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Simple metal salts supported on montmorillonite as recyclable catalysts for intramolecular hydroalkoxylation of double bonds in conventional and VOC-exempt solvents†

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We describe herein an efficient and particularly sustainable catalytic system for the intramolecular hydroalkoxylation of double bonds. A heterogeneous catalyst based on the impregnation of benign metals such as iron and bismuth on montmorillonite was used for a highly atom-economic transformation in DMC, a non-VOC solvent. The transformation allowed the formation of a large range of cyclic ethers from the corresponding unsaturated alcohols and the catalyst could be recycled several times.

Intramolecular hydroalkoxylation of olefins is the most straightforward and atom-economic route to cyclic ethers. The reaction can be promoted or catalysed by a large variety of Brønsted or Lewis acids under homogeneous conditions^{1,2} or with heterogeneous catalysts such as zeolites.^{3,4} The methodology has proven its utility in the synthesis of fine chemicals including bioactives^{5–7} and fragrant molecules,^{8,9} for example (Fig. 1).

To go beyond the laboratory towards large-scale applications, an ideal catalyst should however not only be efficient in terms of yields and selectivity, but also offer recycling capabilities. To achieve this goal, various strategies have been developed including, for example, the use of solid acid catalysts¹⁰ such as zeolites^{11–14} or acidic resins (Amberlyst, Nafion), ^{15,16} attachment of metal complexes or salts to a solid support such as silica^{17,18} or Nafion¹⁹ by physisorption, or by covalent linkage of one or more ligands.²⁰ The non-covalent linkage of catalysts onto solid supports has been developed with success through "catch and release" strategies.²¹ Recently, supported metal nanoparticles (*e.g.* gold nanoparticles) have appeared in the literature as efficient catalysts in various chemical processes.^{22–26}

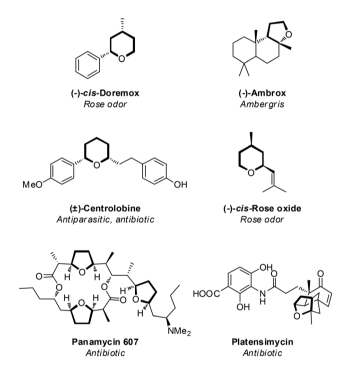


Fig. 1 Fine chemicals featuring cyclic ether frameworks obtained by intramolecular double bond hydroalkoxylation.

We have been looking for a cheap and readily available unprocessed solid material to support simple metal salts and use these supported metals as catalysts for the intramolecular hydroalkoxylation of double bonds. In this regard, montmorillonite (MMT) appeared very appealing to us to provide supported metal catalysts with limited footprints among other advantages as supporting material.^{27,28} We previously reported the use of MMT-supported gallium species as a catalyst for 1,6-enynes cycloisomerisation.²⁹

The catalysts were prepared by impregnation of solutions of sustainable metal salts such as BiCl₃, FeCl₃, and CuCl₂ in MeOH. Under an inert atmosphere, 10 mmol of an anhydrous precursor was added to 10 g of pre-lyophilised MMT and

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50 mL of MeOH. The mixture was stirred vigorously for 4 hours, filtered and then triturated in a minimum amount of methanol. The material was dried under vacuum for 24 h and stored under an inert atmosphere. Characterisation of the catalysts involved ICP-MS titration of the metal contents, indicating 4.87% w/w for Bi-MMT, 2.67% w/w for Fe-MMT, and 2.04% w/w for Cu-MMT. Small angle XRD analysis confirmed the insertion of metal cations in the MMT interlayers. In the case of Bi-MMT, signals attributed to crystals of BiOCl, presumably formed upon hydrolysis within the MMT, could be observed. Those cations are under their most stable oxidation states (Bi³⁺, Fe³⁺/Fe²⁺ (53/47) and Cu²⁺) as witnessed by XPS analysis (see ESI†).

We started our study with a screening of the reaction conditions and solvents including conventional and green solvents such as dimethyl carbonate (DMC) and 2-methyltetrahydrofuran (Table 1). DMC is particularly attractive since it is considered non-VOC thanks to its low vapour pressure (2400 Pa, to be compared to 2340 for water and 29000 for toluene), and have good solvent properties (bp = 90 °C, δ = 3.1, μ = 0.91 D).³⁰

DMC and nitromethane appeared as the most suitable solvents for the model reaction of olefinic alcohol 1a converted with total Markovnikov regioselectivity to cyclic ether 1b. With Bi-MMT, yields beyond 75% in 5 hours at 80 °C were obtained (entries 1 and 2). The reaction in acetonitrile needed prolonged reaction times (24 h, entry 3) to proceed efficiently, while the other tested solvents gave moderate results within the same reaction time (30-70%, entries 4-7).

Table 1 Solvents and conditions of screening^a

Entry	M	Solvent	T (°C)	Time (h)	GC yield ^b	
Effery	111	Волчене	1 (0)	Time (ii)		
1	Bi	DMC	80	5	76%	
2	Bi	CH_3NO_2	80	5	77%	
3	Bi	CH ₃ CN	80	24	76%	
4	Bi	$CH_3(CO)CH_3$	80	24	70%	
5	Bi	2-Me-THF	80	24	20%	
6	Bi	AcOEt	80	24	45%	
7	Bi	DCE	80	24	60%	
8	Fe	DMC	80	7	86%	
9	Fe	CH_3NO_2	80	24	83%	
10	Cu	DMC	80	24	36%	
11	Cu	CH_3NO_2	80	24	81%	
12	Bi	CH_3NO_2	30	24	77%	
13	Fe	CH_3NO_2	30	4	80%	
14	Cu	CH_3NO_2	30	24	7%	
15	MMT	CH_3NO_2	80	7	0%	
16	_	CH_3NO_2	80	24	0%	

^a Reaction conditions: substrate (0.25 mmol), anhydrous and degassed solvent (1 mL) and catalyst (5 mol% metal/substrate ratio). ^b Determined by GC-TCD by external calibration. Trace of hydroarylation product 1b' resulting from the 6-endo-trig addition of one phenyl group across the double bond could be observed.

We only focused on DMC and nitromethane for further catalyst testing and obtained again very good isolated yields of cyclic ether with Fe-MMT (entries 8 and 9). With Cu-MMT, the reaction proceeded poorly in DMC with a low 36% yield, while in nitromethane 81% yield was obtained (entries 10 and 11). In a last set of experiments, the temperature was lowered to 30 °C and the reaction was conducted in nitromethane. Bi-MMT yielded 77% of the product after 24 h (entry 12), but Fe-MMT was the most efficient catalyst under these conditions with 80% isolated yield after only 4 h (entry 13). Cu-MMT did not allow us to reach 10% yield at this temperature (entry 14). Control experiments without metals (MMT alone) and without additives were performed, which showed no background reactions (entries 15 and 16). The performance of these supported catalysts was compared with their homogeneous counterparts. Thus, the reaction of 1a was performed with 5 mol% FeCl₃ and BiCl₃ in DMC at 80 °C. After 7 h, cyclic product **1b** was formed in 76% yield with FeCl3, along with 5% of Friedel-Crafts product 1b'. With BiCl₃, substrate 1a was recovered unchanged, indicating the presence of the active Bi species in the heterogeneous reaction within the solid catalyst and not in the solution phase after leaching from the support. Thus, supported catalysts Fe-MMT and Bi-MMT were equal or superior to their homogeneous equivalents in terms of both activity and selectivity.

Fe-MMT being identified as the most potent catalyst, we further performed a recycling study under our optimised conditions in DMC (Fig. 2). Satisfyingly, the Fe-MMT catalyst could be recycled 4 times and at the 5th cycle still gave 80% yield of cyclised product 1b upon total conversion of 1a. This result suggested that the possible leaching of iron species was not significant in DMC at 80 °C. It is interesting to note that the recycling capacities in CH3NO2 were found to be lower (ESI page S13†).

To further estimate the possibility of leaching in these systems, kinetic plots of reactions of Fe-MMT and Bi-MMT in DMC were recorded and a hot filtration test was performed

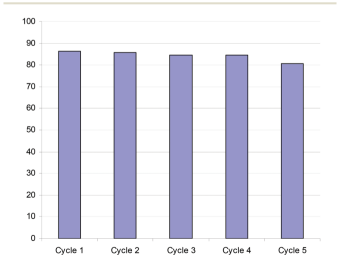


Fig. 2 Recycling studies. Yields of 1b in % obtained after 7 h at 80 °C in DMC with recycled Fe-MMT.

 Table 2
 Substrate scope^a

Table 2	substrate scope					
Entry	Substrate	M	Solvent	<i>T</i> (°C)	Time (h)	Products, isolated yield
1	OH 2a	Fe	DMC	80	7	2b , 98% ^b
2	C ₅ H ₁₁ OH 3a	Fe	DMC	80	7	C ₅ H ₁₁ 3 b , 89%
3	OH 4a	Fe	DMC	80	7	4b, 38%°
4	OH 5a	Fe	DMC	80	7	5b, 89%
5	OH 6a	Fe	DMC	80	7	6b , 63% ^d
6	HO OH 7a	Fe	DMC	80	24	HO 7c (2 dias 1:1), 51% HO 7d (2 dias 1:1), 31%
7	AcO OH 8a	Fe	DMC	80	20	AcO 8c (2 dias 1:1), 59%
8	nC_5H_{11} OH (Z) -9a	Fe	$\mathrm{CH_{3}NO_{2}}$	100	72	nC ₆ H ₁₃
9	Ph OH (Z)-10a	Fe	CH ₃ NO ₂	100	2.5	Ph 0 10b, Quant.
10	(E)- 10a	Fe	$\mathrm{CH_3NO_2}$	100	24	10b, Quant.
11	HO 11a	Fe	DMC	80	48	f
12 13	11a OH (E)-12a	Bi Fe	DMC DMC	80 80	48 5	_f
14 15 16 17	(E)-12a (E)-12a (E)-12a (E)-12a	Bi Fe Bi Bi	$\begin{array}{c} \mathrm{DMC} \\ \mathrm{CH_3NO_2} \\ \mathrm{CH_3NO_2} \\ \mathrm{DCE} \end{array}$	80 80 80	24 5 24 24	12b, 32% + 12b' (trans/cis 2:1), 42% 12b, 21% + 12b' (trans/cis 4:1), 79% 12b' (trans/cis 2:1), 96% 12b' (trans/cis 2:1), 92%
18	OH (E)-13a	Fe	DMC	80	24	13b, 56% ^g
19 20	(E)-13a (E)-13a	Fe Bi	$\mathrm{CH_3NO_2}$ $\mathrm{CH_3NO_2}$	80 80	21 21	13b, 71% Mixture

 $[^]a$ Reaction conditions: substrate (1 mmol), anhydrous and degassed solvent (2 mL) and catalyst (5 mol% metal/substrate ratio). b Reactions performed in DMC-d6 using benzene as the internal standard. c Along with 12% of dehydration product. d Conversion 81%. e Conversion 89%. Traces of a 6-membered ring product were observed (\sim 5%). f No conversion, quantitative recovery of the starting material. g Conversion 63%.

after 75 min reaction time. 31 After removal of the catalyst, the concentration of cyclised product 1b in the filtrate did not change over a period of 220 min, ruling out the possibility of hidden homogeneous catalysis (ESI page S13†).

With this sustainable catalytic system in hand, we next turned our attention to the reaction substrate scope (Table 2). Unsaturated secondary alcohols such as 2a and 3a reacted readily in the presence of 5 mol% Fe-MMT in DMC at 80 °C to yield regioselectively the corresponding 6-membered cyclic ethers 2b and 3b in excellent yields (entries 1 and 2). The introduction of an aryl substituent in geminal position with the hydroxyl group as in 4a was detrimental to the intramolecular hydroalkoxylation and the cyclic ether 4b was formed in a modest 38%, together with 12% of styryl derivatives obtained upon the dehydration of 4a (entry 3). The high mass loss observed was presumably due to a polymerisation reaction. Substrate 5a however, with the phenyl substituent one ethylene away from the alcohol function, could efficiently cyclise to 5b with 89% yield under the same conditions (entry 4). With ortho-prenylated phenol 6a, the conversion was 81% under the same conditions, and the expected cyclic ether 6b was formed in 63% yield (entry 5). Bis-hydroxylated substrate 7a was prepared from biosourced glycerol and afforded a mixture of 5- and 6-membered cyclic ketals 7c and 7d in 51 and 31% yields, respectively (entry 6). The reaction presumably proceeded by isomerisation/cyclisation, involving the formation of an enol ether intermediate, in a process already described with α -methallyloxy carboxylic acids in the presence of Cu(OTf)2 as the catalyst.32 Both products were formed as equimolar mixtures of diastereomers. After conversion of the primary alcohol function into its acetate, the reaction proceeded selectively towards the formation of the 5-membered ring 8c (2 dias 1:1) in 59% yield (entry 7). The TBDMS protecting group was not stable under our conditions (not shown). With a linear substrate such as 9a, featuring an internal disubstituted double bond, the reaction required both high temperatures and longer reaction times in CH3NO2 to reach 89% conversion and to deliver regioselectively the product 9b in 62% yield (entry 8).

Substrate (Z)-10a afforded an example of an internal disubstituted double bond reacting efficiently towards the formation of the 6-membered ring 10b obtained in an excellent 99% yield but required heating in CH₃NO₂ at 100 °C (entry 9). The presence of the phenyl substituent allowed both a perfect regiocontrol of the nucleophilic attack of the hydroxyl group and a sufficient stability of the reactive intermediates. Its stereoisomer (E)-10a reacted similarly, albeit in longer reaction times (entry 10). Substrate 11a featuring a monosubstituted double bond did not react either in the presence of Fe-MMT or Bi-MMT at 80 °C in DMC and was quantitatively recovered unchanged after 48 h (entries 11 and 12). Homogeneous versions of the reaction with this substrate and analogs have been previously reported to require the dual action of an aluminium-based catalyst and a high temperature (250 °C)³³ or the combination of FeCl₃ and silver triflate³⁴ as additives to proceed efficiently.

With this system in hand, we evaluated the possibility to catalyse the tandem reaction involving the formation of C-C and C-O bonds with polyunsaturated alcohols. Firstly, the ortho-geranylated phenol (E)-12a was reacted in the presence of 5 mol% Fe-MMT or Bi-MMT in DMC at 80 °C, and a mixture of the tricyclic product 12b' (in the form of a 2:1 diastereomeric mixture in favor of the trans-fused ring system), resulting from the double cyclisation of 12a, and the monocyclic ether 12b were obtained unselectively (entries 13 and 14). With Fe-MMT in CH₃NO₂, the selectivity in favour of 12b' was increased to 79% (entry 15). Interestingly, the tricyclic product 12b' was the sole product obtained in 96 and 92% isolated yields when the reaction was run with Bi-MMT in CH3NO2 and DCE, respectively (entries 16 and 17). In an effort to obtain the commercial odorant Ambrox®, we tested our system for a tandem reaction with substrate (E)-13a. However, in contrast with the case of substrate (E)-12a, only one single cyclisation was observed to yield the tetrahydrofuran derivative 13b in 56-71% yields (entries 18 and 19). The use of Bi-MMT resulted in the formation of a mixture of isomerised products under the same reaction conditions, but no tandem product (entry 20).

To proceed, the tandem reaction requires the initial activation of the terminal double bond followed by an ene-reaction with the internal double bond, the resulting carbenium ion being trapped intramolecularly by the hydroxyl group. In the reaction of (E)-12a, Bi-MMT, featuring a large cation (ionic radius³⁵ for Bi³⁺ = 0.96–1.17 Å), could only activate the terminal double bond and favour the selective formation of the tandem product, while Fe-MMT, featuring smaller cations (0.49–0.78 Å), could interact with both double bonds and led unselectively to a mixture of products. With (E)-13a, the remote double bond is too hindered, and only the internal double bond is activated by the catalyst followed by the nucleophilic attack of the hydroxyl group leading to the tetrahydrofuranic product.

Conclusions

In summary, we have developed and described herein an efficient and particularly sustainable catalytic system based on the use of benign metals such as iron and bismuth supported on a natural inorganic material, which allow a highly atomeconomic transformation in DMC, a non-VOC solvent. The transformation allowed the formation of a large range of cyclic ethers from the corresponding unsaturated alcohols by intramolecular hydroalkoxylation and the catalyst could be recycled several times.

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