

Cite this: *Chem. Commun.*, 2019, 55, 2293Received 12th December 2018,
Accepted 28th January 2019

DOI: 10.1039/c8cc09853k

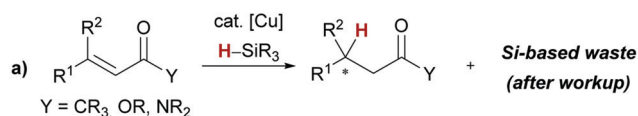
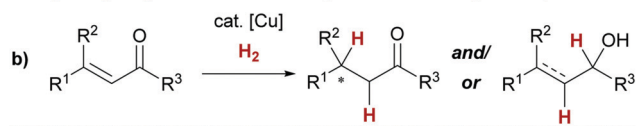
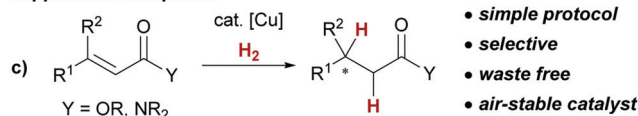
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A simple and air-stable copper(i)/N-heterocyclic carbene complex enables the catalytic hydrogenation of enoates and enamides, hitherto unreactive substrates employing homogeneous copper catalysis and H₂ as a terminal reducing agent. This atom economic transformation replaces commonly employed hydrosilanes and can also be carried out in an asymmetric fashion.

One of the main challenges of contemporary method development for synthetic chemistry is the development of atom economic and sustainable transformations.^{1,2} In this vein, catalytic hydrogenations are much desired reactions, as they serve to replace complex and waste-generating reducing agents such as borohydrides, aluminium hydrides or hydrosilanes.^{3,4} Catalytic reactions involving copper hydride intermediates⁵ serve as a prime example for this challenge: commonly, hydrosilanes are used as stoichiometric reducing agents, whereas the use of dihydrogen (H₂) for the – considerably more atom economic – generation of the desired copper hydride complexes has been much less explored.^{6,7} The hallmark reaction of the so-called copper hydride catalysis⁵ is arguably the conjugate reduction of α,β -unsaturated carbonyl or carboxyl compounds employing hydrosilanes (Scheme 1a). Even though the replacement of hydrosilanes with H₂ has been attempted for the reduction of enones, the chemoselectivity of the 1,2- vs. the 1,4-reduction turned out to be challenging to control (Scheme 1b).⁶ Furthermore, the corresponding enoates emerged as too unreactive in combination with H₂.^{8,9} Enoates are common substrates for the catalytic hydrogenation with precious metals such as rhodium and ruthenium,^{3,10} and replacement with base metal catalysts such as copper complexes is highly desirable.

We herein report the first copper-catalysed conjugate reduction of previously unreactive α,β -unsaturated esters and amides with H₂ employing well-defined and air-stable copper(i)/N-heterocyclic

Catalytic hydrogenation of α,β -unsaturated carboxylic acid derivatives using copper(i)/N-heterocyclic carbene complexes†

Birte M. Zimmermann,^{1b} Sarah C. K. Kobosil and Johannes F. Teichert^{1b*}*previous work:**catalytic conjugate reduction of enones and enoates with hydrosilanes (generates Si-based waste)**catalytic hydrogenation of enoates (chemoselectivity issues)**This work: catalytic hydrogenation of enoates with well-defined copper/NHC complexes*

Scheme 1 Challenges in atom-economic copper-catalysed conjugate reductions.

carbene complexes¹¹ (Scheme 1c). Additionally, this approach circumvents the use of waste-generating hydrosilanes, resulting in a simple and sustainable protocol for the catalytic hydrogenation of α,β -unsaturated esters and amides.

The copper-catalysed hydrogenation of enoates was optimised employing ethyl β -methyl cinnamate (**1**, Table 1). Using sodium *tert*-butanolate (NaOtBu) as additive for the generation of the key Cu–O-bond for H₂ activation,¹² we investigated common copper(i)/NHC complexes **3–7** at 90 bar H₂ and 60 °C in THF (Table 1, entries 1–5). From these experiments, mesityl-derived complexes **3** and **6** emerged as most active, as judged by the conversion of **1** to **2**. Notably, no other side-products were observed. We found that the amount of NaOtBu could be lowered to 30 mol% while maintaining full conversion, however, even smaller amounts of the additive (10 mol%) led to an almost complete halt of the reaction (Table 1, entries 6 and 7). Lowering the H₂ pressure to 50 bar at these limiting conditions led to diminished conversion

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† Electronic supplementary information (ESI) available: Experimental procedures, characterisation and NMR spectra. See DOI: 10.1039/c8cc09853k



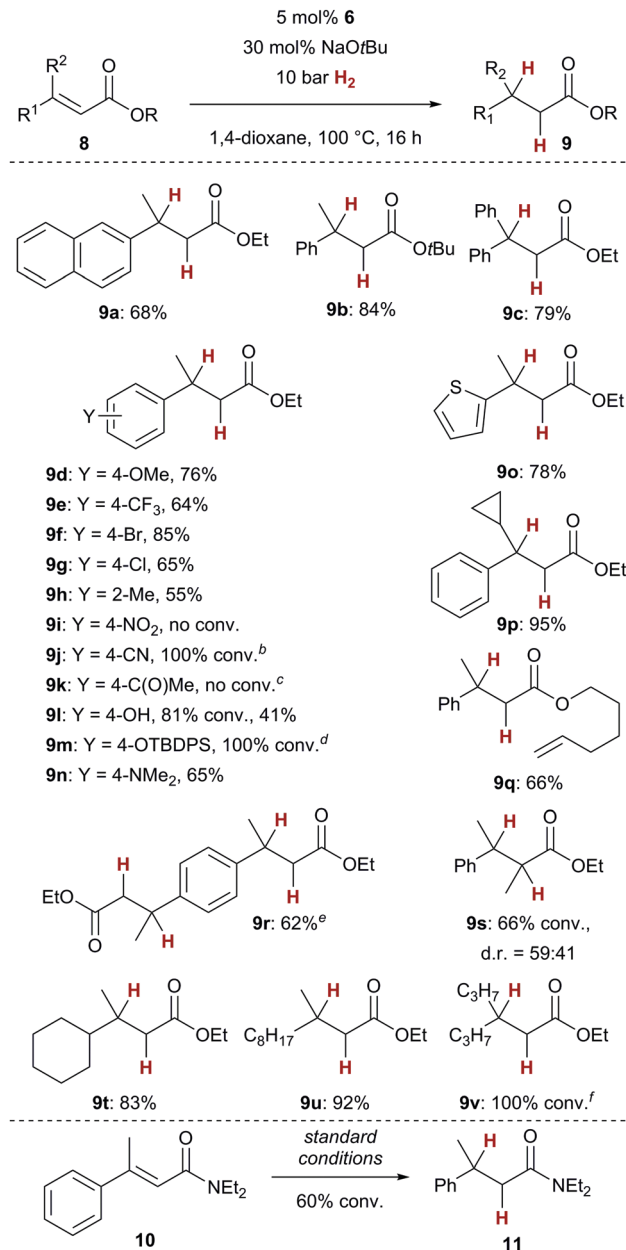
Table 1 Cu-catalysed hydrogenation of enoates, optimisation^a

| Entry | [Cu] | Conditions | Conv. ^b |
|-------|------|---|--------------------|
| 1 | 3 | 110 mol% NaOtBu, 90 bar H ₂ , THF, 60 °C | > 95% |
| 2 | 4 | As entry 1 | 67% |
| 3 | 5 | As entry 1 | 17% |
| 4 | 6 | As entry 1 | > 95% |
| 5 | 7 | As entry 1 | 41% |
| | | | |
| 6 | 3 | 30 mol% NaOtBu, 90 bar H ₂ , THF, 60 °C | > 95% |
| 7 | 3 | 10 mol% NaOtBu, 90 bar H ₂ , THF, 60 °C | 6% |
| 8 | 3 | 30 mol% NaOtBu, 50 bar H ₂ , THF, 60 °C | 32% |
| 9 | 3 | 30 mol% NaOtBu, 50 bar H ₂ , 1,4-dioxane, 60 °C | > 95% |
| 10 | 3 | 30 mol% NaOtBu, 10 bar H ₂ , 1,4-dioxane, 60 °C | 5% |
| 11 | 3 | 30 mol% NaOtBu, 10 bar H ₂ , 1,4-dioxane, 100 °C | > 95% |
| 12 | 6 | 30 mol% NaOtBu, 10 bar H ₂ , 1,4-dioxane, 60 °C | 79% |
| 13 | 6 | 30 mol% NaOtBu, 10 bar H ₂ , 1,4-dioxane, 100 °C | > 95% ^c |
| 14 | 6 | 30 mol% NaOtBu, 1 bar H ₂ , 1,4-dioxane, 100 °C | 32% |

^a Reactions were carried out on a 0.25 mmol scale. ^b Determined by ¹H NMR spectroscopy. ^c 71% isolated yield.

of 32% (Table 1, entry 8). A subsequent solvent optimisation revealed 1,4-dioxane as optimal,¹³ with full conversion of **1** reached at lowered H₂ pressure of 50 bar (Table 1, entry 9). When comparing catalysts **3** and **6** at 10 bar H₂ pressure, imidazolium-based copper complex **6** turned out to be more active (Table 1, entries 10 vs. 12), even though no reactivity difference could be detected at higher temperature of 100 °C. The investigation of the substrate scope was therefore carried out with catalyst **6**. Notably, lowering the H₂ pressure with catalyst **6** to 1 bar still led to a detectable conversion of **1** (32% conv. Table 1, entry 14).

With optimised reaction conditions in hand, we set out to investigate the substrate scope of the copper-catalysed conjugate reduction of α,β -unsaturated carboxylic acid derivatives **8** (Scheme 2). We found that the catalytic hydrogenation could be applied to a variety of aryl or alkyl substituted enoates **8**. Next to the successful generation of simple naphthyl derivate **9a**, also the sterically more demanding *tert*-butyl ester **9b** as well as diphenyl-substituted ester **9c** could be furnished in good yields (68–84%). Both electron donating and withdrawing groups were tolerated as substituents of the cinnamic acid derivatives **9d–9h** with similar results in terms of yield. Notably, no protodehalogenation was observed with bromide **9f** or chloride **9g**. As in previous studies,⁷ the presence of a nitro or a carbonyl group (**9i**, **9k**) led to no or diminished conversion, whereas nitrile derivative **9j** led to full conversion. Of note is the fact that protic groups such as a free phenol (**9l**) were susceptible to the



Scheme 2 Cu-catalysed hydrogenation of enoates and enamides, scope^a. ^a If not noted otherwise, *E*-configured enoates **8** have been employed. *E/Z* ratios: **8f**: 90:10, **8i**: 88:12, **8k**: 87:13, **8m**: 93:7, **8u**: 77:23. ^b The product decomposed during purification, 5% isolated yield. ^c No 1,2-reduction of the ketone was observed. ^d A mixture of the corresponding ethyl and *tert*-butyl ester were observed (83:17). The separation turned out to be tedious, resulting in lower yields (OEt: 40%, OtBu: 7%). ^e 10 mol% **6**, 60 mol% NaOtBu used. ^f Product is volatile.

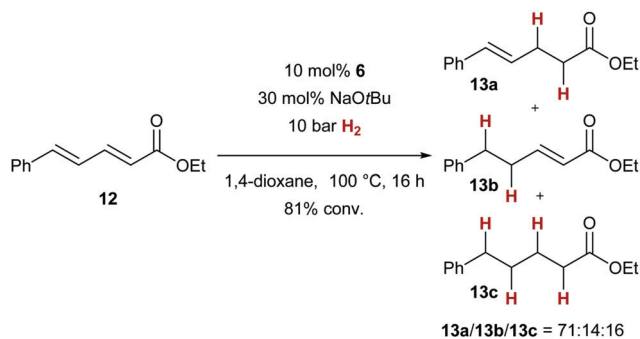
conjugate reduction.¹⁴ Also, the silyl-protected variant (**9m**) could be converted successfully. Dimethylaniline **9n** and thiophene-derived **9o** as possibly coordinating substrates could be successfully hydrogenated. The generation of cyclopropane-substituted **9p** in almost quantitative yield (95%) gives an important indication that no carbon-based radical is involved in the overall process. The clean formation of alkene-substituted ester **9q** underscores the chemoselectivity of the present catalyst,



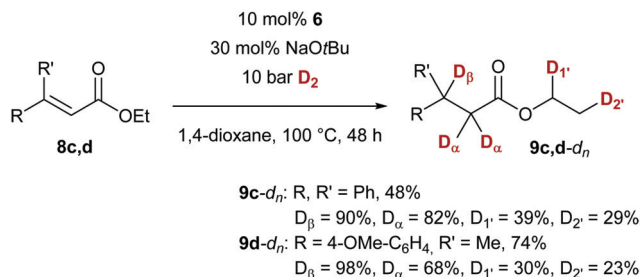
as no alkene hydrogenation was observed.¹⁵ Next to the successful formation of diester **9r**, also **9s**, bearing a methyl group in α -position, which generally slows down the conjugate addition in other copper-catalysed processes,¹⁶ was turned over by the catalyst. Ester **9s** was formed as a 59:41 mixture of diastereomers. Finally, dialkyl-substituted enoates **9t–9v** could be converted with similarly good results in terms of yield. We could demonstrate that even enamide **10**, which generally is too electron-rich for reactivity in other copper-catalysed conjugate addition reactions,¹⁷ displays some reactivity with our catalyst (60% conv., 17% yield for **11**). This result underscores the fact that the copper/NHC complexes employed in this transformation serve a key role for the generation of copper hydride intermediates with higher reactivity in comparison to the commonly used phosphine complexes.⁵ The latter could not realize any conversion of carboxylic acid derivatives.

Sorbic acid derivative **12** gave a mixture of 1,4- and 1,6-addition products, with unsaturated ester **13a** (from 1,4-addition of the hydride nucleophile) as the major product (Scheme 3). The formation of **13b** and **13c** can be explained by prior 1,6-addition of the copper hydride, and, in the case of **13c**, subsequent 1,4-addition.

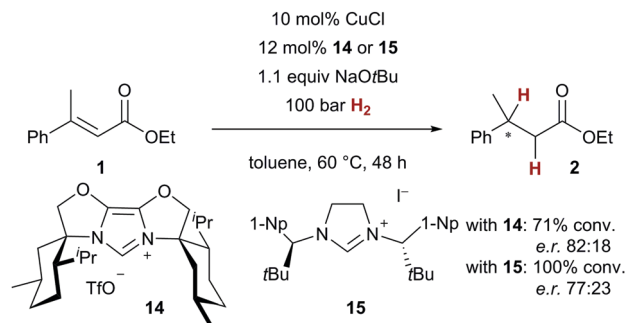
To gain some insight into the mechanism of the present protocol, we carried out the conjugate reduction in the presence of deuterium gas (D_2 , Scheme 4). With ethyl esters **8c** and **8d**, as expected for a hydride transfer reaction, the deuterium incorporation in the β position was high ($\geq 90\%$ D), but also significant isotope labeling in the α position was found, indicating an enolisation process with *t*-BuOD after the conjugate addition. Unexpectedly, also deuterium incorporation was observed in the ethyl ester (23–39% D, D_1' and D_2').¹⁸



Scheme 3 Conjugate reduction of an $\alpha,\beta,\gamma,\delta$ -unsaturated ester.



Scheme 4 Cu-catalysed conjugate reduction with D_2 .



Scheme 5 Asymmetric conjugate reduction of cinnamate **1** (1-Np = 1-naphthyl).

As an asymmetric variant of catalytic conjugate reduction with H_2 is highly attractive,¹⁶ we turned our attention to chiral NHC ligands. In preliminary experiments, we were able to demonstrate that indeed stereoselection with chiral NHC ligands is possible, even though more drastic reaction conditions (100 bar H_2 , 48 h reaction time) had to be employed with chiral NHC precursors **14**¹⁹ and **15**²⁰ (Scheme 5). The corresponding reduced ester **2** was obtained with an enantiomeric ratio of up to 82:18 with chiral NHC precursor **14**, demonstrating the viability of an asymmetric reaction.

In summary, we have developed a simple and efficient copper-catalysed conjugate reduction of enoates and enamides with H_2 . Well-defined and easily accessible copper(I)/NHC complexes are employed as catalysts, giving the desired products which had so far not been accessible with copper/phosphine complexes due to the lower reactivity of these substrates employing H_2 . The present catalytic protocol serves as an atom economic alternative to replace the commonly used and waste-generating hydrosilanes as stoichiometric reducing agents with dihydrogen (H_2). Isotope labelling studies indicate a polar mechanism involving a copper hydride intermediate. In addition, the viability of an asymmetric reaction has been demonstrated.

This work was supported by the German Research Council (DFG, Emmy Noether Fellowship for J. F. T., TE1101/2-1), by the Fonds der Chemischen Industrie (Liebig-Stipendium for J. F. T.) and by the Daimler and Benz Foundation (postdoctoral fellowship for J. F. T.). Prof. Dr Martin Oestreich (TU Berlin) is kindly thanked for generous support.

Conflicts of interest

There are no conflicts to declare.

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