

Elucidating Dramatic Ligand Effects on SET Processes: Iron Hydride versus Iron Borohydride Catalyzed Reductive Radical Cyclization of Unsaturated Organic Halides

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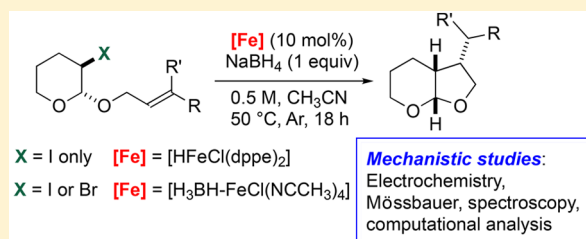
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S Supporting Information

ABSTRACT: An iron(II) borohydride complex ($[(\eta^1\text{-H}_3\text{BH})\text{-FeCl}(\text{NCCH}_3)_4]$) is employed as the precatalyst in iron-catalyzed radical cyclizations of unsaturated organic halides in the presence of NaBH_4 . Mechanistic investigations have established that the ligand bound to the metal center (acetonitrile versus ethylenebis(diphenylphosphine) (dppe)) plays a crucial role in the structure and reactivity of the active anionic iron(I) hydride ($[\text{HFeCl}(\text{dppe})_2]^-$) and borohydride ($[(\eta^1\text{-H}_3\text{BH})\text{FeCl}(\text{NCCH}_3)_4]^-$) with unsaturated haloacetals. This work provides new insights into iron(I) hydride and borohydride species and their potential implication in single-electron processes.



INTRODUCTION

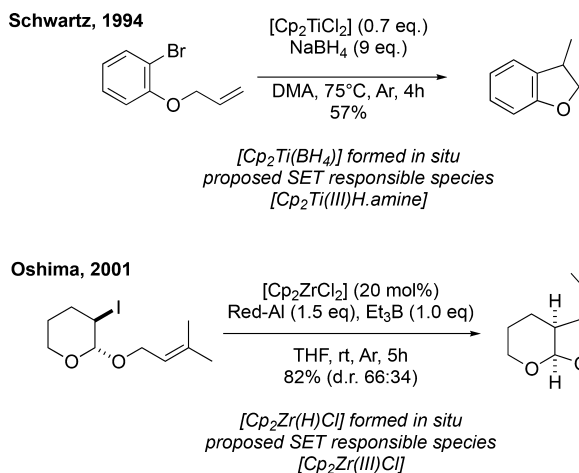
Transition metals have been widely employed in radical chemistry¹ since the pioneering works of Kochi and co-workers, which documented the importance of single-electron-transfer (SET) processes in such metal-catalyzed reactions.² Particularly, transition-metal complexes with low oxidation state readily undergo reduction of organic substrates to form a variety of carbon-centered radicals under the appropriate reaction conditions.³ Among these, it has been proposed that metal hydride complexes, which have proven their efficiency in catalysis and organic synthesis,⁴ generate low-valent transition-metal hydride species under reducing conditions, acting as both mediators and hydrogen donors for reductive radical reactions with organic halides. However, the in situ formation of the transition-metal hydride species, generated from the corresponding metal-halogenated complexes and a stoichiometric amount of hydride source (LiAlH_4 , NaBH_4 , NaBH_3CN , Red-Al), and the subsequent reaction mechanism have both remained largely unexplored. Such radical transformations have been performed with metal complexes of gallium,⁵ indium,⁶ zirconium,⁷ or titanium.⁸ While plausible mechanisms for these reactions have been proposed, tangible evidence is still anecdotal. For instance, Oshima and co-workers used commercially available Schwartz reagent (bis(cyclopentadienyl)zirconium(IV) chloride hydride) as a stoichiometric reducing reagent, or catalytic reaction conditions

of 20 mol % $[\text{Cp}_2\text{ZrCl}_2]$ in the presence of Red-Al, and Et_3B to promote reductive radical cyclization reactions of haloacetals. The authors proposed that the zirconium hydride (Zr-H) species was formed in situ from $[\text{Cp}_2\text{ZrCl}_2]$ in the presence of the strong reducing agent and acted as the precatalyst for the reaction. Hydrogen atom abstraction would give rise to the active zirconium(III) radical which undergoes SET with the substrate.⁷ In one of the earliest examples of this chemistry, Schwartz employed substoichiometric $[\text{Cp}_2\text{TiCl}_2]$ in the presence of a large excess of sodium borohydride for an aryl halide reductive cyclization reaction (Scheme 1). Mechanistic studies were carried out on the titanium borohydride complex $[\text{Cp}_2\text{Ti}(\text{BH}_4)]$ which was presumably formed under the reaction conditions. A titanium hydride amine complex ($\text{Cp}_2\text{Ti}^{\text{III}}\text{H}\cdot\text{amine}$) was suggested to be responsible for electron transfer. Evidence of this species was circumstantial but showed that the rate of cleavage of the Ti-HBH_3 bond varied in the presence of different amines such as pyridine and *N,N*-dimethylethylamine.⁸ These studies led to the conclusion that metal hydride and borohydride complexes can both act as precatalysts of the reduction reaction of organic halides

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Scheme 1. Transition-Metal-Mediated Radical Reactions and Postulated Intermediates Formed in Situ

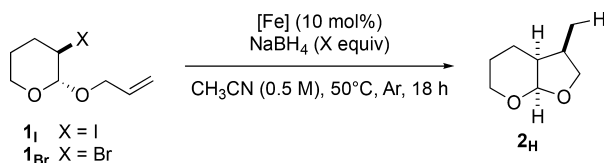


70 depending on the nature of the metal and potentially also the
 71 ligand.

72 Iron is reactive in redox chemistry: in particular, an array of
 73 reduction reactions has been carried out in the presence of
 74 metal reducing agents. Thanks principally to the pioneering
 75 work by Kharash and Kochi, iron catalysts have been exploited
 76 in a broad range of cross-coupling reactions of alkyl halides with
 77 Grignard reagents.⁹ Meunier and co-workers first reported that
 78 a stoichiometric amount of an iron–magnesium complex
 79 ($\text{Cp}(\text{DIPHOS})\text{FeMgBr}$) mediated the 5-exo radical cyclization
 80 reaction of 5-hexenyl bromide.¹⁰ Oshima and co-workers
 81 developed intramolecular radical cyclization reactions of
 82 haloacetals in the presence of 5 mol % of iron(II) dichloride
 83 and a stoichiometric Grignard reagent as the reducing agent.¹¹
 84 Other iron-mediated reduction reactions have also been
 85 developed,¹² and particularly, other mixed iron(II) or iron-
 86 (III)/hydride systems have also been applied to several
 87 different radical transformations.¹³

88 Over the last 50 years, various iron monohydride¹⁴ and
 89 borohydride¹⁵ complexes have been reported in the literature,
 90 but their use in radical synthesis remains scarce.

91 In an initial communication, we reported a novel 5-exo
 92 cyclization of 2-allyloxy-3-halo-tetrahydropyrans ($\mathbf{1}_I$, $\mathbf{1}_{Br}$) in the
 93 presence of iron(II) dichloride and sodium borohydride
 94 (Scheme 2).¹⁶

Scheme 2. Reaction of Haloacetals ($\mathbf{1}$) in the Presence of Iron(II) Complexes and NaBH_4 

95 As shown in Table 1, iron(II) dichloride (10 mol %)
 96 promoted the reaction of haloacetals ($\mathbf{1}_I$, $\mathbf{1}_{Br}$) in the presence of
 97 excess sodium borohydride (1.5 equiv) to give 73% and 78%
 98 yields of the bicyclic product ($\mathbf{2}_H$), respectively (Table 1,
 99 entries 1 and 2). Reaction of iodoacetal ($\mathbf{1}_I$) was also complete
 100 in the presence of $[\text{HFe}^I\text{Cl}(\text{dppe})_2]$ ¹⁷ ($\mathbf{3}$) and sodium
 101 borohydride (1 equiv) and provides support that an iron(II)
 102 hydride complex is a precatalyst for this reaction (Table 1, entry

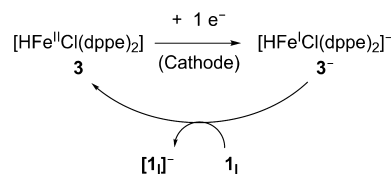
Table 1. Reactivity of $\mathbf{1}$ in the Presence of $[\text{Fe}]$ (10 mol %) and NaBH_4 ^a

entry	substrate	$[\text{Fe}]$	NaBH_4 (equiv)	yield (%)
1	$\mathbf{1}_I$	FeCl_2	1.50	73 ^b
2	$\mathbf{1}_{Br}$	FeCl_2	1.50	78 ^b
3	$\mathbf{1}_I$	$[\text{HFeCl}(\text{dppe})_2]$	1.00	70 ^c
4	$\mathbf{1}_{Br}$	$[\text{HFeCl}(\text{dppe})_2]$	1.00	0

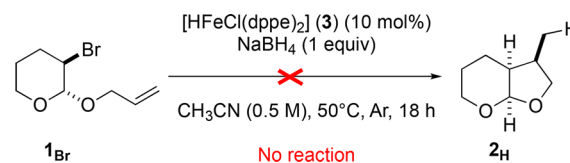
^aReaction conditions: $\mathbf{1}$ (1.0 mmol, 0.5 M in CH_3CN), $[\text{Fe}]$ (10 mol %), NaBH_4 (X equiv), 50 °C, 18 h. ^b_{dr} = 87:13. ^c_{dr} = 90:10.

3). Interestingly, no reaction was observed with the dihydride
 103 counterpart, $[\text{H}_2\text{Fe}(\text{dppe})_2]$. 104

Electrochemical studies of $[\text{HFe}^I\text{Cl}(\text{dppe})_2]$ ($\mathbf{3}$) identified
 105 that an anionic iron(I) hydride species, $[\text{HFe}^I\text{Cl}(\text{dppe})_2]^-$
 106 ($\mathbf{3}^-$), was electrogenerated or formed in the presence of
 107 NaBH_4 . This species ($\mathbf{3}^-$) was the active catalyst for the
 108 reaction, activating iodoacetal ($\mathbf{1}_I$) by electron transfer (SET)
 109 and regenerating the precatalyst in a catalytic cycle (Scheme 3).
 110 s3

Scheme 3. Electrochemical Generation of $[\text{HFe}^I\text{Cl}(\text{dppe})_2]^-$ ($\mathbf{3}^-$) and Electron Transfer (SET) to $\mathbf{1}_I$ 

As shown in Scheme 4, during followup investigations we
 111 s4 discovered that bromoacetal ($\mathbf{1}_{Br}$), in contrast to iodoacetal
 112

Scheme 4. No Reaction of Bromoacetal ($\mathbf{1}_{Br}$) in the Presence of $[\text{HFe}^I\text{Cl}(\text{dppe})_2]$ ($\mathbf{3}$) and NaBH_4 

($\mathbf{1}_I$), was completely unreactive in the presence of $[\text{HFeCl}(\text{dppe})_2]$ ($\mathbf{3}$) (Table 1, entry 4).^{16b} 113

114
 115 Herein, we report in detail the explanation for the lack of
 116 reactivity of $[\text{HFe}^I\text{Cl}(\text{dppe})_2]^-$ with bromoacetal ($\mathbf{1}_{Br}$) and our
 117 investigation to better understand the important factors that
 118 determine the reactivity of iron hydride complexes. We have
 119 elucidated the crucial role played by the ligand, thus giving new
 120 insights into the reactivity of iron hydride complexes to mediate
 121 radical cyclization reactions. To our knowledge, no detailed
 122 mechanistic investigations have been reported on iron-
 123 mediated radical reactions involving the in situ formation of
 124 an iron hydride versus borohydride species in acetonitrile. In
 125 this article we discuss our efforts to expand the understanding
 126 of iron-mediated reductive radical reactions. 126

RESULTS AND DISCUSSION 127

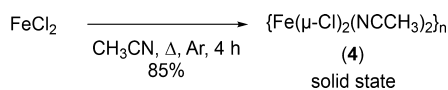
**Synthesis and Characterization of Iron(II) Bis-Chloride
 128 Complexes in Acetonitrile.** As previously mentioned, an
 129 iron(II) hydride complex ($[\text{HFeCl}(\text{dppe})_2]$ ($\mathbf{3}$)) serves as a
 130 precatalyst for reductive 5-exo radical cyclization reactions. This
 131 structurally defined complex can be easily synthesized from
 132

133 iron(II) dichloride upon reaction with sodium borohydride and
 134 dppe.¹⁷ This led us to consider the identity of the precatalyst
 135 formed in situ from the same source of iron(II) in the presence
 136 of sodium borohydride in acetonitrile in the absence of any
 137 phosphine ligand. Acetonitrile is usually considered as a pure σ -
 138 donor ligand and will readily fill vacant metal coordination sites
 139 to provide stability in solution.¹⁸ Indeed, literature reports
 140 supported the formation of iron–acetonitrile species. These
 141 complexes have even been isolated and characterized; however,
 142 their chemistry has not been greatly explored.¹⁹

143 Miller and co-workers have reported the solvation of iron(II)
 144 dichloride with acetonitrile to give an isolated species.²⁰ The
 145 crystal structure revealed the 1D chain polymer dichlorobis-
 146 (acetonitrile)iron(II) ($\{\text{Fe}(\mu\text{-Cl})_2(\text{NCCH}_3)_2\}_n$ (**4**)). This
 147 species was first described in 1964 by Hathaway and Holah
 148 and characterized by IR spectroscopy and titration.^{19a} A
 149 procedure to synthesize the dichlorotetrakis(acetonitrile)iron-
 150 (II) complex $\text{FeCl}_2(\text{NCCH}_3)_4$ is also available.²¹

151 The dichlorobis(acetonitrile)iron(II) complex ($\{\text{Fe}(\mu\text{-}$
 152 $\text{Cl})_2(\text{NCCH}_3)_2\}_n$ (**4**)) was synthesized following the procedure
 153 reported by Miller and co-workers, giving off-white crystals in
 154 85% yield (Scheme 5).²⁰ The X-ray structure matched the

Scheme 5. Synthesis of $\{\text{Fe}(\mu\text{-Cl})_2(\text{NCCH}_3)_2\}_n$ (**4**)



155 literature data. In this chain, an octahedral environment is
 156 accommodated by the iron centers, with bridging chloride
 157 anions and acetonitrile molecules in an apical position.

158 High-resolution ESI mass spectrometry revealed the
 159 presence of $[\text{FeCl}(\text{NCCH}_3)_2]^+$, a fragment resulting from the
 160 loss of one chloride ligand in the unit motif of the monomer
 161 (calcd 172.95634, found 172.95654).

162 The Mössbauer spectrum recorded at 80 K on a powder
 163 sample of $\{\text{Fe}(\mu\text{-Cl})_2(\text{NCCH}_3)_2\}_n$ (**4**) (natural abundance in
 164 ^{57}Fe) reveals three doublets. The major doublet (64%) has
 165 nuclear parameters that are characteristic of high-spin ($S = 2$)
 166 Fe^{II} ($\delta = 1.24 \text{ mm s}^{-1}$, $\Delta E_{\text{Q}} = 2.07 \text{ mm s}^{-1}$). The isomer shift
 167 value is in agreement with that previously determined for an
 168 analogous 1D chain.²² The second doublet ($\delta = 1.20 \text{ mm s}^{-1}$,
 169 $\Delta E_{\text{Q}} = 1.34 \text{ mm s}^{-1}$, contribution 26%) also exhibits nuclear
 170 parameters featuring a high-spin Fe^{II} ion. The two species differ
 171 in the quadrupole splitting values, suggesting that the two
 172 molecules of acetonitrile may be in either the trans or cis
 173 positions. The third doublet ($\delta = 0.34 \text{ mm s}^{-1}$, $\Delta E_{\text{Q}} = 1.12 \text{ mm}$
 174 s^{-1} , contribution 10%) presents nuclear parameters that are
 175 close to those of the tetrahedral ferric ion in $[\text{N}_6\text{Fe}^{\text{III}}-(\mu\text{-O})-$
 176 $\text{Fe}^{\text{III}}\text{Cl}_3]^+$,²³ suggesting a slight contamination by the diferric μ -
 177 oxo species $[\text{Cl}_3\text{Fe}-(\mu\text{-O})-\text{FeCl}_3]^{2+}$.

178 In order to investigate more closely the nature of the
 179 transient iron species generated in situ in the catalytic
 180 cyclization described in this work (Scheme 2), Mössbauer
 181 spectra of frozen solutions containing ^{57}Fe -enriched samples
 182 were recorded. First, $\{\text{Fe}(\mu\text{-Cl})_2(\text{S})_2\}_n$ ($\text{S} = \text{NCCH}_3$) was
 183 synthesized following Miller's procedure, at 0.05 M concen-
 184 tration to mimic the cyclization conditions (Scheme 5). An
 185 aliquot of this solution was frozen without preliminary
 186 purification, and the Mössbauer spectrum was recorded at 80
 187 K. Most of the spectrum (ca. 90% of the total area) could be
 188 simulated with two quadrupole doublets (I and II, Figure 1a)

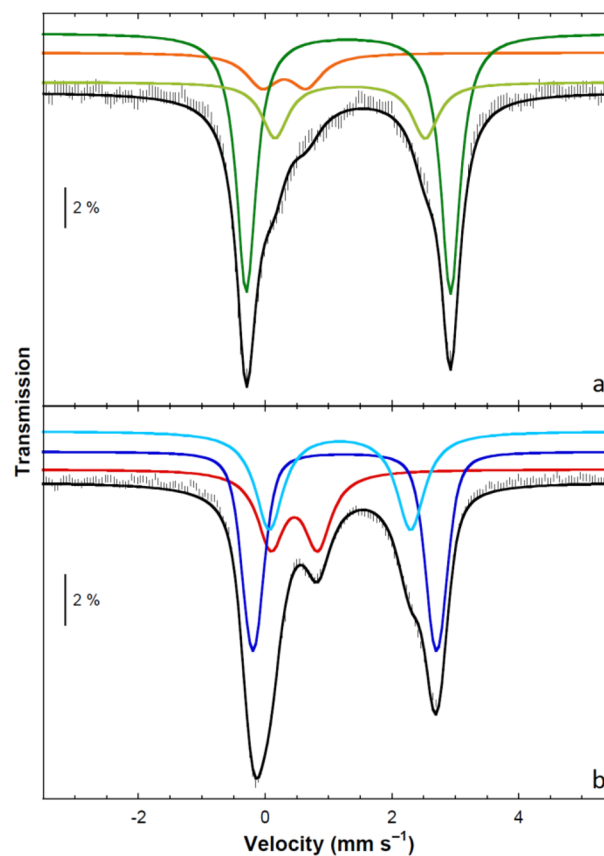


Figure 1. Zero-field Mössbauer spectra recorded at 80 K on acetonitrile solutions of $^{57}\text{FeCl}_2$ before (a) and after (b) the addition of 53 equiv of NaBH_4 . Experimental data are shown as hatched bars, and simulations are overlaid as solid black lines. Contributions are shown in color as upper traces. See text and Table 2 for parameters.

189 characteristic of two high-spin Fe^{II} species ($S = 2$) featuring the
 190 metal in an octahedral environment ($\delta_{\text{I}} = 1.31 \text{ mm s}^{-1}$, $\Delta E_{\text{Q,I}} = 190$
 191 3.22 mm s^{-1} (67%) and $\delta_{\text{II}} = 1.34 \text{ mm s}^{-1}$, $\Delta E_{\text{Q,II}} = 2.37 \text{ mm}$
 192 s^{-1} (19%)), in agreement with literature data.²⁴ Therefore,
 193 these results suggest that at the reaction concentration (0.05
 194 M) the iron ion adopts an octahedral environment in
 195 acetonitrile. In addition, the isomer shift is 0.1 mm s^{-1} higher
 196 than that in the solid state. The same increase has been
 197 previously measured at room temperature on powder samples
 198 between $\text{FeCl}_2 \cdot 2\text{H}_2\text{O}$ presenting the same polymeric 1D chain
 199 ($\delta = 1.12 \text{ mm s}^{-1}$) and $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$, where the ferrous ions are
 200 hexacoordinated by two chloride ions in trans positions and
 201 four water molecules ($\delta = 1.22 \text{ mm s}^{-1}$).²⁵ This strongly
 202 suggests that, in acetonitrile, octahedral ferrous complexes are
 203 formed with two chloride ions and four acetonitrile molecules
 204 coordinated: $[\text{FeCl}_2(\text{S})_4]$ (**4S**) ($\text{S} = \text{NCCH}_3$), which would
 205 putatively coexist as both the trans and cis isomers. The
 206 octahedral dication $[\text{Fe}(\text{S})_6]^{2+}$ was not detected under such
 207 conditions.²⁶

208 The cyclic voltammetry of $[\text{FeCl}_2(\text{S})_4]$ (**4S**; 4 mM) in
 209 acetonitrile (containing $^{\text{t}}\text{Bu}_4\text{NBF}_4$ (0.3 M) as the supporting
 210 electrolyte) exhibited several irreversible reduction peaks
 211 (Figure 2a). One major reduction peak was observed at E_{RI}^{P}
 212 $= -1.41 \text{ V}$ versus the saturated calomel electrode (SCE), along
 213 with two minor reduction peaks at $+0.48$ and -1.92 V . On the
 214 reverse scan, a major oxidation peak was observed at -0.048 V ,
 215 with minor oxidation peaks at -0.35 and $+0.61 \text{ V}$. This result
 can be explained by considering the equilibrium between ionic 216

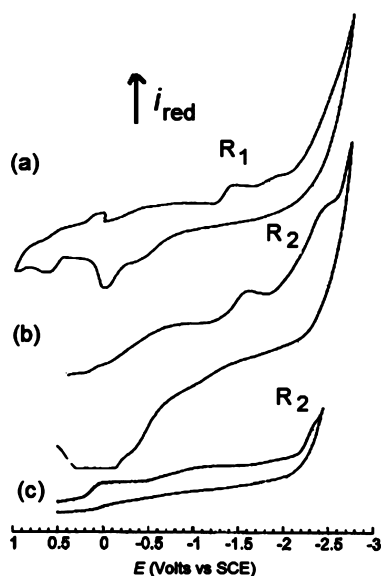
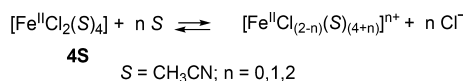


Figure 2. Cyclic voltammetry performed at a gold-disk electrode ($d = 1$ mm) at 22 °C in acetonitrile (S) containing ${}^n\text{Bu}_4\text{NBF}_4$ (0.3 M) as the supporting electrolyte: (a) Reduction of $[\text{Fe}^{\text{II}}\text{Cl}_2(\text{S})_4]$ (4S; 4 mM) at a scan rate of 0.5 V s^{-1} ; (b) $[\text{Fe}^{\text{II}}\text{Cl}_2(\text{S})_4]$ (4S; 4 mM) in the presence of NaBH_4 (4 equiv) at a scan rate of 5 V s^{-1} ; (c) reduction of isolated $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{S})_4]$ (5; 4 mM) at a scan rate of 0.5 V s^{-1} .

217 species formed from $[\text{FeCl}_2(\text{S})_4]$ (4S) in an acetonitrile
 218 solution (Scheme 6).^{27,28} The position of the equilibrium is
 219 expected to be sensitive to the concentration and the ionic
 220 strength induced by the supporting electrolyte.

Scheme 6. Postulated Iron(II) Complexes Formed in Acetonitrile

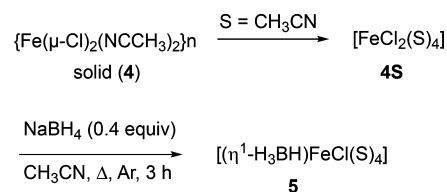


221 **Characterization of the Iron(II) Complex Formed in**
 222 **Situ from the Reaction of $[\text{FeCl}_2(\text{S})_4]$ (4S) with NaBH_4 in**
 223 **Acetonitrile.** The reaction of nonstabilized iron(II) and
 224 iron(III) salts such as FeBr_2 and FeBr_3 by NaBH_4 has been
 225 explored by several groups, and the structure of the resulting
 226 species was remarkably solvent dependent. For example, in the
 227 presence of water in diethyl ether, the reduction of FeBr_2 leads
 228 to the formation of noncrystalline metallic Fe^0 , whereas in dry
 229 diglyme borohydride-ligated iron(II) complexes such as
 230 $[\text{Fe}(\text{BH}_4)_2(\text{S})_n]$ (S = diglyme) are obtained, with no further
 231 reduction of the metal occurring at temperatures lower than 65
 232 °C.²⁹ However, the analogous bis-borohydride complex proved
 233 to be unstable in dry diethyl ether.³⁰ Several well-defined
 234 iron(II) borohydride complexes supported by various (PNP)
 235 pincer ligands have been reported and characterized by X-ray
 236 diffraction.³¹ In some cases, the iron center features η^1
 237 coordination of the borohydride ligand³¹ or bridging $\mu_2, \eta^1: \eta^1$ -
 238 H_2BH_2 moieties.^{31d} Some of these complexes exhibited good
 239 catalytic activities in ketone hydrogenation^{31c} and methanol
 240 dehydrogenation.³²

241 The reaction of $\text{FeCl}_2(\text{S})_4$ (4S) with NaBH_4 was monitored
 242 by cyclic voltammetry. The reduction peaks of $[\text{FeCl}_2(\text{S})_4]$
 243 (4S) (Figure 2a) decreased upon addition of sodium
 244 borohydride (up to 4 equiv). At the same time a new reduction
 245 peak was observed at $E_{\text{R}2}^{\text{p}} = -2.32$ V (Figure 2b). The

reduction peak R_2 was attributed to the formation of a new
 246 complex that has been independently synthesized and
 247 characterized to be the borohydride iron(II) complex $[(\text{BH}_4)\text{-}$
 248 $\text{FeCl}(\text{S})_4]$ (5) (Scheme 7) (vide infra for the characterization
 249 data).
 250

Scheme 7. Synthesis of $[(\eta^1\text{-H}_3\text{BH})\text{FeCl}(\text{NCCH}_3)_4]$ (5) by Reaction of NaBH_4 with $\{\text{Fe}(\mu\text{-Cl})_2(\text{NCCH}_3)_2\}_n$ in Acetonitrile



The isolated complex $[(\eta^1\text{-H}_3\text{BH})\text{FeCl}(\text{S})_4]$ (5) was
 251 prepared as shown in Scheme 7. Reaction of dichlorobis-
 252 (acetonitrile)iron(II) (4) with sodium borohydride (0.4 equiv)
 253 in acetonitrile gave an orange-brown solid. This new complex
 254 (5) was extremely air sensitive and thermally unstable, which
 255 made its complete characterization difficult. If appropriate
 256 precautions were not taken to maintain an inert environment, a
 257 noticeable and rapid color change of the complex occurred,
 258 from dull orange-brown to bright yellow. X-ray analysis
 259 revealed that the bright yellow complex was hexakis-
 260 (acetonitrile)iron(II) (μ -oxo)bis(trichloroiron(III)) ($[\text{Fe}-$
 261 $(\text{NCCH}_3)_6][\text{Cl}_3\text{Fe}(\mu\text{-O})\text{FeCl}_3]$).³³
 262

The isolated complex $[(\eta^1\text{-H}_3\text{BH})\text{FeCl}(\text{S})_4]$ (5) was
 263 characterized by cyclic voltammetry. It exhibited the same
 264 reduction peak at $E_{\text{R}2}^{\text{p}} = -2.37$ V (Figure 2c) as the complex
 265 generated in situ from 4S and NaBH_4 (Figure 2b). The
 266 reduction peak of 5 is irreversible at low scan rate ($\nu = 0.5$ V
 267 s^{-1} , Figure 2c), indicating that the electrogenerated complex 5⁻
 268 is not stable on the time scale of the cyclic voltammetry, in
 269 contrast to $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]^-$ (3⁻).¹⁶ However, at a higher
 270 scan rate ($\nu = 5$ V s^{-1} , shorter time scale), the reduction peak of
 271 5 at R_2 became partially reversible, suggesting the formation of
 272 $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^-$ (5⁻), which is partially stable
 273 on the time scale of this cyclic voltammogram (Figure 2b).
 274

Monitoring the addition of two 1 equiv amounts in
 275 succession of NaBH_4 to $[\text{FeCl}_2(\text{S})_4]$ (4S) in CD_3CN by
 276 ${}^{11}\text{B}\{\text{H}\}$ NMR in the presence of ${}^n\text{Bu}_4\text{NBPh}_4$ as an internal
 277 reference revealed that no boron-containing byproducts (such
 278 as $\text{H}_3\text{B}\cdot\text{NCCH}_3$ adducts) were formed. NaBH_4 itself could not
 279 be detected in the $+500/-500$ ppm range. However, free
 280 NaBH_4 was observed as a singlet at -41 ppm for higher NaBH_4
 281 concentrations (>3 equiv/mol of iron). The reaction of NaBH_4
 282 with $[\text{FeCl}_2(\text{S})_4]$ (4S) in acetonitrile in the absence of any
 283 additional coligand is therefore in stark contrast with the
 284 formation of the diphosphine complex $[\text{HFeCl}(\text{dppe})_2]$ (3)
 285 from NaBH_4 and $[\text{FeCl}_2(\text{dppe})_2]$, which can be synthesized
 286 and isolated. In the latter case, a stable terminal hydride is
 287 indeed obtained, and hydroborane BH_3 is released in the bulk,
 288 as confirmed by the characterization of its dppe-ligated adduct
 289 by ${}^{31}\text{P}$ NMR spectroscopy.^{16b} This strongly suggests that no
 290 free diamagnetic boron byproduct is released in the bulk during
 291 the reaction of NaBH_4 with $[\text{FeCl}_2(\text{S})_4]$ (4S) in acetonitrile
 292 and that at least one borohydride anion could act as a ligand to
 293 a paramagnetic NMR-silent species. This hypothesis is
 294 consistent with the work of Klabunde mentioned earlier, who
 295 reported that FeBr_2 reacted with an excess of NaBH_4 in diglyme
 296

Table 2. Parameters Determined from the Simulations of the Mössbauer Spectra Shown in Figure 1^a

addition of NaBH ₄	dite	color	δ (mm s ⁻¹)	ΔE_Q (mm s ⁻¹)	Γ (mm s ⁻¹) ^b	rel area (%)
no ^c	I	dark green	1.31	3.22	0.37	67
	II	light green	1.34	2.37	0.50	19
	III	orange	0.30	0.70	0.60	13
yes ^d	IV	dark blue	1.25	2.90	-0.35	43
	V	light blue	1.18	2.23	0.53	34
	VI	red	0.46	0.74	0.51	24

^aUncertainties are ± 0.03 mm s⁻¹ on the isomer shift δ , ± 0.05 mm s⁻¹ on the quadrupole splitting ΔE_Q , and $\pm 2\%$ on the relative area. ^bThe Lorentzian line presents a full width at half-maximum (fwhm) denoted as Γ . A negative value indicates a Voigt line shape, which is the convolution of a 0.19 mm s⁻¹ fwhm Lorentzian line with a Gaussian whose fwhm is the absolute value of Γ . ^cMössbauer spectrum shown in Figure 1a. ^dMössbauer spectrum shown in Figure 1b.

to give diglyme-stabilized bis-borohydride complexes such as [Fe(BH₄)₂(S)_n] (S = diglyme).²⁹ Moreover, high-resolution ESI mass spectrometry revealed that, after ionization, [(η^1 -H₃BH)FeCl(S)₄] (5) exhibited the same fragment ion [FeCl(S)₂]⁺ as 4, resulting from the loss of a borohydride fragment and two acetonitrile ligands (calcd 172.95634, found 172.95650).³⁴ Taken together, these results suggest that, in the presence of a moderate excess of NaBH₄ in acetonitrile solution, [FeCl₂(S)₄] (4S) reacts to form an iron(II) borohydride complex such as [(η^1 -H₃BH)FeCl(S)₄] (5) by substitution of a single chloride ligand.

The IR spectrum of [(η^1 -H₃BH)FeCl(S)_n] (5, solid phase, Nujol) confirmed the η^1 coordination mode of the borohydride ligand. Well-defined stretches characteristic of a transition-metal-ligated η^1 -borohydride anion were observed at 2429, 2390, and 2352 cm⁻¹ (ν_{BH_3}) and 2042 cm⁻¹ ($\nu_{\text{Fe-H-B}}$).^{31a} The BH₃ deformation band could not be assigned precisely due to the presence of other signals in the 900–1100 cm⁻¹ area (see the Supporting Information). No signal characteristic of a terminal Fe–H bond vibration in the 1800–2000 cm⁻¹ range (usual range for the M–H bond vibration) could be detected, thus excluding the formation of a terminal iron hydride. These data are in good agreement with previous reports for (η^1 -borohydrido)iron(II) species by Koehne^{31d} and Field.^{31b} However, it should be noted that almost all of the reported iron(II) borohydride or terminal hydride complexes accommodate a low-spin configuration.

More structural insights into the nature of this borohydride species under solvation conditions were gained by Mössbauer spectroscopy on frozen solutions, starting from ⁵⁷Fe-enriched complexes. Figure 1b shows the zero-field Mössbauer spectrum recorded at 80 K on a 0.05 M acetonitrile solution of ⁵⁷FeCl₂ after the addition of excess NaBH₄. The addition of 18 or 53 equiv of NaBH₄ leads to identical spectra. Accordingly, only the spectrum with 53 equiv is shown. The lines are quite broad; however, this spectrum significantly differs from that recorded for [FeCl₂(S)₄] (4S) (Figure 1a). A reduced separation of the two more intense lines, now at 0 and 2.8 mm s⁻¹ versus -0.3 and 3.0 mm s⁻¹ for 4S, is observed. In addition, a new line is detected at 0.7 mm s⁻¹. The slight shift of the high-velocity pattern indicates that the starting Fe^{II} species has reacted with NaBH₄ and that new high-spin Fe^{II} species are generated.

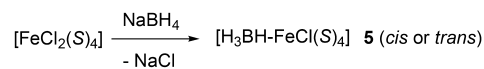
The spectrum shown in Figure 1b was then simulated. In order to reproduce the asymmetric profile of the high-velocity line, three doublets must be considered. The simulation is shown in Figure 1b, and the parameters are given in Table 2. Two doublets (IV and V) present isomeric shift values lying between 1.1 and 1.3 mm s⁻¹ ($\delta_{\text{IV}} = 1.25$ mm s⁻¹, $\Delta E_{\text{Q,IV}} = 2.90$

mm s⁻¹ (43%) and $\delta_{\text{V}} = 1.18$ mm s⁻¹, $\Delta E_{\text{Q,V}} = 2.23$ mm s⁻¹ (34%)), which correspond to high-spin ferrous ions accommodating an octahedral environment, these sites accounting for 77% of the total iron content. The isomeric shift value associated with the third doublet (VI) is much lower ($\delta_{\text{VI}} = 0.46$ mm s⁻¹), which renders this doublet more difficult to assign. Because borohydride was added in great excess, a reasonable hypothesis is that it is associated with a reduced form of iron. This is in agreement with the cyclic voltammogram recorded after addition of 50 equiv of NaBH₄ (see the Supporting Information). A broad oxidation peak is detected at $E^{\text{p}}_{\text{O1}} = +0.2$ V vs SCE, indicating the chemical reduction of Fe^{II} to metallic Fe⁰. The same peak is also observed on the reverse scan in Figure 2a and corresponds to the oxidation of the electrogenerated reduced species adsorbed at the electrode and formed upon electroreduction of [FeCl₂(S)₄] (4S).

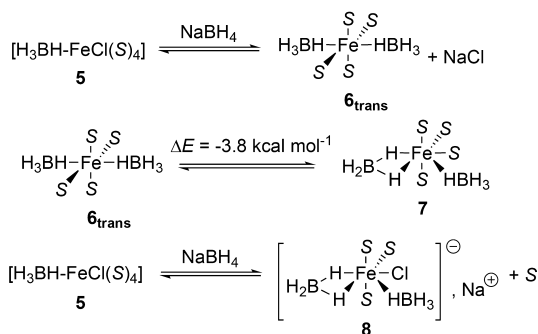
DFT calculations were performed to gain further insights into the high-spin ferrous species 5 generated by the reaction of NaBH₄ with [FeCl₂(S)₄] (4S). Alternative structures were also computed for comparison (Scheme 8). Because the isomer shifts of the doublets labeled IV and V (Table 2) are significantly lower than those of the starting species (doublets I and II, Table 2), it is reasonable to postulate the substitution of one ligand by a stronger σ donor: namely, borohydride or hydride. Calculations were performed on several model complexes (see the Supporting Information). The negatively

Scheme 8. Distribution of (a) Mono- and (b) Bis-Borohydrido Fe^{II} Species Obtained by Reaction of Excess NaBH₄ with [FeCl₂(S)₄] (4S) in Acetonitrile

a) formation of mono-borohydrides:



b) formation of bis-borohydrides:



371 charged ligands (Cl^- , BH_4^- , H^-) may be in the cis or trans
372 positions, and the borohydrido ligand may coordinate in the η^1
373 or κ^2 mode. Since NaBH_4 is present in great excess, the
374 coordination of two borohydride ligands was also considered. A
375 selection of calculated isomer shift values is given in Table 3.

Table 3. Calculated Isomer Shifts (δ) for Several Relevant High-Spin Fe^{II} Borohydrides^a

label	complex	$\rho_0(\text{calcd})$ (e a_0^{-3})	$\delta(\text{calcd})$ (mm s^{-1})
5_{trans}	$\text{trans}-[(\eta^1\text{-H}_3\text{BH})\text{FeCl}(\text{NCCH}_3)_4]$	11816.89	1.20
5_{cis}	$\text{cis}-[(\eta^1\text{-H}_3\text{BH})\text{FeCl}(\text{NCCH}_3)_4]$	11816.85	1.21
6_{trans}	$\text{trans}-[(\eta^1\text{-H}_3\text{BH})_2\text{Fe}(\text{NCCH}_3)_4]$	11817.08	1.13
7	$[(\kappa^2(\text{H},\text{H})\text{-H}_2\text{BH}_2)(\eta^1\text{-H}_3\text{BH})\text{Fe}(\text{NCCH}_3)_4]$	11816.76	1.27
8	$[(\kappa^2(\text{H},\text{H})\text{-H}_2\text{BH}_2)(\eta^1\text{-H}_3\text{BH})\text{FeCl}(\text{NCCH}_3)_3]^-$	11816.69	1.27

^aThe calculated isomer shift values are deduced from the correlation curve established in the Supporting Information.

376 Solvated complex **5** was tentatively assigned as the
377 borohydrido complex $[(\eta^1\text{-H}_3\text{BH})\text{FeCl}(\text{NCCH}_3)_4]$ in the
378 solid phase (Scheme 7). Simulations of the Mössbauer
379 parameters of the trans and cis isomers of $[(\eta^1\text{-H}_3\text{BH})\text{FeCl}$
380 $(\text{NCCH}_3)_4]$ (**5**) were performed (Scheme 8a). Complexes
381 5_{trans} and 5_{cis} were found to be almost isoenergetic ($\Delta E \approx 1$
382 kcal mol^{-1}), with similar nuclear parameters ($\delta = 1.20 \text{ mm s}^{-1}$
383 (5_{trans}); $\delta = 1.21 \text{ mm s}^{-1}$ (5_{cis}); Table 3). It was found that the
384 parent hydrides $[\text{HFeCl}(\text{NCCH}_3)_4]$ led to smaller values of the
385 isomer shift (ca. 1.1 mm s^{-1}), in agreement with H^- having
386 stronger σ donating properties in comparison BH_4^- (see the
387 Supporting Information).

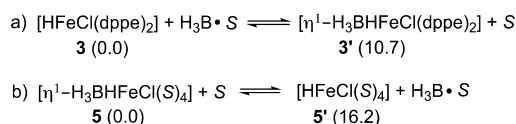
388 Therefore, computed parameters of monoborohydrido
389 complexes 5_{trans} and 5_{cis} (Table 3, and Scheme 8a) reproduce
390 the simulation of doublet V in the experimental spectrum well
391 (Figure 1b and Table 2). Due to the high borohydride to iron
392 ratio (18 or 53 equiv) in the Mössbauer experiments, bis-
393 borohydrido complexes can also be obtained. Complexes $[(\eta^1\text{-}$
394 $\text{H}_3\text{BH})_2\text{Fe}(\text{NCCH}_3)_4]$ (**6**) can be formed by substitution of the
395 remaining chloride anion of **5** by a second borohydride. DFT
396 calculations show that **6** evolve(s) toward the more stable
397 complex **7** $[(\kappa^2(\text{H},\text{H})\text{-H}_2\text{BH}_2)(\eta^1\text{-H}_3\text{BH})\text{Fe}(\text{NCCH}_3)_4]$ ($\Delta E =$
398 $-3.8 \text{ kcal mol}^{-1}$, Scheme 8b). In complex **7**, the steric pressure
399 in the meridional plane containing five ligands leads to longer
400 metal–ligand bonds and therefore to an electronic density at
401 the ^{57}Fe nucleus smaller than that for the octahedral species
402 5_{trans} and 5_{cis} . Consequently, the corresponding isomer shift is
403 higher ($\delta = 1.27 \text{ mm s}^{-1}$, **7**; Table 3). Analogously, $[(\kappa^2(\text{H},\text{H})\text{-}$
404 $\text{H}_2\text{BH}_2)(\eta^1\text{-H}_3\text{BH})\text{FeCl}(\text{NCCH}_3)_3]^-$ (**8**), formed from **5** via
405 substitution of an acetonitrile molecule by BH_4^- , was computed
406 (Scheme 8b). Complex **8** gave a similarly high computed
407 isomer shift ($\delta = 1.27 \text{ mm s}^{-1}$, **8**; Table 3) to **7**. Therefore, the
408 doublet IV in the spectrum (Figure 1b and Table 2) can be
409 explained by the formation of sterically constrained bis-
410 borohydrido species such as **7** and **8**, which exhibit isomer
411 shifts higher than those of **5**.

412 Computation of mono and bis iron(II) borohydride nuclear
413 parameters can explain the general trend observed in the
414 Mössbauer experimental spectrum.³⁵ Moreover, the formation
415 of bis-borohydrido species can explain the fast formation of
416 reduced oxidation states leading to the observation of the
417 doublet VI in Figure 1b. However, due to the broadness of the
418 experimental signals, it can be anticipated that several

structurally similar isomers (or other species with similar
nuclear parameters) coexist in solution.

Iron(II) Hydride versus Iron(II) Borohydride Complexes. As stated in the previous section, there is a major
difference in the electronic properties of the $\text{Fe}^{\text{II}}\text{-H}$ bond in
the well-defined diphosphine-ligated complex $[\text{HFeCl}(\text{dppe})_2]$
(**3**) and in borohydride acetonitrile-ligated complexes such as
 $[(\eta^1\text{-H}_3\text{BH})\text{FeCl}(\text{NCCH}_3)_4]$ (**5**). The former is a low-spin Fe^{II}
terminal hydride, whereas the latter is a high-spin Fe^{II}
borohydride. This difference is well reflected by DFT
calculations. The computation of the enthalpy of formation
of the dppe-stabilized borohydride $\text{trans}-[(\eta^1\text{-H}_3\text{BH})\text{FeCl}$
 $(\text{dppe})_2]$ (**3'**_{trans}) from $\text{trans}-[\text{HFeCl}(\text{dppe})_2]$ (**3**_{trans}) and the
 $\text{H}_3\text{B}\cdot\text{NCCH}_3$ adduct clearly shows that the terminal hydride **3**
is more stable than the corresponding borohydride **3'**_{trans} (ΔE
 $= 10.7 \text{ kcal mol}^{-1}$; Scheme 9a). This equilibrium is the opposite

Scheme 9. DFT-Computed Equilibria between Hydrides and Borohydrides for Complexes (a) **3 and **3'** and (b) **5** and **5'**^a**



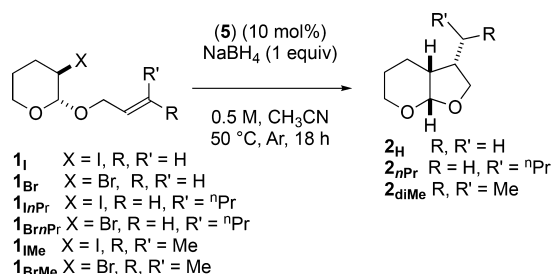
^aAll complexes were computed as trans isomers. Values in parentheses stand for the energies in kcal mol^{-1} .

for the acetonitrile-ligated system, which is shown to be far
more stabilized in the borohydride form, with $\text{trans}-[(\eta^1\text{-}$
 $\text{H}_3\text{BH})\text{FeCl}(\text{NCCH}_3)_4]$ (5_{trans}) stabilized by ca. $\Delta E = 16.2 \text{ kcal}$
 mol^{-1} with respect to the corresponding hydride $\text{trans}-$
 $[\text{HFeCl}(\text{NCCH}_3)_4]$ ($5'_{\text{trans}}$; Scheme 9b).

An analysis of the electronic population of $\text{trans}-[\text{HFeCl}$
 $(\text{dppe})_2]$ (**3**_{trans}) by natural bond orbital (NBO) theory shows
that there is a strong delocalization of the $\sigma(\text{Fe-H})$ molecular
orbital into the antibonding $\sigma^*(\text{Fe-P})$ MOs. The analysis is
drastically different for the acetonitrile-ligated complex $\text{trans}-$
 $[\text{HFeCl}(\text{NCCH}_3)_4]$ ($5'_{\text{trans}}$), where no significant delocaliza-
tion of the $\sigma(\text{Fe-H})$ molecular orbital into the acetonitrile
ligands is observed. Consistent with this observation, all of the
spin density is located on the iron center in $\text{trans}-[\text{HFeCl}$
 $(\text{NCCH}_3)_4]$ ($5'_{\text{trans}}$) (computed Mulliken spin density on iron
4.00, which is the expected theoretical value for a high-spin Fe^{II}
ion), and the iron-ligated acetonitrile ligands remain mostly
neutral (average computed NBO charge for each acetonitrile
ligand $q = +0.04 \text{ lel}$), showing that no charge transfer occurs
from the $\text{Fe}^{\text{II}}\text{-H}$ moiety.

Chlorotetrakis(acetonitrile)iron(II) Borohydride Complex **5 as Precatalyst for Reactions of Unsaturated Haloacetals.** $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ (**5**) was tested as
a suitable precatalyst for reaction with haloacetals (**1**), as shown
in Scheme 10.

Indeed, $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ (**5**) catalyzed the
reaction of both iodoacetal (**1_I**) and bromoacetal (**1_{Br}**)
substrates in the presence of 1.0 equiv of NaBH_4 (Table 4),
in contrast to $[\text{HFeCl}(\text{dppe})_2]$ (**3**), which only mediated the
reaction of iodoacetal (**1_I**). Complete conversion and good
yields were obtained for iodoacetals **1_I** and **1_{InPr}** (entries 1 and
3, Table 4). In the case of **1_{Me}** a moderate yield (46%) of the
bicyclic product (**2_{diMe}**) was obtained along with 19% recovery
of the starting material (entry 5). The unsubstituted
bromoacetal (**1_{Br}**) was completely consumed in the reaction,
and 57% of bicyclic product (**2_H**) was obtained (entry 2, Table 4)

Scheme 10. Reaction of Haloacetals **1 in the Presence of $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ (**5**) and NaBH_4**

Table 4. Reactivity of Haloacetals **1 in the Presence of $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ (**5**) (10 mol %) and Sodium Borohydride (1 equiv)^a**

entry	substrate	yield of 2 (%) (recovered 1 (%) [dr]
1	1 _I	77 (0) [88:12]
2	1 _{Br}	57 (0) [86:14]
3	1 _{ImPr}	70 (0) [82:18]
4	1 _{BrnPr}	38 (38) [81:19]
5	1 _{Ime}	46 (19) [66:34]
6	1 _{BrMe}	23 (55) [65:35]

^aReaction conditions: **1** (1.0 mmol, 0.5 M in CH_3CN), $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ (**5**) (10 mol %), NaBH_4 (1 equiv), 50 °C, 18 h.

471 **4**). Substituted bromoacetals (**1**_{BrnPr} and **1**_{BrMe}) were reactive,
472 albeit with incomplete conversion, giving the bicyclic products
473 in 38% yield (**2**_{nPr}, entry 4, Table 4) and 23% yield (**2**_{diMe}, entry
474 **6**), respectively.

Mechanistic Studies: Exploring Reactivity Differences.

476 This raised an important question about the ability of the
477 iron(II) monohydride or borohydride complexes, ligated by
478 either phosphorus ligands (**3**) or acetonitrile ligands (**5**), to act
479 as precatalysts for the reaction of bromoacetal (**1**_{Br}).

480 Cyclic voltammetry had been used to evidence the catalytic
481 turnover of the active iron(I) anionic hydride catalyst
482 $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]^-$ (**3**⁻) in the presence of iodoacetal (**1**_I).¹⁶
483 Thus, the catalytic activity of $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^-$
484 (**5**⁻) in the presence of haloacetals (**1**_I and **1**_{Br}) was also tested
485 by electrochemistry. The reduction potential of iodoacetal (**1**_I)

in acetonitrile was measured at $E_{\text{red}}^{\text{P}} = -2.5$ V vs. SCE at a scan 486
rate of 0.5 V s⁻¹. In contrast, the reduction potential of 487
bromoacetal (**1**_{Br}) could not be measured, as the substrate was 488
not reduced before the solvent ($E_{\text{red}}^{\text{P}} = -2.7$ V vs. SCE). As 489
shown in Figure 3a, in the presence of iodoacetal (**1**_I), the 490
reduction peak R₂ of $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^-$ (**5**) 491
increases with each aliquot of substrate (0–10 equiv, Figure 492
3a). Therefore, the anionic species $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^-$ 493
(**5**⁻) (generated by electrochemical reduction 494
of $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ (**5**)) activates iodoacetal (**1**_I) 495
by electron transfer. This in turn regenerates $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ 496
(**5**), and a catalytic current was observed. 497
This reveals that the interaction of $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^-$ 498
(**5**⁻) with **1**_I is faster than its decomposition on 499
the time scale of cyclic voltammetry. 500

The procedure was repeated in the presence of bromoacetal 501
(**1**_{Br}). A noticeable catalytic effect in the presence of **1**_{Br} was 502
observed, as shown in Figure 3b (0–12 equiv). This result 503
provides strong evidence that the acetonitrile ligated $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ (**5**) 504
furnishes the catalytic species $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^-$ (**5**⁻) able to activate both **1**_I 506
and **1**_{Br}. The turnover was lower for **1**_{Br} than for **1**_I (compare 507
the catalytic currents in parts b and c of Figures 3, respectively). 508

No catalytic current at R₃ was observed when $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]$ (**3**) 509
was reduced in the presence of increasing amount 510
of **1**_{Br} (1–6 equiv, Figure 3c), in contrast to the case for $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ (**5**) 511
(Figure 3b). Therefore, anionic $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^-$ (**5**⁻) 512
promotes the reduction of **1**_{Br} in contrast to $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]^-$ (**3**⁻), in agreement with 514
the experimental observations. This shows that the ligand 515
structure (hydride versus borohydride) strongly affects the 516
reactivity of the anionic iron(I) complex with **1**_{Br}. 517

These results have interesting mechanistic implications to 518
explain the higher reactivity of bromoacetal (**1**_{Br}) in the 519
presence of $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ (**5**) versus $[\text{HFe}^{\text{II}}\text{Cl}(\text{dppe})_2]$ (**3**). 520
521

The rate of electron transfer of $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^-$ (**5**⁻) or $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]^-$ (**3**⁻) to haloacetals 522
is controlled by the potential gap (ΔE) between the reduction 524
potential of the haloacetals and the oxidation potential of the 525
anionic iron(I) (boro)hydride complexes. This potential gap is 526

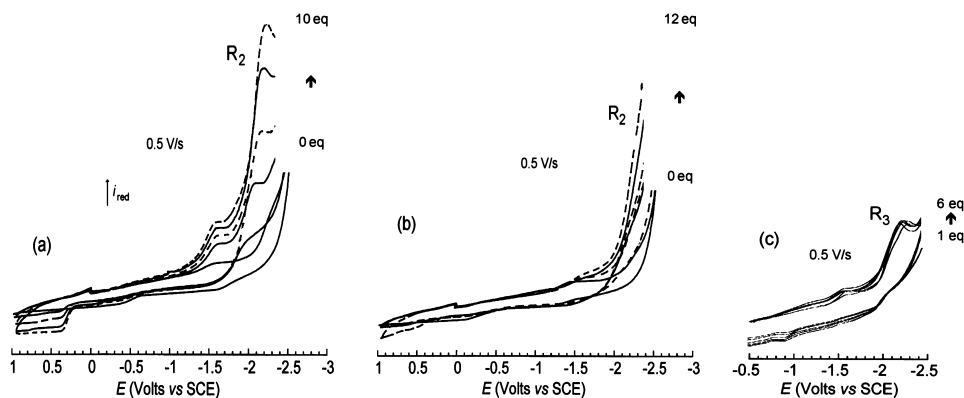


Figure 3. Cyclic voltammetry performed at a gold-disk electrode ($d = 1$ mm) at 22 °C in acetonitrile containing $^n\text{Bu}_4\text{NBF}_4$ (0.3 M) as the supporting electrolyte at a scan rate of 0.5 V s⁻¹: (a) reduction of $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ (4 mM) in the presence of increasing amounts of **1**_I (0 and 4 equiv, solid lines; 6 equiv, dashed line; 8 equiv, solid line; 10 equiv, dashed line); (b) reduction of $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]$ (4 mM) in the presence of increasing amounts of **1**_{Br} (0 and 8 equiv, solid lines; 4 and 12 equiv, dashed lines); (c) reduction of $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]$ (4 mM) in the presence of increasing amounts of **1**_{Br} (1–6 equiv, solid lines).

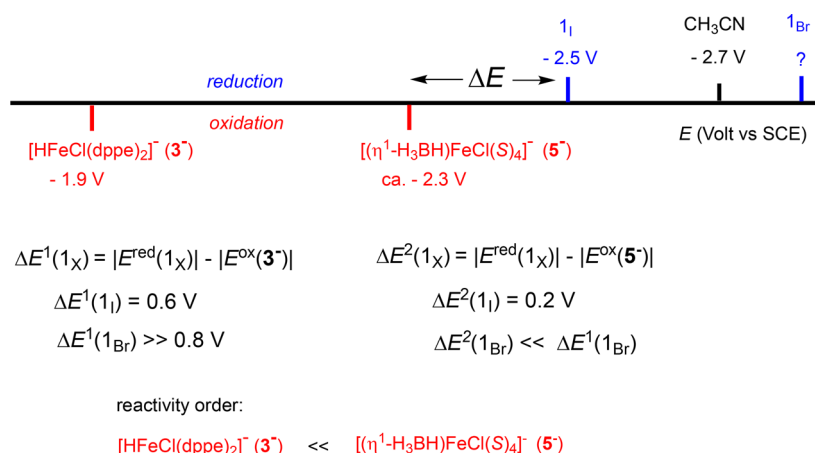


Figure 4. Cyclic voltammetry data used to explain the difference in reactivity of anionic complexes 3^- and 5^- with iodo- and bromoacetals (1_X) ($X = \text{I}, \text{Br}$).

527 crucial to the outcome of the reaction: indeed, the lower the
 528 potential gap, the faster the reaction.³⁶

529 As stated earlier, 1_I is more easily reduced than 1_{Br} at -2.5 V
 530 and more negative than -2.7 V , respectively (Figure 4).
 531 $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]^- (3^-)$ is less easily oxidized (-1.9 V)^{16a} than
 532 $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^- (5^-)$ (estimated to be ca. -2.3
 533 V) (Figure 4).

534 $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]^- (3^-)$ reacts with 1_I because of the low
 535 potential gap, $\Delta E^1(1_I) = 0.6 \text{ V}$, which permits a fast reaction
 536 (Figure 4). The reaction with 1_I is even faster with $[(\eta^1\text{-}$
 537 $\text{H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{S})_4]^- (5^-)$ because the potential gap $\Delta E^2(1_I)$ is
 538 even lower (0.2 V) (Figure 4).

539 The potential gap $\Delta E^1(1_{\text{Br}})$ between $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]^- (3^-)$
 540 and 1_{Br} is larger than 0.8 V . This explains why the catalytic
 541 reaction does not proceed when $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]^- (3)$ is used
 542 as the precatalyst. The potential gap $\Delta E^2(1_{\text{Br}})$ between 5^- and
 543 1_{Br} is smaller than $\Delta E^1(1_{\text{Br}})$ ($\Delta E^2(1_{\text{Br}}) < \Delta E^1(1_{\text{Br}})$, Figure 4),
 544 and therefore $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{S})_4]^- (5^-)$ can activate 1_{Br} and
 545 the catalytic reaction proceeds. In conclusion, the acetonitrile
 546 ligand renders the corresponding catalytically active species
 547 (5^-) far more active than the phosphine ligand (3^-). This
 548 clearly demonstrates the importance of the ligand in
 549 determining the outcome of the catalytic reaction.

550 **Mechanism.** The results of the catalytic reactions and
 551 subsequent mechanistic investigation performed on $[(\eta^1\text{-}$
 552 $\text{H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]^- (5)$ provides evidence for the active
 553 catalyst, the anionic iron(I) borohydride species $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{S})_4]^- (5^-)$ (Figure 5). The latter is formed from
 554 the iron(II) borohydride precatalyst 5 , either by electro-
 555 chemical means or by NaBH_4 . Activation of RX (1_X , $X = \text{I}, \text{Br}$)
 556 by 5^- via electron transfer generates the radical R^\bullet involved in
 557 the cyclization process. Hydrogen atom transfer to form the
 558 final cyclized product $\text{R}'\text{H}$ occurs either directly from BH_4^- or
 559 from $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^- (5)$ (vide infra for the
 560 computational studies that model the final step).³⁷

562 CONCLUSIONS

563 In conclusion it is established that the structure and reactivity of
 564 the active iron catalyst in the cyclization of unsaturated iodo-
 565 and bromoacetals is directly related to the ligand environment
 566 (acetonitrile versus dppe). The iron(II) borohydride $[(\eta^1\text{-}$
 567 $\text{H}_3\text{BH})\text{Fe}^{\text{II}}\text{Cl}(\text{NCCH}_3)_4]^- (5)$ has been synthesized, and its
 568 catalytic and redox properties have been compared to those of
 569 the reported iron(II) hydride $[\text{HFe}^{\text{II}}\text{Cl}(\text{dppe})_2]^- (3)$.¹⁶ As

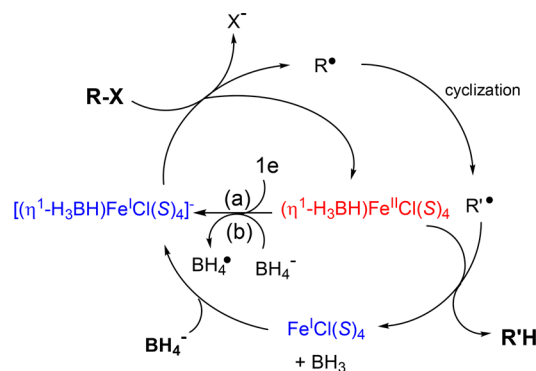


Figure 5. Mechanistic role of the active anionic iron(I) borohydride complex (blue) generated either (a) by electrochemical reduction or (b) by reduction by NaBH_4 of the iron(II) borohydride complex (red).

570 observed by cyclic voltammetry, the anionic iron(I) complex
 571 $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^- (5^-)$ is able to activate both
 572 iodo- and bromoacetals, in contrast to $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]^- (3^-)$,
 573 which can only activate the iodoacetal. This result is in
 574 agreement with the synthetic reactions in which the precatalyst
 575 $[(\eta^1\text{-H}_3\text{BH})\text{Fe}^{\text{I}}\text{Cl}(\text{NCCH}_3)_4]^- (5)$ mediates the cyclization of
 576 the bromoacetal, while $[\text{HFe}^{\text{I}}\text{Cl}(\text{dppe})_2]^- (3)$ cannot. The
 577 potential gap between the reduction potential of the haloacetal
 578 and the oxidation potential of the anionic hydrido (3^-) or
 579 borohydrido (5^-) iron(I) complex is crucial for the reaction
 580 outcome. When the potential gap is too large, the SET is too
 581 slow and the catalytic reaction does not occur. This rationalizes
 582 the role of the ligand which is responsible for the formation of
 583 either hydrido iron(II)/iron(I) complexes (ligand dppe) or
 584 borohydrido iron(II)/iron(I) complexes (ligand acetonitrile),
 585 which possess different redox properties.

586 These findings emphasize new perspectives to be considered
 587 in the design of new radical mediators. The fine tailoring of the
 588 ligand on the metal center is key to the success of such an
 589 approach. Work is underway to use this understanding to
 590 identify potential ligand candidates and validate them as
 591 efficient precatalysts for a wider range of radical reactions.

592 EXPERIMENTAL SECTION

General Considerations. All reactions were carried out in oven-
 593 dried glassware under an argon atmosphere using standard Schlenk 594

595 techniques. Reaction solvents (acetonitrile and dichloromethane) were
596 degassed with argon and dried using a PureSolv Micro Solvent
597 Purification System (Innovative Technology) by percolation through a
598 column packed with neutral alumina under a positive pressure of
599 argon. Other solvents were dried and distilled prior to use by literature
600 methods. All solvents were further degassed via freeze/pump/thaw
601 cycles with argon prior to use. Column chromatography was
602 performed using Merck Geduran SI 60 Å silica gel (35–70 μm).²
603 Purification was performed using Sigma-Aldrich 58 Å neutral alumina
604 (Brockman I, activated) as required. All substrates were filtered
605 through neutral alumina prior to use. Iron(II) dichloride (anhydrous
606 beads, ~10 mesh, 99.99%) and sodium borohydride (99.99%) were
607 purchased from Sigma-Aldrich. These reagents were stored in the
608 glovebox and used as received. NMR spectra were recorded on a
609 Bruker AV400 or Avance III spectrometer fitted with either a QNP or
610 BBFO probe and calibrated using undeuterated acetonitrile (δ_{H} 1.94
611 ppm and δ_{C} 118.26) or chloroform (δ_{H} 7.26 ppm and δ_{C} 77.16 ppm)
612 as internal references. High-resolution ESI mass spectra were recorded
613 using a FT-ICR mass spectrometer (7 T hybrid FTICR Solarix
614 spectrometer, Bruker Daltonik GmbH) combined with an ion funnel
615 geometry to transfer the formed ions. Cyclic voltammetry (CV) was
616 performed with a laboratory-made potentiostat and a PAR Model 175
617 waveform generator. The working electrode was a steady gold disk (d
618 = 1 mm), the counter electrode a platinum wire (ca. 0.2 cm² apparent
619 area), and the reference a saturated calomel electrode. Cyclic
620 voltammograms were recorded with a Nicolet 3091 digital
621 oscilloscope. All experiments were carried out under argon.

622 **Typical Procedure for the Synthesis of Iron Complexes.**
623 *trans*-Hydridochlorobis[1,2-bis(diphenylphosphino)ethane]iron(II),
624 *trans*-[HFeCl(dppe)₂] (3)..^{17,38} Iron(II) dichloride (160 mg, 1.3 mmol)
625 and bis(diphenylphosphino)ethane (1.00 g, 2.5 mmol) were placed in
626 a Schlenk flask in the glovebox, and then ethanol (15 mL) was added
627 under argon. The mixture was stirred vigorously for 10 min and then
628 heated to 50 °C. Sodium borohydride (29 mg, 0.76 mmol) was placed
629 in a separate Schlenk tube in the glovebox, ethanol (12 mL) was added
630 under argon, and the solution was stirred for 10 min. The sodium
631 borohydride solution was added over 15 min to the iron mixture under
632 a counter current of argon. After 2 h the reaction mixture had turned
633 bright red-purple with both red and white precipitates. The reaction
634 mixture was filtered, and the filtrate was washed with distilled water
635 (dried and degassed, 2 mL) and ethanol (dried and degassed, 2 mL)
636 and then dried under vacuum. Benzene (dried and degassed, 20 mL)
637 was added (not all precipitate dissolved) and the solution was filtered
638 under argon. The solvent was removed to give a red-purple solid (550
639 mg, 49%), which was dried under vacuum. The complex was stable in
640 the glovebox for more than 6 months. The title compound had
641 spectral properties identical with those previously reported.

642 *Dichlorobis(acetonitrile)iron(II)* (4).³⁹ Iron(II) chloride (1.00 g, 7.9
643 mmol) was placed in a Schlenk tube in the glovebox. Acetonitrile (5
644 mL) was added under argon, and the reaction mixture was heated to
645 reflux for 3 h. The reaction mixture was cooled to room temperature,
646 resulting in the precipitation of a solid. The solvent was removed
647 under argon, and the solid was washed with toluene (dried and
648 degassed, 2 mL) and hexane (dried and degassed, 2 mL) and then
649 dried under vacuum to give colorless fine crystals (1.40 g, 85%). The
650 complex was stable in the glovebox for more than 6 months. Crystals
651 suitable for single-crystal X-ray crystallography were obtained directly
652 from the mother liquor and corresponded to the data previously
653 reported.

654 *Chlorotetrakis(acetonitrile)iron(II) Borohydride* (5). Dichlorobis-
655 (acetonitrile)iron(II) (4; 360 mg, 1.7 mmol) was placed in a Schlenk
656 flask in the glovebox, acetonitrile (15 mL) was added under argon, and
657 the mixture was stirred for 10 min. Sodium borohydride (30 mg, 0.76
658 mmol) was placed in a separate Schlenk tube in the glovebox,
659 acetonitrile (12 mL) was added under argon, and the solution was
660 stirred vigorously for 10 min. The sodium borohydride solution was
661 added over 15 min to the iron mixture under a counter current of
662 argon. After 3 h the reaction mixture had turned dark orange with
663 some precipitate observed. The reaction mixture was filtered under

argon, the filtrate was concentrated, and the resulting orange-brown
solid (255 mg, 56%) was dried under vacuum and stored under argon.

Typical Procedure for the Synthesis of Haloacetal Substrates. A mixture of *N*-halosuccinimide (76 mmol) and alcohol (74 mmol) in dichloromethane (40 mL) was cooled to -10 °C, and then 3,4-dihydro-2*H*-pyran (6.8 mL, 75 mmol) was added dropwise under argon. The reaction mixture was warmed to room temperature over 3 h and stirred at this temperature overnight. The reaction mixture was diluted with dichloromethane (40 mL) and washed with saturated sodium thiosulfate (3 × 20 mL). The combined aqueous phase was extracted with dichloromethane (3 × 20 mL), the combined organic phase was washed with brine (20 mL) and dried (phase separation paper), and the solvent was removed in vacuo. The residue was purified by column chromatography (20% diethyl ether/80% pentane) to afford the title compound. The product was filtered through neutral alumina prior to use.

Typical Procedure for Cyclization of Haloacetal Substrates. The iron complex (0.1 mmol) and reducing agent were placed in a screw-cap tube in the glovebox (the tube was capped with a Suba-Seal). Acetonitrile (1.5 mL) was added under argon, and the mixture was stirred for ca. 15 min at room temperature. A solution of haloacetal (1.0 mmol) in acetonitrile (0.5 mL) was added under argon, the Suba-Seal was replaced by a screw cap, and the reaction mixture was heated to 50 °C overnight. The reaction mixture was cooled to room temperature and quenched with water (20 mL), and the aqueous phase was extracted with dichloromethane (3 × 20 mL). The combined organic phase was washed with brine (30 mL) and dried (phase separation paper), and the solvent was removed in vacuo. The residue was purified by flash chromatography (5–20% diethyl ether/pentane).

Electrochemical Analyses. Cyclic voltammetry (CV) was performed in a three-electrode cell connected to a Schlenk line (under argon) at room temperature with a laboratory-made potentiostat and a PAR Model 175 waveform generator. The working electrode was a steady gold disk (d = 1 mm) and the counter electrode a platinum wire (ca. 0.2 cm² apparent area). The reference was a saturated calomel electrode (SCE) separated from the solution by a bridge filled with tetrabutylammonium tetrafluoroborate in acetonitrile solution (0.3 M, 2 mL). The same solution (0.3 M, 12 mL) was used as the solvent in the electrochemical cell for all CV experiments reported herein. Cyclic voltammograms were recorded with a Nicolet 3091 digital oscilloscope. All experiments were carried out under argon.

General Procedure A: Determination of Redox Potentials. Substrate (0.048 mmol, 4 mM) was placed in the electrochemical cell containing a solution of tetrabutylammonium tetrafluoroborate in acetonitrile (0.3 M, 12 mL). The CV was performed immediately after mixing, at a scan rate of 0.5 V s⁻¹.

General Procedure B: Electrochemical Reduction of Iron Complexes in the Presence of Sodium Borohydride. The iron(II) complex (0.048 mmol, 4 mM) was placed in the cell, the mixture was stirred briefly, and the CV was performed immediately at a scan rate of 0.5 or 5 V s⁻¹. Sodium borohydride (0.048 mmol) was added, the mixture was stirred briefly, and the CV was performed immediately.

General Procedure C: Electrochemical Reduction of Iron Complexes in the Presence of Haloacetal. The iron complex (0.048 mmol, 4 mM) was placed in the cell, the mixture was stirred briefly, and the CV was performed immediately at a scan rate of 0.5 V s⁻¹. The haloacetal (0.048 mmol) was added, the mixture was stirred briefly, and the CV was performed immediately. Additional aliquots of haloacetal (1 equiv) were added, and the CV was performed immediately, up to a total of 4–12 equiv depending on substrate.

Mössbauer Spectroscopy. Mössbauer spectra were recorded on powder samples of natural-abundance ⁵⁷Fe compounds or on acetonitrile solutions starting with fully enriched ⁵⁷FeCl₂. In the synthesis of isotopically enriched ⁵⁷FeCl₂, elemental 94%-enriched ⁵⁷Fe was heated under a dry in situ generated chlorine atmosphere at 350 °C over 15 min. Dry chlorine was obtained by adding dropwise anhydrous sulfuric acid into a commercial bleach solution; the resulting chlorine gas was then dried by bubbling into sulfuric acid

734 (98%). The experiments were performed using a horizontal trans-
735 mission 4 K closed cycle refrigerator system from Janis and SHI and a
736 100 mCi source of $^{57}\text{Co}(\text{Rh})$ as previously described.⁴⁰ All velocity
737 scales and isomer shifts are referred to the metallic iron standard at
738 room temperature. Analysis of the data was performed with the
739 software WMOSS4 Mössbauer spectral analysis software (www.wmoss.org,
740 2009–2015).

741 **DFT Calculations.** Simulations of the nuclear parameters were
742 performed using the ORCA code (v. 3.0.3) at a DFT level (BP86,
743 Ahlrich's TZVP set for all atoms except Fe, which was treated using
744 the CP(PPP) set). See the [Supporting Information](#) for additional
745 details and the corresponding references. Mössbauer parameters can
746 be easily evaluated using straightforward DFT techniques, which allow
747 the computation of the electronic density at the ^{57}Fe nucleus, denoted
748 ρ_0 (in units of $e a_0^{-3}$, where a_0 is the Bohr radius). A reliable estimation
749 of the isomer shift (δ) can be obtained with a linear extrapolation from
750 ρ_0 .⁴¹ To do so, a benchmark of high-spin and low-spin Fe^{II} complexes
751 has been used to calibrate the chosen level of theory. Computation of
752 the quadrupolar split was not performed herein because this parameter
753 is highly dependent on the symmetry of the occupied electronic levels.
754 For several complexes in the following benchmark, the experimental
755 value of ΔE_{Q} could not be reproduced with a satisfying accuracy.
756 Minor differences between the optimized geometries and the
757 experimental structures can induce strong discrepancies between the
758 experimental and calculated values of ΔE_{Q} .⁴² Computation of the
759 Mössbauer parameters was performed using ORCA 3.0.3 software.⁴¹
760 The BP86 functional⁴³ was used, with Ahlrich's TZVP basis set⁴⁴ for
761 all atoms except Fe, which was described using the CP(PPP) enlarged
762 basis set.⁴⁵

763 ■ ASSOCIATED CONTENT

764 ● Supporting Information

765 The Supporting Information is available free of charge on the
766 ACS Publications website at DOI: [10.1021/acs.organomet.7b00603](https://doi.org/10.1021/acs.organomet.7b00603).

768 Experimental details and procedures, spectral data,
769 electrochemical analysis, Mössbauer spectroscopy, and
770 computational analysis (PDF)

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785 Notes

786 The authors declare no competing financial interest.

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