

Iron(II) Complexes of the Linear *rac*-Tetraphos-1 Ligand as Efficient ² Homogeneous Catalysts for Sodium Bicarbonate Hydrogenation and **Formic Acid Dehydrogenation**

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Supporting Information 7

and are described herein.

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ABSTRACT: The linear tetraphosphine 1,1,4,7,10,10-hexaphenyl-1,4,7,10-tetraphosphadecane 8 9 (tetraphos-1, P4) was used as its rac and meso isomers for the synthesis of both molecularly defined and in situ formed Fe(II) complexes. These were used as precatalysts for sodium 10 bicarbonate hydrogenation to formate and formic acid dehydrogenation to hydrogen and carbon 11 dioxide with moderate to good activities in comparison to those for literature systems based on 12 Fe. Mechanistic details of the reaction pathways were obtained by NMR and HPNMR 13 experiments, highlighting the role of the Fe(II) monohydrido complex $[FeH(rac-P4)]^+$ as a key 14 15 intermediate. X-ray crystal structures of different complexes bearing rac-P4 were also obtained



KEYWORDS: iron phosphine complexes, formic acid dehydrogenation, bicarbonate hydrogenation, X-ray crystallography, 17 HPNMR mechanistic studies 18

INTRODUCTION 19

20 Hydrogen is of crucial importance in the chemical industry and 21 holds great potential as a secondary energy carrier, as a 22 feedstock for direct hydrogen fuel cells.¹ Its generation from 23 renewable sources and its storage in a safe and reversible 24 manner are urgent targets for the widespread application of 25 hydrogen in such technologies. Among the different H₂ storage 26 materials, formic acid (FA) is a nontoxic hydrogen source 27 which can be handled and transported easily and possesses a 28 relatively high H₂ content (4.4 wt %). H₂ generation from 29 formic acid affords $H_2 + CO_2$ mixtures and is therefore an 30 "atom efficient" process, since no hydrogen is wasted in the 31 formation of byproducts (such as H₂O, as in the case of H₂ 32 generation from methanol or methane). In addition, the 33 byproduct CO_2 can be, in the presence of suitable catalysts, 34 rehydrogenated back to FA, affording a zero-carbon footprint 35 cycle for hydrogen storage and release.² The efficient $_{36}$ interconversion of FA to $\mathrm{H_2}$ and CO_2 is of importance for 37 both H₂ storage and release and for the utilization of CO₂ or 38 bicarbonates obtained by its trapping in alkaline water 39 solutions, as a abundant C1 feedstock. In the past decade, 40 there have been a number of reports on selective FA 41 dehydrogenation to produce CO-free H₂, as well as on the 42 hydrogenation of CO₂ or bicarbonates to FA or formate salts. 43 However, most of these catalysts are based on low-abundance 44 noble metals such as ruthenium³ and iridium.⁴ Only recently 45 has this chemistry been extended to non-noble metals such as 46 Fe⁵ and Co.⁶

The most active additive-free Fe-based catalyst system for FA 47 dehydrogenation under mild temperature conditions (40 °C) 48 reported to date was obtained by combining the iron(II) salt 49 $Fe(BF_4)_2 \cdot 6H_2O$ with the tetraphosphine ligand P- 50 (CH₂CH₂PPh₂)₃ (PP₃).^{5e,7} Although the nature of the initial 51 complex formed in this reaction has not been fully ascertained, 52 mechanistic studies indicated that under catalytic conditions 53 (FA in propylene carbonate (PC)) complexes $[FeH(PP_3)]^+$ 54 and $[FeH(\eta^2-H_2)(PP_3)]^+$ are formed. ^{Se,8} This catalytic system 55 was successfully applied to bicarbonate hydrogenation to 56 formates and carbon dioxide valorization to alkyl formates 57 and formamides.^{5a} In continuation of this work, efficient iron- 58 catalyzed hydrogenation of carbon dioxide and bicarbonates 59 was achieved using $Fe(BF_4)_2 \cdot 6H_2O$ and $P^{Ph}P_3$ ($P^{Ph}P_3 = tris(2-60)$ (diphenylphosphino)phenylphosphine). In this case, metal 61 complexation afforded the well-defined complex $[FeF(P^{Ph}P_3)]^+$ 62 via F-BF3 activation. Mechanistic studies established that this 63 complex reacts with H₂ to give $[FeH(\eta^2-H_2)(P^{Ph}P_3)]^+$. High- 64 pressure HPNMR CO₂ hydrogenation experiments in the 65 presence of NEt₃ suggested the formation of the known 66 dihydride complex [Fe(H)₂(P^{Ph}P₃)].^{5f}

In recent years, our group has been interested in FA 68 dehydrogenation and CO₂ hydrogenation, so far using Ru⁹ and 69 Ir¹⁰ homogeneous catalysts. In an effort to develop novel, non- 70 noble-metal-based catalysts for such transformations, we 71

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72 became eager to explore the potential of Fe(II) complexes of 73 other tetradentate phosphines.

The linear tetradentate phosphine ligand 1,1,4,7,10,10-75 hexaphenyl-1,4,7,10-tetraphosphadecane (tetraphos-1, P4) ex-76 ists as a mixture of *rac* (*S*,*S*;*R*,*R*) and *meso* (*S*,*R*) 77 diastereoisomers (hereafter *rac*-P4 and *meso*-P4, respectively), 78 which can be separated by fractional crystallization.^{11,12} Despite 79 the fact that the existence of these stereoisomers was 80 recognized as early as 1974,¹³ the importance of this isomerism 81 was not fully appreciated until the work of Brown and 82 Canning.¹¹ The configurations that these diastereoisomers can 83 adopt in an octahedral complex are denoted as cis- α , cis- β , and 84 trans (Chart 1). While the *meso* isomer can adopt only a trans

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ss or cis- β configuration, all three configurations are physically 86 possible for the rac isomer. Nevertheless, the rac isomer is ⁸⁷ known for its propensity to form cis- α complexes.^{11,14} Since the 88 original preparation of tetraphos-1 by King and co-workers,¹⁵ 89 there have been a number of reports on its coordination 90 behavior.¹¹⁻¹⁶ By a close perusal of the available literature, we 91 noticed that the chemistry of the meso isomer is far more 92 developed than that of the *rac* isomer. Only complexes 93 $[FeBr(P4)][BPh_4]^{16d}$ and *trans*- $[FeH(N_2)(P4)]^{16c}$ have been 94 characterized crystallographically, and in both the ligand 95 exhibits a meso configuration. This was probably due to the 96 fact that the authors used commercial tetraphos-1, which is 97 richer in the *meso* isomer. The syntheses of [FeH(P4)]X, 98 $[Fe(NCS)_2(P4)]$, [FeH(NCS)(P4)]X, and [FeH(CO)(P4)]99 (X = Br, I) were also described, but no indication of the 100 configuration of the P4 ligand was provided.^{16e} Morris and co-101 workers reported on the hydrogen exchange between η^2 -H₂ and 102 hydride ligands in *trans*-[FeH(η^2 -H₂)(*meso*-P4)]BF₄, obtained ¹⁰³ by protonation of the corresponding dihydride complex *trans*-¹⁰⁴ $[Fe(H)_2(meso-P4)]$.^{12,14} To the best of our knowledge, a full 105 exploration of the coordination chemistry of rac-P4 to Fe(II) 106 and the reactivity of the complexes so obtained has never been 107 reported.

In this work, we describe the synthesis of novel Fe(II)109 complexes bearing *rac*-P4, their reactivity toward H₂ and CO₂, 110 and their application as efficient catalysts for FA dehydrogen-111 ation and sodium bicarbonate hydrogenation to sodium 112 formate. The catalytic data are complemented by mechanistic 113 details obtained by model stoichiometric reactions and in 114 operando high-pressure HPNMR experiments. 115

RESULTS AND DISCUSSION

Syntheses and Characterization of Fe(II) Complexes. 116 At first, rac-P4 and meso-P4 were obtained in pure isomeric 117 form from the commercial ligand P4, containing a rac:meso 118 ratio of 1:3, by fractional crystallization as described in the 119 literature.¹⁴ In order to test the coordination abilities of the two 120 isomers with suitable iron(II) sources, the commercially 121 available salt $Fe(BF_4)_2 \cdot 6H_2O$ and the easily accessible 122 complex¹⁷ [Fe(CH₃CN)₆](BF₄)₂ were used as metal precur- 123 sors. The reaction of $Fe(BF_4)_2 \cdot 6H_2O$ with rac-P4 (1:1) was 124 rather sluggish in a variety of common solvents, whereas it 125 proceeded smoothly in propylene carbonate (PC), affording a 126 deep purple solution. The ³¹P{¹H} NMR spectrum of this 127 solution (C_6D_6 insert) showed two broad signals at δ_P 99.9 and 128 60.9 ppm, indicative of Fe(II) complexation by the ligand. 129 19 F{ 1 H} NMR analysis at room temperature showed only a $_{130}$ single, sharp peak for the BF₄ anion, suggesting that the 131 complex [FeF(rac-P4)](BF₄), expected to arise upon F-BF₃ 132 bond activation,^{5f} had not formed. Due to the known 133 propensity of Fe(P4) complexes to adopt a pentacoordinate ¹³⁴ geometry, often completed by halide ligands, ^{16c,d,f} we propose ¹³⁵ that under these conditions the complex $[Fe(\eta^1 - FBF_2)](rac-136)$ P4)](BF₄) (1) has formed, where one of the BF₄ counterions 137acts as a weakly coordinating ligand (Scheme 1).¹⁸ This 138 s1





complex is likely to be fluxional in solution, and the loosely 139 bound BF₄ can be easily replaced by a coordinating solvent X 140 $(X = H_2O, MeOH)$,¹⁹ giving complexes such as *cis-α*-141 $[FeX_2(rac-P4)](BF_4)_2$. This was proven by addition of 142 CD₃OD to a solution of 1 in PC, where a new species formed, 143 showing a ³¹P{¹H} NMR pattern composed of two triplets at δ_p 144 107.6 and 73.8 (²J_{PP} = 29.9 Hz), which we attribute to the 145 solvento species *cis-α*-[Fe(CD₃OD)_x(*rac*-P4)](BF₄)₂ (1'; *x* = 1, 146 2). To date, all our attempts to obtain crystals of either 1 or 1' 147 failed. A similar reactivity was observed upon reacting 148 Fe(BF₄)₂·6H₂O with *meso*-P4, which resulted in the formation 149 of a brown solution containing the putative complex [Fe(η^{1} - 150 FBF₃)(*meso*-P4)](BF₄) (1"), also characterized by two broad 151 signals in the ³¹P{¹H} NMR at δ_p 104.8 and 70.8 ppm. 152

In contrast, the reaction of *rac*-P4 with $[Fe(CH_3CN)_6]^{-153}$ (BF₄)₂ resulted in the quantitative formation of the well-defined 154 complex *cis*- α -[Fe(CH₃CN)₂(*rac*-P4)](BF₄)₂ (**2**) as the sole 155 product (Scheme 2). The ³¹P{¹H} NMR spectrum exhibits two 156 s2 triplets at 100.7 and 65.6 ppm in CD₃CN, which reflect an 157 AA'XX' coupling pattern with equivalent *cis*-P,P coupling 158 constants (²J_{PP} = 31.7 Hz). These values are in close analogy 159 with those attributed by Habeck et al. to *cis*- α -[Fe- 160 (NCS)₂(*rac*-^{pr}P4)] (*rac*-^{pr}P4 = 1,1,4,8,11,11-hexaphenyl- 161 Scheme 2. Synthesis of *rac*-P4 and *meso*-P4 Complexes Starting from $[Fe(CH_3CN)_6](BF_4)_2$.



162 1,4,8,11-tetraphosphaundecane).²⁰ Crystals suitable for X-ray 163 diffraction analysis were grown by adding *n*-pentane to a 164 solution of 2 in acetonitrile/methanol (Figure 1). Complex 2



Figure 1. Molecular structure for the cationic portion of **2**. Ellipsoids are set at the 50% probability level. Hydrogen atoms are omitted for clarity. Selected bond lengths (Å) and bond angles (deg): Fe1–N1, 1.943(3); Fe1–N2, 1.955(4); Fe1–P1, 2.2869(13); Fe1–P2, 2.2137(12); Fe1–P3, 2.2246(12); Fe1–P4, 2.2981(13); N1–Fe1–N2, 88.70(15); N1–Fe1–P1, 90.36(11); N1–Fe1–P2, 92.50(11); N1–Fe1–P3, 174.50(12); N1–Fe1–P4, 90.06(11); N2–Fe1–P1, 91.57(12); N2–Fe1–P2, 117.12(12); N2–Fe1–P3, 93.76(11); N2–Fe1–P4, 88.25(12).

165 crystallizes in the C2/*c* space group and has an octahedral 166 coordination geometry at the Fe(II) center, with Fe–P_{ax} 167 distances (Fe1–P1 = 2.2868(13) Å and Fe1–P4 = 168 2.2982(13) Å) that are longer than the Fe–P_{eq} distances 169 (Fe1–P2 2.2138(13) and Fe1–P3 2.2247(12) Å). Notably, 170 complex **2** was stable in air as a solid and solutions in 171 acetonitrile/methanol could be stored under nitrogen for over 172 1 month without any appreciable decomposition. In contrast, 173 the reaction of *meso*-P4 with [Fe(CH₃CN)₆](BF₄)₂ was not 174 selective and afforded a mixture of two products in an 175 approximately 2:1 ratio, which were identified on the basis of 176 characteristic ³¹P{¹H} NMR resonances^{12,20} (see the <u>Support-</u> 177 <u>ing Information</u>) as the *trans-* and *cis-β* isomers of [Fe-178 (CH₃CN)₂(*meso*-P4)](BF₄)₂, respectively.

Syntheses of Fe(*rac***-P4) Hydrido Complexes.** Due to the relevance of Fe–hydrido complexes to FA dehydrogenation and bicarbonate hydrogenation reactions, we targeted the syntheses of the so far unknown mono- and dihydride iron complexes of *rac*-P4. The analogues of the *meso* isomer have 183 been previously reported.^{12,16e} The monohydrido complex 184 [FeH(*rac*-P4)][BPh₄] (3·BPh₄) was obtained upon reacting 185 *rac*-P4, anhydrous FeCl₂, NaBPh₄, and NaBH₄ in stoichio- 186 metric amounts in THF/MeOH and was characterized by 187 NMR and X-ray diffraction studies upon growing suitable 188 crystals from these solutions. The ³¹P{¹H} NMR spectrum of 189 3·BPh₄ in *d*₈-THF showed two triplets at δ_p 119.4 and 99.4 190 ppm, reflecting an AA'XX' coupling pattern with an observed 191 splitting of 24.5 Hz, while in the corresponding ¹H NMR 192 spectrum, the hydride signal appeared as a broad triplet at 193 -9.16 ppm (²J_{HP} = 24.0 Hz). The crystal structure of 3·BPh₄ 194 displays a pseudo-octahedral geometry, with the hydride ligand 195 occupying two sites in the crystal: i.e., alternatively one or the 196 other cis position in 50% occupancy (Figure 2). The distortion 197 f2



Figure 2. Molecular structure for the cationic portion of $3 \cdot BPh_4$. Ellipsoids are set at the 50% probability level. Hydrogen atoms are omitted for clarity, except for the hydrido ligand. Selected bond lengths (Å) and bond angles (deg): Fe1–P1, 2.2189(7); Fe1–P2, 2.1961(7); Fe1–P3, 2.2207(7); Fe1–P4, 2.1993(8); Fe1–H1h, 1.61(5); Fe1–H1h', 1.41(5); H1h–Fe1–H1h', 90(3); P1–Fe1–P2, 85.38(3); P1–Fe1–P3, 102.58(3); P1–Fe1–P4, 170.41(3); P2–Fe1–P3, 85.74(3); P2–Fe1–P4, 96.87(3); P3–Fe1–P4, 86.91(3); H1h–Fe1–P1, 89.0(2); H1h–Fe1–P2, 172.4(18); H1h–Fe1–P3, 100.6(18); H1h–Fe1–P4, 87.(2); H1h'–Fe1–P1, 83.(2); H1h'–Fe1–P2, 83.(2); H1h'–Fe1–P3, 168(2); H1h'–Fe1–P4, 87 (2).

from the ideal octahedral geometry is evident from the P1- 198 Fe1-P4 angle (170.4°), which is significantly bent in 199 comparison to the analogous P1-Fe1-P4 angle in 2 200 (179.5°), whereas the P2-Fe1-P3 angles are comparable in 201 2 and 3 (85.3° vs 85.7°).

The neutral dihydrido complex $cis-\alpha$ -[Fe(H)₂(rac-P4)] (4) 203 was synthesized from rac-P4, anhydrous FeCl₂, and excess 204 NaBH₄ under reflux conditions in a THF/EtOH mixture. The 205 ³¹P{¹H} NMR spectrum of 4 in d_8 -THF displayed two triplets 206 at δ_p 123.8 and 113.1 ppm with ² J_{PP} = 13.5 Hz due to cis-P,P 207 coupling, whereas the two hydride ligands gave a complex 208 multiplet centered at -11.7 ppm (apparent double septuplet; 209 see the <u>Supporting Information</u>). Crystals of 4 suitable for X- 210 ray analysis were grown by diffusion of MeOH into the solution 211 which resulted from the reaction mixture, after filtration and 212 partial evaporation of the solvent. The solid-state molecular 213 structure of 4 displays a significantly distorted octahedral 214 coordination geometry at the Fe(II) center with the *rac*-P4 215 f3 f3

216 ligand adopting a cis- α configuration (Figure 3). The P1–Fe1– 217 P4 angle in 4 (159.6°) is significantly more distorted than that



Figure 3. Molecular structure of 4. Ellipsoids are set at the 50% probability level. Hydrogen atoms are omitted for clarity, except for hydrido ligands. Selected bond lengths (Å) and bond angles (deg): Fe1–P1, 2.1249(7); Fe1–P2, 2.1510(7); Fe1–P3, 2.1654(7); Fe1–P4, 2.1303(7); Fe1–H1h, 1.55(3); Fe1–H1j, 1.57(2); H1h–Fe1–H1j, 90.6(13); P1–Fe1–P2, 89.00(3); P1–Fe1–P3, 106.14(3); P1–Fe1–P4, 159.62(3); P2–Fe1–P3, 86.05(2); P2–Fe1–P4, 106.21(3); P3–Fe1–P4, 88.71(3); H1h–Fe1–P1, 85.0(9); H1h–Fe1–P2, 91.8(9); H1h–Fe1–P3, 168.6(9); H1h–Fe1–P4, 81.1(9); H1j–Fe1–P1, 82.3(9); H1j–Fe1–P2, 93.3(9); H1j–Fe1–P3, 171.8(9); H1j–Fe1–P4, 83.0(9).

218 in $3 \cdot \text{BPh}_4$ (170.4°). Furthermore, all Fe–P bond distances are 219 significantly shorter (all <2.17 Å) with respect to those 220 observed in 2 (2.19–2.22 Å) and $3 \cdot \text{BPh}_4$ (2.21–2.30 Å).

Reactivity of 1 and 2 toward H₂. To verify the potential 222 of **1** and **2** as hydrogenation catalyst precursors, we investigated 223 at first their reactivity toward molecular H₂ in model reactions 224 under HPNMR conditions. A solution of $Fe(BF_4)_2$.6H₂O and 225 *rac*-P4 in PC (1.5 mL, 0.01 M) was initially transferred into a 226 10 mm medium-pressure HPNMR sapphire tube. ³¹P{¹H} 227 NMR analysis at room temperature under Ar atmosphere 228 showed, as expected, broad signals due to **1**.

Upon addition of CD_3OD for a deuterium lock (0.5 mL), the 229 $_{230}\ ^{31}P\{^{1}H\}$ NMR pattern due to 1' appeared, while no hydride ²³¹ signals were observed in the corresponding ¹H NMR spectrum. 232 The tube was then pressurized at room temperature with 30 bar 233 of H_{2} , which resulted in the quantitative conversion of 1 and 1' 234 into a new species that we identified as $3 \cdot BF_4$ on the basis of its $^{31}\text{P}\{^1\text{H}\}$ NMR pattern being identical with that of the isolated 235 236 monohydride $3 \cdot BPh_4$.²¹ Due to HPNMR conditions and 237 possible H/D exchange, the Fe-H hydrido ligand appeared $_{238}$ as a broad signal centered at ca. -9.16 ppm. In the temperature 239 range 233-353 K, no changes in the spectra were observed, suggesting that a putative hydrido-dihydrogen complex such as 240 $[FeH(\eta^2-H_2)(rac-P4)]^+$ does not form under these conditions, 241 242 in analogy to what was previously described for [FeH(meso-243 P4)]⁺.¹⁴ This was further verified by repeating the experiment 244 using a 0.025 M solution of 1 in PC/CD₃OH (3/1, total 245 volume 2.0 mL) and measuring the longitudinal relaxation time 246 (T_1) at 293 K, respectively, giving values of ca. 900 ms with 247 good exponential fitting of the data, in line with the values 248 expected for a classical hydride.

Complex 2 was remarkably less reactive toward H₂ than 1. 249 Complex 2 was dissolved in CD_3OD and reacted with H_2 (30 250 bar) under HPNMR conditions (see the Supporting 251 Information). At room temperature, in addition to the peaks 252 of unreacted 2, four distinct ³¹P{¹H} NMR resonances were 253 observed to appear at $\delta_{\rm P}$ 121.7 (br s), 104.0 (br d), 101.2 (br 254 d), and 96.3 (br s). The corresponding ¹H NMR spectrum 255 showed an apparent doublet of quartets centered at $\delta_{\rm H}$ –8.5 256 ppm (dq, ${}^{2}J_{\text{HPris,ax1}} = 36.7$ Hz, ${}^{2}J_{\text{HPcis,eq}} = 51.3$ Hz, ${}^{2}J_{\text{HPcis,ax1}} = 257$ ${}^{2}J_{\text{HPcis,ax2}} = 51.1$ Hz). This pattern, indicative of nonequivalent 258 phosphorus atoms typical of an octahedral Fe complex, was 259 attributed to the formation of $cis-\alpha$ -[FeH(NCMe)(rac-260P4)](BF₄) (5). The resonances due to $3 \cdot BF_4$ appeared at 313 261 K. The temperature was then further increased to 333 and 353 262 K. The signals due to 3.BF4 and 5 were observed to increase, 263 reaching almost complete conversion of 2 with a final 1:3 ratio 264 of ca. 1:3 between $3 \cdot BF_4$ and 5. The reaction is reversible, as 265 cooling to 293 K gave back the same pattern initially observed 266 (Scheme 3 and the Supporting Information). The experiment 267 s3





was repeated in the presence of an added base (NEt₃), affording 268 at first a mixture of $3 \cdot BF_4$ and 5 upon heating, and then 5 as the 269 only product after 20 h of standing at 293 K. Addition of Et₂O/ 270 pentane to the reaction mixture resulted in the precipitation of 271 yellow crystals of 5. The corresponding X-ray crystal structure, 272 albeit highly disordered, was however useful to confirm the 273 proposed formula (see the Supporting Information). 274

Reactivity of 3·BPh₄ and 4 toward CO₂. In the next step, 275 we explored the reactivity of the mono- and dihydrides 3·BPh₄ 276 and 4 toward CO₂. Beller et al. showed that insertion of CO₂ 277 into the Fe–H bond of the complex [FeH(PP₃)]⁺ could be 278 achieved under 10 atm of gas pressure, giving the 279 corresponding formate complex.^{5a} In an NMR-scale experi- 280 ment, we reacted the monohydride 3·BPh₄ with CO₂ (1 atm) 281 in *d*₈-THF, obtaining as expected the formate complex *cis-α*- 282 [Fe(η^2 -O₂CH)(*rac*-P4)](BPh₄) (6·BPh₄; Scheme 4), having 283 s4 J³¹P{¹H} NMR signals at δ_P 106.0 (t) and 76.5 (t, ²*J*_{PP} = 29.5 284 Hz). In the corresponding ¹³C{¹H} NMR spectrum, apart from 285 the signal at δ_C 162.4 ppm due to BPh₄⁻, a broad singlet at 286 174.6 ppm compatible with a coordinated formate anion was 287







Figure 4. Molecular structure for the cationic part of $\{\mu_2 - [Fe(MeOH)_4] \cdot \kappa^{10} - [Fe(\eta^2 - O_2CO)(rac - P4)]_2\}$ (BPh₄)₂ (7'). Ellipsoids are set at the 50% probability level. Hydrogen atoms are omitted for clarity. Selected bond lengths (Å) and bond angles (deg): Fe1-O1, 2.035(5); Fe1-O2, 2.027(5); Fe1-P1, 2.286(3); Fe1-P2, 2.205(3); Fe1-P3, 2.197(2); Fe1-P4, 2.297(3); Fe2-O4, 2.032(5); Fe2-O5, 2.046(5); Fe2-P5, 2.282(2); Fe2-P6, 2.217(3); Fe2-P7, 2.204(2); Fe2-P8, 2.260(2); Fe3-O3, 2.133(6); Fe3-O6, 2.076(6); O1-Fe1-O2, 65.0(2); O1-Fe1-P1, 96.25(17); O1-Fe1-P2, 106.26(19); O1-Fe1-P3, 165.40(18); O1-Fe1-P4, 86.72(17); O2-Fe1-P1, 87.63(17); O2-Fe1-P2, 167.59(17); O2-Fe1-P3, 104.43(18); O2-Fe1-P4, 94.20(17); O4-Fe2-O5, 64.8(2); O4-Fe2-P5, 93.80(16); O4-Fe2-P6, 104.65(17); O4-Fe2-P7, 166.98(19); O4-Fe2-P8, 86.48(16); O5-Fe2-P5, 85.36(16); O5-Fe2-P6, 164.85(16); O5-Fe2-P7, 107.37(16); O5-Fe2-P8, 92.69(17); O3-Fe3-O6, 178.2(2).

Table 1. Hydrogenation of Sodium Bicarbonate using either in Situ Formed or Defined Molecular Fe(II) Precatalysts^a

entry	catalyst precursor	substrate:catalyst	T (°C)	$p(H_2)$ (bar)	TON ^{<i>i</i>,<i>k</i>}	yield (%) ^{j,k}
1^b	i	1000	80	60	154 (±4)	15 (±0)
2^{c}	1″	1000	80	60	62 (±16)	6 (±2)
3^d	1	1000	80	60	575 (±52)	58 (±5)
4^d	1	1000	100	60	588 (±74)	59 (±7)
5^d	1	1000	60	60	186 (±14)	19 (±1)
6^d	1	1000	80	30	620 (±36)	62 (±4)
7^d	1	1000	80	10	398 (±14)	40 (±1)
8^e	1	10000	80	60	83 (±27)	1 (±0)
$9^{d_i f}$	1	3000	80	60	723 (±40)	24 (±1)
10^g	2	1000	80	60	762 (±105)	76 (±11)
11^g	2	1000	100	60	555 (±15)	55 (±1)
12^g	2	1000	60	60	161 (±6)	16 (±1)
13^g	2	1000	80	30	766 (±81)	71 (±14)
14^h	2	10000	80	60	1229 (±18)	12 (±0)

^{*a*}General reaction conditions: catalyst precursor (0.01 mmol); NaHCO₃ (10 mmol); MeOH (20 mL); H₂ pressure; 24 h. ^{*b*}Catalyst precursor i: 1 mL of a 0.01 M stock solution of commercial P4 and Fe(BF₄)₂·6H₂O (1:1). ^{*c*}Catalyst precursor 1": 1 mL of a 0.01 M stock solution of 1 in PC. ^{*d*}Catalyst precursor 1: 1 mL of a 0.01 M stock solution of 1 in PC. ^{*e*}O.1 mL of a 0.01 M stock solution of NaHCO₃. ^{*s*}Complex 2 (0.01 mmol) was added to the autoclave from a CH₃CN stock solution, from which the solvent was subsequently removed (0.02 M, 0.5 mL, see the Experimental Section for details). ^{*h*}Complex 2 (0.001 mmol) was added to the autoclave from a CH₃CN stock solution formate)/(mmol of catalyst). ^{*j*}Yields calculated from the integration of ¹H NMR signals due to NaHCO₂, using THF as internal standard. ^{*k*}Values of yields and TONs were calculated as averages from the analysis of two to four samples. The largest deviations are reported in parentheses; selected experiments were repeated to ensure reproducibility.

288 observed. By repetition of the test using in situ generated 3·BF₄ 289 and ¹³CO₂, the singlet at 174.6 ppm turned as expected into a 290 doublet with ¹*J*_{CH} = 208.8 Hz in the corresponding proton-291 coupled ¹³C NMR spectrum. ^{5a,22} Unfortunately, the ¹H NMR 292 signal expected in the range 8.2–8.5 ppm for the formate 293 ligand, diagnostic for η^1 vs η^2 coordination, was lying under the 294 ligand aromatic proton multiplet. After 24 h, the ³¹P{¹H} NMR 295 spectrum showed signals of a new complex with triplets at δ_p 296 106.6 and 73.2 ppm (²*J*_{PP} = 30.4 Hz), which we assigned to the 297 neutral carbonate complex *cis-α*-[Fe(η^2 -O₂CO)(*rac*-P4)] (7). The corresponding ${}^{13}C{}^{1}H$ NMR signal was determined from ${}_{298}$ the experiment run using ${}^{13}CO_2$, giving a singlet at 158.1 ppm. ${}_{299}$ The attribution was confirmed by independent synthesis of 7 ${}_{300}$ by reaction of 1 with an excess of K₂CO₃ in PC. In addition, the ${}_{301}$ formation of complex 7 was observed also in HPNMR ${}_{302}$ experiments upon reacting 2 with NaHCO₃ (vide infra). ${}_{303}$

MeOH diffusion into the d_8 -THF solution recovered after $_{304}$ the NMR experiment described above afforded a few purple $_{305}$ crystals which were found to be suitable for X-ray diffraction $_{306}$ data collection. Quite surprisingly, the solid-state structure $_{307}$

±1

t1

308 revealed a trimetallic unit in which a central $[Fe(MeOH)_4]^{2+}$ 309 moiety bridges two $[Fe(O_2CO)(rac-P4)]$ moieties via the two 310 carbonate ligands by η^{1} -O coordination, as shown in Figure 4. 311 Despite the fact that the formation of the complex { μ_2 -312 $[Fe(MeOH)_4]$ - κ^1O - $[Fe(\eta^2$ - $O_2CO)(rac-P4)]_2$ }(BPh_4)_2 (7') 313 may be accidental, its solid-state structure confirmed the 314 presence of CO₃²⁻ ligands. Carbonate is likely to form by 315 reductive disproportionation of CO₂ into CO₃²⁻ and CO, 316 promoted by 3·BPh_4.²³ This reaction, occurring via a formate 317 intermediate, has been previously described with Fe(II) hydrido 318 complexes such as trans- $[Fe(H)_2(dppe)_2]$ and [Fe- $319 (H)_2(PP_3)].^{23,24}$

³²⁰ Complex 4 was also tested for reactivity with CO₂, to check ³²¹ for the possible formation of Fe hydrido formate complexes, ³²² similarly to what proposed by Beller et al. for [Fe-³²³ (H)₂(P^{Ph}P₃)].^{5f} No reaction was observed under the conditions ³²⁴ described above (i.e., 1 atm of CO₂ in d_8 -THF, room ³²⁵ temperature). The experiment was repeated under a moderate ³²⁶ pressure of CO₂ (7 bar) under HPNMR conditions, but again ³²⁷ no reaction occurred.

Fe-Catalyzed Sodium Bicarbonate Hydrogenation. 328 329 The added base-free hydrogenation of sodium bicarbonate to 330 formate in MeOH was tested in stainless steel autoclaves at different H₂ pressures and temperatures. In a preliminary 331 332 experiment, we tested the activity of a combination of commercial tetraphos-1 (P4) and $Fe(BF_4)_2 \cdot 6H_2O$ (0.01 333 mmol, 1:1 ratio) in the hydrogenation of sodium bicarbonate 334 335 in MeOH. To our delight, at 80 °C under 60 bar H₂, sodium 336 formate was formed with TON = 154 (entry 1). The activity of 337 1 and 1" was then tested to check for ligand effects. The in situ 338 formed precatalysts were obtained from stock solutions made 339 from $Fe(BF_4)_2 \cdot 6H_2O$ and either rac-P4 or meso-P4 (0.01 M in 340 PC). The solutions were analyzed by ${}^{31}P{}^{1}H$ NMR before use 341 to confirm the formation of the corresponding Fe(II) 342 complexes 1 and 1". Catalyst precursor 1" gave a rather poor 343 catalytic performance, reaching a TON value of 62 after 24 h 344 under 60 bar of H₂ and 80 °C using 0.1 mol % catalyst (Table 345 1; entry 2). In contrast, 1 was rather active in the catalytic 346 hydrogenation of NaHCO₃ in MeOH. Under 60 bar of H₂ 347 pressure, using 0.1 mol % of atalyst, rather good yields (58% 348 and 59%) and TONs (575 and 588) were achieved at 80 and 349 100 °C, respectively (Table 1; entries 3 and 4). TON values are 350 on the same order of magnitude as those obtained by Beller et 351 al. with the $Fe(BF_4)_2/PP_3$ system under comparable con-352 ditions.^{5a} At 60 °C (entry 5) the TON decreased to 186 with a 353 formate yield of 19%. The effect of H₂ pressure on the 354 productivity of the reaction was also tested. TON and yield 355 were not affected at 80 °C in passing from 60 to 30 bar (entry 356 6), whereas at 10 bar the yield of formate was slightly reduced 357 (entry 7). Using a catalyst to substrate ratio of 1:10000, 358 significantly lower TON and yield were obtained (entry 8). At 359 an intermediate catalyst to substrate ratio (1:3000, obtained by 360 increasing the substrate concentration) good activity was observed with TON = 723 and 24% yield in formate (entry 361 9). The hydrogenation of NaHCO₂ to NaHCO₂ using the well-362 363 defined molecular complex 2 as the catalyst precursor (0.1 mol %) proceeded smoothly at 80 °C, affording sodium formate in 364 365 excellent yields (76 and 71%) and good TONs (762 and 766) 366 under 60 and 30 bar of H_2 pressure (entries 10 and 13, 367 respectively). When the catalyst loading was lowered to 0.01 368 mol %, an increased TON = 1229 was measured, albeit with a 369 lower yield in formate (12%) (entry 14). At this catalyst to 370 substrate ratio, 2 performed better than 1 (1.2 mmol of sodium

formate vs ca. 0.1 mmol obtained; entries 14 and 8, 371 respectively). Finally, at higher (100 °C) or lower temperatures 372 (60 °C) in the presence of **2** (0.1 mol %), lower yields of 373 formate were obtained (entries 11 and 12). On the basis of 374 these results, we propose that the better performance of the *rac* 375 vs *meso* systems can be related to the preferred *cis* conformation 376 of the former, suitable for an inner-sphere mechanism (vide 377 infra). In the case of *meso*-P4, different isomers can form in 378 solution (Scheme 1), hence decreasing the concentration of the 379 likely active form, i.e. *cis-β*-**2** (Scheme 2).

Mechanistic Studies. To gain mechanistic insights into the 381 iron-catalyzed hydrogenation of NaHCO3 to sodium formate in 382 the presence of 1 and 2, we monitored catalyst evolution by 383 HPNMR spectroscopy under in operando conditions. In detail, 384 a 10 mm HPNMR sapphire tube was initially charged with a 385 0.01 M solution of 1 in PC (1.5 mL), CD₃OD (0.5 mL), and 386 NaHCO3 (84 mg; 1.0 mmol, 100 equiv). The ³¹P{¹H} NMR 387 pattern showed the presence of 1 (25%), 1' (25%), and the 388 new species 8 (50%) (percentages are based on integrals), 389 characterized by two triplets at δ_p 107.1 and 72.9 ppm (${}^2J_{PP}$ = 390 30.3 Hz).²⁵ Pressurization of the reaction mixture with H_2 (30 391 bar) resulted in the formation of the monohydride complex 3. 392 BF_4 (34%) at room temperature. The mixture composition 393 evolved fully to $3 \cdot BF_4$ in less than 2 h upon slow heating to 60 394 °C, as confirmed by ³¹P{¹H} NMR spectra. Further heating to 395 80 °C did not result in further changes of the NMR patterns. A 396 similar experiment was carried out using 2 (0.01 mmol) and 397 NaHCO₃ (100 equiv) in CD₃OD (2 mL). The initial mixture 398 prepared under an Ar atmosphere showed in the corresponding 399 ${}^{31}P{}^{1}H$ NMR the presence of unreacted 2 (84%), 1′ (7%), and ${}_{400}$ 7 (9%). Upon standing at room temperature for 75 min, the 401 resonances observed for 1' and 7 increased significantly (up to 402 34% and 27%), by slow reaction of 2 with NaHCO₃. The slow 403 ligand exchange from CH₃CN to CO₃²⁻ mirrors the reactivity 404 of 2 with H₂ described above. By pressurization of the HPNMR 405 tube with H_2 (30 bar), the resonances due to 5 appeared in the 406 $^{31}P{^{1}H}$ and ^{1}H NMR spectra, already at room temperature. At $_{407}$ 80 °C, the signals of 2, 1', and 7 disappeared, with concomitant 408 formation of 3.BF4 and 5 and free sodium formate (broad 409 signals at 8.6–8.9 ppm in the ¹H NMR spectrum).²⁶ 410

The experimental results clearly indicate in $3 \cdot BF_4$ the key 411 intermediate in the catalytic hydrogenation of NaHCO₃ with 1 412 and 2, similarly to what was described by Beller and co-workers 413 in the case of CO₂ hydrogenation by $[Fe(H)(PP_3)]^{+,5a}$ Despite 414 the fact that we could not observe other catalytic intermediates 415 in addition to 3 under HPNMR conditions, an outer-sphere 416 mechanism involving intermolecular hydride transfer is 417 unlikely, as it would not account for the different catalytic 418 activities observed for *rac* and *meso* systems. In contrast, we 419 suggest that an inner-sphere mechanism requiring two available 420 cis positions would be more likely and consistent with the 421 better catalytic activity observed using 1. A proposed 422 mechanism for NaHCO₃ hydrogenation centered on 3 is 423 shown in Scheme 5.

Formic Acid Dehydrogenation. FA dehydrogenation to 425 H_2/CO_2 gas mixtures was tested in the presence of the in situ 426 and preformed catalysts described above, using an inert solvent 427 (PC) under isobaric conditions (1 atm) and in the absence of 428 added base, the development of gas during the reaction being 429 measured with a manual gas buret. The gas mixtures were 430 analyzed off-line by FT-IR spectroscopy, showing the absence 431 of CO for all tests (detection limit 0.02%).²⁷ Much to our 432 surprise, the well-defined catalyst precursor **2** (0.1 mol %) was 433

t2

Scheme 5. Proposed Mechanism for the Catalytic Hydrogenation of NaHCO₃ in the Presence of 3



434 inactive in the dehydrogenation of FA in PC at 40 °C. Thus, we 435 targeted the use of in situ catalysts formed using rac- and meso-436 P4. Initially, we checked the activity of commercial P4 (0.01 437 mol %, meso/rac = ca. 3) under the same conditions described 438 above and observed a FA conversion of 4% after 6 h, 439 corresponding to TON = 444 (Table 2; entry 1). When pure 440 rac-P4 was used, generating in situ catalyst 1 (0.1 mol %), FA 441 dehydrogenation proceeded with good conversions, reaching 442 TON = 604 after 8 h at 40 °C (entry 2). As reported for the 443 PP₃/Fe(BF₄)₂.6H₂O catalyst system, ^{Se} higher ligand/Fe ratios 444 are beneficial to reach high reaction turnovers. Using a Fe/rac-445 P4 = 1/2 ratio,^{5e} as expected the catalyst performance 446 improved significantly, affording full conversion of FA in ca. 6 447 h (TON = 1000; entry 3). Using a catalyst to substrate ratio of 448 1:10000 at 40 °C, low conversions (11%) were obtained after 6 449 h, with TON = 1081 (entry 5). Using the same catalyst to 450 substrate ratio at 60 °C gave a higher TON value of 3088 after 451 6 h (entry 6). Using a higher Fe to ligand ratio (1:4) at 60 °C, 452 considerably enhanced catalytic activity was achieved (TON = 453 6061, 6 h; entry 9). In contrast, precatalysts obtained from $454 \text{ Fe}(BF_4)_2 \cdot 6H_2O$ and meso-P4 showed worse catalytic activities 455 (generally ca. 33% lower) in comparison to rac-P4 (entries 4, 7, 456 8, and 10). Also in this case, the exclusive cis geometry forced 457 by rac-P4 is the most suitable to convey a catalytically active 458 species, in comparison to meso-P4, for which different 459 geometrical isomers are possible. Selected results are 460 summarized in Table 2. Selected reaction profiles (volumes

vs time) of catalytic runs obtained at a catalyst to substrate ratio $_{461}$ of 1:10000, at various Fe to ligand ratios and temperatures, are $_{462}$ shown in Figure 5. Disappointingly, recycling experiments with $_{463}$ fs catalyst:substrate = 1:1000, Fe:ligand = 1:2, and 40 °C showed $_{464}$ a severe drop in activity from the first to the third cycle, namely $_{465}$ from TON = 1000 to 295 after 6 h.

Mechanistic Studies. The reactivity of the different 467 precatalysts with FA was studied by monitoring stoichiometric 468 reactions by NMR and by HPNMR under in operando 469 conditions. A solution of complex 2 (0.7 mL, 0.012 M in PC, 470 C_6D_6 insert) was reacted with FA (1 equiv) for 1 h in a NMR 471 tube. No changes in the ³¹P{¹H} and ¹H NMR spectra were 472 observed even after heating to 60 °C, confirming that 2 is not 473 reactive under these conditions, probably due to stable 474 coordination of MeCN ligands to the Fe center.

In contrast, addition of 1 equiv of FA to a solution of 1 in PC 476 in a NMR tube (0.7 mL, 0.042 M, C₆D₆ insert) at room 477 temperature resulted in the formation of the monohydride 478 $[FeH(rac-P4)](BF_4)$ (3·BF₄) and of the formate complex 479 $[Fe(\eta^2-O_2CH)(rac-P4)](BF_4)$ (6·BF₄), initially in a 6:1 ratio. 480 Heating to 40 °C for 1 h and then leaving the tube overnight at 481 25 °C gave almost complete conversion to 3.BF4. The 482 experiment was repeated in the presence of a large excess of 483 FA (100 equiv), with catalyst evolution monitored by HPNMR 484 spectroscopy. A 10 mm HPNMR sapphire tube was thus 485 charged with a solution of 1 in PC (1.8 mL; 0.012 M), to which 486 CD₃OD (0.4 mL) was added for deuterium lock. Upon 487 addition of FA at room temperature, complexes 3.BF4 and 6. 488 BF4 were observed to form in a 1:6 ratio. The probe head was 489 then heated to 40 °C. After 1 h, the reaction mixture evolved 490 further with formation of a new species (9), characterized by 491 four structured signals in the ³¹P{¹H} NMR (see the 492 Experimental Section) and by a complex high-field resonance 493 signal (ddd; $\delta_{\rm H}$ –9.55 ppm, $^2J_{\rm PP}$ = 25.5, 46.5, 70.7 Hz; 1H, 494 FeH) in the corresponding ¹H NMR spectrum, indicative of 495 the formation of an octahedral [FeHL(rac-P4)] complex with 496 cis- α configuration. Prolonged heating resulted in complete 497 conversion to 9, affording a yellow solution. Further multi- 498 nuclear NMR analysis and ESI-MS spectroscopy data obtained 499 from aliquots of the final solution allowed us to identify 500 complex 9 as the Fe carbonyl hydrido complex cis- α - 501 $[FeH(CO)(rac-P4)](BF_4)$ (L = CO; for details see the 502 Experimental Section). 503

Table 2. Formic Acid Dehydrogenation Catalyzed using in Situ Fe(II) Precatalysts
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entry	ligand	substrate:catalyst	Fe/ligand	$T(^{\circ}C)$	$V_{1\mathrm{h}}~(\mathrm{mL})^d$	TON _{1 h} ^e	$\mathrm{TOF}_{10\mathrm{min}}^{f}$	$V_{\rm final}~({ m mL})^d$	$\mathrm{TON}_{\mathrm{final}}^{e}$	total conversn (%)
1^a	$P4^c$	10000	1:2	40	25	97	232	115	444 (6 h)	4
2 ^{<i>a</i>}	rac-P4	1000	1:1	40	220	85	35	1560	604 (8 h)	60
3 ^{<i>a</i>}	rac-P4	1000	1:2	40	345	133	139	2570	1000 (6 h)	100
4 ^{<i>a</i>}	meso-P4	1000	1:2	40	165	64	151	810	313 (8 h)	31
5^b	rac-P4	10000	1:2	40	45	174	347	280	1081 (6 h)	11
6^b	rac-P4	10000	1:2	60	215	830	1853	800	3088 (6 h)	31
7^{b}	meso-P4	10000	1:2	40	15	58	116	90	348 (6 h)	3
8^b	meso-P4	10000	1:2	60	70	270	579	260	1003 (6 h)	10
9^b	rac-P4	10000	1:4	60	400	1544	1737	1570	6061 (6 h)	61
10^{b}	meso-P4	10000	1:4	60	140	540	579	590	2278 (8 h)	23

^{*a*}Reaction conditions: $Fe(BF_4)_2 \cdot 6H_2O$, 5.3 mmol; ligand, 1–4 equiv with respect to Fe; HCOOH, 5.3 mol (2 mL); PC, 5 mL. ^{*b*}Reaction conditions as in footnote *a*, except for the following: $Fe(BF_4)_2 \cdot 6H_2O$, 5.3 μ mol. ^{*c*}Commercial tetraphos-1 (P4) ligand, *meso*-P4:*rac*-P4 = 3. ^{*d*}Gas evolution measured by manual gas buret, based on two to four tests, error ±10%. Gas mixture analyzed off-line by FTIR spectroscopy. ^{*c*}Defined as (mmol of gas produced)/((mmol of catalyst) h), calculated at conversions observed after 10 min.



Figure 5. Reaction profiles of selected FA dehydrogenation catalytic runs using a catalyst to substrate ration of 1:10000 at different temperatures and Fe:P4 ratios. For legends and conditions, see Table 2.

As CO may result from a competitive FA decomposition 504 pathway, i.e. dehydration to H₂O and CO, we thought it was of 505 interest to investigate further the reaction of 1 with FA. Some 506 hints were given from the experimental data described above. 507 First, CO was never detected in the gas mixtures resulting from 508 the catalytic runs by off-line FTIR measurements (see the 509 Supporting Information for a representative spectrum).²⁷ 510 becond, complex 9 was never obtained in the NMR experiment 511 arried out using an FA to 1 ratio of 1:1. Third, 9 was formed 512 under isochoric conditions (HPNMR) in the presence of 100 513 quiv of FA. Under these conditions, it is likely that the CO₂ 514 ressure built up in the HPNMR tube during the course of the 515 experiment may have undergone partial reductive disproportio-516 $_{517}$ nation to CO and CO₃²⁻, as previously observed upon prolonged reaction of the monohydride 3.BPh₄ with CO₂. 518

To confirm this hypothesis, we repeated the experiment in 519 520 the glass reactor (isobaric conditions) normally used for the catalytic runs. Under the same conditions applied for the 521 522 HPNMR experiment, gas evolution was complete after 20 min and again no CO was detected in the gas mixture. Furthermore, 523 the mixture remained purple throughout the run, whereas a 524 bright yellow should be expected upon formation of 9 in high 525 concentrations. As further confirmation, NMR analysis of the 52.6 catalytic mixture at the end of the run showed the typical 527 ${}^{1}P{}^{1}H{}$ NMR resonances of $3 \cdot BF_{4}$ and $6 \cdot BF_{4}$ in a 1:1 ratio, 528 while signals due to 9 were not observed. On the basis of these 529 data, although we cannot rule out that at low catalyst 530 concentrations (0.01 mol %) catalyst deactivation may occur 531 by formation of 9, we propose that in closed (isochoric) vessels 532 Fe-catalyzed CO₂ reductive disproportionation becomes a 533 competing pathway, and CO coordination to 3.BF4 gives the 534 stable (and catalytically inactive) octahedral 9. 535

The pathway for the base-free FA catalytic dehydrogenation 536 reaction is thus proposed as shown in Scheme 6. In step (i), the 537 catalyst precursor 1, formed in situ from $Fe(BF_4)_2 \cdot 6H_2O$ and 538 *rac*-P4, reacts with FA to give the formate complex [Fe(η^2 -539 O_2CH (rac-P4) (BF₄) (6·BF₄), which after a $\eta^2 \rightarrow \eta^1$ 540 coordination shift from 6 to 6' (ii) and rearrangement (iii) 541 undergoes β -hydride elimination to give back 3·BF₄ and CO₂ 542 (iv). Protonation of $3 \cdot BF_4$ by FA results in the fast elimination 543 544 of H₂ and regeneration of the formate complex $\mathbf{6}$ ·BF₄ (v).





CONCLUSIONS

In summary, the coordination chemistry of the *rac* and *meso* 546 isomers of the linear tetraphosphine 1,1,4,7,10,10-hexaphenyl- 547 1,4,7,10-tetraphosphadecane (tetraphos-1, P4) toward Fe(II) 548 was explored in detail, giving novel complexes which were 549 applied as catalysts for base-free H_2/CO_2 generation from 550 formic acid and for the hydrogenation of sodium bicarbonate to 551 formate under mild conditions, showing a higher activity in the 552 case of Fe/*rac*-P4 systems. Mechanistic studies highlighted the 553 pivotal role of the monohydride $[FeH(rac-P4)]^+$ in both 554 reactions and showed that CO_2 reductive disproportionation 555 should not be underestimated as a competing pathway in the 556 case of Fe(II)/polyphosphine systems. A full DFT study of 557 both catalytic reactions promoted by Fe/tetraphos-1 is 558 currently under way.

EXPERIMENTAL SECTION

General Methods and Materials. All syntheses were 561 performed using standard Schlenk techniques under an 562 atmosphere of dry nitrogen or argon. Solvents were freshly 563 distilled over appropriate drying agents, collected over Linde 564 type 3A or 4A molecular sieves under nitrogen, and degassed 565 with nitrogen or argon gas. The ligand 1,1,4,7,10,10- 566

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⁵⁶⁷ hexaphenyl-1,4,7,10-tetraphosphadecane (tetraphos-1, P4) was ⁵⁶⁸ supplied by Pressure Chemicals Inc., Pittsburgh, PA. ¹³C-⁵⁶⁹ labeled carbon dioxide (99 atom % ¹³C) was purchased from ⁵⁷⁰ Sigma-Aldrich. [Fe(MeCN)₆](BF₄)₂ was synthesized according ⁵⁷¹ to literature methods.¹⁷ Anhydrous FeCl₂, Fe(BF₄)₂·6H₂O, and ⁵⁷² propylene carbonate (PC) were purchased from commercial ⁵⁷³ suppliers and used without further purification.

Synthetic Procedures. Reaction of rac-P4 with $Fe(BF_{a})_{2}$. 574 575 6H2O. The ligand rac-P4 (67 mg, 0.1 mmol) was dissolved in 576 propylene carbonate (PC; 2.0 mL) with gentle heating (40-50 °C) to afford complete dissolution. One equivalent of 577 578 $Fe(BF_4)_2 \cdot 6H_2O$ (34 mg, 0.1 mmol) was added to the colorless solution, which immediately turned deep purple. ³¹P{¹H} NMR 579 analysis showed quantitative formation of a single product. The 580 purple product could be precipitated by adding a large amount 581 582 of Et₂O (at least 8.0 mL). The decanted solid was recovered by removing the colorless solution via cannula and washed with 583 584 Et₂O to remove all propylene carbonate, yielding the analytically pure complex $[Fe(\eta^1 - FBF_3)(rac - P4)](BF_4)$ (1). 585 Due to the poor stability of 1 as an isolated solid, we chose to 586 use stock solutions of 1 in PC for both catalytic and NMR 587 experiments. Yield: 78 mg (94%). ³¹P{¹H} NMR (121.49 MHz, 588 589 PC + C₆D₆ capillary): $\delta_{\rm p}$ 99.8 (br s; 2P, PPh), 60.9 ppm (br s; 2P, PPh₂). ¹⁹F{¹H} NMR (376.15 MHz, PC + C₆D₆ insert): δ_P 590 591 154 ppm (s; 4F, BF₄).

Reaction of meso-P4 with $Fe(BF_4)_2 \cdot 6H_2O$. In a 5 mm NMR 592 593 tube, rac-P4 (20 mg, 0.03 mmol) was dissolved in propylene carbonate (PC; 0.7 mL). Gentle heating (40-50 °C) was 594 595 needed to afford complete dissolution of the ligand. One s96 equivalent of $Fe(BF_4)_2 \cdot 6H_2O$ (10 mg, 0.03 mmol) was added to the colorless solution, which immediately turned brown and 597 598 then yellow. ³¹P{¹H} NMR analysis showed the formation of $[Fe(\eta^1-FBF_3)(meso-P4)](BF_4)$ (1") as a single product. No 599 600 attempts were made to isolate the product. ${}^{31}P{}^{1}H$ NMR (121.49 MHz, PC + C_6D_6 insert): δ_P 104.2 (br s; 2P, PPh), 601 70.5 ppm (br s; 2P, PPh₂). 602

Synthesis of cis- α -[Fe(MeCN)₂(rac-P4)](BF₄)₂ (**2**). The ligand 603 604 rac-P4 (134 mg, 0.2 mmol) was suspended in MeCN (10.0 605 mL), and the mixture was vigorously stirred until the 606 tetraphosphine turned into a thin powder. One equivalent of $[Fe(MeCN)_6](BF_4)_2$ (95 mg, 0.2 mmol) was added to the 607 white suspension, affording a bright orange solution. The 608 reaction mixture was stirred until a clear solution was obtained 609 610 and was subsequently stirred 1 h more. The solution was then 611 concentrated under vacuum to remove all volatiles. The 612 resulting orange solid was then dissolved in a minimum 613 volume of acetonitrile (ca. 0.5 mL). Addition of pentane 614 resulted in the precipitation of analytically pure 2 as a 615 crystalline, orange solid. Yield: 170 mg (87%). Crystals of 2 616 suitable for X-ray diffraction data collection were grown by 617 adding pentane (4.0 mL) to an acetonitrile/methanol solution (0.5 + 1.0 mL) of 2. ³¹P{¹H} NMR (121.49 MHz, CD₃CN): $\delta_{\rm P}$ 618 619 100.7 (t, ${}^{2}J_{PP}$ = 31.7 Hz; 2P, PPh), 65.6 ppm (t, ${}^{2}J_{PP}$ = 31.7 Hz; 2P, PPh₂). ESI-MS: calcd for ${}^{12}C_{46}{}^{1}H_{48}{}^{14}N_{2}{}^{56}Fe^{31}P_{4}$ ([M]⁺) m/ 620 621 z 404.10532, found m/z 404.10474.

Reaction of meso-P4 with $[Fe(MeCN)_6](BF_4)_2$. In an NMRc23 scale experiment, meso-P4 (13 mg, 0.02 mmol) was placed into c24 an NMR tube, to which 0.5 mL of CD₃CN was added. The c25 NMR tube was shaken vigorously to help dissolution of the c26 ligand, and subsequently $[Fe(MeCN)_6](BF_4)_2$ (ca. 10 mg, 0.02 c27 mmol) was added, resulting in an immediate color change to c28 red-orange. The reaction mixture was analyzed by ${}^{31}P{}^{1}H{}$ c29 NMR, which showed the formation of *trans*-[Fe-

Synthesis of [FeH(rac-P4)](BPh₄) (3·BPh₄). In a flame-dried 636 Schlenk tube kept under argon, rac-P4 (67 mg, 0.1 mmol) was 637 dissolved in 3.0 mL of THF. A stoichiometric amount of 638 anhydrous FeCl₂ (13 mg, 0.1 mmol) was added as a solid, and 639 the resulting deep blue solution was stirred for 5 min at room 640 temperature. NaBPh₄ (35 mg; 0.01 mmol) and MeOH (1.5 641 mL) were added to the reaction mixture, which was then stirred 642 vigorously for about 10 min. NaBH4 (4 mg, 0.1 mmol) was 643 then added to the reaction mixture as a solid, and a vigorous 644 reaction took place, affording an intense red mixture. All 645 volatiles were removed under vacuum, and the solid residue was 646 redissolved in THF (8.0 mL). The resulting suspension was 647 filtered via cannula into a second Schlenk tube kept under 648 argon, affording a limpid red solution, from which all volatiles 649 were removed under vacuum, affording NMR-pure 3.BPh4. 650 Yield: 103 mg (95%). Crystals suitable for X-ray diffraction data 651 collection were obtained by adding MeOH to a THF solution 652 of 3·BPh₄. ³¹P{¹H} NMR (d_8 -THF, 121.49 MHz): δ_P 119.4 (t, 653 ${}^{2}J_{PP} = 24.5$ Hz; 2P, PPh), 99.4 (t, ${}^{2}J_{PP} = 24.5$ Hz; 2P, PPh₂). ¹H ₆₅₄ NMR (d_8 -THF, 300.13 MHz, negative region): $\delta_{\rm H}$ –9.16 (t, 655 ${}^{2}J_{\rm HP} = 24.0$ Hz; 1H, FeH).

Synthesis of cis- α -[Fe(H)₂(rac-P4)] (4). The synthetic 657 procedure described for the synthesis of trans-[Fe(H)₂(meso- 658 P4)] was adapted with slight modifications.¹⁴ A three-necked 659 round-bottom flask equipped with a reflux condenser was 660 charged under argon with rac-P4 (67 mg, 0.1 mmol) and dry 661 THF (2.5 mL). A solution of anhydrous FeCl₂ (13 mg, 0.1 662 mmol) in THF (2.5 mL) was added via cannula, and the 663 resulting mixture was stirred for 10 min. NaBH₄ (20 mg, 0.55 664 mmol) was added as a solid, and the dark blue reaction mixture 665 that was obtained was heated to reflux. As no visible changes 666 occurred, additional THF (3.0 mL) was added, followed by 667 another aliquot of NaBH₄ (10 mg, 0.27 mmol) and absolute 668 EtOH (0.5 mL). As EtOH was added, a vigorous reaction took 669 place and the deep blue mixture turned orange. After gas 670 evolution had ceased, additional NaBH₄ (10 mg, 0.27 mmol) 671 and absolute EtOH (0.5 mL) were added, and again, gas 672 evolution was observed. The orange mixture was refluxed for 673 about 10 min after gas evolution had ceased, after which it was 674 cooled to room temperature and filtered via cannula. The 675 volume of the solution was partially reduced under vacuum, and 676 dry methanol was subsequently layered on top of the orange 677 solution, from which bright yellow crystals formed. Yield: 53 678 mg (72%). ³¹P{¹H} NMR (d_8 -THF, 121.49 MHz): δ_P 123.8 (t, 679 ${}^{2}J_{\rm PP}$ = 13.5 Hz; 2P, PPh), 113.1 (t, ${}^{2}J_{\rm PP}$ = 13.5; 2P, PPh₂). ${}^{1}H_{680}$ NMR (d_8 -THF, 300.13 MHz, negative region): $\delta_{\rm H}$ –11.7 (m; 681 2H, Fe(H)₂). ESI-MS: calcd for ${}^{12}C_{42}{}^{1}H_{43}{}^{56}Fe^{31}P_4$ ([M – H]⁺) 682 m/z 727.16592, found m/z 727.16523.

Reaction of **3**·BPh₄ *with* CO₂. A few crystals of **3**·BPh₄ (ca. 684 10 mg) were placed in an NMR tube under argon and dissolved 685 in d_8 -THF (0.5 mL). CO₂ (1 atm) was then bubbled through 686 the solution, which then turned light purple. NMR analysis 687 revealed quantitative formation of the expected formate 688 complex [Fe(η^2 -OCHO)(*rac*-P4)](BPh₄) (**6**·BPh₄). ³¹P{¹H} 689 NMR for **6**·BPh₄ (d_8 -THF, 161.99 MHz): δ_P 106.0 (t, ² J_{PP} = 690 28.5 Hz; 2P, PPh), 76.5 (t, ² J_{PP} = 26.1 Hz; 2P, PPh₂). ¹³C{¹H} 691 ⁶⁹² NMR for 6·BPh₄ (d_8 -THF, 100.6 MHz): δ_C 162.4 (dd; ² J_{PP} = ⁶⁹³ 49.4 Hz, ² J_{PP} = 98.7 Hz, BPh₄), 174.6 (br s, Fe(O₂CH)).

After 24 h acquisition, the ³¹P{¹H} NMR spectrum revealed 695 the formation of the carbonate complex [Fe(η^2 -O₂CO)(*rac*-696 P4)] (7). On the basis of ³¹P{¹H} NMR integration, complexes 697 **6** and 7 resulted in an approximately 1:0.6 ratio. ³¹P{¹H} NMR 698 for 7 (d_8 -THF, 161.99 MHz): δ_P 106.6 (t, ² J_{PP} = 30.4 Hz; 2P, 699 PPh), 73.2 (t, ² J_{PP} = 30.4 Hz; 2P, PPh₂). No ¹³C{¹H} NMR 700 resonance was observed for the carbonate O₂CO carbon atom 701 of 7. The experiment was repeated using **3**·BF₄ and ¹³CO₂, 702 showing the same ³¹P{¹H} NMR and ¹³C NMR (proton 703 coupled) signals at 174.6 (d, ¹ J_{CH} = 208.8 Hz) and 158.1 ppm 704 (s) for **6** and 7, respectively.

⁷⁰⁵ A few purple crystals suitable for X-ray diffraction data ⁷⁰⁶ collection were obtained by layering MeOH on top of the d_{8} -⁷⁰⁷ THF solution and standing for 1 day. The X-ray crystal ⁷⁰⁸ structure revealed the serendipitous formation of the trimetallic ⁷⁰⁹ complex { μ_2 -[Fe(MeOH)₄]- κ^1 O-[Fe(η^2 -O₂CO)(*rac*-P4)]_2}-⁷¹⁰ (BPh₄)₂ (7').

Reaction of 1 with K_2CO_3 . A 0.5 mL portion of a 0.01 M 711 712 stock solution of 1 in PC were placed in a 5 mm NMR tube 713 under argon. Solid K₂CO₃ (7.0 mg, 0.05 mmol) was then 714 added. The solution in the NMR tube was stirred with a small 715 stirring bar, and the purple solution turned initially bright pink 716 and then bright red. d_8 -Toluene (0.2 mL) was added for deuterium lock, and the red solution was analyzed by $^{31}\text{P}\{^1\text{H}\}$ 717 NMR and ¹³C{¹H} NMR. ³¹P{¹H} NMR analysis showed 718 719 formation of carbonate complex 7, whereas no ${}^{13}\dot{C}{}^{1}H$ NMR 720 resonance was observed for the carboxylic O₂CO carbon of 7. ³¹P{¹H} NMR for 7 (PC + d_8 -toluene, 121.49 MHz): δ_P 105.1 721 722 (t, ${}^{2}J_{PP}$ = 30.4 Hz; 2P, PPh), 69.0 (t, ${}^{2}J_{PP}$ = 30.4 Hz; 2P, PPh₂). Reaction of 1 with FA under HPNMR Conditions and 723 724 Formation of cis- α -[FeH(CO)(rac-P4)](BF₄) (9). A 10 mm 725 HPNMR sapphire tube was charged with a solution of 726 Fe(BF₄)₂·6H₂O (14 mg; 0.04 mmol) and rac-P4 (28 mg; 727 0.04 mmol) in propylene carbonate (1.8 mL) under argon. 728 CD₃OD (0.4 mL) was then added for deuterium lock, followed 729 by HCOOH (0.15 mL, 4.15 mmol; 100 equiv with respect to 730 Fe). The tube was closed and placed in the NMR probe. The 731 probe head was gradually heated to 60 °C, and the reaction was 732 monitored by ${}^{31}P{}^{1}H{}$ NMR (see the Supporting Information). 733 The tube was left at 60 °C overnight, resulting in a yellow 734 solution. ³¹P{¹H} and ¹H NMR analysis revealed the 735 quantitative formation of $cis-\alpha$ -[Fe(H)(CO)(rac-P4)](BF₄) 736 (9). ³¹P{¹H} NMR (121.49 MHz, CD₃OD): δ 114.6 (dt, ²J_{PP} $_{737} = 23.5, {}^{2}J_{PP} = 38.6, 1P), 105.1 \text{ (br dd, } {}^{2}J_{PP} = 8.6, {}^{2}J_{PP} = 21.9;$ 738 1P), 100.9 (ddd, ${}^{2}J_{PP} = 10.5$, ${}^{2}J_{PP} = 39.3$, ${}^{2}J_{PP} = 68.7$; 1P), 92.3 $(dd, {}^{2}J_{PP} = 37.9, {}^{2}J_{PP} = 68.5; 1P).$ ¹H NMR (300.13 MHz, 739 CD₃OD, negative region): δ –9.6 (ddd, ²*J*_{PP} = 25.5, ²*J*_{PP} = 46.5, 740 ${}^{2}J_{PP} = 70.7$; 1H, FeH). ${}^{13}C{}^{1}H{}$ NMR (75.47 MHz, CD₃OD, 741 742 carbonyl region): δ 162.77 (s; CO). A sharp singlet of higher 743 intensity was also observed at δ 162.1 ppm for HCOOH. ESI-744 MS: calcd for ${}^{12}C_{43}{}^{1}H_{43}{}^{56}Fe^{16}O^{31}P_4$ ([M]+) m/z 753.16550, 745 found *m*/*z* 753.16517.

746 **Catalytic Tests.** *Catalytic Sodium Bicarbonate Hydro*-747 *genation Tests.* In a typical experiment, a 40 mL magnetically 748 stirred stainless steel autoclave built at CNR-ICCOM was 749 charged under an inert atmosphere (glovebox) with NaHCO₃ 750 (typically 840 mg, 10 mmol) and the catalyst (0.01–0.001 751 mmol as solid or stock solution in PC). The autoclave was then 752 closed and thoroughly purged through several vacuum/argon 753 cycles. MeOH (20.0 mL) was then added to the autoclave by 754 suction. Finally the autoclave was pressurized with H₂ gas at the desired pressure. The autoclave was then placed into an oil bath 755 preheated to the desired temperature and stirred for the set 756 reaction time. After the run, the autoclave was cooled in an ice/ 757 water bath and depressurized, and the catalytic mixture was 758 transferred to a flask and concentrated under vacuum at room 759 temperature. The formate content was determined by analyzing 760 aliquots (ca. 30 mg) of the solid mixture dissolved in D_2O (0.5 761 mL) by ¹H NMR, using dry THF (20 μ L) as internal standard 762 with a relaxation delay of 20 s.

Catalytic Formic Acid Dehydrogenation Tests. In a typical 764 experiment, a solution of catalyst (typically 5.3 mmol) in 765 propylene carbonate (5 mL) was placed under an argon 766 atmosphere in a magnetically stirred glass reaction vessel 767 thermostated by external liquid circulation and connected to a 768 reflux condenser and gas buret (2 mL scale). After the solution 769 was heated to 40–60 °C, HCOOH (2.0 mL) was added and 770 the experiment started. The gas evolution was monitored 771 throughout the experiment by reading the values reached on 772 the burets. The gas mixture was analyzed off-line by FTIR 773 spectroscopy using a 10 cm gas-phase cell (KBr windows) to 774 check for CO formation (detection limit 0.02%).

ASSOCIATED CONTENT

776 777

S Supporting Information

The following files are available free of charge on the ACS 778 Publications website at DOI: 10.1021/cs501998t. 779

General methods and equipment, NMR spectra and 780 details of HPNMR experiments, information on FA 781 dehydrogenation tests, and details of the X-ray structure 782 determinations (PDF) 783 Crystallographic data for the X-ray crystal structure of 2 784 (CIF) 785 Crystallographic data for the X-ray crystal structure of 3. 786 BPh_4 (CIF) 787 Crystallographic data for the X-ray crystal structure of 4 788 (\underline{CIF}) 789 Crystallographic data for the X-ray crystal structure of 5 790 (CIF) 791 Crystallographic data for the X-ray crystal structure of 7' 792 (CIF) 793 AUTHOR INFORMATION 794

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Notes	797
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