (0.1 mL, 1.12 mmol) was added into the blend. A given mass of H2O2 was added at last for beginning the reaction, and in the meanwhile the blend was agitated for a couple of hours. The research was conducted in atmospheric pressure. Examinations of GC (9890B) equipped by a FID sensor as well as a capillary pillar (OV-1701; 30 m × 0.25 mm × 0.25 μm) were conducted to investigate the product composition. TPD analysis was performed, from 70 °C to 180 °C by a variation of 10 °C, and then from 180 °C to 280 °C by a variation of 30 °C.

Synthesis of μ-(SCH(CH2CH3)CH2S)–Fe3(CO)6 (1). The complex 1 was prepared according to the published procedures with a modified procedure.11,12 1,2-Butanedithiol (1.39 g, 11.4 mmol) was added to the solution of 1,2-butanedithiol (1.39 g,
11.4 mmol) in 30 mL toluene. The mixture was stirred at reflux for 72 h under an atmosphere of nitrogen. The solvent was evaporated under 110 °C. Then the residue was purified by a silica gel column and eluted with hexane. Single crystals of product 1 were obtained by slow diffusion of hexane into CH2Cl2 solution at -20 °C. Yield: 1.1 g (24%). C13H11O4S2Fe2 (analytical calc.): C 30.03, H 2.02, O 24.00; found: C 29.98, H 2.01, O 23.89.

IR (hexane, cm⁻¹): ν(CO) 2076(s), 2035(m), 2060(s), 1922(m), 1938(m). 1H NMR (400 MHz, CD2Cl2): δ (ppm) 1.31 (d, 3H, Ar-H), 0.81 (d, 3H, cis), 0.79 (d, 3H, trans), 0.78 (d, 3H, cis), 1.09 (m, 2H, trans), 1.46 (m, 2H, cis), 1.59 (m, 2H, cis), 1.61 (m, 2H, trans), 1.63 (m, 2H, cis), 1.69 (m, 2H, trans), 1.90 (m, 2H, cis), 1.99 (m, 2H, trans), 2.13 (m, 2H, cis), 2.26 (m, 2H, trans), 2.43 (m, 2H, cis), 2.59 (m, 2H, trans), 2.64 (m, 2H, cis), 2.85 (m, 2H, trans), 3.60 (m, 2H, cis), 3.62 (m, 2H, trans), 6.29 (m, 2H, cis), 6.23 (m, 2H, trans), 6.29 (m, 2H, cis), 6.23 (m, 2H, trans), 7.66 (m, 15H, Ar). Synthesis of μ-(SCH(CH2CH3)CH2S)-Fe2(CO)5PCy3 (2). The complex μ-(SCH(CH2CH3)CH2S)-Fe2(CO)5PCy3 (2) was prepared according to the published procedures with a modified procedure.

The mixing of complex 1 178.8 mg (0.44 mmol) and Me2NO 34.8 mg (0.46 mmol) in 5 mL THF was carried out at 0 °C for 30 min under N2, the color turned into black from red. Then PCy3 (30.8 mg, 0.11 mmol) was added in the THF solution. The mixture was stirred for 2 h at room temperature under N2 and then the mixture solution was evaporated to dryness in vacuum. The residue was purified by a silica gel column and eluted with hexane. The second red band eluted with ethyl acetate gave a red product of complex 2 (129.8 mg, 45%). Single crystals of product 2 were obtained by slow diffusion of hexane into CH2Cl2 solution at room temperature. Anal. calc. (%) for C27H23Fe2O5PS2: C 51.13, H 3.66, O 12.61. Found (%): C 51.09, H 2.01, O 23.89. NMR (CD2Cl2): δ CO = 208.82, δ dalkyl = 13.69, 30.18, 41.86, 56.44. 1H NMR (400 MHz, CD2Cl2): δ (ppm) 2.41, 2.39, 2.38, 2.29, 2.25, 2.24, 2.20, 2.18 (m, 2H, SCH2).

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Notes and references