



Woof, C. R., Durand, D. J., Fey, N., Richards, E., & Webster, R. L. (2021). Iron Catalyzed Double Bond Isomerization: Evidence for an Fel/FeIII Catalytic Cycle. *Chemistry - A European Journal*, *27*(19), 5972-5977. https://doi.org/10.1002/chem.202004980

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Catalysis

Iron Catalyzed Double Bond Isomerization: Evidence for an Fe^I/Fe^{III} Catalytic Cycle

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Abstract: Iron-catalyzed isomerization of alkenes is reported using an iron(II) β -diketiminate pre-catalyst. The reaction proceeds with a catalytic amount of a hydride source, such as pinacol borane (HBpin) or ammonia borane (H₃N·BH₃). Reactivity with both allyl arenes and aliphatic alkenes has been studied. The catalytic mechanism was investigated by a variety of means, including deuteration studies, Density Func-

Introduction

Two-electron chemistry within the field of iron catalysis is incredibly rare.^[1] We envisioned that if a controlled one electron reduction of an Fe(II) species could be carried out with a welldefined iron pre-catalyst, then it might be feasible to access an oxidative addition (OA)/reductive elimination (RE) (i.e. Fe^I/Fe^{III}) catalytic cycle to afford activity similar to that typically observed with precious metal congeners. Previous catalytic studies within our group using an iron(II) β -diketiminate complex (hereafter referred to as complex 1) have focused on bond forming transformations (for example hydrophosphination, hydrogenation and dehydrocoupling, Scheme 1 a). However, double bond isomerization gives the ideal opportunity to explore a unimolecular transformation that relies on a 1,3-hydro-

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U	https://doi.org/10.1002/chem.202004980
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© 2021 The Authors. Chemistry - A European Journal published by Wiley-VCH GmbH. This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited. tional Theory (DFT) and Electron Paramagnetic Resonance (EPR) spectroscopy. The data obtained support a pre-catalyst activation step that gives access to an η^2 -coordinated alkene Fe^I complex, followed by oxidative addition of the alkene to give an Fe^{III} intermediate, which then undergoes reductive elimination to allow release of the isomerization product.

gen shift,^[2] the simplicity of which may give increased likelihood of OA/RE chemistry (Scheme 1 b). Carbon–carbon double bonds are important in synthetic chemistry: they are used to introduce diverse functionality to organic molecules, including pharmaceuticals, and they are also used extensively in the petrochemical-based polymer industry. Double bond isomerization can change the properties of a molecule without the need for stoichiometric exogenous reagents.^[3]

Although there are several elegant examples of earth abundant metal-catalyzed isomerization/functionalization^[4] (e.g. hydroboration,^[5] hydrosilylation,^[6] allylic alcohol to ketone^[7]), which ultimately provide an energetically favorable driving force for the isomerization step, there are far fewer examples of the use of abundant metal pre-catalysts to only invoke iso-

a) Previous catalytic studies using pre-catalyst 1



Scheme 1. a) Previous catalytic reactions undertaken using 1; b) this work.

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merization. Isomerization-only processes catalyzed by iron are dominated by thermodynamically favored cis-to-trans and/or terminal to internal double bond isomerization. Some of the earliest examples of iron-catalyzed reactions date back to the 1960's when Frankel, Emken and Davison reported the use of Fe(CO)₅ as a catalyst in double bond isomerization in fatty acids and esters.^[8] Triiron dodecacarbonyl was used by Casey and Cyr to undertake isomerization of 3-ethyl-1-pentene and detailed mechanistic studies suggested that the reaction likely proceeded via an Fe(n³-allyl)(H) intermediate.^[9] Beyond catalysis performed using iron carbonyl complexes,^[10] von Wangelin has reported isomerization using Fe(acac)₃ with a Grignard additive at room temperature.^[11] More recently, Koh and co-workers have reported an elegant regiodivergent isomerization process that uses B₂pin₂ or PhMe₂SiBpin in conjunction with an iron catalyst and LiOtBu. Mechanistic studies revealed that the reaction is likely to proceed via an iron-hydride with subsequent olefin insertion and β -hydride elimination.^[12] Pertinent to this study, Smith has employed an Fe^I species, exploiting the propensity for spin crossover, to achieve iron catalyzed double bond isomerization via a two-electron catalytic cycle.^[13]

From the synthetic studies we have already disclosed in hydrofunctionalization chemistry,^[14] we postulated that access to a two-electron isomerization catalytic cycle would be possible by limiting the quantity of hydride source employed in order to access Fe^I, meanwhile preventing competitive Fe^{II} hydroboration. We herein report evidence to support this hypothesis, using pre-catalyst **1**.

Results and Discussion

Following a short optimization procedure we found that a cocatalytic quantity (10 mol%) of pinacol borane (HBpin), ammonia borane (H₃N·BH₃) or dimethylamine borane (Me₂HN·BH₃) are all similarly competent at transforming allyl benzene (**2** a) into β -methylstyrene (**3** a) with 5 mol% **1** at 60°C in 16 h (Table 1, also see Supporting Information for further optimization).

Our previous synthetic and theoretical studies with 1 have indicated that hydroboration^[14] and transfer hydrogenation^[15]

Table 1. Optimization of iron catalyzed double bond isomerization using allylbenzene as the substrate.					
	Catalyst (5 mol%), additive (10 mol%)	Conversion to 3 a (%)	trans:cis		
1	1, none	0	0		
2	None, HBpin	0	0		
3	1, HBpin	93	7.6:1		
4	1 , H ₃ N·BH ₃	99	25:1		
5 ^[a]	1, H ₃ N·BH ₃	12	8:1		
6	1, $Me_2HN\cdot BH_3$	99	33:1		
7 ^[b]	1, TMP·BH ₃	73	3.6:1		
8	1, PhH ₂ N·BH ₃	44	2.9:1		

Standard reaction conditions: C_6D_6 (0.6 mL), allylbenzene (0.5 mmol), catalyst (0.025 mmol), additive (0.05 mmol), 60 °C, 16 h. Conversion and selectivity determined by in situ ¹H NMR spectroscopy. [a] RT. [b] TMP = 2,2,6,6-tetramethylpiperidine. 80 °C, 48 h.

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reactions, amongst others, proceed in a redox-neutral manner, in particular operating via an Fe^{II} hydride. In light of this, we sought to rule out an alkyl-based mechanism where HBpin or an alternative reagent serves as a hydride source, forming an iron(II) hydride which can consequently undergo addition across a double bond and subsequent β -hydride elimination at the more substituted position. Further investigation was able to discount this mechanism taking place. Firstly, the iron(II) hydride dimeric species **4** is poor in catalysis (16% conversion to **3a**,16 h, 60°C as opposed to 93% **3a** with **1** and HBpin, see Scheme 2d), contrasting with our dehydrocoupling work.^[16]



Scheme 2. a)-c) Deuterium labelling studies; d) Discounted redox-neutral catalytic isomerization cycle.

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the lack of NMR evidence for **4** forming in situ suggests it is not formed as part of a pathway to an active species. Secondly, deuteration studies give outcomes contradicting what would be expected for a redox neutral mechanism (Scheme 2).

Firstly, there is no ²H-incorporation in substrates when using deuterated hydride sources (Scheme 2a), and full isoretention is observed in the transformation of p-labelled substrate $2a-d_2$ (Scheme 2b) to $3a-d_2$. If 1 is activated to form an Fe-hydride (or deuteride) then a small amount of H/D exchange would be expected owing to the intermolecular nature of catalyst activation and H transfer.

Secondly, in a competitive reaction of **2a**-**d**₂ and **2b**, isoretention is again observed (Scheme 2 c), indicating hydride transfer is intramolecular rather than intermolecular. At this stage, the deuterium labelling studies closely match the results one would expect from an η^3 -allyl mechanism.^[3] Qualitatively, the reactions are observed to change from the dark yellow of 1 to a vivid red; a color that is synonymous with the formation of Fe¹. Furthermore, upon investigating our proposed catalytic cycle with DFT calculations, we were able to demonstrate that a redox-neutral Fe^{II} catalytic cycle of the form shown in Scheme 2 d is disfavored due to an exceptionally high energy barrier for the β -hydride elimination step for the more stable, high-spin quintet surface ($\Delta G^{\pm} = 37.4 \text{ kcal mol}^{-1}$, see Supporting Information for details). Evans method NMR analysis^[17] gives $\mu_{obs} = 5.7$ for the pre-catalyst 1, which is consistent with measurements from Hessen of similar Fe^{II} complexes.^[18] However, addition of HBpin, H₃N·BH₃ or Me₂HN·BH₃ to 1 results in a decrease in the magnetic moment to $\mu_{obs} = 5.1$. We tentatively attribute this to a shift from pure Fe^{II} to a mix of oxidation/ spin states being present. Adding 2a does not appear to affect μ_{obs} further. At this stage, we feel we have reasonable evidence to rule out the entirely redox-neutral cycle of the form shown in Scheme 2 d. No reaction is observed with the radical clock 6bromo-1-hexene, therefore we are confident that there are no organic radicals present or propagating in solution.

To support the NMR results, X-band CW EPR studies were recorded in benzene:toluene (200:10 µL) frozen solution. A solution of 1 in the absence of substrate indicates the presence of a small amount of high-spin Fe^{III}, which we assign to the side formation of LFeCl₂ during synthesis of the pre-catalyst (see Supporting Information). Direct addition of 2a to 1 did not lead to any changes in the EPR spectrum (see Supporting Information), indicating no change in oxidation state induced by the alkene. However, upon addition of HBpin or H₃N·BH₃ to 1, the appearance of a well-resolved rhombic signal without any observable hyperfine coupling is observed, consistent with a low spin S = 1/2 iron d⁷ center (Figure 1). Addition of **2a** to this mixture does not lead to further spectral changes. Computer simulation of this species revealed the spin Hamiltonian parameters characterized by g=[1.984 2.018 2.200], which is analogous to Holland's characterization of an Fe¹ species interacting with benzene, and our previous report of Fe¹ complexes formed upon reaction of FeCl3 with aryl/alkyl Grignard reagents.^[19] The Fe^I signal is superimposed on a broad signal associated with a high-spin Fe^{III} center, vide infra (also see Supporting Information for details). Cyclic Voltammetry was also



Figure 1. i) CW X-band EPR spectrum [T = 140 K] of 1 + HBPin. ii) Experimental (a), and simulation (a') of CW X-band EPR spectrum [T = 140 K] of 1 + HBPin, focusing on the center-field region. The broad signal originating from high-spin Fe^{III} (seen in i) and also in Figure S1) has been background subtracted. See Supporting Information for corresponding simulations and discussion.

used to determine whether HBpin, H₃N·BH₃ or Me₂HN·BH₃ are strong enough reductants to convert 1 into an Fe¹ complex (see Supporting Information for details). Cyclic Voltammetry studies of 1 show two strong reduction peaks. We assign the first to the reduction of our Fe^{II} pre-catalyst to an Fe^I species, which we determine to be -1.48 vs. Fc/Fc⁺. The lack of a large peak in the oxidative direction suggests the irreversibility of the Fe^{II}/Fe^I reduction process, potentially ruling out a oneelectron oxidation/reduction catalytic cycle. The second peak has a potential of -1.92 vs. Fc/Fc⁺, and we believe it is ligandbased and is unlikely to have significant bearing on the catalytic pathway. The relatively strong reducing potentials of both HBpin and H₃N·BH₃ indicate that such a reduction is thermodynamically feasible.^[20] Because of this, we propose that the active species in catalysis is initially an Fe¹ species. This complex may be stabilized by either a molecule of solvent, as previously reported, or by a molecule of substrate. Substitution that, once formed, the η^2 -allylbenzene complex $\mathbf{1}_{\mathbf{B}}$ is more stable, and we therefore believe that if $\mathbf{1}_{A}$ forms, displacement to form $\mathbf{1}_{B}$ is rapid and facile. Wide sweep width NMR spectroscopy of both stoichiometric and catalytic reactions (using HBpin or H₃N·BH₃) give complex spectra that we attribute to off-cycle or catalytically inactive Fe^{II} complexes or Fe^{III} formed

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Scheme 3. a) **1** reacts with borane to generate $1_{A'}$ which can then undergo further stabilization with alkene; b) proposed Fe^I/Fe^{III} catalytic cycle for the isomerization of allylbenzene.

by disproportionation of Fe^{II} into Fe^{II} and Fe^{III} (see Supporting Information for extended discussion). Unfortunately, we have not been able to isolate any key intermediates from reaction mixtures^[22] and characterize them crystallographically, and detailed kinetics studies have been hampered by the reaction temperature coupled to the paramagnetic iron center, which have made in situ NMR monitoring challenging. However, in situ UV-vis spectroscopy measurements give $\lambda_{max} = 494$ and 552 nm (for catalysis in the presence of H₃N·BH₃) or $\lambda_{max} = 497$ and 547 nm (for catalysis in the presence of HBpin); wavelengths that have been linked to both Fe^{I} and $Fe^{III} \beta$ -diketiminate species.^[21,23] Using the method reported by Scheer for the synthesis of the η^6 -toluene analogue of $\mathbf{1}_{A'}^{[24]}$ we obtain 87% **3a** (7.5:1 *trans:cis*) after 16 h at 60 $^{\circ}$ C when 5 mol% of this Fe^I species is used in catalysis (31% in a 6.3:1 ratio after 2 h); these results are in-line with those obtained using the same conditions with pre-catalyst 1 (5 mol% 1 and 10 mol% HBpin, 27%, 5.7:1 ratio after 2 h; and 93%, 7.6:1 ratio after 16 h) and further support the proposal that the reaction proceeds via an Fe¹ species.

To summarize the experimental results thus far: 1 appears to undergo a change in oxidation state in the presence of a hydride source and this is supported by NMR and EPR studies. Electrochemistry indicates that the hydride source can reduce 1 to an Fe¹ species and work from Holland has demonstrated that alkenes are able to form η^2 -Fe¹ complexes (i.e. 1_A and 1_B). An η^6 -arene Fe¹ complex gives similar catalytic results to 1. Deuterium labelling indicates that the hydride is not transferred to the isomerized product and therefore it appears to be acting as a pre-catalyst activator only. Having ruled out a redox-neutral Fe^{II} pathway both experimentally and computationally, we further investigated the isomerization of allylbenzene to β -methylstyrene using DFT calculations. We propose that the reaction proceeds from our active Fe^I η^2 -alkene species $\mathbf{1}_{B}$, in the doublet spin-state, via a selectivity-determining oxidative addition of one of the phenyl adjacent C–H bonds to form either of Fe^{III} η^3 -allyl species, $\mathbf{1}_{C\ trans}$ or $\mathbf{1}_{C\ cis}$. These intermediates then undergo a reductive elimination of the proton onto the terminal carbon to form the Fe^{II} allyl species, $\mathbf{1}_{D\ trans}$ or $\mathbf{1}_{D\ cis}$. A ligand exchange with the substrate $\mathbf{2}\mathbf{a}$ then yields the major product, *trans*- β -methylstyrene or the minor product, *cis*- β -methylstyrene, while also regenerating $\mathbf{1}_{B}$ (Scheme 3 b).

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We suggest that the doublet spin state of the complex is largely preserved throughout the catalytic cycle, although we have also investigated the other spin states, with a particular focus on the *trans*-selective route. While the quartet 1_B was calculated to be more stable than the doublet by 21.0 kcal mol⁻¹, we found, similarly to Smith,^[13] that the activation barrier for the *trans* selective oxidative addition on the quartet surface ($\Delta G^{\pm} = 45.0 \text{ kcal mol}^{-1}$) was far in excess of what might be considered accessible under the experimental conditions used (see Supporting Information). In addition to this, our EPR results do not support the presence of a quartet Fe^{III} species.

Starting from our initial species 1_B, rotation of the Ph-C bond to give the appropriate orientation for each oxidative addition is a low-energy process ($< 1 \text{ kcal mol}^{-1}$), with the subsequent activation barriers ($\Delta G^{\pm} = 25.7 \text{ kcal mol}^{-1}$ for *trans*, $\Delta G^{\pm} = 28.1 \text{ kcal mol}^{-1}$ for *cis*) supporting the experimentally observed trans selectivity of the reaction (Table 2). Due to steric interactions arising from the proximity of the substrate phenyl ring to the 2,6-diisopropylphenyl groups on the β -diketiminate ligand, the *cis* η^3 -allyl product of the oxidative addition, intermediate $\mathbf{1}_{c\ cis}$, generates an overall positive ΔG for that step (+19.7 kcal mol⁻¹ relative to $\mathbf{1}_{B}$), while the less hindered $\mathbf{1}_{C trans}$ is 2.7 kcal mol⁻¹ more stable than $\mathbf{1}_{C cis}$ and results in a slightly reduced free energy difference, $\Delta G = +18.0$ kcal mol⁻¹, making the formation of 1_{C trans} slightly less reversible than 1_{C cis}. From there, both isomers have relatively small barriers for the reductive elimination ($\Delta G^{\pm} = 2.6 \text{ kcal mol}^{-1}$ for *trans* and 4.8 kcal mol⁻¹ for *cis*), the rate determining barrier for both is the oxidative addition, supported by a secondary kinetic isotope effect ($K_D/K_H = 1.03 \pm 0.06$) observed experimentally. Both the trans (-4.6 kcal mol⁻¹) and cis selective (-3.3 kcal mol⁻¹) isomerization of allylbenzene are calculated to be exothermic overall.

The sextet Fe^{III} intermediate $\mathbf{1}_{C \ trans}$ lies below the doublet electronic configuration at our chosen level of theory. As

Table 2. Calculated activation barriers ΔG^{+} (kcal mol ⁻¹) for the isomerization of allylbenzene for the proposed Fe ^I /Fe ^{III} catalytic cycle.			
Selectivity	Oxidative addition	Reductive elimination	
trans cis	25.7 28.1	2.6 4.8	

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Figure 2. Calculated free energy surface for the *trans*-selective isomerization of allylbenzene in doublet (black line) and sextet (red line) electronic configurations, relative to the initial on-cycle species $1_{B neutral}$. B3LYP-D3/6-31G(d), PCM = benzene, 298 K.

shown on the energy surfaces plotted in Figure 2, the highspin Fe^{III} species detected by EPR is likely to be $1_{C \ trans}$ (3.1 kcal mol⁻¹ more stable than the doublet), although we concede that computational method effects might affect this prediction. The observed reaction outcome could thus be explained either by considering only the low spin species, or by spin crossovers onto the high spin surface near the oxidative addition step and crossing back onto the low spin pathway near the reductive elimination. Unfortunately, we were unable to locate minimum energy crossing points (MECPs) between these surfaces with the method described by Harvey et al. (see Supporting Information for further details).^[25]

To prove utility beyond allyl benzene we have explored a range of substrates (Scheme 4). We have opted to use H₃N·BH₃ as our hydride source as it is an easy to handle, inexpensive reagent with good sustainability credentials. In most cases there is good selectivity for the trans-styrene product (3a to 3d). However, when the aromatic ring is functionalized with strongly electron donating (3e and 3f) or withdrawing groups (3g and 3h), there is a drop-off in trans-selectivity and the cis product becomes more favorable. We link this, and the dropoff in trans selectivity observed with HBpin (see Table 1), to rate of reaction: reactions of electron donating or withdrawing substrates, or those mediated by HBpin, are slower, meaning there is less opportunity for cis-to-trans isomerization to take place prior to product isolation. Good turnover is observed for the double bond isomerization of β - to α -pinene (3 i) and a modest amount of exo- to endo-double bond isomerization is observed for valencene (2j). Unfortunately, no reactivity is observed with secondary and tertiary amine-containing alkenes (diallylamine, N-allylaniline and 4-methyl-diallylaniline), or with 1,1-disubstituted alkenes such as (2-methyl-2-propenyl)benzene or multiple double bond containing terpenes such as β farnesene (which showed good reactivity during hydroboration studies).^[14] We have explored the selectivity obtained for the isomerization of hexene (see Supporting Information) but more forcing reaction conditions are required (80 °C, 48 h, in comparison to 60 °C, 16 h).

Finally, to demonstrate the applicability of this isomerization catalysis to sustainable chemistry, we have utilized our catalysis to afford a key synthetic fragrance and flavor molecule (Scheme 4b). In a reaction sequence involving iron-catalyzed double bond isomerization followed by ozonolysis, we have

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Scheme 4. a) Double bond isomerization substrate scope. [a] $80 \,^{\circ}$ C; b) Preparation of fragrance molecule piperonal using iron catalyzed double bond isomerization followed by ozonolysis.

transformed safrole into piperonal, a floral/vanilla scented compound with both fragrance and synthetic applications.

Conclusions

In summary, we have reported a rare example of iron catalysis likely proceeding through a two-electron oxidation/reduction mechanistic pathway. A catalytic amount of HBpin or H₃N·BH₃, rather than act as a source of hydride, acts to reduce our Fe^{II} pre-catalyst to an Fe^I species, a proposal supported by EPR and electrochemical studies. An η^3 -allyl-type mechanism, proceeding via an Fe^I/Fe^{III} redox active cycle, is supported by deuterium labelling and DFT studies, leading to terminal alkenes being isomerized to more substituted products in a *trans* selective manner. The presence of a redox active catalytic cycle

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raises the question of whether this chemistry is limited to double bond isomerization, or whether we can exploit this to other two-electron transformations.

Acknowledgements

The EPSRC Centre for Doctoral Training in Catalysis (EP/L016443/1) is thanked for funding CRW and DJD. We thank the EPSRC UK National Mass Spectrometry Facility at Swansea University for MS analysis and the Centre for Computational Chemistry at the University of Bristol for providing access to computing resources. The authors would also like to thank Dr. Josh Tibbetts for assistance with ozonolysis experiments, Professor Frank Marken and Mr. Jack Stewart for helpful discussion on Cyclic Voltammetry studies, and Dr. Danila Gasperini for the preparation of the Fe¹ η^6 -toluene analogue of $\mathbf{1}_A$.

Conflict of interest

The authors declare no conflict of interest.

Keywords: homogeneous catalysis • iron • isomerization • reaction mechanisms • redox chemistry

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Manuscript received: November 17, 2020 Revised manuscript received: January 19, 2021 Accepted manuscript online: January 25, 2021 Version of record online:

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FULL PAPER

Fe(I)/(III) cycle

- d-labelling
- EPR
- Electrochemistry
- DFT

Using deuterium labelling, electrochemistry, EPR and DFT, an iron catalyzed double bond isomerization reaction was disclosed that is likely to proceed through on-cycle Fe¹ and Fe^{III} intermediates. The reaction uses an Fe^{II} pre-

Ph

-e(I)



catalyst that undergoes a one-electron reduction to get to the first on-cycle species. These studies on mechanism, along with synthetic scope of the reaction, are presented.

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Catalysis

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Iron Catalyzed Double Bond Isomerization: Evidence for an Fe^I/Fe^{III} Catalytic Cycle