Green Chemistry



View Article Online





Cite this: Green Chem., 2015, 17, 2398

Facile route to conformal hydrotalcite coatings over complex architectures: a hierarchically ordered nanoporous base catalyst for FAME production+

Julia J. Creasey, a Christopher M. A. Parlett, b Jinesh C. Manayil, b Mark A. Isaacs, b K. Wilson^b and Adam F. Lee*^b

An alkali- and nitrate-free hydrotalcite coating has been grafted onto the surface of a hierarchically ordered macroporous-mesoporous SBA-15 template via stepwise growth of conformal alumina adlayers and their subsequent reaction with magnesium methoxide. The resulting low dimensional hydrotalcite crystallites exhibit excellent per site activity for the base catalysed transesterification of glyceryl triolein with methanol for FAME production.

Received 1st September 2014, Accepted 27th January 2015 DOI: 10.1039/c4ac01689k

www.rsc.org/greenchem

Rising global energy demand over the next 25 years, notably among emergent economies, is driving the quest for sustainable routes to low cost, liquid transportation fuels from biomass feedstocks.² Around 9% of transportation energy needs are predicted to be met via liquid biofuels by 2030.3 The past decade has seen much criticism of first-generation biofuels derived from edible plant materials which are attributed to significant land use changes and deforestation in South East Asia.4 In order for advanced bio-fuels to be considered truly sustainable, they must be sourced from non-edible crop components, forestry waste, alternative non-food plants such as switchgrass, Miscanthus or Jatropha curcas⁵ which require minimal cultivation and do not compete with traditional arable land or drive deforestation, algal sources or the lignocellulosic components of municipal waste such as packaging materials.

Although there is burgeoning interest in extracting bio-oils from aquatic biomass, which can yield 80-180 times the annual volume of oil per hectare than plants, 6 process scale-up and the availability of nutrient resources remains challenging.⁷ The biorefinery concept affords biomass the simplest and most popular approach to drop-in transportation fuels,8 based upon carbohydrate pyrolysis and subsequent hydrodeoxygenation (HDO)9 of the resulting bio-oils or their gasification and subsequent Fischer-Tropsch processing10 to gasoline and diesel,¹¹ or lipid transesterification to biodiesel.¹² Catalytic

depolymerisation of lignin may also unlock opportunities for the co-production of phenolics and related aromatic compounds via biorefineries for fine chemical and pharmaceutical applications¹³ improving their cost-effectiveness.

Biodiesel is a clean burning and biodegradable fuel¹⁴ which remains popular for meeting transportation energy requirements in Europe, 15 Asia, 16 the Americas 17 and Africa. 18 Commercial biodiesel is produced almost entirely via the liquid base catalysed transesterification of C14-C20 triacylglyceride (TAG) components of lipids with C₁-C₂ alcohols¹⁹ into fatty acid methyl esters (FAMEs) which constitute biodiesel. Higher alcohols have also been exploited20 as they offer a less corrosive FAME with improved physical characteristics.²¹ Isolation of the desired biodiesel product from homogeneous base catalysts (and unreacted mono- and di-alkyl glycerides and glycerol by-product) is necessary to circumvent saponification and emulsification side reactions and produce a high quality biofuel.6 Heterogeneous, solid acid22-24 and base catalysts offer facile FAME separation, eliminating the requirement for quenching steps and permitting continuous biodiesel production, 25 and a purer glycerol by-product stream for use as a commodity chemical in the food and cosmetics industry. Among solid base catalysts, hydrotalcites, 26-28 alkaline earth oxides²⁹⁻³³ and alkali-doped mesoporous silicas³⁴ are good potential candidates for biodiesel formation under mild conditions. Hydrotalcites (of general formula $[M(II)_{1-x}M(III)_x$ - $(OH)_2^{x+}(A_{n-x/n})\cdot mH_2O)$ are a subset of microporous layered double hydroxides, 35 conventionally synthesised via co-precipitation from nitrates36 in the presence of alkalis which are problematic due to NOx emissions³⁷ and in situ alkali leaching and consequent FAME contamination.38-40 We recently reported an alkali/nitrate-free route to tunable Mg-Al hydro-

^aSchool of Chemistry, Cardiff University, Cardiff CF10 3AT, UK

^bEuropean Bioenergy Research Institute, Aston University, Birmingham B4 7ET, UK. E-mail: a.f.lee@aston.ac.uk; Tel: +44 (0)121 2044036

[†] Electronic supplementary information (ESI) available: Full material synthesis and characterisation. See DOI: 10.1039/c4gc01689k

talcite coatings via the direct reaction of Mg(OCH₃)₂ with a conventional, bulk alumina support.⁴¹ While the resulting materials exhibited excellent turnover frequencies (TOFs) towards TAG transesterification, they suffer from a number of important drawbacks, namely: poor specific activity per unit mass towards bulky C₁₈ substrates (0.042 mmol min⁻¹ g⁻¹), low surface areas, and restricted (and disordered) pore architectures available through the use of pure alumina templates. In contrast, *hydrothermally stable* silica frameworks can be readily synthesised, offering diverse pore interconnectivities and bi- or tri-modal pore networks.⁴²⁻⁴⁴

Here we extend our previous methodology to create crystalline, catalytically active hydrotalcite coatings via a versatile two-step methodology, permitting (i) the first genesis of an ultrathin alumina adlayer over a complex (hierarchically ordered) template, and (ii) facilitating its subsequent reaction stoichiometric $Mg(OCH_3)_2$ to form a MM-SBA-15 hydrotalcite catalyst. This novel methodology opens the way to a new class of solid bases built upon the tunable interconnectivity and porosity afforded by underlying silica architectures. The resulting nanocomposite combines the high surface area and excellent mass-transport characteristics of the parent silica, and solid basicity and transesterification performance of a pure hydrotalcite.

Catalyst synthesis

Synthesis of macroporous-mesoporous SBA-15 (MM-SBA-15)

An hierarchical macroporous–mesoporous SBA-15 silica was prepared following the method of Dhainaut $et~al.^{22}$ Briefly polystyrene beads synthesised via the emulsion polymerisation approach of Vaudreuil and co-workers were added to an acidified, aqueous solution of Pluronic® P123 surfactant prior to the addition of tetramethoxysilane. The resulting gel was hydrothermally aged without agitation, and the solid obtained filtered, washed and dried at room temperature before calcination at 550 °C for 6 h in air.

Alumina grafting onto MM-SBA-15 (Al-MM-SBA-15)

The Al-MM-SBA-15 hierarchical framework employed the method of Landau and co-workers developed for MCM-41.46 Aluminium-tri-sec-butoxide (14.5 g) was dissolved in anhydrous toluene (100 cm³) at 85 °C with stirring. Triethylamine (2.1 cm³) was added to this solution, followed by dried MM-SBA-15 (1 g). After 6 h stirring at 85 °C the solution was filtered under vacuum (~0.1 bar), with the recovered solid washed three times in toluene (100 cm³). The alumina surface was then hydrolysed in ethanol (318 cm³) containing water (1.6 cm³) for 24 h at 25 °C with stirring, and the resulting solid recovered by vacuum filtration and washed with ethanol (300 cm³) before drying at 80 °C in a vacuum oven overnight. A three-step calcination sequence was utilised to form an alumina monolayer: the material was first heated to 250 °C for 1 h, then 400 °C for 1 h and finally 500 °C for 4 h (each ramp rate 1 °C min⁻¹). Consecutive grafting cycles were undertaken

employing an identical protocol in order to progressively build-up alumina monolayers over the silica surface, adjusting the quantities to maintain the initial Al:Si stoichiometry.

Synthesis of hydrotalcite-coated MM-SBA-15 (HT/MM-SBA-15)

Magnesium methoxide solution (8-10 wt% in methanol) was added to Al-MM-SBA-15 (400 mg, dried for 1 h at 80 °C), at the minimum quantity to form a homogeneous paste on mixing. After stirring for 15 min, the mixture was dried under vacuum at 80 °C for 1 h to remove excess methanol. The surface Mg: Al atomic ratio was tuned by adjusting the volume of magnesium methoxide (10.8 cm³ for the MM-SBA-15). The resulting material was calcined at 450 °C for 15 h under 20 cm³ min⁻¹ O_2 (ramp rate 1 °C min⁻¹). After cooling to room temperature under N₂ (20 cm³ min⁻¹), the powder was added to distilled water (50 cm³ for every 300 mg of powder) in a 100 cm³ roundbottomed pressure vessel and heated to 125 °C with stirring for 21 h. After cooling to room temperature, the final HT/ MM-SBA-15 sample was filtered, washed with deionised water and dried in a vacuum oven overnight at 80 °C, before storage in a desiccator. This synthesis proved successful on the multigram scale. A conventional hydrotalcite reference material was prepared via our alkali-free, co-precipitation method from Mg-(NO₃)₂·6H₂O and Al(NO₃)₃·9H₂O precursors, ²⁶ with the Mg: Al atomic ratio tuned to match that of the MM-SBA-15.

Materials characterisation

Nitrogen porosimetry was undertaken on a Quantachrome Nova 1200 porosimeter. Multi-point BET surface areas were calculated over the relative pressure range 0.01-0.3. Pore diameters and volumes were calculated applying either the t-plot or BJH methods to the desorption isotherm. Powder XRD patterns were recorded on a PANalytical X'pertPro diffractometer fitted with an X'celerator detector and Cu Kα source; the Scherrer equation was used to calculate HT crystallite sizes. XPS was performed on a Kratos Axis HSi X-ray photoelectron spectrometer fitted with a charge neutraliser and magnetic focusing lens employing Al K_{α} monochromated radiation (1486.7 eV). Spectral fitting was performed using CasaXPS version 2.3.15. Base site densities were measured via CO2 pulse chemisorption and subsequent temperature programmed desorption (TPD) on a Quantachrome ChemBET 3000 system coupled to an MKS Minilab QMS. SEM analysis was carried out on a Carl Zeiss EVO SEM fitted with an Oxford Instruments energy dispersive X-ray (EDX) analyser employing Oxford Instruments Inca Software. TGA was performed using a Stanton Redcroft STA780 thermal analyser.

Transesterification

The HT/MM-SBA-15 and conventional HT materials were tested as catalysts in the transesterification of triolein to form methyl trioleate (FAME) using a Radleys Starfish parallel

Paper

reactor. Briefly, 50 mg of catalyst was added to 10 mmol of triolein using a 30:14:1 methanol:butanol:oil ratio; butanol was added as a co-solvent to help solubilise the triglyceride. In light of the significant differences in HT content between our conventional and SBA-15 coated materials, a common total mass of catalyst (rather than mass of hydrotalcite) was employed to ensure identical mixing characteristics within the reaction vessel. Reactions were carried out at 90 °C in a modified ACETM 50 cm³ round bottom pressure flask, with aliquots removed periodically from the reaction mixture for analysis on a Varian 450 GC with 8400 autosampler (programmable oncolumn injection onto a Phenomenex ZB-1HT column (15 m × $0.53~\text{mm} \times 0.15~\mu\text{m}$ film thickness). Initial rates were calculated from the linear portion of the reaction profile during the first 60 min of the reaction. Turnover frequencies (TOFs) were determined by normalising rates to the total base site density from CO₂ chemisorption.

Results and discussion

Characterisation of Al-MM-SBA-15

Alumina grafted silica (Al-MM-SBA-15) was first prepared as support for subsequent conversion to a high area, hierarchically ordered hydrotalcite coating. The alumina grafting process was repeated four times to obtain a uniform multilayer interface, with textural properties characterised after each grafting in order to examine the evolution of the aluminasilica interface. Low angle XRD and TEM of the parent MM-SBA-15 and the sequentially alumina grafted variants (Fig. S1-2†) confirmed the presence of ordered mesopores indicative of SBA-15.47 Characteristic (100), (110) and (200) reflections were observed for all materials, indicative of the p6mm space group expected for hexagonally arranged mesoporous channels.⁴⁸ Macropore incorporation shifted these reflections to higher angle relative to conventional mesoporous SBA-15, associated with a small contraction in the mesopore lattice parameter.²² This contraction is attributed to curvature of the mesopore channels as they coalesce around the polystyrene bead template due to strong electrostatic interactions between the beads, block co-polymer and silica precursor. Long range, hexagonally ordered mesopores remained present for Al-MM-SBA-15 even following four consecutive grafting cycles.

Nitrogen porosimetry of the parent MM-SBA-15 and alumina grafted analogues confirmed that the mesoporosity intrinsic to the SBA-15 framework is maintained after each grafting cycle (Fig. S3a-b \dagger). However, the BET surface area (and interconnecting micropore area from t-plot analysis), mean mesopore diameter, and total mesopore and micropore volumes decreased progressively with each grafting cycle (Table 1), consistent with an increasing thickness of conformal alumina overlayer uniformly distributed throughout the pore network.

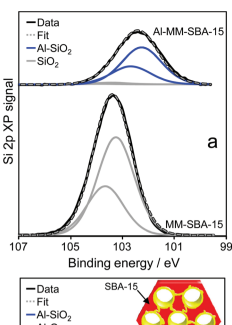
The surface of Al-MM-SBA-15 was subsequently investigated by XPS. Successful alumina grafting was confirmed by the presence of surface Al, with the Al: Si atomic ratio increasing monotonically with each cycle, reaching 22.3 wt% Al after four cycles. Fig. 1 compares the Al and Si 2p chemical environments for the parent and alumina grafted mesoporous silicas. The pure MM-SBA-15 exhibited a single Si 2p spin-orbit split doublet centred with Si 2p3/2 component at 103.1 eV binding energy associated with pure silica. Alumina grafting significantly attenuated the substrate signal consistent with a conformal (Frank-van der Merwe) growth mode, rather than the formation of 3-dimensional alumina islands. This attenuation was accompanied by the emergence of a second low binding energy doublet at 102.2 eV for Al-MM-SBA-15, which can only be associated with a new, interfacial silicon species. This hypothesis is supported by the observation of two distinct Al 2p spin-orbit doublets, at 73.8 eV and 74.7 eV. The former is consistent with pure alumina, and its intensity increases monotonically with grafting cycle relative to the high binding energy state, precisely as expected if the latter was associated with an interfacial alumina species. The opposing binding energy shifts for the interfacial Al and Si species are similar to those previously observed for alumina grafted SBA-15,49 and can be understood in terms of their different Pauling electronegativities and associated induced dipoles mediated via the Al-O-Si bridges which increase and decrease the local initial state charge on interfacial Si and Al atoms respectively. An estimate of the alumina film thickness may be obtained by comparing the experimentally determined Al surface density (derived from porosimetry and XPS) with that for a crystalline alumina phase such as α-Al₂O₃, which exhibits a rhombohedral (4.75 Å \times 4.75 Å) surface unit mesh containing three Al atoms within the (006) plane as shown in Scheme 1.50 With a total surface area of 473 m² g⁻¹, a single α-Al₂O₃ monolayer covering the entire silica pore network would contain

Table 1 Porosimetry of Al-MM-SBA-15 as a function of grafting cycle

Material	Surface area ^a /m ² g ⁻¹	Mesopore volume ^b / cm ³ g ⁻¹	Mesopore diameter ^b /nm	Micropore volume ^c /cm ³ g ⁻¹	Micropore area ^c /m ² g ⁻¹
MM-SBA-15	473	0.66	3.9	0.079	166
Al-MM-SBA-15-1	332	0.48	4.0	0.024	58
Al-MM-SBA-15-2	271	0.47	4.0	0.013	33
Al-MM-SBA-15-3	240	0.46	3.7	0.007	19
Al-MM-SBA-15-4	201	0.45	3.7	0.004	13

^a Multi-point BET method. ^b BJH mesopore volume and pore diameter from the desorption isotherm. ^c Calculated by t-plot method.

Green Chemistry



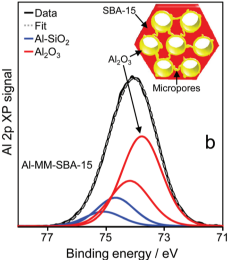
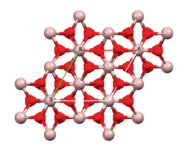


Fig. 1 (a) Si and (b) Al 2p XP spectra of parent MM-SBA-15 and Al-MM-SBA-15 following four alumina grafting cycles.



Scheme 1 Crystal structure of $\alpha\text{-alumina}$ with primitive cell highlighted. 50

0.0009 mol Al, equating to an Al loading of 24.3 wt%. This is close to the observed value of 22.3 wt%, and indicates that an alumina film approximately 0.7 monolayers thick (~0.17 nm) is formed following four grafting cycles, which would constrict the mesopores by 0.34 nm relative to the parent MM-SBA-15,

in excellent agreement with the observed pore diameter decrease of 0.3 nm seen in Table 1.

Characterisation of HT/MM-SBA-15

Powder XRD diffraction patterns for the methoxide functionalised Al-MM-SBA-15 material prepared via four alumina grafting cycles (HT/MM-SBA-15), and a reference bulk HT sample prepared by conventional alkali-free co-precipitation (ConvHT) are shown in Fig. 2. The HT/MM-SBA-15 sample shows a diffraction pattern characteristic of a pure HT phase, very similar to that observed for the ConvHT standard, but with broader reflections indicative of significantly smaller crystallite sizes (as anticipated in light of the highly dispersed alumina substrate, which by inference appears to undergo little restructuring during the crystallisation process) and turbostratic disorder.51 There was no evidence for brucite52 and only a single weak reflection likely associated with trace MgO. This confirms the successful synthesis of a hydrotalcite phase through direct reaction of a pre-formed, ultrathin alumina film and magnesium methoxide from solution.

Crystallite sizes determined using the Scherrer equation, interlayer spacings, lattice parameters and Mg: Al ratios determined using Vegard's law (Fig. S4†) are reported in Table 2. The composition, lattice parameter and interlayer spacings of the HT/MM-SBA-15 material were almost identical to that of the ConvHT, confirming that the hydrotalcite phase formed at the surface of the hierarchical silica support was essentially indistinguishable from that of obtained by traditional synthetic methods, but with a surface area around five times higher. However, the significant difference in microporous crystallites size is expected to hinder accessibility of reactants

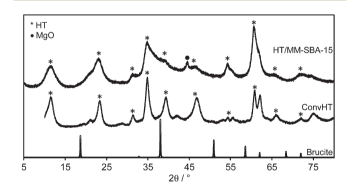


Fig. 2 Powder XRD patterns for HT/MM-SBA-15 and a conventional HT standard (offset for clarity).

Table 2 Crystallographic properties of HT/MM-SBA-15 and ConvHT

Material	Crystallite	Interlayer	Lattice	Mg : Al
	size ^a /nm	spacing ^a /nm	parameter ^a /nm	ratio ^b
HT/MM-SBA-15	1.18	0.76	0.305	2.21:1
ConvHT	6.2	0.77	0.305	1.95:1

 a Calculated using Scherrer equation. b Determined from lattice parameter using Vegard's Law.

Paper

to active sites within the ConvHT interlayers relative to the HT/

MM-SBA-15 sample whose dimensions suggest a hydrotalcite bilayer wherein a far greater proportion of base sites reside on exposed surfaces.

Textural properties of the HT/MM-SBA-15 material are compared with those of the Al-MM-SBA-15 precursor in Fig. 3. Nitrogen porosimetry evidences retention of mesopore and macropore character within the adsorption/desorption isotherms following HT crystallisation, although their demarcation is not as clear as for Al-MM-SBA-15 (Fig. S3†), while Table S1† shows virtually no change in either the mesopore volume or mean mesopore diameter upon reaction of the alumina adlayer with Mg(OCH₃)₂. This suggests that either extremely thin HT crystallites are formed throughout the bimodal pore network (consistent with XRD), or that hydrotalcite formation is confined to the macropores. The latter

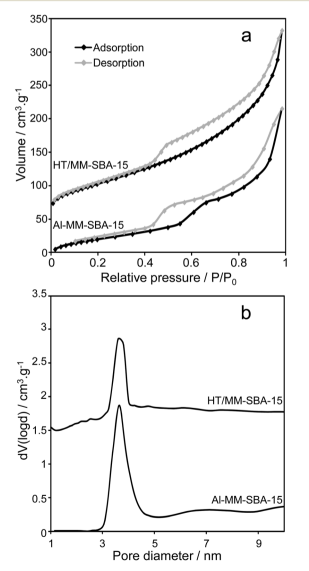


Fig. 3 (a) N₂ adsorption-desorption isotherms and (b) BJH pore size distributions for Al-MM-SBA-15 precursor and HT/MM-SBA-15 (offset for clarity).

would be expected to hinder accessibility of the mesopores (for which macropores serve as the principal conduits), and hence reduce both the mesopore volume and total surface area, in contrast to the observed values reported in Table S1.†

SEM of the HT/MM-SBA-15 (Fig. 4) confirms the macropore network present within the parent MM-SBA-15 support is retained throughout the material after hydrothermal treatment, a measure of the excellent stability of silica frameworks towards high temperature water, and conditions that a comparable pure hierarchical alumina structure would be unlikely to survive. TEM shows macropores are decorated with high aspect ratio hydrotalcite nanocrystallites. Thermogravimetric analysis confirms the excellent thermal stability of the HT/MMSBA-15 (Fig. S5†), with only a small 10% weight loss between 70 and 220 °C, associated with the desorption of physisorbed water from the HT surface and water from within the interlayers, 53 and a 5% loss between 250 and 350 °C attributed to hydroxide anions in the brucite-like layers.⁵⁴ EDX elemental analysis of the HT/MM-SBA-15 yields an overall Mg: Al atomic ratio of 2.2:1, in good agreement with that derived from Vegard's law in Table 2, and a total Mg content of 19.8 wt% i.e. a quarter that of a bulk hydrotalcite of comparable Mg:Al ratio,26 consistent with the formation of hydrotalcite nanocrystals approximately 1 nm thick relative to silica walls around 4–5 nm thick in the MM-SBA-15 support. 49

Surface base properties of the HT/MM-SBA-15 and ConvHT bulk reference materials were assessed by temperature-programmed desorption of CO2-saturated samples, presented in Fig. S6.† The desorption profile of HT/MM-SBA-15 mirrors that of the bulk hydrotalcite, both exhibiting a single desorption feature around 340 °C indicative of moderate strength base sites, albeit spanning a broader temperature range and hence distribution of basicity for the HT/MM-SBA-15. Total base site densities for the HT/MM-SBA-15 and ConvHT materials were 6.4×10^{18} and 8.6×10^{19} m⁻² respectively, although one should recall that a large proportion of sites present within the inter-

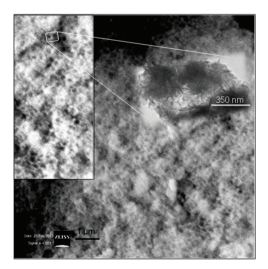


Fig. 4 SEM and TEM micrographs of HT/MM-SBA-15. Insets highlight macropore network and hydrotalcite crystallites decorating macropores.

Green Chemistry Paper

layers of the bulk hydrotalcite structure may be inaccessible to sterically-demanding substrates. These results confirm that high aspect ratio hydrotalcite crystallites formed over the hierarchical silica support possess similar intrinsic basicity to conventional co-precipitated analogues.

Surface analysis of HT/MM-SBA-15 vielded a Mg: Al atomic ratio of 2.21 and Mg content of 16.7 wt%, both very similar to values determined by EDX, evidencing uniform incorporation of Mg into the alumina film throughout the pore network of the Al-MM-SBA-15 precursor. Si 2p XP spectra shown in Fig. 5a reveal that hydrotalcite formation was accompanied by attenuation of the interfacial alumina species, and concomitant appearance of a new low binding energy chemical environment at 101.5 eV. The latter suggests that interfacial silicon atoms are now bound (through oxygen bridges) to a less polarising adlayer relative to alumina, consistent with the exchange of Al3+ for Mg2+ cations. Fig. 5b shows analogous changes in the Al chemical environment, with attenuation of the pure (and interfacial) alumina adlayer, and emergence of a high energy Al state ~74 eV, consistent with the introduction of Mg²⁺ cations into the grafted alumina film during hydrotalcite formation. The corresponding Mg 2s XP spectrum of HT/ MM-SBA-15 presents a single chemical environment around 88.5 eV. In summary, over 75% of the MM-SBA-15 silica surface is contacted with a hydrotalcite phase, and a similar proportion of the initially grafted alumina adlayer in Al-MM-SBA-15 is converted into hydrotalcite.

Transesterification of triglycerides

In order to establish the catalytic efficacy of the HT film encapsulating the hierarchical silica template, the HT/ MM-SBA-15 material was screened in the transesterification of glyceryl triolein, a bulky C18 triglyceride that is a major component on oilseed feedstocks, with methanol for FAME (biodiesel) production. The resulting reaction profiles for HT/ MM-SBA-15 and the co-precipitated ConvHT analogue are shown in Fig. 6. Transesterification proceeded rapidly over both catalysts during the first hour of reaction before slowing dramatically, to give limiting conversions of 34% and 64% for HT/MM-SBA-15 and ConvHT respectively. While the absolute FAME productivity of the bulk hydrotalcite is clearly superior, it is important to recall that the HT/MM-SBA-15 only contains a thin hydrotalcite coating and the majority of this catalyst is composed of inert silica. A fairer comparison of the relative catalytic performance is obtained from their initial rates of triolein conversion and turnover frequencies (TOFs) normalised per base site utilising the CO2 TPD measurements. This reveals a common initial rate of 1 mmol g_{catalyst}⁻¹ min⁻¹, however one must recall that the HT/MM-SBA-15 material only contains one quarter of the amount of hydrotalcite present within the bulk ConvHT material, hence the rate normalised per mass of hydrotalcite is four times higher for HT/ MM-SBA-15 catalyst. Since the base site density of the coated hydrotalcite is also ~34% lower than that of its bulk counterpart, the rate enhancement per base site of the coated material is higher still, translating to TOFs of 7.6 min⁻¹ for the bulk

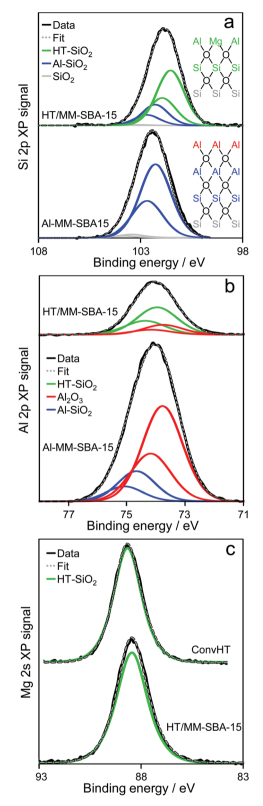


Fig. 5 (a) Si 2p and (b) Al 2p, and (c) Mg 2s XP spectra of Al-MM-SBA-15 precursor and HT/MM-SBA-15.

ConvHT standard versus 66 min⁻¹ for HT/MM-SBA-15. Hydrotalcites prepared via conventional co-precipitation are among the most widely-used catalysts for triglyceride transesterifica**Paper**

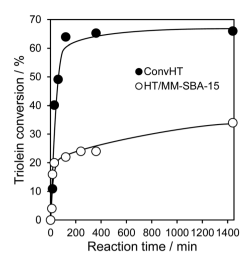


Fig. 6 Triolein conversion to methyl trioleate *via* transesterification with methanol (1:30 molar ratio) at 90 °C over HT-MM-SBA-15 and a conventional HT standard

tion to FAME, hence the nine-fold rate enhancement observed for our HT/MM-SBA-15 material provides a striking benchmark of its exceptional performance. While the magnitude of this enhancement does fall at longer reaction times, likely due to partial deactivation of the coating, the HT/MM-SBA-15 remains three times as active per base site as the bulk hydrotalcite, even after 1400 min reaction.

Since the intrinsic base strength of active sites within the conventional and hierarchical hydrotalcite catalysts is the same (common CO₂ desorption temperatures, Fig. S6†), we attribute this nine-fold rate enhancement of HT/MM-SBA-15 to superior mass-transport characteristics of the macroporous-mesoporous architecture. Indeed, the magnitude of the HT/MM-SBA-15 enhancement with respect to the ConvHT standard is comparable to that previously reported for a macroporous pure HT material, ²⁸ but affords a far more flexible and hydrothermally stable framework than the latter synthesis.

Conclusions

Sequential, wet-chemical surface modification of nanostructured silicas with Al and Mg precursors offers a versatile route to the preparation of high area, tailored solid base hydrotalcite catalysts. Stepwise grafting and thermal processing of aluminium-tri-sec-butoxide results in a uniform alumina monolayer throughout the bimodal macropore–mesopore network. Subsequent reaction with Mg(OCH₃)₂ affords stoichiometric incorporation of aluminium from the alumina adlayer into \sim 1 nm Mg₂Al hydrotalcite crystallites, which possess identical basicity as a co-precipitated, bulk hydrotalcite. In contrast to bulk (monomodal) alumina templates, the development of a silica based methodology results in HT/MM-SBA-15 catalyst exhibits similar specific mass activity in the transesterification of glyceryl triolein with methanol as a bulk hydrotalcite, despite containing only a small fraction of the number of

active sites, indicating far greater active site accessibility to the bulky TAG reactant. The latter conclusion is supported by a nine-fold enhancement in the TOF per base site for the hierarchical hydrotalcite, indicating the majority of base sites in HT/MM-SBA-15 reside at the external surface of nanoscale crystallites within the meso- and macropores, rather than within the microporous interlayers of conventional hydrotalcites. Our methodology is readily extendable to diverse silica architectures and other metal oxides, opening up opportunities for the facile introduction of hydrotalcite solid basicity into complex two- or three-dimensional materials, *e.g.* membranes and monoliths, for catalysis and sorption applications.

Acknowledgements

We thank the EPSRC (EP/G007594/4) for financial support and a Leadership Fellowship (AFL) and studentship (JJC), and the Royal Society for the award of an Industry Fellowship (KW).

Notes and references

- 1 International Energy Outlook 2013, Report DOE/EIA-0484 (2013), 2013.
- 2 R. Luque, L. Herrero-Davila, J. M. Campelo, J. H. Clark, J. M. Hidalgo, D. Luna, J. M. Marinas and A. A. Romero, *Energy Environ. Sci.*, 2008, **1**, 542–564.
- 3 BP, BP Energy Outlook 2030, 2011.
- 4 F. Danielsen, H. Beukema, N. D. Burgess, F. Parish, C. A. BrÜHl, P. F. Donald, D. Murdiyarso, B. E. N. Phalan, L. Reijnders, M. Struebig and E. B. Fitzherbert, *Conserv. Biol.*, 2009, 23, 348–358.
- 5 W. M. J. Achten, L. Verchot, Y. J. Franken, E. Mathijs, V. P. Singh, R. Aerts and B. Muys, *Biomass Bioenergy*, 2008, 32, 1063–1084.
- 6 T. M. Mata, A. A. Martins and N. S. Caetano, *Renewable Sustainable Energy Rev.*, 2010, 14, 217–232.
- 7 R. Pate, G. Klise and B. Wu, *Appl. Energy*, 2011, **88**, 3377–3388
- 8 A. Demirbas, Energy Policy, 2007, 35, 4661–4670.
- 9 P. M. Mortensen, J. D. Grunwaldt, P. A. Jensen, K. G. Knudsen and A. D. Jensen, *Appl. Catal.*, *A*, 2011, **407**, 1–19.
- 10 H. Jahangiri, J. Bennett, P. Mahjoubi, K. Wilson and S. Gu, *Catal. Sci. Technol.*, 2014, 4, 2210–2229.
- 11 F. Trippe, M. Fröhling, F. Schultmann, R. Stahl, E. Henrich and A. Dalai, *Fuel Process. Technol.*, 2013, **106**, 577–586.
- 12 C. S. K. Lin, L. A. Pfaltzgraff, L. Herrero-Davila, E. B. Mubofu, S. Abderrahim, J. H. Clark, A. A. Koutinas, N. Kopsahelis, K. Stamatelatou, F. Dickson, S. Thankappan, Z. Mohamed, R. Brocklesby and R. Luque, *Energy Environ. Sci.*, 2013, 6, 426–464.
- 13 M. P. Pandey and C. S. Kim, *Chem. Eng. Technol.*, 2011, 34, 29-41
- 14 G. Knothe, Top. Catal., 2010, 53, 714-720.

15 K. Bozbas, Renewable Sustainable Energy Rev., 2008, 12, 542–552.

Green Chemistry

- 16 M. H. M. Ashnani, A. Johari, H. Hashim and E. Hasani, *Renewable Sustainable Energy Rev.*, 2014, 35, 244–257.
- 17 J. C. Bergmann, D. D. Tupinambá, O. Y. A. Costa, J. R. M. Almeida, C. C. Barreto and B. F. Quirino, *Renewable Sustainable Energy Rev.*, 2013, 21, 411–420.
- 18 B. Amigun, J. K. Musango and W. Stafford, *Renewable Sustainable Energy Rev.*, 2011, **15**, 1360–1372.
- 19 A. F. Lee, J. A. Bennett, J. C. Manayil and K. Wilson, *Chem. Soc. Rev.*, 2014, **43**, 7887–7916.
- 20 J. Geuens, J. M. Kremsner, B. A. Nebel, S. Schober, R. A. Dommisse, M. Mittelbach, S. Tavernier, C. O. Kappe and B. U. W. Maes, *Energy Fuels*, 2007, 22, 643–645.
- 21 G. Knothe, Fuel Process. Technol., 2005, 86, 1059–1070.
- 22 J. Dhainaut, J.-P. Dacquin, A. F. Lee and K. Wilson, *Green Chem.*, 2010, 12, 296–303.
- 23 J. P. Dacquin, A. F. Lee, C. Pirez and K. Wilson, *Chem. Commun.*, 2012, **48**, 212–214.
- 24 Zillillah, T. A. Ngu and Z. Li, *Green Chem.*, 2014, **16**, 1202–1210.
- 25 K. Wilson and A. F. Lee, *Catal. Sci. Technol.*, 2012, **2**, 884–897.
- 26 D. G. Cantrell, L. J. Gillie, A. F. Lee and K. Wilson, *Appl. Catal.*, A, 2005, 287, 183–190.
- 27 Y. Liu, E. Lotero, J. G. Goodwin and X. Mo, *Appl. Catal.*, *A*, 2007, 33, 138–148.
- 28 J. J. Woodford, J.-P. Dacquin, K. Wilson and A. F. Lee, *Energy Environ. Sci.*, 2012, 5, 6145–6150.
- 29 R. S. Watkins, A. F. Lee and K. Wilson, *Green Chem.*, 2004, 6, 335–340.
- 30 K. Wilson, C. Hardacre, A. F. Lee, J. M. Montero and L. Shellard, *Green Chem.*, 2008, **10**, 654–659.
- 31 J. M. Montero, P. Gai, K. Wilson and A. F. Lee, *Green Chem.*, 2009, **11**, 265–268.
- 32 M. Verziu, B. Cojocaru, J. Hu, R. Richards, C. Ciuculescu, P. Filip and V. I. Parvulescu, *Green Chem.*, 2008, **10**, 373–381.
- 33 J. J. Woodford, C. M. A. Parlett, J.-P. Dacquin, G. Cibin, A. Dent, J. Montero, K. Wilson and A. F. Lee, *J. Chem. Technol. Biotechnol.*, 2014, **89**, 73–80.
- 34 M. C. G. Albuquerque, I. Jimenez-Urbistondo, J. Santamaria-Gonzalez, J. M. Merida-Robles, R. Moreno-Tost, E. Rodriguez-Castellon, A. Jimenez-Lopez, D. C. S. Azevedo, C. L. Cavalcante Jr. and P. Maireles-Torres, Appl. Catal., A, 2008, 334, 35–43.

- 35 S. J. Mills, A. G. Christy, J. M. R. Genin, T. Kameda and F. Colombo, *Mineral. Mag.*, 2012, **76**, 1289–1336.
- 36 Y. Xi and R. J. Davis, J. Catal., 2009, 268, 307-317.
- 37 M. Behrens, K. F. Girsgdies, I. Kasatkin, F. Hermerschmidt, K. Mette, H. Ruland, M. Muhler and R. Schlogl, *Chem. Commun.*, 2011, 47, 1701–1703.
- 38 D. M. Alonso, R. Mariscal, M. L. Granados and P. Maireles-Torres, *Catal. Today*, 2009, **143**, 167–171.
- 39 M. Di Serio, R. Tesser, L. Casale, A. D'Angelo, M. Trifuoggi and E. Santacesaria, *Top. Catal.*, 2010, 53, 811–819.
- 40 Y. C. Sharma, B. Singh and J. Korstad, *Fuel*, 2011, **90**, 1309–1324
- 41 J. J. Creasey, A. Chieregato, J. C. Manayil, C. M. A. Parlett, K. Wilson and A. F. Lee, *Catal. Sci. Technol.*, 2014, 4, 861–870
- 42 Y. Wan and D. Zhao, Chem. Rev., 2007, 107, 2821-2860.
- 43 Z. Li, X. Wei, T. Ming, J. Wang and T. Ngai, *Chem. Commun.*, 2010, **46**, 8767–8769.
- 44 C. M. A. Parlett, K. Wilson and A. F. Lee, *Chem. Soc. Rev.*, 2013, 42, 3876–3893.
- 45 S. Vaudreuil, M. Bousmina, S. Kaliaguine and L. Bonneviot, *Adv. Mater.*, 2001, 13, 1310–1312.
- 46 M. V. Landau, E. Dafa, M. L. Kaliya, T. Sen and M. Herskowitz, *Microporous Mesoporous Mater.*, 2001, **49**, 65–81.
- 47 A. Y. Khodakov, V. L. Zholobenko, R. Bechara and D. Durand, *Microporous Mesoporous Mater.*, 2005, **79**, 29–39.
- 48 D. Zhao, Q. Huo, J. Feng, B. F. Chmelka and G. D. Stucky, *J. Am. Chem. Soc.*, 1998, **120**, 6024–6036.
- 49 C. M. A. Parlett, L. J. Durndell, A. Machado, G. Cibin, D. W. Bruce, N. S. Hondow, K. Wilson and A. F. Lee, *Catal. Today*, 2014, 229, 46–55.
- 50 E. N. Maslen, V. A. Streltsov, N. R. Streltsova, N. Ishizawa and Y. Satow, *Acta Crystallogr., Sect. B: Struct. Sci.*, 1993, **49**, 973–980.
- 51 G. Thomas and P. V. Kamath, *J. Chem. Sci.*, 2006, **118**, 127–133.
- 52 A. V. G. Chizmeshya, M. J. McKelvy, R. Sharma, R. W. Carpenter and H. Bearat, *Mater. Chem. Phys.*, 2003, 77, 416–425.
- 53 H. C. Greenwell, P. J. Holliman, W. Jones and B. V. Velasco, *Catal. Today*, 2006, **114**, 397–402.
- 54 S. Abelló, F. Medina, D. Tichit, J. Pérez-Ramírez, J. C. Groen, J. E. Sueiras, P. Salagre and Y. Cesteros, *Chem. Eur. J.*, 2005, **11**, 728–739.