Peripheral mechanism of a carbonyl hydrosilylation catalysed by an SiNSi iron pincer complex†

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Combined experimental and theoretical analysis of the carbonyl hydrosilylation catalysed by an iron(0) pincer complex reveals an unprecedented mechanism of action. The iron(0) complex is in fact a precatalyst that is converted into an iron(II) catalyst through oxidative addition of a hydrosilane. Neither the hydrogen atom nor the silicon atom bound to the iron site are subsequently transferred onto the carbonyl acceptor, instead remaining at the sterically inaccessible iron atom throughout the catalytic cycle. A series of labelling, crossover and competition experiments as well as the use of a silicon-stereogenic hydrosilane as a stereochemical probe suggest that the iron site is not directly involved in the hydrosilylation. Strikingly, it is the silyl ligand attached to the iron atom that acts as a Lewis acid for carbonyl activation in this catalysis. The whole catalytic process occurs on the periphery of the transition metal. Computation of the new peripheral as well as plausible alternative inner and outer sphere mechanisms support the experimental findings.

Introduction

Iron-catalysed carbonyl hydrosilylation can be traced back to seminal reports by Brunner1 but these contributions were decades ahead of their time. With few exceptions, catalyst development was focused on complexes of rare transition metals while limited progress had been made involving iron-based catalysts for carbonyl hydrosilylation with different ligand designs were introduced in recent years.2 The advent of these new catalysts immediately poses the question whether the mechanisms are similar to those established for rare transition-metal complexes or totally unprecedented.3 However, little detail is known about the mechanisms of action of iron complexes in catalytic hydrosilylation.

Mechanisms of transition-metal-catalysed hydrosilylations exhibit a wide variety of modes of activation.4 However, the known mechanisms are characterised as either inner sphere5,6 or outer sphere7,8 where both the substrate and the hydrosilane are directly in contact with the metal or outer sphere9,8 where only one of the two is in contact with the metal centre. The proposed mechanisms for iron-catalysed hydrosilylations range from inner sphere mechanisms with σ-bond-metathesis-type Si–H bond cleavage at an iron–oxygen bond10,11 to outer sphere mechanisms with iron acting as a Lewis acid,12 either activating the hydrosilane or the carbonyl group.

Driess and co-workers recently introduced silylenes as σ-donor ligands in iron-based catalysis, and iron(0) complexes 1 and 2 (Fig. 1) were applied to carbonyl hydrosilylation.9 Cooperativity between the iron(0) atom and the silicon atom in the hydrosilane 1 was postulated to be relevant in the catalytic cycle.6 The SiNSi iron(0) pincer complex 2 was, in turn, believed to be a pre-catalyst10 but a detailed mechanistic analysis remained challenging. We report here the disclosure of a unique mechanism of a transition-metal-catalysed carbonyl hydrosilylation that takes place neither inner nor outer sphere but on the periphery of the metal centre without its direct involvement.

Fig. 1 Iron(0) complexes 1 (ref. 9a) and 2 (ref. 9b) introduced by Driess and co-workers.
Results and discussion

Optimisation and scope

The SiNSi pincer complex 2 (2.5 mol%) was found to catalyse the hydrosilylation of various acetophenones 3a–3i with silane 4a at elevated temperatures11 (Table 1, entries 1–9; see the ESI† for the optimisation of the reaction conditions). Both electron-donating (entries 1 and 3) as well as -withdrawing (entry 6) substituents at the aryl group were tolerated with the exception of a Et$_2$N group in the para position (entry 2). The reaction was, however, sensitive toward steric hindrance. Substituents in the ortho position significantly lowered the yield (entries 7–9), and a 2,6-disubstituted substrate did not react (3j, entry 10). Benzo-phenone (3k, entry 11) reacted readily while propiophenone and isobutyrophenone (3l and 3m, entries 12 and 13) afforded 18 and 16% yield, respectively. Hydrosilylation of cyclopropyl substituted ketone (3n, entry 14) proceeded efficiently12 (see the ESI† for full scope).

Isolation of the active catalyst

To gain insight into the mechanism, we investigated the reaction between iron(0) complex 2 and hydrosilanes 4a–c (2 → 7a–c, Scheme 1). Heating at 70 °C, a new set of distinct signals appeared in the $^1$H as well as in the $^{29}$Si and $^{31}$P NMR spectra. The $^1$H NMR spectrum clearly indicated the formation of an iron hydride, and detailed NMR analysis revealed that the hydride was likely to be trans to the apical phosphine ligand. The silyl group was assigned by 2D NMR experiments to be in the equatorial position trans to the pyridine ligand. We then obtained single crystals of 7b (Si = Me$_3$P) suitable for X-ray diffraction analysis, and that confirmed the molecular structure deduced from the NMR analysis. The structure shows a distorted octahedral iron(ii) coordination environment. The hydride was located tilted toward one of the silylene donor arms deviated from the trans coordination to the Me$_3$P ligand (P–Fe–H$_1$ 170.20(5)°), a situation similar to that of known iron(ii) hydride pincer complexes.13 The Fe–Si bond distances 2.1509(7)/2.1715(7) Å for Fe–Si$_1$/Si$_2$ and 2.2986(8) Å for Fe–Si$_3$ are within the range of iron silylene and silyl complexes.14,15 The t-Bu groups encage the iron hydride with a Si$_1$–Fe–Si$_2$ angle far from the linearity, 144.54(3)°.

To validate whether the thus formed iron(ii) complex 7 is the active catalyst, we measured the kinetic profiles for the hydrosilylation of 3a with hydrosilane 4a catalysed by 2 or 7a (Scheme 2). Conversion with 2 was only 15% after 1 h while the reaction had reached 74% with 7a. The reaction with 7a continued with significantly higher rate reaching 86% at 4 h compared to only 53% with 2. After 22 h, nearly full conversion

<table>
<thead>
<tr>
<th>Entry</th>
<th>$R_1$</th>
<th>$R_2$</th>
<th>Yield of $^5$ (%)</th>
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<tr>
<td>1</td>
<td>3a</td>
<td>X = MeO</td>
<td>Me</td>
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<tr>
<td>2</td>
<td>3b</td>
<td>X = Et$_2$N</td>
<td>Me</td>
</tr>
<tr>
<td>3</td>
<td>3c</td>
<td>X = Me</td>
<td>Me</td>
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<tr>
<td>4</td>
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<td>Me</td>
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<td>5</td>
<td>3e</td>
<td>X = H</td>
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<td>3f</td>
<td>X = CF$_3$</td>
<td>Me</td>
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<tr>
<td>7</td>
<td>3g</td>
<td>X = MeO</td>
<td>Me</td>
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<tr>
<td>8</td>
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<td>X = Me</td>
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<td>9</td>
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<td>X = Cl</td>
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<tr>
<td>14</td>
<td>3n</td>
<td>c-Pr</td>
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Reactions were performed on 0.10 mmol scale employing precatalyst 2 (2.5 mol%) and (EtO)$_2$SiH (4a 1.5 equiv). Average yield from two runs determined by GLC-MS analysis and $^1$H NMR spectroscopy using anisole as internal standard.

Scheme 1. Identification of the catalytically active iron(ii) complexes 7 from iron(0) precatalyst 2 and molecular structure of 7b. Hydrogen atoms except for the iron hydride are omitted for clarity.

Scheme 2. Kinetic profiles of the iron(0) and iron(ii) complexes 2 and 7a in hydrosilylation.
is obtained for both. The greater initial rate of the catalysis with 7a strongly supports the assignment of the iron(II) complex 7 as the active catalyst.

**Stoichiometric experiments: hydride transfer**

With the iron(II) hydrides 7 in hand, we had a closer look at the hydride transfer. Maintaining 7b and deuterium-labelled hydrosilane 4b-d₁ in THF at 70 °C resulted in slow H/D exchange, visible both at the silicon and iron atoms (Scheme 3, left). The stoichiometric reaction between iron(II) hydride 7b, hydrosilane 4b-d₁ (ca. 95% deuteration grade), and ketone 3e was puzzling though (Scheme 3, right). Initially, the H : D ratio at the methine position of silyl ether 8eb is nearly 50 : 50. However, it quickly decreases to 36 : 64 at 25% conversion within 6 hours and then gradually increases again, returning to 50 : 50 at full conversion after a few days. Meanwhile, the corresponding reaction with partially deuterated 4b-d₁ (ca. 50% deuteration grade) yielded 8eb with little deuterium incorporation at 19% conversion (H/D = 90 : 10). That ratio subsequently decreases to 78 : 22 to reach equilibrium after 24 hours. These results reveal that even though the hydride at the silicon atom in 4 is exchanging with the iron-bound hydride in 7, hydride transfer to the carbonyl carbon atom of 3 most likely occurs from the hydrosilane 4 and not from complex 7. Also, the reaction with partially deuterated hydrosilane indicates that the kinetic isotope effect (KIE) of the hydride transfer is significant. Precise value of the KIE could not be measured due to competing H/D exchanges (vide infra).

The possible H/D exchange at the methine position of silyl ether 8eb was verified using 8eb-d₁ (Scheme 4, top). Treatment of 8eb-d₁ with equimolar amounts of the iron(II) hydride 7b indeed led to H/D scrambling. Conversely, no erosion of the enantiomeric purity was seen when subjecting enantiopure silyl ether (S)-8eb to the typical protocol (precatalyst 2 and hydrosilane 4b generate catalyst 7b, Scheme 4, bottom). The configurational stability of (S)-8eb suggests that the hydride transfer itself is irreversible, and a concerted mechanism involving frontside attack at the asymmetrically substituted carbon atom is needed to explain the hydrogen atom exchange between the catalyst and the product.

These unusual scramblings were then investigated by DFT calculations. Both were found to proceed via a silylene-assisted concerted mechanism (9a⁺ for Si–H and 10a⁺ for C–H, Scheme 5) where the hydride on the iron atom is first shifted to the silicon atom of the adjacent donor-stabilised silylene ligand forming a pentacoordinate silicon atom while the Si–H or C–H bond interact with the now accessible iron centre. Both transition states are paired with their corresponding isomer where the second silylene ligand accepts the hydride. Attempts to locate the transient intermediates between the two degenerate conformations were not successful.

After the transition state, the silylene-bound hydrogen atom migrates back to the silicon and carbon atom, respectively. The activation barriers of the scrambling reactions (18.5 kcal mol⁻¹ for Si–H and 20.1 kcal mol⁻¹ for C–H) are energetically accessible under the reaction conditions.

**Stoichiometric experiments: phosphine dissociation**

To probe the potential lability of the phosphine, we performed phosphine crossover experiments. Slow exchange of the phosphine ligand with (CD₃)₃P(6-d₆) was observed (Scheme 6, 9a⁺, left) and the methine carbon atom (10a⁺, right); Gibbs free energies given in parentheses in kcal mol⁻¹; Si = Si(OEt)₂.
When 7b was subjected to repeated reflux/freeze-pump–thaw cycles, we detected a new iron hydride species that was tentatively assigned as the expected phosphine-dissociated iron silyl hydride 11b–d6, with C6D6 as an 11b–d6 ligand (Scheme 6, top right). The generation of this new iron compound was accompanied by formation of disilane 12b and, with longer reaction times, decomposition into a complex mixture. The role of the disilane 12b in the formation of the complex 11b remains unexplained. Unfortunately, attempts to isolate 11b–d6 were unsuccessful. Its generation in the presence of acephenethone (3e) did not lead to the formation of the silyl ether 8eb (see the ESI† for details), providing further evidence against the role of 11 as an intermediate in the catalytic reaction. In fact, the dissociation of the Me3P (6) is significantly slower in the presence of ketone 3e. The formation 11 was also investigated computationally (Scheme 6, bottom). Phosphine dissociation from 7a gives energetically highly unfavoured intermediate cis-14a that readily coordinates benzene to form adduct 11a.

It must be noted here that catalysis with Guan’s related iron(n) POCOP-pincer complex is thwarted by additional Me3P (6), indicating dissociation of one of the phosphine ligands as part of the catalytic cycle.38 When we added 25 mol% of Me3P (6, 10 equiv./catalyst) to the reaction mixture, the reaction was unaffected (Scheme 7, cf. Table 1, entry 5).

As is to be expected from the above observations, the hydride complex 7b did not produce any silyl ether 8eb when reacted stoichiometrically with ketone 3e in the absence of a hydrosilane (Scheme 8, top left). What was fascinating though is that the silyl ligand in 7b also remains untouched throughout the catalysis: 7b derived from Me2PhSiH (4b) catalyses the hydroisilylation of 3e with MePh2SiH (4c) with hardly any incorporation of the Me3PhSi moiety into the product; silyl ether 8ec rather than 8eb is formed almost exclusively (Scheme 8, right). This crossover experiment unequivocally proves that iron(n) complexes 7 are the actual catalysts, originating from oxidative addition of hydrosilanes 3 to the iron(0) complex 2; 2 is a pre-catalyst. During the crossover experiment no changes in the characteristic signals of complex 7b in the 1H and 31P NMR spectra were detected. However, when the assumed inability of 7b to exchange their silyl groups was examined with another control experiment (Scheme 8, bottom left), we observed slow exchange with ca. 36% conversion of 7b to 7c after 24 h. The Me2PhSi/MePh3Si scrambling was accompanied with formation of phosphate-dissociated, benzene-stabilised compounds 11b and 11c. Only traces of Me2PhSiH (4b) were observed, indicating that the exchange (7b to 7c) is in fact a side product of the decomposition rather than simple scrambling of the silyl groups.

Hydroisilylation with a silicon-stereogenic hydrosilane

With sufficient knowledge of the active catalyst, we decided to analyse the stereochemical course at the silicon atom of the reacting hydrosilane (Scheme 9).18 Catalyst 7b promoted the reaction between highly enantioenriched hydrosilane (9S)-4d (e.r. > 95 : 5) and ketone 3e but conversion was slow as expected from the data obtained with achiral triorganoisilane 4b (see Table S1, entry 12 in the ESI†). After 6 days, we were able to

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Scheme 6 Phosphine scrambling and dissociation. Double-ended arrows in 11b–d6 show H, 29Si HMQC NMR correlation. Gibbs free energies given in parentheses in kcal mol⁻¹ [Si = SiMe2Ph].

Scheme 7 Effect of excess Me3P on the catalytic activity.

Scheme 8 Stoichiometric control and crossover experiments.
isolate the silyl ether \((\text{Si} \ R)\text{-8ed}\) in 31% yield; diastereoselection was poor. The enantiomeric ratio of unreacted \((\text{Si} \ S)\text{-4d}\) was found to be unaffected. Subsequent reductive cleavage of the Si–O bond in \((\text{Si} \ R)\text{-8ed}\) (known to proceed with stereoretention at silicon atom\(^{19}\)) liberated \((\text{Si} \ S)\text{-4d}\) with overall retention of the stereochemistry at the silicon atom (e.r. > 95 : 5). Hence, the hydrosilylation step involves frontside attack at the silicon atom, and that makes a mechanism involving Lewis-acid activation of the hydrosilane unlikely.\(^{18}\)

DFT calculation of conventional inner and outer sphere mechanisms\(^{23}\)

Based on the combined findings, we propose that the active iron(II) catalyst 7 is generated from the iron(0) precatalyst 2 by oxidative addition of hydrosilane 4 to the zero-valent iron atom (Scheme 1). As expected due to the steric congestion around the iron(II) centre in 7a (grey box), we did not locate any structure resulting from direct insertion of the ketone C=O group into the iron hydride 7a or the silylene ligands (not shown). Instead, we were able to find a minimum structure for the phosphine-dissociated complex \(\text{cis-14a}\) (Scheme 10, left). In agreement with the experiments, \(\text{cis-14a}\) is however significantly higher in energy (29.2 kcal mol\(^{-1}\) relative to 7a). The intermediate \(\text{cis-14a}\) readily coordinates THF forming the adduct 17a. This intermediate is however a resting state if not a “mechanistic dead-end”. Ketone coordination to the iron centre of \(\text{cis-14a}\) gives intermediate 19oa with activated carbonyl group. The catalytic cycle is closed by an outer sphere concerted hydrosilane addition 20oa\(^{2}\) to the ketone with an activation barrier of 33.7 kcal mol\(^{-1}\).

Isomerisation of \(\text{cis-14a}\) to \(\text{trans-14a}\) was found to be strongly disfavoured, but again, ketone coordination to \(\text{trans-14a}\) lowered the energy significantly (Scheme 10, right). The following hydride transfer from 23oa passes through 24oa\(^{\ddagger}\) (30.9 kcal mol\(^{-1}\)) to afford the iron alkoxide 25oa. The silylated alcohol is released by an inner sphere silylation through 26oa\(^{2}\) with an energy barrier of 34.9 kcal mol\(^{-1}\). An alternative inner sphere mechanism could be a reductive elimination from the intermediate 25oa. However, the energy barrier for the transition state 27oa\(^{2}\) was found to be high, and the resulting iron(0) complex 28 is energetically disfavoured. Recoordination of phosphine 6 gives iron(0) complex 29 which oxidatively adds to a silane 4a to form 7a.

Peripheral mechanism: support from DFT calculations

In addition to the high energy barriers, neither outer nor inner sphere mechanisms give satisfactory fits to the experimental evidence. Hence, we looked for an adduct of 7a and 3o wherein
the silyl group would act as a Lewis acid (Scheme 11). Coordination of the ketone to the silyl group via low energy transition state $300a^\ddagger$ (8.1 kcal mol$^{-1}$) led to intermediate $310a$ (2.8 kcal mol$^{-1}$) with a pentacoordinate silicon atom, and the C=O double bond being significantly elongated compared to its equilibrium distance from 1.211 to 1.251 Å, indicating activation. Lewis pair formation is followed by coordination of hydrosilane $4a$ to the carbonyl group in $310a$, and the hydrosilylation event releases $80a$ through transition state $320a^\ddagger$ with retention at the silicon atom. In accordance with our labelling experiments (cf. Scheme 3), this is the rate-determining step (14.3 kcal mol$^{-1}$). To further validate this, we conducted a competition experiment between electron-rich $3a$ and electron-deficient $3f$ (Scheme 12). The $para$ substitution in $3$ exerts a pronounced electronic effect, and $F_3C$-substituted $3f$ was consumed significantly faster than MeO-substituted $3a$. This reactivity pattern is not unprecedented, and it has been seen previously in the activation of carbonyl compounds with silicon-based Lewis acids. The carbonyl carbon atom in $3f$ ($X = CF_3$) is more positively polarised accelerating the hydride transfer, than that of donor-substituted $3a$ ($X = \text{OMe}$). The reactivity is also in agreement with the proposed KIE based on the control reactions with deuterated silane $4b$-$d$ (Scheme 3).

**Conclusions**

The value of the present study is that it demonstrates, to our knowledge for the first time, an unusual case where the transition metal of a catalyst complex is not directly involved in the catalytic process. Activation of both substrate and reagent as well as the bond-forming and -breaking events happen in the ligand sphere, i.e., on the periphery of the transition metal. Coincidentally, the mechanism becomes outer sphere at silicon. The more conventional inner or outer sphere mechanisms do not apply to this unique catalyst.

**Acknowledgements**

This research was supported by the Cluster of Excellence UniCat of the Deutsche Forschungsgemeinschaft (EXC 314/2). T.S. is grateful to The New Széchenyi Plan TAMOP-4.2.2/B-10/1-2010-0009. M.O. is indebted to the Einstein Foundation Berlin for an endowed professorship.

**Notes and references**

6. For selected examples of inner sphere hydrosilylation, see: (a) I. Ojima, M. Nihonyanagi, T. Kogure, M. Kumagai,
For reviews of the related outer sphere


The complex cis-14a was calculated to be diamagnetic with vacant centre on the iron. The singlet-triplet gap was calculated to be only 5.1 kcal mol⁻¹.

The adduct 31 was not observed by the ¹H or ³¹P NMR spectroscopy, even at lower temperatures. Also attempts to perform a Gutmann–Beckett analysis with Et₃PO led to only slight broadening of the ³¹P NMR resonance of the phosphine oxide.

(a) K. Müther and M. Oestreich, Chem. Commun., 2011, 47, 334–336; (b) Ref. 20.