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Introduction

Metal-catalysed C–H functionalisation has emerged as a powerful synthetic method for the direct derivatisation of typically inert C–H bonds and obviating the need for prefunctionalised substrates.¹ While the application of C–H functionalisation strategies has expanded, these methods generally rely on precious metal catalysts. Examples using Earthabundant metal,² and even non-metal,³ catalysts have emerged as potential alternatives, however their use and development has been comparably limited.

A number of iron-catalysed C–H functionalisation methods have been developed,^{2a} but the isolation and characterisation of well-defined, catalytically relevant species has remained limited to only a few studies, particularly where C–H metallation, sometimes referred to as C–H activation, has been proposed.⁴ Holland reported the observation and isolation of an iron-aryl species arising from C–H metallation of benzene using a low oxidation-state iron β -diketiminate complex (Scheme 1a).^{4a}

Iron-catalysed alkene and heteroarene H/D exchange by reversible protonation of iron-hydride intermediates†

Luke Britton,^a Jamie H. Docherty, ^b*^a Jan Sklyaruk,^a Jessica Cooney,^a Gary S. Nichol, ^a Andrew P. Dominey^b and Stephen P. Thomas ^b*^a

C-H functionalisation reactions offer a sustainable method for molecular construction and diversification. These reactions however remain dominated by precious metal catalysis. While significant interest in ironcatalysed C-H activation reactions has emerged, the isolation, characterisation and mechanistic understanding of these processes remain lacking. Herein the iron-catalysed C(sp²)-H bond hydrogen/ deuterium exchange reaction using CD₃OD is reported for both heterocycles and, for the first time, alkenes (38 examples). Isolation and characterisation, including by single-crystal X-ray diffraction, of the key iron-aryl and iron-alkenyl C-H metallation intermediates provided evidence for a reversible protonation of the active iron hydride catalyst. Good chemoselectivity was observed for both substrate classes. The developed procedure is orthogonal to previous iron-catalysed H/D exchange methods which used C_6D_6 , D_2 , or D_2O as the deuterium source, and uses only bench-stable reagents, including the iron(II) pre-catalyst. Further, a new mechanism of iron-hydride formation is reported in which β hydride elimination from an alcohol generates the iron hydride. The ability to produce, isolate and characterise the organometallic products arising from C-H activation presents a basis for future discovery and development.

> Similarly, Ackermann reported the C–H metallation of pivalophenone using a low oxidation-state iron complex $[Fe(PMe_3)_4]$ that allowed for the isolation and characterisation of the key iron-aryl species responsible for allene hydroarylation (Scheme 1a).^{4b,c} Additional iron-aryl complexes have been isolated and characterised through independent synthesis, usually from organometallic aryl reagents.^{4d-g}

> The iron bisdiphosphino complex, $[dmpe_2FeH_2] 2 (dmpe = Me_2PCH_2CH_2PMe_2)$, has been shown to undergo stoichiometric C-H bond metallation of benzene in solution-phase studies.⁵ It was therefore questioned whether the metallation of $C(sp^2)$ -H bonds could be made general and harnessed to generate an iron-aryl species applicable to H/D exchange. In order to achieve H/D exchange, photoirradiation of $[dmpe_2FeH_2] 2$ would be used to trigger H₂ elimination to give $[dmpe_2Fe^0]$. This would undergo $C(sp^2)$ -H bond oxidative addition (metallation) to give an aryl(hydrido) iron species, $[dmpe_2Fe(H)Ar]$, from which exchange of the hydrido-ligand to the deuteride isotopologue, would give $[dmpe_2Fe(D)Ar]$, and leave only a final reductive elimination to complete a potential catalytic cycle.

Commonly, direct H/D exchange reactions have involved the use of precious metal catalysts, with only a few Earth-abundant systems reported. Chirik reported the H/D exchange of arenes using iron-pincer complexes $[(MesCNC)Fe(N_2)_2]$ and deuterium gas, as well as $[(MesCNC)Fe(CH_2SiMe_3)_2(N_2)]$ and C_6D_6 (Scheme 1b).⁶ de Ruiter reported the C_6D_6 -mediated H/D exchange

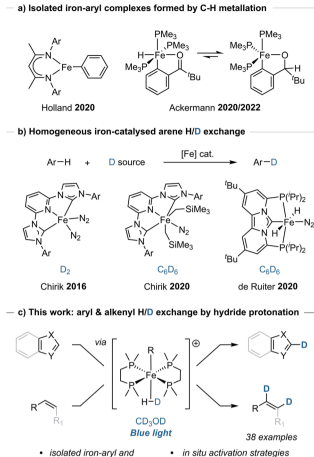
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[&]quot;EaStCHEM School of Chemistry, University of Edinburgh, Edinburgh EH9 3FJ, UK. E-mail: jamie.docherty@ed.ac.uk; stephen.thomas@ed.ac.uk

^bGSK Medicines Research Centre, Stevenage SG1 2NY, UK

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iron-alkenyl intermediates to iron-hydrides

Scheme 1 a) Previous examples of isolated iron-aryl complexes by direct C-H metallation. (b) Previous homogeneous iron-catalysed H/ D exchange catalysts. (c) This work: arene and alkene C-H metallation and H/D exchange using CD₃OD and a bench-stable iron(II) precatalyst.

reactions of (hetero)arenes using an alternative iron-complex [(PCNHCP)Fe(H)₂(N₂)] (Scheme 1b).⁷ Beller and Lei reported the H/D exchange reaction of arenes bearing electron-donating groups using D₂O and a heterogeneous iron catalyst.⁸ Additional examples of Earth-abundant metal-catalysed HIE reactions of arenes have been reported which rely on pre-installed directing groups.9 Of note to this manuscript, Hartwig used a Ag₂CO₃/JohnPhos catalyst and CH₃OD for the H/D exchange reaction of 5-membered heteroarenes.10

Results and discussion

Investigations began by exploring the viability of H/D exchange with a selection of related iron pre-catalysts, deuterium sources and catalyst activation methods (see ESI, Tables S1-S3[†]). Precatalyst activation with alkoxide salts has been shown to generate catalytically active metal-hydride species when combined with commercial silane or borane reagents such as pinacolborane (HBpin).11 This removes the need to generate and handle the highly air- and moisture sensitive iron

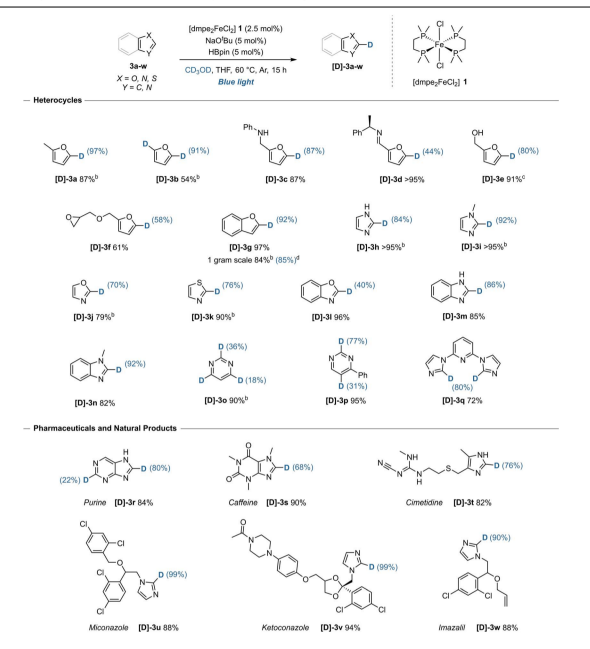
dihydride. To assess the catalytic feasibility of this strategy in *situ* generated $[dmpe_2FeH_2]$ 2 (from $[dmpe_2FeCl_2]$ 1 + NaO^tBu + HBpin) was used in combination with 2-methylfuran 3a, blue light irradiation (450 nm) and a variety of deuterium sources. Deuterium incorporation was achieved using deuterated alcohols as the deuterium source, with CD₃OD giving the highest deuterium incorporation and exclusive C-5 regioselectivity. Control reactions showed the necessity for continuous light irradiation, iron complex, phosphine ligand, and the alkoxide activator (see ESI, Table S4[†]). Notably, good levels of deuterium incorporation were observed even in the absence of HBpin (vida infra). Optimisation of the light source using monochromatic LEDs in the range of 365-525 nm, showed the blue region (450 nm) of the spectrum to be optimal (see ESI, Table S5[†]). The substitution of CD₃OD for CD₃OH resulted in no deuterium incorporation, indicating that the most acidic, alcohol, hydrogen was responsible for deuterium-exchange. The state-ofthe-art precious metal equivalent for non-directed H/D exchange of arenes using ROD-type deuterium sources resulted in global C-H deuteration and showed no chemoselectivity for alkene versus arene C-H bonds (reaction conditions: $Pd(OAc)_2$ (10 mol%), ligand (15-30 mol%), HFIP : $D_2O = 3:7$, 80-120 °C, 18-48 h).12

Catalytic H/D exchange of heteroarenes

The generality of the method with respect to substrate compatibility and scope was then assessed (Table 1). Furan 3b, furan derivatives bearing a free secondary-amine 3c and imine 3d all underwent successful, chemoselective deuteration to give the monodeuterated furans [D]-3b, [D]-3c and [D]-3d, respectively, with good to excellent deuterium incorporation and complete control of regioselectivity (C-5) in all cases. Pinacol boronic ester protected alcohol 3e reacted to give the deuterated isotopologue [D]-3e, albeit with alcohol deprotection upon work up. Benzofuran 3g underwent site-selective deuteration at the C-2 position, with no detectable deuteration at any of the other five available $C(sp^2)$ -H bonds. Unprotected imidazole **3h** and methyl-protected imidazole 3i underwent exclusive C-2 selective deuteration to give the deuterated imidazoles [D]-3h and [D]-3i, respectively. This is notable as both previously reported ironcatalysts gave unselective deuteration of N-methyl imidazole with exchange at at all the C(sp²)–H bonds.^{6,7} Using this system, oxazole 3j, 1,3-thiazole 3k and benzoxazole 3l all underwent deuteration with mild to good deuterium incorporation, again at a single site. 6-Membered heterocycle pyrimidine 30 proved amenable to deuteration at C-2, C-4 and C-6, albeit with only moderate incorporation. When 4-phenylpyrimidine 3p was used, deuteration occurred at both C-2 and C-5 positions with significantly greater inclusion than the unsubstituted analogue.

To illustrate the utility and tolerance of this system with respect to complex targets, the deuteration procedure was successfully applied to several pharmaceutical and natural product structures (Table 1, lower). Purine 3r, a compound ubiquitous to biological systems and common foodstuffs, reacted efficiently with incorporation at both C-2 and C-8. A further range of biologically active molecules including caffeine

Table 1 Iron-catalysed Site-selective Heteroarene C(sp²)-H H/D Exchange^a



^{*a*} Reaction conditions: arene (0.33 mmol), [dmpe₂FeCl₂] **1** (2.5 mol%), NaO^{*t*}Bu (5 mol%), HBpin (5 mol%), THF (0.2 mL), CD₃OD (0.2 mL), blue-light irradiation, 15 h. Reported yields denote quantity of material recovered post-reaction. Deuterium incorporation determined by ¹H and/or ¹³C NMR spectroscopy. ^{*b*} Quantity of recovered product determined using 1,3,5-trimethoxybenzene as an internal standard. ^{*c*} Starting material **3e** = 4,4,5,5-tetramethyl-2-(furan-2-ylmethoxy)-1,3,2-dioxaborolane, free alcohol [**D**]-**3e** isolated following deprotection. ^{*d*} Performed on a 9.3 mmol scale, 48 h.

3s, histamine receptor antagonist cimetidine **3t** and antifungal compounds; miconazole **3u**, ketoconazole **3v** and imazalil **3w**, all reacted efficiently with predictable deuterium incorporation at the most acidic $C(sp^2)$ -H site. While the imidazole unit of **3w** underwent H/D exchange under the reaction conditions to give **[D]-3w**, the allylic ether was not deuterated. The functional group compatibility and regioselectivity shown illustrates the system is suitable for potential late-stage site-selective isotopic exchange of heteroarene units.

While high levels of deuterium incorporation were obtained using this system, it must be noted that an alternative alkoxidemediated reaction exists for substrates with highly acidic $C(sp^2)$ -H bonds. Excluding benzoxazole **31**, which achieved greater levels of deuterium incorporation in the absence of precatalyst and HBpin,¹³ all substrates underwent greater deuterium incorporation and selectivity under the iron-catalysed reaction conditions (see ESI, Table S8†).

Catalytic H/D exchange of alkenes

While H/D exchange reactions of alkenes with non-reductive sources of deuterium, i.e. CD₃OD, D₂O or C₆D₆, have been previously reported using precious metals catalysts (iridium,14 rhodium,15 ruthenium,16 and palladium17), Earth-abundant metal-mediated H/D exchange reactions of alkenes have so far been limited to either stoichiometric studies,18 or unwanted side reactions.19 Using this iron-catalysed system, styrene derivatives 4a-f bearing both electron-donating- and electron-withdrawing groups were efficiently deuterated at all alkene positions with no detection of deuterium incorporation at any of the arene $C(sp^2)$ -H bonds. This represents the first reported example of an iron-catalysed H/D exchange reaction of alkenes (Table 2). It was observed that for the deuteration of styrene derivatives there was a slight trend for greater deuterium incorporation at the α position relative to both β positions, contrasting with results often seen with precious metal alternatives.14b,15b,17

2,3-Dihydrofuran 4g underwent deuteration at both alkene $C(sp^2)$ -H bonds while the 6-membered 2,3-dihydropyran 4h was selectively deuterated at the 3-position. Monosubstituted acyclic alkenes undecene 4i and O-tosyl alkene 4j both underwent moderate deuterium incorporation. Naturally occurring allylic alcohols Linalool 4k and anti-cancer agent Sclareol 4m, both of which contain free alcohol groups, were deuterated with moderate deuterium incorporation. Cinchona alkaloid quinine 4n and diastereoisomer quinidine 4o were both successfully reacted and with exclusive deuteration of the alkene over the other five available C(sp²)-H sites. Minimal reactivity was displayed towards carbon substituted 1,2-cis alkenes including cyclohexene and cyclooctene, and no reactivity was observed for trans-, 1,1-disubstituted-, trisubstitued- or activated (e.g. acrylates or acrylamides) alkenes. Background, base-mediated H/D exchange was not observed for any alkene substrate (see ESI, Table S8[†]).²⁰





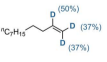
[D]-4f 86%

0

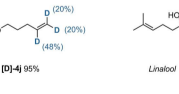


'n

[D]-4a 85%



[D]-4i 61%



b

[D]-4c 89%

(61%)

(62%)

[D]-4a 77%

(62%)

Linalool [D]-4k 96%

[dmpe2FeCl2] *

(63%)

[D]-4d 58%b

[D]-4h 84%b

p ^(10%)

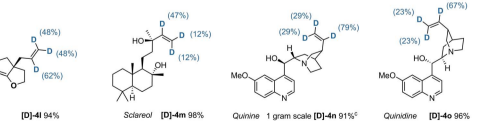
(36%)

D (10%)

(40%)

D (53%)

(53%)

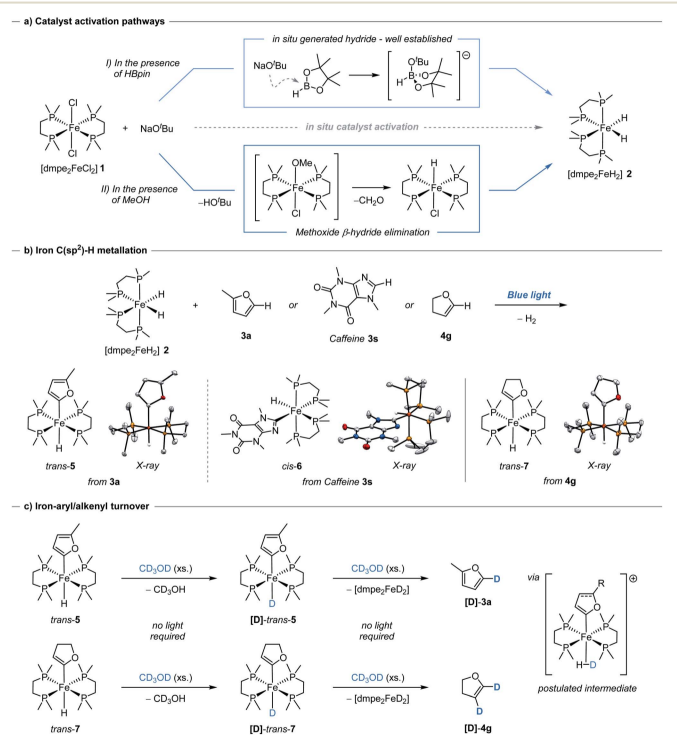


^a Reaction conditions: alkene (0.33 mmol), [dmpe₂FeCl₂] 1 (2.5 mol%), NaO^tBu (5 mol%), HBpin (5 mol%), THF (0.2 mL), CD₃OD (0.2 mL), bluelight irradiation, 15 h. Reported yields denote quantity of material recovered post-reaction. Deuterium incorporation determined by ¹H and/or ¹³C NMR spectroscopy. ^b Quantity of recovered product determined using 1,3,5-trimethoxybenzene as an internal standard. ^c Performed on a 3.4 mmol scale (1.0 g recovered), 48 h.

Mechanistic investigations

Having demonstrated catalytic H/D exchange across arenes and alkenes, the fundamental catalytic steps were investigated. The generation of the photoactive iron hydride [dmpe₂FeH₂] 2,

through the *in situ* activation of [dmpe₂FeCl₂] **1** using an alkoxide salt and HBpin, has been previously reported (Scheme 2a, I),^{11b} but did present a question of how significant deuterium incorporation of 2-methylfuran **3a** was achieved in the absence of HBpin (see ESI, Table S4[†]). Methanol solutions



Scheme 2 Mechanistic Studies for Iron-catalysed H/D Exchange. (a) *in situ* Activation pathways of $[dmpe_2FeCl_2]$ **1** pre-catalyst; (I) in the presence of HBpin; (II) in the presence of CH₃OH. (b) C(sp²)–H bond activation to give intermediate *trans*-5, *cis*-6, and *trans*-7. X-ray ellipsoids are shown at 50% probability. Hydrogen atoms, with the exception of Fe-hydrides, have been omitted for clarity. (c) Fe-hydrides (*trans*-5 and *trans*-7) to Fe-deuterides (**[D]**-*trans*-7), exchange using CD₃OD in the absence of light, postulated intermediate species shown to the right. Generation of **[D]**-3a and **[D]**-4g through slow exchange between CD₃OD and Fe-deuterides **[D]**-*trans*-7, respectively.

of [dmpe₂FeCl₂] **1** were observed to exist as a 1 : 1 electrolyte of [dmpe₂Fe(HOMe)Cl]⁺ and Cl⁻, as seen by ³¹P NMR spectroscopy.²¹ Upon the addition of a stoichiometric amount of sodium *tert*-butoxide, the immediate generation of iron-hydride species, including [dmpe₂FeHCl] and [(dmpe₂FeH)₂(µ-dmpe)]²⁺, were observed by ¹H and ³¹P NMR spectroscopy. Following blue light irradiation, [dmpe₂FeH₂] **2** was also observed in solution alongside an additional unknown iron hydride species. This unknown iron-hydride was not observed under standard activation conditions; sodium *tert*-butoxide and HBpin (see ESI,[†] Part 12).

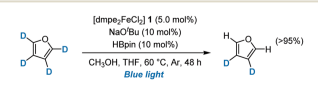
The formation of these catalytically active iron-hydrido complexes was proposed to proceed by tert-butoxide-mediated deprotonation of [dmpe₂Fe(HOMe)Cl]⁺ to generate an unstable iron methoxide complex, [dmpe₂Fe(OMe)Cl], which underwent β-hydride elimination releasing formaldehyde and generating [dmpe₂FeHCl] (Scheme 2a, II). The formation of [dmpe₂FeHCl] was also observed by ¹H and ³¹P NMR spectroscopy when the reaction was performed in ethanol and benzyl alcohol, with benzaldehyde being formed and observed by ¹³C NMR spectroscopy in the latter case. When the reaction was performed in CD₃OH, [dmpe₂FeDCl], along with other irondeuteride complexes, were observed by ³¹P NMR spectroscopy (see ESI,[†] Part 12). The use of alcohols lacking available β hydrogens, tert-butanol and phenol, resulted in no iron-hydride species being observed. Similar reactivity has been proposed for the analogous ruthenium-dmpe complexes.22 It should be noted that although activation to the catalytically active ironhydride complexes can be achieved in the absence of HBpin, this activation was less controlled in terms of clean formation of [dmpe₂FeH₂] under catalytic conditions.

[trans-dmpe₂Fe(H)(2-methylfuryl)] trans-5, has previously been characterised in solution.^{11b} By careful tuning of the reaction and crystallization conditions, it was characterised in the solid-state by single-crystal X-ray diffraction (Scheme 2b, left). Significantly this reactivity was not limited to furan derivatives and was extended to the reaction of caffeine 3s to give the corresponding complex, [cis-dmpe₂Fe(H)(caffeine)] cis-6 (Scheme 2b, center). Although these species were crystallised as the trans- and cis-isomers, respectively, solution phase cistrans isomerisation was reasonably proposed.23 Rapid and quantitative exchange of hydride for deuteride was observed upon addition of excess CD₃OD (approx. 250 eq.) to a THF solution of [trans-dmpe2Fe(H)(2-methylfuryl)] trans-5 to give the deuterated isotopologue [trans-dmpe2Fe(D)(2-methylfuran)] [D]trans-5 in the absence of blue light irradiation. Exchange of hydride for deuteride occurring by protonation of [transdmpe₂Fe(H)(2-methylfuryl)] trans-5 with CD₃OD,²⁴ to give the cationic complex [dmpe₂Fe(HD)(2-methylfuryl)]⁺ which reversibly released CD₃OH (Scheme 2c, upper). Further monitoring of this solution by ¹H and ³¹P NMR spectroscopy showed the slow generation of [dmpe₂FeD₂] [D]-2 and 2-methyl-5D-furan [D]-3a at room temperature in the absence of blue light irradiation. Additionally, no further intermediary species were observed, and similar reactivity was not observed when using ^tBuOH. Presumably, turnover proceeded either through a stepwise cationic protonation and release of the arene, or by a concerted

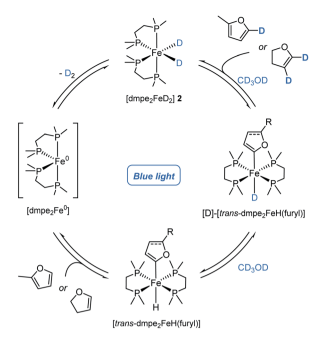
sigma bond metathesis reaction with CD_3OD , both resulting in an [dmpe₂Fe(D)(OMe)] intermediate which collapsed by β hydride elimination to generate the photoactive complex [dmpe₂FeD₂] **[D]-2** and formaldehyde (*vida ante*).

The reversible nature of deuterium incorporation led to it being questioned whether this process could be used for selective protodedeuteration to give selectively deuterated products. d_4 -Furan was regioselectively protodedeuterated at the 2- and 5-positions to give d_2 -[3D,4D]-furan with >95% incorporation using 5 mol% of Fe(II) pre-catalyst 2 and MeOH (Scheme 3). The synthesis of d_2 -[3D,4D]-furan has only been previously prepared using a multistep sequence.²⁵

Considering the H/D exchange of alkenes, blue light irradiation of [dmpe₂FeH₂] 2 in the presence of 2,3-dihydrofuran, led exclusively to the C-2C(sp^2)-H metallation product, [transdmpe₂Fe(H)(2,3-dihydrofuryl)] trans-7 (Scheme 2b, right). Despite reports of Fe-alkenyl C(sp²)-H metallation products being observed in the solution phase,^{5b} complex trans-7 represents the first example of an Fe-alkenyl complex characterised in the solid-state by single-crystal X-ray diffraction (Fe-C bond distance 1.338(25) Å, C=C bond distance 1.338(2) Å). This was in contrast to previous reports using the analogous bis(diethylphosphino)ethane ligand where C-3 metallation was proposed based on solution phase ¹H and ³¹P NMR spectroscopy.²⁶ In analogy to aryl iron hydride trans-7, the alkenyl iron complex trans-7 displayed similar reactivity in the presence of CD₃OD; quantitative conversion to the iron-deuteride isotopologue, [D]-trans-7, was observed upon the addition of excess CD₃OD in the absence of blue light irradiation. Slow conversion of [trans-dmpe₂Fe(H)(2,3-dihydrofuryl)] trans-7 to [dmpe₂FeD₂] [D]-2 with the release of 2,3-dihydro-4D,5D-furan [D]-4g was observed by ¹H and ³¹P NMR spectroscopy in the absence of blue light irradiation. The irradiation of [dmpe₂FeH₂] 2 in the presence of 4-tert-butylstyrene 4a, led to a complex mixture of products observed by ¹H and ³¹P NMR spectroscopy. When performing the deuteration of *tert*-butylstyrene in the absence of blue light irradiation at 60 or 80 °C, a greater α : β ratio was observed to that under standard reaction conditions (see ESI, Table S6[†]). When the allylic ethers, 2,5-dihydrofuran and allyl benzyl ether, were reacted under standard conditions, alkene isomerisation to the vinyl ethers was observed producing 2,3dihydrofuran (62%) and the benzyl propenyl ether (62%, 46 : 54 Z: E), respectively. Similarly, 4-allylanisole showed small amounts of isomerisation to (Z)-anethole (<10%) (see ESI,† Part 12). Control reactions ruled out any acid- or base-mediated



Scheme 3 Protodedeuteration of d_4 -furan. Reaction conditions: d_4 -furan (0.33 mmol), [dmpe₂FeCl₂] 1 (5.0 mol%), NaO^tBu (10 mol%), HBpin (10 mol%), THF (0.2 mL), CH₃OH (0.2 mL), blue-light irradiation, 48 h. Deuterium incorporation determined by ¹H and ¹³C NMR spectroscopy.



Scheme 4 Proposed reaction mechanism for the iron-catalysed H/D exchange reaction of heteroarenes and alkenes.

isomerisation. The aforementioned observations, alongside previous work,^{5b,27} would suggest that the deuteration of alkenes is occurring through a combination of (direct) C–H metallation and hydrometallation mechanisms, with the latter observed to mediate alkene isomerisation alongside exchange (Scheme 4).

Conclusions

In summary, direct C-H metallation of heteroarenes and alkenes using an iron(II) hydride species has enabled the development of an iron-catalysed H/D exchange reaction using a bench-stable iron(II) pre-catalyst, and CD₃OD as the deuterium source. This method is mechanistically different from and complementary to previously described iron-catalysed strategies, and allows for precise and predictable formation of C-D bonds without the need for directing groups. Regioselectivity was obtained and controlled through highly selective C-H metallation by the proposed active catalyst. Significantly, the use of a non-reductive deuterium source (CD₃OD) has enabled extension to exchange of alkene C-H bonds and to molecules containing reducible functional groups. Isolation and characterisation, including single-crystal X-ray diffraction, of the key iron-aryl and iron-alkenyl intermediates allowed for mechanistic investigations that provided evidence for a reversible protonation of these iron C-H activation products. Further, a new mechanism of iron-hydride formation is presented in alcohol solvents.

Data availability

Experimental data is provided in the ESI.†

Author contributions

J. H. D. and S. P. T. conceived and discovered the reaction. L. B., J. H. D., J. S. and J. C. conducted the experimental work. G. S. N. conducted the X-ray crystallographic analysis. S. P. T. and A. P. D. advised investigations. All authors contributed to the manuscript.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- (a) I. V. Seregin and V. Gevorgyan, Chem. Soc. Rev., 2007, 36, 1173-1193; (b) M. M. Díaz-Requejo and P. J. Pérez, Chem. Rev., 2008, 108, 3379-3394; (c) J. Yamaguchi, A. D. Yamaguchi and K. Itami, Angew. Chem., Int. Ed., 2012, 51, 8960-9009; (d) Z. Huang, H. N. Lim, F. Mo, M. C. Young and G. Dong, Chem. Soc. Rev., 2015, 44, 7764-7786; (e) F. Roudesly, J. Oble and G. Poli, J. Mol. Catal. A: Chem., 2017, 426, 275-296; (f) D. J. Abrams, P. A. Provencher and E. J. Sorensen, Chem. Soc. Rev., 2018, 47, 8925-8967; (g) B. Liu, L. Yang, P. Li, F. Wang and X. Li, Org. Chem. Front., 2021, 8, 1085-1101.
- 2 (a) R. Shang, L. Ilies and E. Nakamura, *Chem. Rev.*, 2017, 117, 9086–9139; (b) P. Gandeepan, T. Muller, D. Zell, G. Cera, S. Warratz and L. Ackermann, *Chem. Rev.*, 2019, 119, 2192–2452; (c) L. Ilies, *Bull. Chem. Soc. Jpn.*, 2021, 94, 404–417; (d) J. Loup, U. Dhawa, F. Pesciaioli, J. Wencel-Delord and L. Ackermann, *Angew. Chem., Int. Ed.*, 2019, 58, 12803–12818.
- 3 (a) M.-A. Légaré, M.-A. Courtemanche, É. Rochette and F.-G. Fontaine, *Science*, 2015, 349, 513–516; (b)
 K. Chernichenko, M. Lindqvist, B. Kótai, M. Nieger, K. Sorochkina, I. Pápai and T. Repo, *J. Am. Chem. Soc.*, 2016, 138, 4860–4868; (c) J. Lv, X. Chen, X.-S. Xue, B. Zhao, Y. Liang, M. Wang, L. Jin, Y. Yuan, Y. Han, Y. Zhao, Y. Lu, J. Zhao, W.-Y. Sun, K. N. Houk and Z. Shi, *Nature*, 2019, 575, 336–340; (d) T. He, H. F. T. Klare and M. Oestreich, *J. Am. Chem. Soc.*, 2022, 144, 4734–4738.
- 4 (a) S. F. McWilliams, D. L. J. Broere, C. J. V. Halliday,
 S. M. Bhutto, B. Q. Mercado and P. L. Holland, *Nature*,
 2020, 584, 221–226; (b) A. M. Messinis, L. H. Finger, L. Hu
 and L. Ackermann, *J. Am. Chem. Soc.*, 2020, 142, 13102–
 13111; (c) A. M. Messinis, J. C. A. Oliveira, A. C. Stückl and
 L. Ackermann, *ACS Catal.*, 2022, 12, 4947–4960; (d)
 S. L. Daifuku, M. H. Al-Afyouni, B. E. R. Snyder,

J. L. Kneebone and M. L. Neidig, *J. Am. Chem. Soc.*, 2014, **136**, 9132–9143; (e) R. B. Bedford, P. B. Brenner, E. Carter, J. Clifton, P. M. Cogswell, N. J. Gower, M. F. Haddow, J. N. Harvey, J. A. Kehl, D. M. Murphy, E. C. Neeve, M. L. Neidig, J. Nunn, B. E. R. Snyder and J. Taylor, *Organometallics*, 2014, **33**, 5767–5780; (f) Y. Liu, M. Shi and L. Deng, *Organometallics*, 2014, **33**, 5660–5669; (g) T. E. Boddie, S. H. Carpenter, T. M. Baker, J. C. DeMuth, G. Cera, W. W. Brennessel, L. Ackermann and M. L. Neidig, *J. Am. Chem. Soc.*, 2019, **141**, 12338–12345.

- 5 (a) S. D. Ittel, C. A. Tolman, A. D. English and J. P. Jesson, J. Am. Chem. Soc., 1976, 98, 6073–6075; (b) M. V. Baker and L. D. Field, J. Am. Chem. Soc., 1986, 108, 7433–7434.
- 6 (a) R. Pony Yu, D. Hesk, N. Rivera, I. Pelczer and P. J. Chirik, *Nature*, 2016, **529**, 195–199; (b) J. Corpas, P. Viereck and P. J. Chirik, *ACS Catal.*, 2020, **10**, 8640–8647.
- 7 S. Garhwal, A. Kaushansky, N. Fridman, L. J. W. Shimon and G. d. Ruiter, *J. Am. Chem. Soc.*, 2020, **142**, 17131–17139.
- 8 W. Li, J. Rabeah, F. Bourriquen, D. Yang, C. Kreyenschulte, N. Rockstroh, H. Lund, S. Bartling, A.-E. Surkus, K. Junge, A. Brückner, A. Lei and M. Beller, *Nat. Chem.*, 2022, 14, 334–341.
- 9 (a) J. Zhang, S. Zhang, T. Gogula and H. Zou, ACS Catal., 2020, 10, 7486–7494; (b) S. Kopf, H. Neumann and M. Beller, Chem. Commun., 2021, 57, 1137–1140; (c) F. Bourriquen, N. Rockstroh, S. Bartling, K. Junge and M. Beller, Angew. Chem., Int. Ed., 2022, 61, e202202423.
- 10 A. Tlahuext-Aca and J. F. Hartwig, ACS Catal., 2021, 11, 1119– 1127.
- 11 (a) J. H. Docherty, J. Peng, A. P. Dominey and S. P. Thomas, *Nat. Chem.*, 2017, 9, 595–600; (b) L. Britton, J. H. Docherty, A. P. Dominey and S. P. Thomas, *Molecules*, 2020, 25, 905; (c) L. Britton, M. Skrodzki, G. S. Nichol, A. P. Dominey, P. Pawluć, J. H. Docherty and S. P. Thomas, *ACS Catal.*, 2021, 11, 6857–6864.
- 12 M. Farizyan, A. Mondal, S. Mal, F. Deufel and M. van Gemmeren, *J. Am. Chem. Soc.*, 2021, **143**, 16370–16376.
- 13 Presumed to be a result of a lower concentration of base present in the reaction mixture.

- 14 (a) J. Zhou and J. F. Hartwig, Angew. Chem., Int. Ed., 2008, 47, 5783–5787; (b) M. Hatano, T. Nishimura and H. Yorimitsu, Org. Lett., 2016, 18, 3674–3677.
- 15 (a) B. Rybtchinski, R. Cohen, Y. Ben-David, J. M. L. Martin and D. Milstein, *J. Am. Chem. Soc.*, 2003, **125**, 11041–11050;
 (b) A. Di Giuseppe, R. Castarlenas, J. J. Pérez-Torrente, F. J. Lahoz and L. A. Oro, *Chem.-Eur. J.*, 2014, **20**, 8391–8403.
- 16 (a) G. Erdogan and D. B. Grotjahn, J. Am. Chem. Soc., 2009,
 131, 10354–10355; (b) S. K. S. Tse, P. Xue, Z. Lin and G. Jia,
 Adv. Synth. Catal., 2010, 352, 1512–1522; (c) A. Bechtoldt and L. Ackermann, ChemCatChem, 2019, 11, 435–438.
- 17 N. Camedda, A. Serafino, R. Maggi, F. Bigi, G. Cera and G. Maestri, *Synthesis*, 2020, 52, 1762–1772.
- 18 (a) C. P. Lenges, M. Brookhart and B. E. Grant, *J. Organomet. Chem.*, 1997, **528**, 199–203; (b) J. Choi, L. Tang and J. R. Norton, *J. Am. Chem. Soc.*, 2007, **129**, 234–240.
- 19 (a) H. Fong, M.-E. Moret, Y. Lee and J. C. Peters, Organometallics, 2013, 32, 3053-3062; (b) D. Zell, M. Bursch, V. Müller, S. Grimme and L. Ackermann, Angew. Chem., Int. Ed., 2017, 56, 10378-10382; (c) J.-F. Li, Z.-Z. Wei, Y.-Q. Wang and M. Ye, Green Chem., 2017, 19, 4498-4502.
- 20 T. R. Puleo, A. J. Strong and J. S. Bandar, *J. Am. Chem. Soc.*, 2019, **141**, 1467–1472.
- 21 J. M. Bellerby, M. J. Mays and P. L. Sears, *Dalton Trans.*, 1976, 1232–1236.
- 22 (a) J. R. Fulton, S. Sklenak, M. W. Bouwkamp and R. G. Bergman, *J. Am. Chem. Soc.*, 2002, **124**, 4722–4737; (b) X.-J. Dai and C.-J. Li, *J. Am. Chem. Soc.*, 2016, **138**, 5433–5440.
- 23 T. Dombray, C. G. Werncke, S. Jiang, M. Grellier, L. Vendier, S. Bontemps, J. B. Sortais, S. Sabo-Etienne and C. Darcel, *J. Am. Chem. Soc.*, 2015, 137, 4062–4065.
- 24 M. V. Baker, L. D. Field and D. J. Young, *Chem. Commun.*, 1988, 546–548.
- 25 R. M. Spycher, L. Hausherr-Primo, G. Grassi and A. Bauder, J. Mol. Struct., 1995, 351, 7–17.
- 26 T. Morikita, M. Hirano, A. Sasaki and S. Komiya, *Inorg. Chim. Acta*, 1999, **291**, 341–354.
- 27 S. Komiya, N. Oyasato and T. Furukawa, *Bull. Chem. Soc. Jpn.*, 1989, **62**, 4078–4079.