Controlled deposition of titanium oxide overcoats by non-hydrolytic sol-gel for improved catalyst selectivity and stability

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Advances in the synthetic control of surface nanostructures could improve the activity, selectivity and stability of heterogeneous catalysts. Here, we present a technique for the controlled deposition of TiO2 overcoats based on non-hydrolytic sol-gel chemistry. Continuous injection of Ti( iPrO)4 and TiCl4 mixtures led to the formation of conformal TiO2 overcoats with a growth rate of 0.4 nm/injected monolayer on several materials including high surface area SBA-15. Deposition of TiO2 on SBA-15 generated medium-strength Lewis acid sites, which catalyzed 1-phenylethanol dehydration at high selectivities and decreased deactivation rates compared to typically used HZSM-5. When supported metal nanoparticles were similarly overcoated, the intimate contact between the metal and acid sites at the support-overcoat interface significantly increased propylcyclohexane selectivity during the deoxygenation of lignin-derived propyl guaiacol (89% at 90% conversion compared to 30% for the uncoated catalyst). For both materials, the surface reactivity could be tuned with the overcoat thickness.

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1. Introduction

Metal oxides with uniform composition are omnipresent in catalysts and catalyst supports due to their ease of production, but their use can lead to low selectivity and stability due to the lack of precise design of their surface functionalities. Deposition of thin metal oxide layers on the surface offers the opportunity to create optimal environment on the material surface [1–6]. Multiple studies demonstrate that liquid phase sol-gel based techniques can be used for the controlled growth of oxide films [7–11]. The mild operative conditions and relatively simple equipment needed for sol-gel applications make these synthetic routes appealing, especially as scale-up is considered. However, while high levels of control have been reported for SiO2 deposition, which benefits from kinetically slow hydrolysis/condensation [13], highly reactive titanium oxide precursors prevent controlled deposition of thin oxide layers (<50 nm). Although remarkable results were achieved by controlling the kinetics of condensation with catalysis [14] or chelating agents [15], deposition of multiple oxide layers on challenging substrates such as high surface area mesoporous materials or supported metal nanoparticles remains a challenge [16–20].

The most versatile and powerful techniques for thin film deposition on high surface area catalyst powders such as Atomic Layer Deposition (ALD) rely at present on complex instrumentation and gas phase operations. ALD [21] is the most prominent of these techniques and has been employed to improve selectivity [22] and stabilize supported nanoparticles against sintering [2,3,23]. However, ALD requires cumbersome instrumentation and is difficult to scale up, especially on high surface area materials and powders. In addition, gas phase synthesis offers limited possibilities for tuning material porosity and structure compared to liquid phase synthesis. Tilley and coworkers developed a thermolytic molecular approach, which allowed the generation of multicomponent oxides such as Ti-Si [24] as well as various surface modifications [25], but deposition was limited to one molecular layer [26]. In addition to overcoating strategies, high-temperature calcination and reduction were also employed to cover Co particles with a titanium oxide layer, leading to improvements in catalyst stability and changes in selectivity thanks to strong metal-support interaction during aqueous-phase hydrogenation of furfuryl alcohol [27].

Non-hydrolytic sol-gel (NHSG) chemistry is a well-known technique for material synthesis and has been previously used for catalyst synthesis [28,29]. However, we have found limited
use of sol-gel chemistry for the deposition of thin oxide films and its effect on catalysis. NHSS precursors were used for the layer-by-layer deposition of TiO$_2$ on SBA-15 [30]. However, the alternated manual precursor addition for each half layer limited the control on the overcoat deposition and no effects on catalytic properties were reported. Similarly, ALD was used to deposit TiO$_2$ films over Co/TiO$_2$ catalysts to prevent sintering and leaching of cobalt nanoparticles [31]. However, no selectivity or reactivity effects were demonstrated beyond reduced deactivation.

In this study, we introduce a new titanium oxide thin film deposition methodology based on non-hydrolytic sol-gel (NHSS) chemistry. Our deposition is carried out in a single liquid phase step involving the continuous injection of a mixture of NHSS precursors and results in uniform coatings that are comparable in conformality to those obtained with ALD, even for high surface area materials. More importantly, we demonstrate that the selectivity of both a high-surface area homogeneous oxide catalyst (SBA-15) and a supported metal catalyst (Pt/SiO$_2$) can be significantly enhanced and controlled with the thickness of the overcoat for several reactions.

2. Materials and methods

2.1. Chemicals and materials

All reagents and solvents were analytical grade, obtained from commercial suppliers and used without further purification, unless otherwise noted. Air and moisture-sensitive reagents were handled using a nitrogen filled glove box and Schlenk techniques. Titanium isopropoxide (98%) was obtained from Acros and distilled under reduced pressure. Titanium tetrachloride (99%), tetramethyl orthosilicate (TMOS, 99%), Pluronic 123 (P123, average $M_n$ 7.0 g/mol) and 1-phenylethanol (98%) were obtained from Sigma-Aldrich. Ethanol (99%) was obtained from Fisher Scientific. Orthosilicate (TMOS, 99%), Pluronic 123 (P123, average $M_n$ 7.0 g/mol) and 1-phenylethanol (98%) were obtained from Sigma-Aldrich. Ethanol (99%) was obtained from Fisher Scientific. 1,4-dioxiane (99.8%) was obtained from ABCR. Hydrogen hexachlororoloplatinate(IV) hexahydrate (H$_2$PtCl$_6$·6H$_2$O, 99.9%) was obtained from Strem. Aerosil TiO$_2$ P25 was obtained from Evonik. TiO$_2$ PC500 was provided by Crystal. Siral40 was provided by Sasol. Toluene (VWR, 99.5%) and diethyl ether (99%, Fisher Scientific) were purchased from Strem. Aerosil TiO$_2$ P25 was obtained from Evonik. TiO$_2$ PC500 was provided by Crystal. Siral40 was provided by Sasol. Toluene (VWR, 99.5%) and diethyl ether (99%, Fisher Scientific) were purchased from Strem. Aerosil TiO$_2$ P25 was obtained from Evonik.

2.2. Materials synthesis

2.2.1. Cuprous oxide nanoparticles synthesis

Cu$_2$O nanoparticles were prepared by mixing aqueous solutions 1 M of CuSO$_4$·5H$_2$O with 1 M of reducing agent Na$_2$S$_2$O$_3$ in volumetric proportion of 1:4. The mixture was then poured into an Erlenmeyer flask containing equimolar amount of 1 M sodium hydroxide. A bright orange precipitate appeared almost instantaneously, indicating that the copper was precipitating under the form of cuprous oxide. The solution was vigorously stirred for 30 min. The mixture was centrifuged and the collected particles washed four times with water and once with ethanol. The sample was then dried overnight under vacuum at 65 °C.

The materials were annealed at 200 °C (1.5 °C/min ramp) for 2 h under inert gas (N$_2$) flow to increase their stability toward air oxidation. This synthetic route reportedly yields 1 1 1 plane-oriented cuprous oxide nanoparticles [33].

2.2.2. Supported platinum nanoparticle synthesis

Pt/SiO$_2$ and Pt/TiO$_2$ were prepared by incipient wetness impregnation of the support with a 0.1 M HCl aqueous solution of H$_2$PtCl$_6$·6H$_2$O, calcination at 500 °C in a muffle furnace and reduction under H$_2$ flow (100 mL/min) at 300 °C (2 °C/min ramp) for 10 h.

2.2.3. SBA-15 synthesis

The synthesis of mesoporous silica SBA-15 was based on methods found in literature [34–37]. Pluronic 123 (7.0 g) was dissolved in a 1.6 M aqueous HCl solution (262.5 g) in a round bottom flask and heated to 35 °C in oil bath. Then, TMOS (10 mL) was added to start the synthesis, stirring the solution vigorously for 1 min followed by moderate stirring for 24 h. Afterward, a hydrothermal treatment under static conditions at 140 °C was performed for an additional 24 h. This was followed by filtration, and a day-long Soxhlet extraction with ethanol. The sample was then dried in a ventilated oven at 90 °C for 12 h. Finally, the acquired solid was calcined in a muffle furnace for 6 h at 500 °C (1 °C/min ramp).

2.2.4. Overcoating representative procedure

The substrate (0.5 g) was loaded into a double neck round bottom flask and dried for 12 h at 150 °C under vacuum (<10$^{-5}$ mbar). After refilling the flask with nitrogen, the solid was suspended in anhydrous toluene (20 mL) and sonicated for 10 min to ensure uniform particle dispersion. A reflux condenser was fitted to the flask and the suspension was stirred at 110 °C. The precursor solution was prepared under inert atmosphere by dissolving appropriate amounts of TiCl$_4$ and freshly distilled Ti(O$i$Pr)$_4$ in anhydrous toluene to yield a molar ratio [Ti]/[Ti$(O$i$Pr)$_4$] = 1. Concentrations were adjusted to achieve one monolayer per mL, calculated from the ratio of the titania precursor’s projected surface area (determined from its Van der Waals volume projection as described in our previous work [38]) and the substrate surface area. The precursor solution was added to the refluxing suspension through a rubber septum using a gas-tight glass syringe (Hamilton, US) with a PTFE plunger and an automatic syringe pump (KDS 100 Legacy). The needle was kept at a distance of at least 10 cm from the mixture to avoid premature TiO$_2$ condensation on the needle tip (Fig. S2). The addition rate was typically set to one monolayer per hour, or 0.5 monolayer per hour for high surface area substrates (>100 m$^2$/g). After completing the precursor addition, the mixture was stirred for 2 h and cooled to 80 °C. Then, 20 mL of anhydrous diethyl ether were added to quench unreacted Ti–Cl bonds and the mixture was stirred for another 12 h. The solid was collected by centrifugation (5 min, 4500 rpm) and washed three times with 50 mL diethyl ether, twice with 50 mL ethanol, before being redispersed in H$_2$O. Aging of the as-synthesized overcoat was performed by stirring the material for 12 h at 70 °C in an aqueous solution of 0.2 M NH$_4$OH. Finally, the solid was isolated by filtration, dried under vacuum at 80 °C and calcined for 5 h at 500 °C (2 °C/min ramp) under flow of synthetic air.

2.3. Material characterization

2.3.1. Inductively coupled plasma optical emission spectrometry (ICP-OES)

Elemental analyses were performed at the EPFL Central Environmental Laboratory using inductively coupled plasma optical emission spectrometry (ICP-OES) on an ICP-OES Multitype
Shimadzu instrument. Metals were dissolved using a 1:2 volumetric mixture of concentrated aqueous nitric acid (ca. 70 wt.%) and hydrochloric acid (ca. 37 wt.%) solutions.

2.3.2. Transmission electron microscopy (TEM)

The overcoat morphology was characterized by Transmission Electron Microscopy (TEM) and High-Angle Annular Dark-Field Scanning Transmission Electron Microscopy (HAADF-STEM) on a FEI Talos with 200 kV acceleration voltage in the mode resulting in atomic number contrast (Z contrast). Lacey carbon grids were prepared by directly “dipping” the grid into the sample powder. For thickness and particle size distribution, at least 50 measurements were taken. Energy-Dispersive X-ray spectroscopy (EDXS) analysis was performed using Bruker Espirit software. Average thickness and standard deviations were calculated from STEM images using a minimum of 60 equally spaced random measurements on at least 10 different particles using ImageJ software. The resulting data was analyzed with Origin2016Pro and Igor Pro software.

2.3.3. Magic angle spinning (MAS) solid-state nuclear magnetic resonance (ssNMR)

MAS ssNMR spectra were recorded on a 900 MHz Bruker Avance spectrometer equipped with a conventional double resonance CP-MAS probe using a 3.2 mm zirconia rotor with Vespel caps at a spinning speed of 24 kHz. 

2.3.4. N\textsubscript{2} physisorption

Brunauer-Emmett-Teller (BET) surface, Barrett-Joyner-Halenda (BJH) mesoporous and DFT microporous volumes were calculated from N\textsubscript{2}-physisorption measurements on a Micromeritics 3Flex apparatus at liquid nitrogen temperature between 10^{-5} and 0.99 relative N\textsubscript{2} pressure. Samples (ca. 100 mg) were dried at 120 °C (temperature reached with a ramp of 2 °C/min) under vacuum (<10^{-3} mbar) for 4 h and a leak seal was performed prior to analysis.

2.3.5. X-ray diffraction (XRD)

Small angle XRD measurements were performed in a PANalytical Empyrean system (Theta-Theta, 240 mm) equipped with parallel beam mirror optics with Cu Kα as a radiation source.

2.3.6. NH\textsubscript{3} temperature programmed desorption

The concentration of acid sites was calculated from NH\textsubscript{3} temperature programmed desorption (TPD) performed on a Micromeritics Autochem 2920 II instrument. Typically, the sample (ca. 100 mg) was loaded to a U-shaped cell and dried for 4 h under He flow (50 mL/min) at 300 °C (2 °C/min). After cooling down to 50 °C, the flow was switched to a 1:99 (volumetric ratio) NH\textsubscript{3}:He mixture for 1 h to saturate the sample with NH\textsubscript{3}. The sample was then flushed with He (50 mL/min) to remove physisorbed NH\textsubscript{3} and the temperature was ramped to 500 °C at a rate of 10 °C/min. During this process, ammonia desorption was monitored using a calibrated thermal conductivity detector as well as MKS Cirrus II mass spectrometer calibrated to mass 16.

2.3.7. Diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS)

DRIFT spectra were recorded using a high temperature Harrick DRIFT cell on a Perkin Elmer Frontier spectrometer equipped with a mercury cadmium telluride detector. Spectra were typically collected with 32 scans at a resolution of 4 cm\textsuperscript{-1}. Helium (100 mL/min) was passed through a bubbler filled with pyridine to saturate the samples with pyridine vapor (30 min, 150 °C). Physisorbed pyridine was eliminated by flowing pure helium (30 min, 150 °C) through the cell.

2.4. Catalytic testing

2.4.1. 1-Phenylethanol dehydration

1-Phenylethanol dehydration was carried out in 10 mL Alltech glass reactors (Grace) with silicon sealing disks and PTFE screw caps. Typically, reactors were loaded with 5 mL of 1,4-dioxane and 15 mg of catalyst, 200 μL of 1-phenylethanol and 50 μL of dodecane (internal standard). Reactors were sonicated for 15 min to ensure catalyst dispersion and the reaction was started by placing reactors into the holes of an aluminum block heated by a hot plate. Low stirring speeds around 200 rpm were employed to avoid catalyst deposition to the walls above the solvent level. 50 μL aliquots were collected after cooling down the reactor in an ice bath. Samples were diluted in 1,4-dioxane (10x) and filtered. Reagent and products concentrations were determined with an Agilent Technologies 7890A Gas Chromatography (GC) apparatus equipped with a flame ionization detector (FID) and a HP-5 capillary column (50 m, 0.32 mm). Compound identification was performed on an Agilent Technologies 7890B series GC equipped with an HP5-MS capillary column and an Agilent 5977A series Mass Spectroscopy detector. 1-phenylethanol, dodecane and styrene were quantified by means of a calibration curve. The effective carbon number method was used to quantify ethers and dimers by using the detector response of the peak of interest compared to that of an internal standard. Details for this method, first reported and verified in our previous work [39] are given in the supporting information.

2.4.2. 3-Propyl-guaiacol hydrodeoxygenation

The catalyst (50 mg) was loaded into a 25 mL stainless-steel reactor (Parr) and a 10 mL isooctane solution containing 1 mmol of substrate was added. After purging 3 times with 50 bars of H\textsubscript{2}, the H\textsubscript{2} pressure was set to 15 bar and the temperature was ramped to 200 °C using a heating plate and heating tape controlled by an Omega PID controller. After reaction, the reactor was cooled down under a flow of air and a 50 μL fraction was collected once room temperature was reached and the vessel depressurized. The aliquot was diluted 10 times with isooctane and analyzed by gas chromatography. Reaction time and substrate amount were varied to achieve different levels of conversion.

3. Result and discussion

3.1. Design of the optimal titanium oxide precursor

The first objective was to develop a liquid phase overcoating technique resulting in a conformal titanium oxide layer of controlled thickness while avoiding homogeneous nucleation. To do so, quantitative reaction of the metal oxide precursor on the substrate surface was required to maintain control over the deposition. In addition, reactivity of the precursor had to be limited to the surface. Finally, relatively fast condensation kinetics were sought to allow deposition in reasonable reaction times.

Our approach was based on the pioneering work of Vioux et al., who introduced non-hydrolytic sol-gel condensation routes to metal oxides in anhydrous conditions [40–44]. Transition metal oxides can be formed by etherealysis or alcoholysis of metal halides with ethers, alcohols or metal alkoxides. These condensations are characterized by oxo-bridging between metal atoms by
elimination of small molecules. The alcoholysis route was not considered because condensation kinetics were too slow with metal alkoxides and too fast with metal halides. Furthermore, water generated during the condensation step would likely yield by-products and increase rates. Etherolysis involves an intermediate alkoxide formation step that is difficult to control [41], and hence was also not chosen. To obtain the targeted reactivity, we focused on the direct condensation of metal halide TiCl₄ with titanium alkoxides, while screening various reaction conditions. We achieved optimal control on the condensation for the reaction between TiCl₄ and Ti(OiPr)₄ (titanium isopropoxide). Fast ligand recombination is known to occur in metal halides and metal alkoxides mixtures [41] but no condensation was observed at room temperature between TiCl₄ and Ti(OiPr)₄ even after several days. Condensation kinetics increased drastically with temperature and a quantitative reaction between titanium precursors was observed within 30 min at 110 °C, in agreement with the report by Arnal et al. [41]. Toluene was selected as a solvent because its reflux conditions were close to this temperature and no undesired reactivity with the precursors was observed.

Next, we optimized the [PrO]/[Ti] molar ratio as the relative amounts of metal halide and metal alkoxide substantially influenced the condensation kinetics. A ratio of [PrO]/[Ti] = 1—achieved with a 3:1 TiCl₄/Ti(OiPr)₄ molar mixture—formed primarily Ti(PrO)Cl₃ as reported by Arnal et al. [40]; we observed a sol-gel induction period as low as 10 min at 110 °C (refluxing toluene) that fulfilled our requirement for relatively fast condensation. We investigated the reactivity further using higher ratios ([PrO]/[Ti] > 1), where gelation and subsequent condensation to titanium oxide took several hours to a day, without reaching full conversion. Conversely, employing lower ratios ([PrO]/[Ti] < 1) resulted in low final precursor conversion due to a lack of bridging oxygen atoms. As depicted in Fig. S1, Ti(PrO)Cl₃ readily undergoes self-condensation catalyzed by its strong Lewis acidity [45]. Hence, Ti(PrO)Cl₃ acted as both the reactive and catalytic species during titanium oxide formation.

### 3.2. Deposition of TiO₂ overcoats

The reactivity of metal precursors during the deposition of the first layer differs from the self-condensation presented earlier, because substrate surface hydroxyl groups are involved. Due to the lower reactivity of self-condensation compared to hydrolytic cleavage, we expected the precursor to react preferentially with the substrate surface. As Ti-Cl bonds are more reactive than Ti-O’Pr toward hydrolysis [46,47], we postulated a surface condensation model where surface hydroxyls reacted primarily with the chloride ligands, leading to the formation of HCl. Each titanium center could react with one or two hydroxyl groups, yielding a mono or bis-grafted species (Fig. 1); the coordination to 3 or 4 hydroxyl groups was not considered, as it would yield unrealistic bond strain [48,49]. We calculated the projected surface area of Ti(OiPr)Cl₃ according to Van der Waals radius to be 0.45 nm², while the surface of a metal oxide such as SiO₂ is populated by ca. 4 OH/nm² after dehydroxylation under vacuum at 300 °C [50]. Therefore, surface hydroxyl groups were likely not fully consumed. Consequently, monolayer coverage was calculated by using the ratio of the precursor projected surface area and the substrate surface area. Here, we define a “layer” by analogy with ALD—i.e., the amount of Ti precursor required to saturate the surface. Similarly to ALD, the first monolayer will not cover the surface with one continuous layer of TiO₂ due to the space left behind by the precursor ligands [51].

In contrast, precursor dosing according to OH density, as performed by Yan et al. [30] resulted in undesired excess TiO₂ formation and poor deposition control. In contrast, the formation of subsequent layers likely proceeded by direct condensation between surface titanium and precursor molecules in solution by elimination of a propyl chloride molecule.

Typical overcoating procedures such as ALD require alternated steps of substrate exposure to precursors. Here, the use of a single mixture allows us to use continuous precursor addition with an automated syringe pump (Fig. S2). Continuous addition maintained a low precursor concentration in solution, which prevented homogeneous self-condensation. Moreover, the automated process does not require the presence of an operator during deposition. Preliminary tests showed that a high degree of condensation was achieved after 40 min of reaction. Hence, addition rates were set to one monolayer/hour and 0.5 monolayer/hour for substrates with surface areas below and above 100 m²/g, respectively, to ensure full condensation of the precursor on the substrate surface. After deposition of the last metal oxide layer, highly reactive Ti-Cl

![Fig. 1. Precursor formation and cycle deposition on silica surface.](image-url)
bonds remained within the coating network due to stoichiometric excess of chloride ligands during condensation. To avoid the formation of agglomerates by uncontrolled hydrolysis/condensation reactions after synthesis, the material was first quenched by refluxing in boiling anhydrous diethyl ether to exchange highly reactive chloride ligands with relatively more stable ethoxides. Further washing with non-anhydrous diethyl ether and ethanol removed the condensation by-products iPrCl and HCl and progressively hydrolyzed ethoxide ligands to hydroxyl groups and oxo-bridges. Finally, full cross-linking within the metal oxide network was ensured by aging the materials in a dilute aqueous ammonia solution. Residual chlorine was detected by Energy-Dispersive X-ray Spectroscopy (EDXS) (Fig. S3) but its quantitative removal was achieved after calcination under air flow at 500 °C (Fig. S4).

3.2.1. Deposition on model Cu₂O and SiO₂ substrates

Cu₂O and SiO₂ spheres were selected as model oxide materials because their spherical shape allows precise thickness determination by TEM. High Angle Annular Dark Field Scanning Transmission Electron Microscopy (HAADF STEM) together with EDXS mapping pictures reveal the formation of conformal overcoats after depositing TiO₂ on both SiO₂ and Cu₂O (Fig. 2A and C). The deposition of TiO₂ on Cu₂O illustrates the compatibility of our method with substrates presenting low hydroxyl group density [52]. Coverage calculations suggest the lower limit for the surface hydroxyl density is 2 OH/nm². A lower density would lead to less than full coverage and could modify the resulting deposition. The in-depth line scan profile for 30 TiO₂@Cu₂O suggests the presence of a dense overcoat located only at the surface of the substrate. The TiO₂ signal sharply and symmetrically decreases to noise at the material interface and with distance from the substrate (Fig. 2B).

TiO₂@Cu₂O overcoat thickness could be precisely adjusted by the number of deposited monolayers with a growth rate of 0.4 nm/monolayer (Fig. 2D). In addition, BET surface areas (S_BET) calculated from N₂ physisorption isotherms revealed the formation of a porous overcoat with increasing surface area for increasing overcoat thicknesses (Fig. 2D). The 20 first layers show a steep S_BET increase from 18 m²/g for the initial Cu₂O spheres to 50 m²/g. From 20 layers, we observed a decrease of S_BET growth rate per layer. We attributed this non-linearity to an initial surface chemistry modification effect, which gradually decreases as further layers are eventually grafted on a surface that is identical to a pure metal oxide. Furthermore, we did not observe any decrease in surface area that may be caused by particle agglomeration for high layer numbers (>20). Titania loading ranged from 5.5 wt.% for 10 TiO₂@Cu₂O to 19 wt.% for 40 TiO₂@Cu₂O (see Tables S1 and S2 for all values). The porosity and structure of the coating can be further modified by ageing and calcination. While deposition of 15 TiO₂ layers on SiO₂ spheres does not influence pore size distribution even after calcination at 500 °C, mesoporosity around 5 nm is generated by ageing in basic aqueous solution (0.1 M NH₃) for 12 h at 60 °C (Fig. S6A). Generation of mesoporosity is accompanied by an increase of BET surface area and total pore volume (Fig. S6B). For coated samples with and without ageing, calcination causes a decrease in pore volume and surface area (Fig. S6B). Generation of mesopores is expected during TiO₂ NHSG synthesis [53], we suspect the mesopores are formed after ageing due to removal of residual ligands that were blocking these pores.

3.2.2. Deposition on high-surface area SBA-15

To illustrate the applicability of the overcoating technique to high surface area materials, we investigated the deposition of

Fig. 2. TEM images and corresponding EDXS mapping for (A) 30 TiO₂@Cu₂O (Cu: green; Ti: red); (B) In depth profile for 30 TiO₂@Cu₂O elemental mapping along the white line drawn in (A); (C) 10 TiO₂@SiO₂ (Si: green; Ti: red); (D) BET surface area and average thickness for increasing TiO₂ overcoat layers on Cu₂O spheres. Selected representative TEM pictures used for thickness measurements are shown in Fig. S5. The thickness for the 10-layers sample is not shown, because the coatings were too thin for reliable thickness measurements.
TiO₂ layers within the mesoporous structure of SBA-15 silica. We overcoated mesoporous silica with 1, 2, 4, 6, 8 and 10 layers TiO₂, using the technique described above.

Conformal overcoat deposition is typically challenging on such materials, but EDXS mapping after deposition of 10 TiO₂ layers clearly reveals uniform distribution of titanium into the mesopores of SBA-15, while no TiO₂ aggregates were observed on SBA-15 outer surface (Fig. 3A–C). In addition, no evidence of homogeneous nucleation was detected (see Fig. S7 for additional STEM pictures and EDXS mapping at lower magnification). HAADF STEM images show retention of the 2D hexagonal structure with a unit-cell parameter of 10.5 nm. EDXS line scan across the pore channels revealed coherence of Si and Ti intensities, suggesting that TiO₂ layers formed during overcoating are inside SiO₂ pore walls (Fig. 3D). Due to high surface area of SBA-15 (690 m²/g), titania loading reached 68 wt.% after deposition of 10 TiO₂ layers (see Table S3 for all values).

Diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) shows a progressive red-shift of the hydroxyl group band upon deposition of TiO₂ layers from 3745 cm⁻¹ (SBA-15) to 3735 cm⁻¹ (10TiO₂@SBA-15, Fig. S8). This illustrates a transition from a surface covered by isolated silanol groups to the creation of a population of Ti-OH groups. ¹H magic angle spinning (MAS) solid-state nuclear magnetic resonance (ssNMR) was investigated, as chemical shifts are very sensitive to the nature of surface hydroxyl groups [54]. The spectra of pure SBA-15 displayed a single peak at 2 ppm, which is characteristic of surface silanols. This peak was no longer observed after deposition of just one monolayer equivalent of TiO₂, which was consistent with the full coverage of these silanols after deposition of the first layer (Fig. 3E). Furthermore, an additional peak was observed at 2.3 ppm for 1TiO₂@SBA-15, attributed to =Si(OH)OTi species formed by the reaction of geminal silanol =Si(OH)₂ groups with the TiO₂ precursor. This peak was no longer observed after deposition of 8 TiO₂ layers, consistent with the formation of a thicker TiO₂ overcoat. Finally, hydroxyl groups bound to tetrahedrally coordinated Ti⁴⁺ were observed around 3.3 ppm after deposition of TiO₂ [55]. Those species are characteristic of amorphous titania, while hydroxyl groups on 6-coordinated Ti⁴⁺ sites from anatase and rutile TiO₂ phases are expected below 2.5 ppm [55].

¹H,²⁹Si cross polarization (CP) ssNMR was investigated to probe modification of surface Si groups after TiO₂ deposition. As opposed to single pulse ssNMR, CP ssNMR conditions highlight the surface species because this technique is more sensitive to ²⁹Si that are spatially close to surface silanols compared to bulk species. The ¹H-²⁹Si CP ssNMR spectrum of pure SBA-15 (Fig. 3F) presented the 3 characteristic ²⁹Si resonances for bulk siloxanes (Q₄; =Si–=Si: –111 ppm), silanols (Q₃; =SiOH: –100 ppm) and geminal silanols (Q₂; =Si(OH)₂: –90 ppm). After deposition of TiO₂, the intensity of the Q₂ peak decreased due to the transformation of geminal silanols into silanols (=Si(OH)OTi) (Q₃), in agreement with ¹H NMR results. The latter Q₃(Ti) peak, expected around ~98 ppm [56], was most likely contributing to the broad resonance at ~100 ppm. Deposition of TiO₂ was also characterized by the appearance of a new peak at ~108 ppm attributed to the formation of Q₃(Ti) (=SiOTi) by reaction of surface =SiOH (Q₃) with the TiO₂ precursor [56]. Overall, MAS ssNMR confirmed that the silica surface was fully covered and that hydroxyl groups were formed on amorphous Ti⁴⁺ tetrahedral sites, in agreement with the absence of the crystalline TiO₂ pattern in the X-ray diffractogram (Fig. S9).

Retention of the SBA-15 long-range order was further investigated by small-angle X-ray diffraction (Fig. S10). Typical SBA-15 diffraction peaks were observed at 2₀ = 0.86, 1.5 and 1.75°, corresponding to the (1 0 0), (1 1 0) and (2 0 0) reflections, respectively, consistent with p6mm symmetry of the silica mesoporous structure [57]. The intensity of the main diffraction peak at 0.86° decreased gradually with the increasing number of layers. The other reflection peaks were considerably weaker and were not retained after the TiO₂ overcoat deposition. Finally, d-spacing calculated from Bragg equation [58] on the main peak decreased from 10.3 nm (SBA-15) to 9.6 nm and 9.5 nm after deposition of 6 and

Fig. 3. HAADF STEM picture (A) and EDXS maps of Ti (B) and Si (C) for 10TiO₂@SBA-15; (D) line scan showing abundance of Si and Ti along the line visible on (A) (smoothing by average was performed to minimize experimental noise); ¹H (E) and ¹H-²⁹Si CP (F) MAS ssNMR spectrum of SBA-15 and after deposition of 1, 4, 6, 8 and 10 layers TiO₂.
10 layers, respectively, in agreement with STEM measurements (Fig. 3A). Perturbation of the SBA-15 diffraction pattern was expected upon overcoating, but the absence of major shift of the (1 0 0) reflection confirms retention of the SBA-15 structure.

Nitrogen physisorption revealed a decreasing surface area upon deposition of TiO$_2$ layer onto SBA-15, with a drop from 480 m$^2$/g for SBA-15 to 275 m$^2$/g after deposition of 10 layers (Fig. 4C). The BET surface area dropped significantly after the deposition of the first two layers, which was attributed to rapid filling of micropores. DFT micropore analysis from N$_2$ physisorption confirmed a significant decrease of microporosity compared to regular SBA-15 (Fig. S11). Gradual pores filling did not alter the pore structure, as the type H1 hysteresis loop observed in the isotherms was retained and is consistent with capillary pore filling associated with a mesoporous channel. Specifically, the parallel adsorption and desorption branches excluded the formation of TiO$_2$ islands or other effects that would modify the shape of the channels (Fig. 4A). The hysteresis loop gradually lost verticality, consistent with a widening of the mesopore size distribution (Fig. 4B). We observed a light shift of the hysteresis loop toward higher partial pressure upon deposition of 2 and 4 layers, consistent with pores widening (Fig. 4B). This effect could be attributed to an increase of mean hydraulic ratio (i.e. pores volume to pore internal surface area) caused by micropore filling (i.e. sharp decrease surface area) and minimal mesopores shrinking upon deposition of first 4 layers [59]. This explanation is consistent with the line scan shown in Fig. 3D that showed incorporation of TiO$_2$ within the channel walls. Depositon of further layers (>6) lead to a mean pore diameter decrease, which was more in line with what is expected after deposition (Fig. 4A and B) [12].

3.3. Formation of acid sites: TiO$_2$@SBA-15

While SBA-15 does not have significant acidity, acidity can be generated by the formation of mixed oxide at the titania-silica interface [60]. We observed controlled acid site formation during overcoating. Ti-SBA-15 materials prepared by co-condensation of tetraethoxysilane and titanium isopropoxide have been reported [61,62], but this synthesis route does not allow control of the nature of surface sites in the same way as our overcoating approach does.

Acid sites were titrated using ammonia Temperature Programmed Desorption (NH$_3$ TPD), which revealed the presence of acid sites characterized by a desorption peak centered around 200 °C, characteristic of medium acid strength (Fig. 5A). Deposition of 2 layers TiO$_2$ created 33 μmol acid sites per gram and a maximum was reached for 6 TiO$_2$ layers with 83 μmol acid sites per gram (Fig. 5B). The number of acid sites dropped after deposition of 10 TiO$_2$ layers (35 μmol/g), which is consistent with the creation of a pure TiO$_2$ phase. As a comparison, we measured 517 μmol acid sites per gram on HZSM-5 with a second desorption peak at 300 °C attributed to strong acid sites (Fig. 5A), whereas pure TiO$_2$ shows a broad distribution of acid strength. Pyridine adsorption monitored by Fourier Transformed Infra-Red spectroscopy was used to further identify the nature of acid sites. The use of Diffuse Reflectance (DRIFT) does not allow precise acid site quantification but density trends can be observed in addition to qualitative sites identification. While negligible interaction was observed after exposure of SBA-15 to pyridine vapor, two bands characteristic of Lewis acid sites were observed after deposition of 2 TiO$_2$ layers at 1445 and 1605 cm$^{-1}$ (Fig. 5C). A third band at 1490 cm$^{-1}$ can be attributed to both Lewis and Bronsted acid sites but the absence of bands at 1545 or 1640 cm$^{-1}$ reveals the absence of significant Bronsted acidity on all materials [63,64]. Since Bronsted acidity in silica-supported TiO$_2$ materials has been recently attributed to the presence of chlorides and alkoxides impurities [60,65], those data further confirmed the efficient elimination of any precursor residue from the metal oxide framework during post-synthetic treatment. The maximum intensity for the peak attributed to Lewis acid sites was observed for 6 TiO$_2$ layers where hydrogen bonding was observed as well with a band at 1590 cm$^{-1}$. Beyond 6 TiO$_2$ layers, the band intensity decreased which is consistent with the formation of a pure TiO$_2$ surface, in agreement with FTIR and ssNMR spectroscopy results. In summary, the TiO$_2$ overcoat created a tunable number of medium strength Lewis acid sites on SBA-15 with a narrow acid strength distribution.

In view of the tunable number of medium strength Lewis acid sites generated on TiO$_2$@SBA-15, we investigated its use for the catalytic 1-phenylethanol dehydration, which suffers from poor yields and catalyst deactivation issues. In the past, zeolites such as HZSM-5 have been indicated as most suitable metal free catalysts for this reaction, but the presence of strong acid sites leads to fast deactivation and low selectivity due to styrene polymerization [66–68].

The reactivity of TiO$_2$@SBA-15 was investigated at 150 °C in 1,4-dioxane and compared to the reference HZSM-5. The zeolite could achieve 60% selectivity at 4% conversion but selectivity dropped quickly with increasing conversion due to the formation of dimers and styrene oligomers, reaching only 12% at 99% conversion (Fig. 6A). Oligomers are not detected by gas chromatography but...
In comparison, the use of 6TiO2@SBA-15 as a catalyst was attributed to strong acid sites characterized by a desorption peak centered at 400 °C in NH3 TPD (Fig. 5A). Styrene polymerization on HZSM-5 can also be limited by lowering temperature to 95 °C and using cyclohexane as the solvent, as reported by Bertero et al. (reaching a 63% styrene yield compared to 57% here) [68]. Conversely, the controlled acidity on 6TiO2@SBA-15 was sufficient to dehydrate phenylethanol but did not lead to fast styrene degradation.

The reactivity of TiO2@SBA-15 was strongly related to acidity and dehydration kinetics, and gradually increased with thicker TiO2 overcoats in line with the increased number of acid sites (Table 1). The use of pure SBA-15 led to no measurable 1-phenylethanol conversion. While 2TiO2@SBA-15 was not very active, activity increased with thicker layer reaching 6.7 mmol PHE converted per gram of catalyst per min for 6TiO2@SBA-15. Selectivity was also affected by the number of deposited layers, reaching a maximum for 8 TiO2 layers (57% at 99% conversion, Fig. 6C). Finally, TiO2 P25 has slightly lower selectivity at high conversion (50% at 99% conversion) but its activity was about 2–3 orders of magnitude lower (Table 1).

Contrary to HZSM-5, the mesoporous structure of TiO2@SBA-15 does not provide shape selectivity, but the formation of targeted acidity maximized activity while preventing further condensation reactions on strong acid sites. This controlled acidity led to higher selectivity and limited catalyst deactivation. Our results demonstrate that NHSG overcoating can improve catalytic selectivity by controlling the metal oxide surface functionalities.

3.4. Metal-acid bifunctional catalysis

We also explored the overcoating of supported metal particle catalysts to create metal and acid sites in close proximity at the interface between metal, overcoat and support. Interfacial effects are of great interest for various catalytic applications [69–71]. Specifically, we investigated the compatibility of the NHSG TiO2 deposition with the presence of Pt nanoparticles (NPs) supported on SiO2 spheres. First, platinum NPs were deposited on SiO2 spheres to form Pt/SiO2, with a mean nanoparticle size of 3.3 ± 2.1 nm (2.0 wt.% Pt; Fig. 7A). After deposition of 10 TiO2 layers onto Pt/SiO2, STEM (Fig. 7B and C) and EDXS mapping (Fig. 7D) showed the formation of a conformal overcoat (6.1 wt.% TiO2, see Table S4 for all values). After deposition of 20 TiO2 layers (Fig. 7E), the coating was still conformal but became coarser. The platinum nanoparticles size distribution was unaffected by deposition of TiO2 (3.6 ± 1.3 nm for 10TiO2@Pt/SiO2; see histograms in Fig. S15), attesting to the compatibility of our deposition method with the presence of metal NPs. Furthermore, the platinum particles were systematically located at the interface between the SiO2 sphere surface and the overcoat (Fig. 7C). The surface area of the materials increased linearly with the number of TiO2 layers (Fig. S16). As a control to determine the importance of having platinum located at the interfacial acidic sites, platinum NPs were also deposited on SiO2 overcoated with 10 TiO2 layers (denoted as Pt/10TiO2@SiO2), where we expected the platinum NPs to be located on the top of the TiO2 overcoat instead of at the interface between overcoat and support (Fig. 7F). In this case, the likely stronger metal-support interaction after deposition of Pt on the acidic 10TiO2@SiO2 compared to pure SiO2 led to smaller NPs size (2.0 ± 1.0 nm). Finally, 10 TiO2 layers were deposited on Pt/TiO2 to explore the importance of mixed oxide formation (sample denoted as 10TiO2@Pt/TiO2).

We investigated the performances of our bifunctional platinum/acid sites in the hydrodeoxygenation of oxygenated aromatics derived from the lignin fraction of biomass [72–83]. The presence of their presence is correlated to the decreased carbon balance and consistent with the brown color of the catalyst after reaction (Fig. S12). During a recycling study, HZSM-5 deactivated quickly and 55% selectivity at 99% conversion (Fig. 6B). Stability of styrene under the reaction conditions was also investigated and its conversion reached 48% within 3 h using the zeolite while almost 2 days were needed to reach 44% conversion with 6TiO2@SBA-15 (Fig. S13). The fast polymerization of styrene on HZSM-5 was attributed to strong acid sites characterized by a desorption peak centered at 400 °C in NH3 TPD (Fig. 5A). Stability of styrene on HZSM-5 can also be limited by lowering temperature to 95 °C and using cyclohexane as the solvent, as reported by Bertero et al. (reaching a 63% styrene yield compared to 57% here) [68].

The NHSG TiO2 deposition with the presence of Pt nanoparticles (NPs) supported on SiO2 led to smaller NPs size (3.3 ± 2.1 nm for 2.0 wt.% Pt; Fig. 7A). After deposition of 10 TiO2 layers onto Pt/SiO2, STEM (Fig. 7B and C) and EDXS mapping (Fig. 7D) showed the formation of a conformal overcoat (6.1 wt.% TiO2, see Table S4 for all values). After deposition of 20 TiO2 layers (Fig. 7E), the coating was still conformal but became coarser. The platinum nanoparticles size distribution was unaffected by deposition of TiO2 (3.6 ± 1.3 nm for 10TiO2@Pt/SiO2; see histograms in Fig. S15), attesting to the compatibility of our deposition method with the presence of metal NPs. Furthermore, the platinum particles were systematically located at the interface between the SiO2 sphere surface and the overcoat (Fig. 7C). The surface area of the materials increased linearly with the number of TiO2 layers (Fig. S16). As a control to determine the importance of having platinum located at the interfacial acidic sites, platinum NPs were also deposited on SiO2 overcoated with 10 TiO2 layers (denoted as Pt/10TiO2@SiO2), where we expected the platinum NPs to be located on the top of the TiO2 overcoat instead of at the interface between overcoat and support (Fig. 7F). In this case, the likely stronger metal-support interaction after deposition of Pt on the acidic 10TiO2@SiO2 compared to pure SiO2 led to smaller NPs size (2.0 ± 1.0 nm). Finally, 10 TiO2 layers were deposited on Pt/TiO2 to explore the importance of mixed oxide formation (sample denoted as 10TiO2@Pt/TiO2).

We investigated the performances of our bifunctional platinum/acid sites in the hydrodeoxygenation of oxygenated aromatics derived from the lignin fraction of biomass [72–83]. The presence of
of acid sites close to metal nanoparticles has been shown to favor C—O bond cleavage and therefore deoxygenation. We studied hydrodeoxygenation of 3-propylguaiacol, one of the major monomers derived from lignin depolymerisation [39] and targeted 100% selectivity of propylcyclohexane instead of a mixture of partially deoxygenated products such as substituted phenol and cyclohexanol (Fig. 8A). A reaction at 200 °C and 15 bar hydrogen in isooctane using either Pt/SiO$_2$ or Pt/TiO$_2$ yielded low selectivity to propylcyclohexane (5 and 20% for 20 and 23% conversion, respectively) with an important formation of oxygenated product at various conversions (Fig. 8A). Selectivity to propylcyclohexane is significantly higher for 10TiO$_2$@Pt/SiO$_2$ (reaching 90% at 95% conversion) even at low conversion (selectivity of 40% at 20% conversion), clearly illustrating the benefit from the presence of Lewis acid sites in close proximity with the platinum particles. The highest selectivity was achieved for 5-10TiO$_2$@Pt/SiO$_2$ (depending on conversion) while deposition of 2 TiO$_2$ layers only marginally improved selectivity at low conversion but was equivalent to other overcoated materials at high conversion (Fig. 8B). Interestingly, Pt/10TiO$_2$@SiO$_2$ and Pt/ASA (platinum particles supported on amorphous silica-alumina) led to low selectivities that were similar to the uncoated catalysts. This result emphasizes the importance of the close proximity of the metal and the Lewis acid sites, which can be achieved only when supported metal nanoparticles are overcoated (Fig. 8C). Deposition of 10 TiO$_2$ layers onto Pt/TiO$_2$ (10TiO$_2$@Pt/TiO$_2$) did not improve selectivity due to the absence of mixed oxide at the interface. It should be noted that the activity of the overcoated materials was significantly reduced due to partial blockage of surface metal sites by the TiO$_2$ overcoat, dropping from 10.7 μmol h$^{-1}$ g$_{\text{cat}}^{-1}$ for Pt/SiO$_2$ to 0.3 μmol h$^{-1}$ g$_{\text{cat}}^{-1}$ for 2TiO$_2$@Pt/SiO$_2$ (Fig. S18). The amount of accessible Pt surface sites was titrated using volumetric CO chemisorption, and showed

![Fig. 6. Selectivity and conversion vs time for 1-phenylethanol dehydration for (A) HZSM-5 and (B) 6TiO$_2$@SBA-15; (C) Styrene selectivities at 99% conversion and initial activities (mmol PHE converted cm$^{-2}$ min$^{-1}$); (D) recycling test for 6TiO$_2$@SBA-15 and HZSM-5 performing 1-phenylethanol dehydration. Conversion and selectivity plots for all materials are provided in Fig. S14.](image-url)

![Fig. S14](image-url)

**Table 1**

<table>
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<tr>
<th>Entry</th>
<th>Material</th>
<th>(S_{\text{BET}}) (m$^2$/g)</th>
<th>Initial activity (mmol$_{\text{PHE}}$ converted per g cat per min)$^a$</th>
<th>Initial activity (mmol$_{\text{PHE}}$ per cm$^2$ cat per min)</th>
<th>Acid sites (μmol/g)$^b$</th>
<th>Initial TOF (mol PHE per mol acid site per min)</th>
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$^a$ Determined using the initial slope.

$^b$ Acid sites quantified by NH$_3$ TPD; all selectivity and conversion plots are presented in Fig. S14 as well as styrene selectivities at 80% conversion, Fig. S14g.
a 5-fold decrease in the ratio of accessible Pt sites after deposition of 2 TiO$_2$ layers (Fig. S19). The extent of metal coverage by the overcoat is comparable with reported ALD overcoated materials \[2,23\], but the possibility of tuning overcoat porosity enabled by our solution phase synthesis is currently under investigation to improve metal access. Our previous work involving liquid phase overcoating with alumina further suggests that TiO$_2$ overcoat may also be able to suppress catalyst deactivation by nanoparticle sintering \[38\].

4. Conclusions

By using NHSG chemistry, we were able to create highly conformal TiO$_2$ overcoats over a diverse range of metal oxide and metal
nanoparticle catalysts. The coverage of the materials, including high-surface-area SBA-15 was comparable to ALD despite being an easily scalable 1-step process in liquid phase conditions. These overcoats can be used to create acid sites at the interface with the substrate with a strength and density that can be controlled by varying the overcoating thickness. Overcoating supported metal nanoparticle catalysts led to similar sites that, because they were in close proximity to metal sites, greatly increased the catalyst's deoxygenation ability. Since we demonstrated that this method is compatible with a wide range of metal oxides, including high surface area materials and supported metal nanoparticle catalysts, we anticipate that it could be used for the post-synthesis modification of a wide variety of important catalytic materials. Finally, we propose that, because this overcoating method leads to control over the selectivity and activity of particular catalytic site, it could be especially valuable for many increasingly important biomass conversion reactions where the breadth of functional groups routinely leads to unwanted side reactions.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version, at https://doi.org/10.1016/j.jcat.2017.11.023.

References


