Green Chemistry



COMMUNICATION

View Article Online
View Journal | View Issue



Cite this: *Green Chem.*, 2016, **18**, 686

Received 21st August 2015, Accepted 21st September 2015

DOI: 10.1039/c5gc01965f

www.rsc.org/greenchem

Attractive aerobic access to the α , β -unsaturated acyl azolium intermediate: oxidative NHC catalysis *via* multistep electron transfer†

L. Ta,‡ A. Axelsson‡ and H. Sundén*

To replace high molecular weight oxidants with air (O_2) we introduce multistep electron transfer NHC catalysis. The method provides a general and selective oxidation of the α,β -unsaturated aldehyde derived homoenolate to the synthetically useful α,β -unsaturated acyl azolium intermediate. Several independent oxidative NHC-catalyzed reactions are viable with this strategy and the products can be isolated in high to excellent yields.

In the vibrant field of N-heterocyclic carbene (NHC) catalysis¹ oxidative reaction paths have gained widespread attention in recent years. Several synthetic methodologies have emerged relying on the addition of a stoichiometric oxidant.² Of particular importance is the *in situ* oxidation of the homoenolate to the α,β -unsaturated acyl azolium (Scheme 1).

As a reaction intermediate, the unsaturated acyl azolium has been exploited in, for example, oxidative esterifications,³ macrocyclization, annulation reactions and cyclo-additions creating possibilities for broad molecular complexity. However, the majority of these reactions rely on the addition of a high molecular weight oxidant such as Kharasch oxidant 1.7,8 Taking into account the price, separation and disposal of waste (hydroquinone) of the oxidant it is clear that oxidants, such as 1, present a significant obstacle in terms of scale-up, sustainability and economy. A more atom efficient strategy would be to use environmentally friendly oxidants such as air (O2).9 It is highly desirable to use O2 as a terminal oxidant since it is inexpensive and non-toxic, and the reaction shows a high efficiency per weight of oxidant, with formation of water as the sole byproduct. In oxidative NHC catalysis, direct oxidations with O2 involving α,β-unsaturated aldehydes are reported to give carboxylic acids (Scheme 2a). 10 Furthermore, introduction of a catalytic oxidant, FeCl₃ 11 or Ru(bpz)₃(PF₆)₂, 12 enables aerobic oxidative esterifications (Scheme 2b). However, reactive carbons

reactive carbons

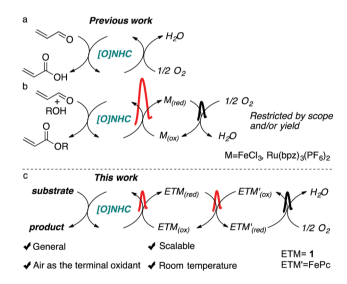
reactive carbons

reactive carbons

reactive carbons

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Scheme 1 Oxidation of the homoenolate to the $\alpha,\beta\text{-unsaturated}$ acyl azolium.



Scheme 2 a) Direct oxidation with O_2 . (b) High energy barrier for the catalytic aerobic oxidative esterification. (c) Aerobic multistep electron transfer NHC catalysis, reliant on the combination of ETMs, lowers the energy barrier for oxidations with O_2 .

Chemistry and Chemical Engineering, Chalmers University of Technology, Kemivägen 10, 412 96 Göteborg, Sweden. E-mail: sundenh@chalmers.se † Electronic supplementary information (ESI) available: Experimental procedures, and ¹H NMR, ¹³C NMR, and ¹⁹F NMR data. See DOI: 10.1039/c5gc01965f

‡These authors contributed equally

these reactions are hampered by a high energy barrier for the oxidation of the NHC-aldehyde-adduct to the acyl azolium, reflected in high reaction temperatures, long reaction times or **Green Chemistry** Communication

low reactivity. Clearly, mild and efficient protocols for aerobic oxidations accessing the unsaturated acvl azolium intermediate would provide new possibilities for green chemistry and would therefore be most desirable.

High activation barriers are a general problem for direct catalytic oxidation with air or pure O2. This often results in unselective reactions with kinetic byproducts, 13 thus making the general use of oxygen problematic as a terminal oxidant. A possible way to circumvent this is to mimic the respiratory chain and to create a low energy path for electrons to flow from the substrate to oxygen.14 This has been envisioned and accomplished in transition metal-catalysis by introducing one or several electron transport mediators (ETMs) between the substrate specific catalyst and O2. The most prominent example is the Wacker oxidation where CuCl2 acts as an ETM, shuffling electrons from Pd⁰ to O₂. ^{13,15} Bäckvall and coworkers¹³ have extended the generality of this strategy to a broad range of aerobic transition metal-catalyzed reactions such as 1,4-oxidation of 1,3-dienes, 16 oxidative carbocyclization of allene-substituted olefins¹⁷ and aerobic oxidations of alcohols.18 Others have adapted this principle for a variety of oxidation reactions.19

Here we present, as a part of our interest in NHC catalysis, 20 a strategy for oxidative NHC-catalyzed reactions relying on an electron transport chain involving two ETMs that enables efficient aerobic access to the synthetically useful α,β-unsaturated acyl azolium intermediate (Scheme 2c).

As a model reaction we choose to investigate the oxidative esterification of α,β -unsaturated aldehydes (Table 1).²¹

α,β-Unsaturated aldehydes, in the presence of NHCs, are known to undergo several distinct reactions to give, for example, saturated esters (4),22 unsaturated carboxylic acids (5), 10a,b and γ -butyrolactones (6). 23 Thus, selectivity towards the unsaturated ester 3 would give a good measure of the selectivity in the oxidation event. The initial screening, in open reaction vessels, revealed that quinone 1 alone was indeed possible to use in substoichiometric quantities²⁴ in the oxidative esterification of aldehyde 2. The reaction gave 3 in 24% yield after 7 h along with several byproducts from competing reaction paths (entry 2). To test our hypothesis of the possibility to combine oxidative NHC catalysis and a coupled system of ETMs for electron transfer, we investigated the effect of an additional ETM. A drastic improvement was achieved when performing the reaction in the presence of 1 and iron phthalocyanine (FePc [0.0055 equiv.]) with almost full conversion after 4 h (entry 3). Notably, the reaction proceeds with a slightly better efficiency compared to the stoichiometric reaction (entry 1). Further screening revealed that quinone 1 could be replaced with 2,6-di-tert-butylphenol (7) with equivalent efficiency (85% yield). Phenol 7 is the precursor in the synthesis of 1 and under our reaction conditions 1 is readily formed in situ. Additionally, 7 is a much cheaper reagent than 1 and therefore became the ETM-precursor of choice (entry 4). Thereafter, different NHC precatalysts were investigated (Table 1, entries 5-7) with A proving superior as a precatalyst. Furthermore, the loading of the NHC precatalyst is important:

Table 1 NHC-catalyzed esterification aerobic oxidative cinnamaldehyde^a

Entry	NHC	ETM	ETM' (FePc, eq.)	Time (h)	Yield ^b (%)	Selectivity ^c (%)
1	Α	1 (1 eq.)	_	4	74	78
2	A	1 (0.01 eq.)	_	7	24	40
3	A	1 (0.01 eq.)	0.0055	4	80	88
4	A	7 (0.02 eq.)	0.0055	4	85	89
5	В	7 (0.02 eq.)	0.0055	_	_	_
6	C	7 (0.02 eq.)	0.0055	22	65	65
7	D	7 (0.02 eq.)	0.0055	4	68	68
8	\mathbf{A}^d	7 (0.02 eq.)	0.0055	4	19	19
9	\mathbf{A}^e	7 (0.02 eq.)	0.0055	4	67	74
A N) B	S c N	Mes L-Cl Mes	+ Mes N - Cl 7	tBu OH	∕tBu

^aThe reactions were performed in open reaction vessels at room temperature (r.t) in MeCN (1 mL) with 2 (66.1 mg, 0.5 mmol, 1 eq.), TBD (34.8 mg, 0.25 mmol, 0.5 eq.), MeOH (0.0809 ml, 2 mmol, 4 eq.), NHC (2.3 mg, 0.01 mmol, 0.02 eq.), ETM (see Table) and ETM' (see table). b Yield determined by NMR against the internal standard. c Yield over conversion. d 0.2 eq. of NHC was used. e 0.05 eq. of NHC was used. TBD = 1,5,7-triazabicyclo[4.4.0]dec-5-ene.

loadings higher than 0.02 equiv. result in a lowered reaction efficiency (compare entry 4 with entries 8 and 9).

Mechanistically, elimination experiments show that reactions performed without 7 stop after 100 minutes at 40% yield (Fig. 1). If FePc is removed from the reactions a similar trend is observed. Without 7 or FePc the reaction reaches 30% yield and then stops. In all control experiments full conversion of the starting material can be noted indicating competing background reactions. In the reactions performed without 7 a possible background reaction is the formation of a peracid. Indirect evidence of this is the isolation of the carboxylic acid after aqueous work up. This is in agreement with previous reports that suggest that the homoenolate intermediate (Fig. 2, II) is oxidized in the presence of O₂ to a peroxo-species (III) rather than to the acyl-azolium intermediate (IV). 10,12 Conversely, in the presence of both 7 and FePc, a low energy path is created so that the homoenolate intermediate (II) is oxidized by quinone 1 to the acyl azolium (IV). Intermediate IV reacts with the alcohol to give the unsaturated ester V regenerating the NHC. The proposed redox cascade is rationalized by elimination experiments and the decreasing redox potential of the three oxidants, O_2 , FePc $(E = +0.74 \text{ V } \text{vs. SCE})^{25}$ and 1 $(E = -0.52, -0.89 \text{ V } \nu s. \text{ SCE}).^{26,27}$

Having found our optimal reaction conditions for the oxidative esterification of α,β-unsaturated aldehydes the scope

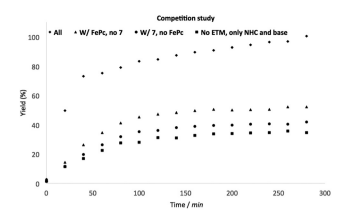


Fig. 1 Kinetic profile of the oxidative esterification of 2 to 3. See the ESI† for experimental details.

Fig. 2 Catalytic cycle for the aerobic oxidative esterification of aldehydes

of the reaction was investigated (Table 2). The reaction proceeds in good to excellent yields for a range of methyl cinnamates incorporating both electron donating and electron withdrawing substituents (entries 1-7). For example, halogen substituents are well tolerated on the aromatic ring and both 4-chloro and 4-fluoro methyl cinnamates can be isolated in 87% and 73% yield, respectively (entries 2 and 3). The reaction is also compatible with a broad range of alcohols yielding the corresponding α,β-unsaturated esters in high to excellent yields (72-98% yield, entries 8-16). For example, allyl and benzyl alcohols give their corresponding esters in 78% and 98% yield, respectively (entries 10 and 11). Moreover, methoxycinnamates such as amiloxate 19 and octinoxate 20, two out of several sunscreening agents that are based on a 4-methoxycinnamate core-structure, were both isolated in 72% yield.²⁸ A diol such as 1,4-butanediol was efficiently converted to the unsaturated ester 22 in 74% yield with only traces of the diacylated byproduct found in the reaction mixture. Aliphatic $\alpha,\!\beta\text{-unsaturated}$ aldehydes proved to be difficult substrates and 2-hexenal and citral gave sluggish reactions. Nonetheless, conjugated aldehydes worked well and sorbic aldehyde provided ester 23 in 69% yield. Furthermore, the reaction is scalable and compound 12 could be isolated in 1.5 g, 95% yield

Table 2 Reaction scope of the oxidative esterification

$$R^{1}$$
 O + $R^{2}OH$ A , 7, FePc, TBD R^{1} OR^{2}

Entry	Product	R^1	\mathbb{R}^2	Yield ^b
1	3	, r. r.		87%
2	8	`zst CI	-ξ-Me	87%
3	9	F	-ξ-Me	73%
4	10	in the second se	-ξ-Me	72%
5	11	OMe	-ξ-Me	88%
6	12	O ₂ N	-ξ-Me	64%
7	13	جَحْ (NO ₂	-ξ-Me	79%
8	14	, ref	·\$<	89%
9	15	OMe		93%
10	16	OMe	72~//	78%
11	17) Z ^S OMe	^{ZZ} Ph	98%, 95% ^c
12	18	`z ^z	¸₹ ^{\$} ∕∕^Ph	96%
13	19	`z ^z	`z ^z .	72%
14	20	OMe	zs.	72%
15	21) Jef	; s O O	76%
16	22	re de la companya de	Ş [₹] ∕∕∕OH	74%
17	23	`\$ ^{\$}	جُحْبِ Ph	69%

^a See the ESI for experimental details. ^b Isolated yields. ^c 1.5 g isolated yield.

(entry 11). In addition, with regard to the oxidants, this reaction requires only 26.7 mg of 7 and 18.9 mg of FePc. In comparison, anaerobic stoichiometric oxidation would need at least 2.48 g of quinone 1 to achieve a similar result. The oxidative esterifications run very clean, none of the expected byproducts can be detected in crude NMR, which simplifies purification. It also signifies the very high selectivity in each catalytic step of this multi-component reaction. Moreover, the method provides a mild direct access to unsaturated esters from aldehydes, which circumvents oxidation to the carboxylic acid that traditionally is the starting point in ester formation. Recycling experiments showed that addition of new starting materials to the reaction vessel was successful and compound 3 could be obtained in average 71% yield over three cycles (see the ESI†). These results confirm that the catalytic system is not degraded over the course of the reaction and that it would be possible to design a system for recover and reuse.

Encouraged by the efficiency of our simple aerobic multistep electron transfer NHC catalysis the reaction was further examined for compatibility with other reactions such as conjugate additions. Accordingly, 1,3-ketoesters and 1,3-diketones were investigated as nucleophiles in combination with cinnamaldehyde (Scheme 3). The approach proved viable and lactones 24–26 could be isolated in 92, 80 and 82% yield, respectively. These reactions require a slightly higher loading of both ETMs and NHC but instead the loading of the base can be reduced.

In summary we have developed a new strategy relying on ETMs for aerobic oxidative NHC-catalyzed reactions. The *in situ* oxidation of the homoenolate to the unsaturated acyl azolium is here exemplified with oxidative esterifications and oxidative Michael-additions of 1,3-dicarbonyl compounds to α,β -unsaturated aldehydes. The reactions have a broad substrate scope and the products can be isolated in good to excellent yields. The use of O_2 in the form of air as the terminal oxidant offers an environmentally friendly and inexpensive way to scale-up the important chemistry of oxidative NHC catalysis. In the lab we are currently exploring other forms of aerobic oxidative multistep electron transfer NHC-catalyzed reactions.

Scheme 3 Syntheses of lactones **24–26**. See the ESI† for experimental details. Yields refer to the isolated product.

Acknowledgements

This work was generously supported by the Swedish Research Council VR and FORMAS, Magnus Bergvalls stiftelse and Chalmers Areas of Advance Nano.

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