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Modelling atomic layer deposition overcoating formation on a porous heterogeneous catalyst[†]

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Atomic layer deposition (ALD) was used to deposit a protective overcoating (Al₂O₃) on an industrially relevant Co-based Fischer–Tropsch catalyst. A trimethylaluminium/water (TMA/H₂O) ALD process was used to prepare ~ 0.7 –2.2 nm overcoatings on an incipient wetness impregnated Co–Pt/TiO₂ catalyst. A diffusion–reaction differential equation model was used to predict precursor transport and the resulting deposited overcoating surface coverage inside a catalyst particle. The model was validated against transmission electron (TEM) and scanning electron (SEM) microscopy studies. The prepared model utilised catalyst physical properties and ALD process parameters to estimate achieved overcoating thickness for 20 and 30 deposition cycles (1.36 and 2.04 nm respectively). The TEM analysis supported these estimates, with 1.29 \pm 0.16 and 2.15 \pm 0.29 nm average layer thicknesses. In addition to layer thickness estimation, the model was used to predict overcoating penetration into the porous catalyst. The model estimated a penetration depth of $\sim 19~\mu m$, and cross-sectional scanning electron microscopy supported the prediction with a deepest penetration of 15–18 μm . The model successfully estimated the deepest penetration, however, the microscopy study showed penetration depth fluctuation between 0–18 μm , having an average of 9.6 μm .

1 Introduction

Atomic layer deposition (ALD) presents an interesting path to modify heterogeneous catalysts with active or inert inorganic compounds.¹ ALD overcoating provides a versatile toolbox for catalyst active site modification and protection. ALD techniques enable precise control and conformality over other methods, such as solution-based overcoatings.²-⁴ Several publications give examples of ALD overcoatings preventing catalyst deactivation through leaching and sintering,⁴-⁶ where catalysts are modified through repeated self-limiting reaction cycles to achieve desired overcoating thicknesses with atomic-level precision. Due to these properties, ALD process is also relevant for very high aspect ratio materials and nanoporous structures, such as catalysts.⁻٬8 After the deposition, overcoating structures can be modified to specific needs with temperature¹٬9,¹¹⁰ or reactive treatments.¹¹ Although ALD gives a wide variety of tools to modify different materials, the

Many of the successful examples 9,10,12 present the ALD overcoating deposition on a catalyst surface and overcoating formation around a specific active metal site. However, less information is available on the precursor penetration into a porous heterogeneous catalyst. Therefore, it is important to understand the ALD process and precursor parameter effect on the achieved penetration depth. In literature, this has been addressed with models to predict a single parameter, such as precursor sticking probability, 13 growth modes (growthper-cycle behaviour)14-16 as well as experimental methods to select ALD process conditions (temperature, pressure and exposure time). 17 Others present holistic models to estimate a full deposition process for varying precursors, experimental conditions and substrate materials. 18-20 An especially interesting approach for catalytic application has been related to porous material deposition modelling. 7,21-24 These examples present a foundation for modelling precursor transport and required exposure time to achieve the desired precursor penetration through the porous substrate. However, the previous modelling studies have not extensively been connected to microscopical examination and resulting catalytic performance. Our work presents a coupled diffusion-reaction differential equation model to estimate ALD overcoating penetration into a Co-Pt/TiO2 porous catalyst, similar to commercial

precursor transport into the porous substrates is no trivial question.

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catalysts listed by Rytter and Holmen.²⁵ The model is used to estimate deposition thicknesses with a given set of catalyst structural parameters and precursor parameters. The presented model is compared against microscopy (TEM and SEM) measurement data.

2 Materials and methods

2.1 Atomic layer deposition precursor diffusion model

To address atomic layer deposition precursor transport into the selected TiO₂ catalyst structure, a diffusion-reaction partial differential equation model was prepared based on the work of Yanguas-Gil and Elam, 7,18,22 Keuter et al. 23 and Ylilammi et al. 19 In the presented model, the precursor was considered as an ideal gas with homogeneous concentration over the catalyst surface. The porous substrate (catalyst) was characterised by mean porosity, mean tortuosity and mean pore size, resulting in the utilisation of the mean diffusion coefficient (Knudsen diffusion coefficient, D_{ki}). As the precursor diffuses through the porous material, the precursor density decreased because of interaction with surface reactive sites, following secondorder reaction kinetics. A practical simplification was made to assume reactive sites having uniform distribution on the porous substrate, having equal reaction probability with the precursor molecule. After the surface reaction, the precursor molecule had no ability for desorption or surface migration.

The prepared model consists of two coupled diffusion-reaction differential equations: 7,23

$$\frac{\partial n_{\rm P}(t,z)}{\partial t} = D_{\rm ei} \frac{\partial^2 n_{\rm P}(t,z)}{\partial z^2} - \bar{s}_{4}^1 v_{\rm th} \beta_0 \cdot n_{\rm P}(t,z) \cdot \Theta(t,z) \tag{1}$$

$$\frac{\partial \Theta(t,z)}{\partial t} = -s_0 \frac{1}{4} v_{\text{th}} \beta_0 \cdot n_{\text{P}}(t,z) \cdot \Theta(t,z)$$
 (2)

Eqn (1) gives the volumetric precursor density (n_P) , dependent upon the time (t) and the depth (z) inside the porous catalyst particle. As the extent of surface reaction depends on both precursor density and available reaction sites, the consumed precursor density is related to the ratio (\bar{s}) of surface area (A_O) and pore volume (V_P) . The β_0 is a simplified estimation with the Langmuir equation for precursor sticking probability to the coated surface. The second coupled differential equation (eqn (2)) considers the fraction of available reaction sites $\Theta(t,z)$, where s_0 is the average surface area of adsorption sites, also including the shielding effect of ligands (steric hindrance) on the precursor molecule surface coverage. The degree of surface coverage is determined as $1 - \Theta(t,z)$. The gas phase diffusion

coefficient in porous structures can be divided into two parts, *i.e.*, into molecular diffusion and Knudsen diffusion (eqn (3)).

$$\frac{1}{D_{i}} = \frac{1}{D_{mi}} + \frac{1}{D_{ki}} \tag{3}$$

where the gas phase diffusion coefficient is given by D_i , molecular diffusion coefficient by D_{mi} and Knudsen diffusion coefficient by D_{li} .

Due to catalyst pore dimensions and TMA partial pressure (1.3 kPa at 293 K), the transport of precursor molecules is dictated by Knudsen diffusion and therefore, ${D_{\rm mi}}^{-1}$ in eqn (3) can be neglected:

$$\frac{1}{D_{\rm i}} = \frac{1}{D_{\rm ki}} \to D_{\rm i} = D_{\rm ki} \tag{4}$$

The estimated TMA mean free path ($\sim 18~\mu m$) was significantly larger than the catalyst average pore radius of 28 nm. In the case of porous materials, it should be considered that the whole volume of a catalyst particle is not available for the diffusion. In order to take this into account, the voidage of a porous particle is expressed with porosity value ε , and tortuosity (labyrinth factor) is expressed by value τ . The Knudsen diffusion coefficient considers molecular movement and interaction with pore walls in a perfectly straight and cylindrical structure. Therefore, the determination of effective diffusion coefficient is expanded to eqn (5):

$$D_{\rm ei} = D_{\rm eki} = \frac{\epsilon}{\tau} D_{\rm ki} \tag{5}$$

where $D_{\rm ki}$ is multiplied with a simplification factor of the tortuous, discontinuous and complex pore structure. Then, the effective diffusion coefficient ($D_{\rm ei}$) for the studied porous catalytic material is based on Knudsen diffusion and is given as:

$$D_{\rm ei} = \frac{\epsilon}{\tau} \frac{2}{3} r \sqrt{\left(\frac{8k_{\rm B}T_{\rm P}}{\pi m_{\rm P}}\right)} \tag{6}$$

Values of parameters in eqn (6), *i.e.*, mean pore radius (r), porosity (ϵ) and tortuosity (τ) are presented in Table 1. In addition to the catalyst structural parameters, Table 2 gives trimethylaluminium (TMA) precursor parameters required for the used model. Fig. 1 presents a schematic for model domains and eqn (7)–(10) give the initial and boundary conditions applied to solve partial differential equations (eqn 1 and 2). The table presents only precursor parameters for TMA, neglecting the second half-reaction compound (H₂O). This was due to the smaller mass and molecular dimensions of water, enabling easier and faster diffusion into the porous material. Therefore,

Table 1 Catalyst structural parameters

Parameter	Value	Source
Porosity, ε Tortuosity, τ Mean pore radius, r BET surface area, $A_{\rm O}$ Pore volume, $V_{\rm P}$	$\begin{array}{c} 0.213 \\ 32.1 \\ 2.77 \times 10^{-8} \text{ m} \\ 38.3 \text{ m}^2 g_{\text{cat}}^{-1} \\ 2.65 \times 10^{-7} \text{m}^3 {g_{\text{cat}}}^{-1} \end{array}$	${ m TiO_2}$ support manufacturer ${ m TiO_2}$ support manufacturer BJH measurement BET measurement BJH measurement

Trimethylaluminium (TMA) precursor parameters

Parameter	Value	Source
Temperature, $T_{\rm P}$	423.15 K	_
Precursor mass, $m_{\rm P}$	$72.082 \times 1.66 \times 10^{-27} \text{ kg}$	_
Maximum density of particles adhering to the surface, $\sigma_{\rm P}$	$5 \times 10^{18} \text{ m}^{-2}$	8, 26–28
Precursor density outside the porous material, $n_P(t, z = 0)$	$3.21 \times 10^{23} \text{ m}^{-3}$	a
Sticking probability, eta_0	2×10^{-3}	19, 27, 29, 30

^a Determined from TMA vapour pressure (1.3 kPa at 293 K). ³¹

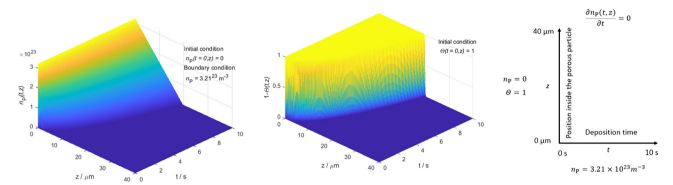


Fig. 1 Precursor depletion profile in depth and time domains of the model (left) with a set of initial and boundary conditions. Surface coverage profile in depth and time domains of the model (middle) with initial condition. A schematic illustration of the used model domains (right)

TMA half-cycle was considered as the limiting factor in overcoating formation. 19,23

2.1.1 Initial conditions

$$\Theta(t=0,z)=1, \forall z \in [0,40 \,\mu\text{m}] \tag{7}$$

$$n_{\rm P}(t=0,z)=0, \forall z \in [0,40 \ \mu {\rm m}]$$
 (8)

2.1.2 Boundary conditions

$$n_{\rm p}(t, z=0) = 3.21 \times 10^{23} \,\mathrm{m}^{-3}, \,\forall \, t \ge 0$$
 (9)

$$\left. \frac{\partial n_{\rm P}(t,z)}{\partial z} \right|_{z=40\,\mu{\rm m}} = 0, \forall t \ge 0$$
 (10)

2.2 Solution of partial differential equation system

The partial differential equation system was solved by applying Matlab software. The second derivative of spatial coordinate z (depth in the porous catalyst particle) was discretised using the finite differences method. This was done using a Matlab modification of the dss044 algorithm developed by Schiesser.³² In this algorithm, the second derivative is discretised applying five-point central difference formulas, based on fourth-order approximation. The formed system of ordinary differential equations (ODEs) was solved using the Matlab algorithm ode15s, suitable for stiff ODE systems.

2.3 Catalyst preparation

Stepwise co-impregnation was used to prepare the Co-Pt/TiO₂ catalyst. In the first step, a TiO2 support (ACCU® SPHERE 0.4 mm ST32244, $S_{\rm BET}$ 52 m₂ g⁻¹, $V_{\rm pore}$ 0.37 mL g⁻¹ and $d_{\rm pore}$ 28.9 nm) was dried at 100 °C under a vacuum for 1 h. The support was then co-impregnated twice with an aqueous solution of cobalt nitrate (Co(NO₃)₂·6H₂O) and platinum(IV) nitrate (Pt(NO₃)₄). Between impregnation steps, the catalyst was dried in a rotary vacuum evaporator and calcined at 300 °C for 4 h. The resulting catalyst contained 21.0 wt% cobalt and 0.2 wt% platinum. The ALD (trimethylaluminium (TMA)/ water) process was carried out using Picosun R-200 ALD equipment. Nitrogen (purity 99.999%) was used as a carrier gas and TMA deposition cycles (10, 20 and 30) were performed at 150 °C (TMA purity 99.999%, Sigma-Aldrich), with a stop-flow deposition and purge cycles of 10 s/80 s/10 s/80 s. The stop-flow cycle had a 0.7 s precursor pulse and 9.3 s equilibrium time. Deionised water was used as a second half-reaction reagent (with stop flow equilibrium time). The TMA and water were evaporated at room temperature. After the ALD process, the resulting catalyst samples required heat treatment to re-open catalyst active sites beneath alumina overcoating. 10,33 The heat treatment was conducted in the tubular reactor prior to FT reaction. For catalyst characterisation experiments, the heat treatment was performed with a Micromeritics 3Flex 3500 instrument. The heat treatment was done in an N_2 flow (40 N mL min⁻¹ g_{cat}^{-1}), having a heating ramp rate 4 °C min⁻¹ from 25 °C to 420 °C at atmospheric pressure.

Catalyst characterisation

2.4.1 Nitrogen adsorption and desorption. The N₂ physisorption experiments were carried out with a Micromeritics 3Flex 3500 instrument. A catalyst sample (~300 mg) was placed in a VacPrep degassing station and kept at 150 °C for 12 h under a vacuum (10⁻² mbar). After degassing, the tube was mounted on the measuring instrument. The catalyst surface

area was estimated using the Brunauer-Emmett-Teller (BET)34 equation, and the Barrett-Joyner-Halenda (BJH)35 method was used for total pore volume and average pore diameter determination. The average pore diameter was evaluated from the nitrogen desorption branch.

2.4.2 Transmission electron microscopy (TEM). The transmission electron microscopy studies were performed in a Jeol IEM 2800 analytical HR-TEM at an accelerating voltage of 200 kV. The samples are prepared by crushing the catalyst particles in a mortar and pestle. The powder is redispersed in acetone and sonicated to achieve suitable dispersion. Of this suspension, 10 µL is drop casted on an EMR holey carbon 400 square mesh Cu grid and air dried. Prior to TEM investigation, the grids are preserved under rotary vacuum conditions.

2.4.3 Scanning electron microscopy-energy-dispersive X-ray spectrometer (SEM-EDS). Scanning electron microscopy (SEM) and energy-dispersive X-ray spectrometry (EDS) examinations were conducted with a Zeiss Crossbeam scanning electron microscope, having an EDAX (energy-dispersive X-ray) spectrometer. Sample casts were prepared by the addition of epoxy resin (~ 23.3 g, EpoFix, Struers), mixing with ~ 10 mg of carbon powder (VULCAN XC72R GP-3875, CABOT) and stirred by hand to achieve a homogeneous mixture. Then, 2.8 g of hardener (EpoFix hardener, Struers) was added into the cast mixture. The mixture was poured into a mould after inserting catalyst sample particles into the mould. The resin mixture was cured for 24 hours, and cured cast was ground to expose crosssections of the catalyst particles. Before the SEM experiment, the cured cast was sputtered with Pt to induce a signal from the sample. Electron acceleration voltage was 15 keV.

3 Results and discussion

3.1 Overcoating penetration to the porous catalyst structure

Fig. 2 presents a SEM image of overcoated catalyst sample particles. Catalyst support manufacturer (Saint-Gobain Norpro, ACCU® SPHERE ST32244) reported measured particle size distribution for catalyst support as 396 \pm 23 μ m. In order to address ALD overcoating penetration experimentally, catalyst particles (similar to Fig. 2) were casted in resin, sanded and polished to hemispherical shape.

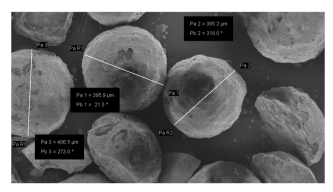


Fig. 2 Example SEM image of catalyst particles with measured diameters.

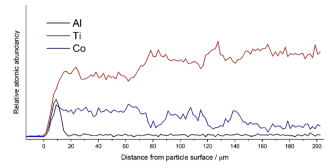


Fig. 3 SEM-EDS line scan from hemispherical catalyst particle having 30 cycle ALD overcoating. Atomic abundancies for Al, Ti and Co from the surface of the particle to the particle centre at 202 µm.

Fig. 3 gives the SEM-EDS line scan measurement spectra starting from the outside of the catalyst particle until the middle of the measured particle (~200 μm). Alumina overcoating had a clear response at the surface of the catalyst particle, ranging from 0-18 µm. In addition to the ALD overcoating, a clear boundary was observed for Co at \sim 75 µm. At a similar depth, the Ti response increased as relative atomic abundancy was increased from Co to the Ti response.

Line scan (Fig. 3) presents information only from the selected crosscut. Therefore, to address ALD overcoating penetration uniformity, Fig. 4 presents SEM-EDS elemental mapping images for two catalyst particles. Measured from these elemental mapping images (with ImageJ version 1.53 k), the overcoating depth had an average of 9.5 and 9.6 µm for fig. a and c respectively, while the

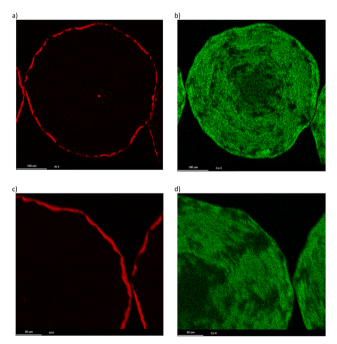


Fig. 4 SEM-EDS elemental mapping conducted on hemispherical catalyst particles having 30 cycle ALD overcoating. Elemental mapping channels for Al images (a and c), for cobalt images (b and d). In fig. (a), Al response at the middle of the particle resulted from the grinding process, leaving a small quantity of Al in this unexpected area.

deepest penetration was between 15–18 μ m. The cobalt penetration ranged between 80–140 μ m. The cobalt missing from the middle of the particles was an indication of successful grinding close to the assumed hemispherical shape. Interestingly, both cobalt elemental mapping and line scan show varying cobalt loading for distinct locations of the catalyst particle. This unevenness might have resulted from varying support densities having both anatase (80%) and rutile (20%) TiO₂ crystallite phases or non-uniform tortuosity and porosity throughout the catalyst particle. With the precursor diffusion–reaction model, the expected overcoating penetration depth was 19 μ m, corresponding well with Fig. 4, a measured maximum penetration depth of 15.2 μ m.

Fig. 5 presents a diffusion-reaction model estimate for the degree of surface coverage $(\Theta[t, z])$ as a function of the position inside the porous particle. With the used catalyst structural parameters (Table 1), deposition process and precursor parameters (Table 2), the estimated degree of surface coverage remains constant until ~19.1 μm, whereafter the precursor coverage depleted rapidly ($\Theta = 0$ at 19.9 µm). In addition to the model prediction, Fig. 5 presents the measured SEM-EDS line scan data. When measuring the line scan from the deepest observed penetration depth, the penetration profile corresponds closely to the model prediction. The Gaussian-like distribution with the SEM-EDS measurement data resulted from X-ray scattering at the measured surface. Due to the scattering effect, the Al signal slowly increased until 4-9 µm and started to decrease after 6-10 µm. The deepest penetration depth was similar to SEM-EDS elemental mapping with 15-18 μm. Therefore, the diffusion-reaction model is slightly overestimating the penetration depth. The overestimation might result from several factors (discussed later in detail in the section "Diffusion-reaction model parameter sensitivity"). However, the most relevant misinterpretation might be related

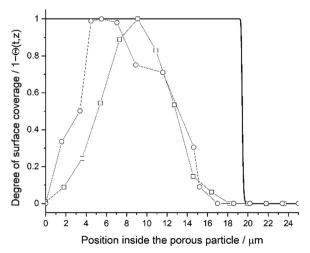


Fig. 5 Degree of surface coverage as a function of position inside the porous particle. Solid black line for model prediction overlayed with two separate SEM-EDS line scan measurements. First SEM-EDS line scan measurement with round markers and a repetition measurement from different location with square markers.

to the used precursor density outside the porous particle. The precursor density (m^{-3}) was calculated from TMA vapour pressure (1.3 kPa at 293 K), thus giving the ideal maximum precursor density outside the catalyst particle.

3.2 Overcoating thickness and conformality

The previous section discussed the overcoating penetration depth into the catalyst particle. In addition to the penetration profile estimation, the presented diffusion-reaction model was utilised to estimate overcoating thickness on the catalyst surface. The model predictions were compared to experimental measurements. Fig. 6 gives the HR-TEM image of the TiO₂ particle having a 30-cycle amorphous overcoating. The overcoating in Fig. 6 presents a rather uniform thickness of 2.93 \pm 0.21 nm (0.98 Å per cycle). However, when considering several microscopy images from separate TiO2 particles, the average overcoating thickness was 2.15 ± 0.29 nm (0.72 Å per cycle). In literature, the growth-per-cycle of 0.98 Å per cycle for the TMA/ H₂O process has been reported for lateral high-aspect-ratio (LHAR) test structures at a deposition temperature 300 °C.²⁰ Similar values have been reported by Ott et al.36 and Puurunen,16 with GPC decreasing linearly with deposition temperature from 1.2 Å per cycle (180 °C) to 0.9 Å per cycle (300 °C). The deposition temperature in our experiment was 150 °C; thus, a lower GPC was achieved compared to the silicabased test structures.

To present overcoating thickness variation, Fig. 7 gives the thickness measurement results as a histogram with a Gaussian distribution. Interestingly, a rather wide distribution of overcoating thicknesses were measured. The 30c sample deviation was clearly increased compared to the 20c sample. The overcoating distribution for the 30c sample was ranging from 1.25 to 3.75 nm. Table 3 summarises the measured thicknesses for 20 and 30 cycle ALD overcoating with experimental error and sample size. The same table also presents the model prediction thickness values for corresponding overcoated catalysts.

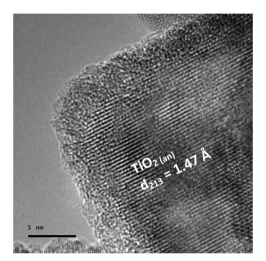


Fig. 6 $\,$ TiO $_2$ support particle TEM image and amorphous 30 cycle ALD overcoating with measured overcoating thickness 2.93 \pm 0.21 nm.

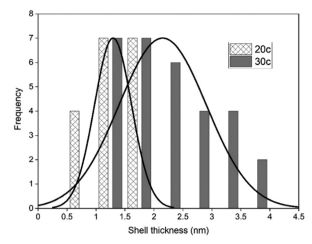


Fig. 7 Histogram of TEM image analysis for overcoating thickness. Frequency of individual measurements are counted in 0.5 nm intervals.

According to Lu et al.,37 the TMA/H2O process on Pd nanoparticles, 4 different growth regions exist depending on the deposition cycles. With deposition cycles ≥ 30 , the growthper-cycle (GPC) is constant. The other three growth regions are the first deposition cycle, 2-12 cycles and 13-30 cycles. The first cycle yielded 1.6 times the mass gain with compared to the constant GPC region; while the second cycle mass gain was only 0.8 of the constant GPC region. After the second cycle, GPC increased stepwise again to 1.6 at cycle 12, and after that, stabilised slowly to constant GPC at cycle 30. Our findings in Table 3 are in contradiction with these results, as 20 deposition

cycles resulted in a slightly lower GPC compared to 30 cycles. However, when considering the experimental error, there is no significant difference between the GPCs. Therefore, making conclusions on the different growth regions was not possible with the given results. To determine overcoating thickness with the diffusion-reaction model, an average of the measured GPC's was used (GPC = 0.681). If the presented model would be used to estimate overcoating thicknesses for higher amount of deposition cycles, the 30c sample GPC should be used in the calculation.

In addition to the prepared diffusion-reaction model and characterization experimental work, the overcoated samples were studied in the Fischer-Tropsch reaction. These results are reported in the supporting information, where 20 and 30 cycle ALD overcoated samples present a decreased rate of deactivation. Although the deactivation rate could be decreased (see Fig. S1, ESI†), this benefit comes with a price of promoted methanation activity and decreasing chain propagation α-value (see Table S2 and Fig. S2, ESI†). The chain propagation α -value was decreasing linearly with respect to overcoating thickness from 0.917 (non-overcoated catalyst) to 0.908 (30 deposition cycles).

3.3 Diffusion-reaction model parameter sensitivity

Parameter sensitivity analysis was used to determine each variable effect on the achieved penetration depth. In the presented diffusion-reaction model, there are two sets of parameters: catalyst structural (Table 1) and ALD precursor parameters (Table 2). From these parameters, the most significant effects

Table 3 Measured (HR-TEM) and predicted overcoating thickness. Diffusion-reaction model uses an average GPC (0.685 Å per cycle) from the TEM measurements to give overcoating thicknesses

ALD cycles	HR-TEM results			Growth-per-cycle	Diffusion-reaction model
	Overcoating thickness (nm)	Experimental error $\pm 2\sigma$	Sample size (n)	(Å per cycle)	Overcoating thickness (nm)
10	_	_	_	_	0.68
20	1.29	0.16	18	0.65	1.36
30	2.15	0.29	30	0.72	2.04

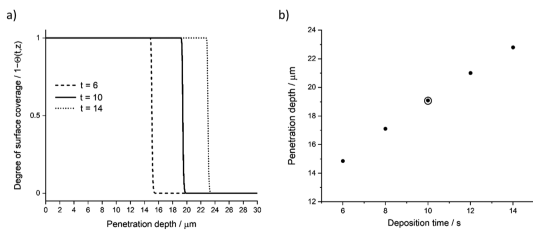


Fig. 8 Parameter sensitivity analysis, (a) degree of surface coverage profile variation with three exposure time (t) values, (b) achieved penetration depth (deepest penetration depth at $\Theta = 1$) with varying deposition time (t). Used model parameter marked with surrounding circle.

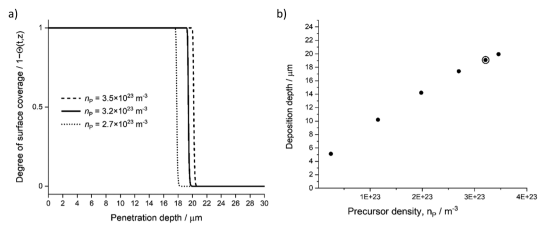


Fig. 9 Parameter sensitivity analysis, (a) degree of surface coverage profile variation with three n_P values, (b) achieved penetration depth (deepest penetration depth at $\Theta = 1$) with varying n_P . Used model parameter marked with surrounding circle. Corresponding TMA partial pressures were 100 Pa (penetration depth 5 µm), 460, 800, 1100, 1300, and 1400 Pa.

were related to catalyst porosity (ε) , catalyst tortuosity (τ) , deposition time (t), precursor density outside the porous material (n_P) , maximum density of particles adhering to the surface $(\sigma_{\rm P})$ and deposition temperature $(T_{\rm P})$.

3.3.1 Sensitivity related to deposition process and precursor parameters. Fig. 8 presents achieved penetration depth for varying exposure time (t, seconds). In the fig. a, the slope at the adsorption front stays constant, and the exposure time mainly affects the penetration depth. This is as expected when β_0 , n_P and σ_P remained constant. Although deposition time is an easy parameter to modify, the full ALD process duration will increase rapidly (see Fig. 8b) if no other parameters are changed to achieve deeper penetration into the catalyst particle.

In addition to the deposition cycle time (single half-reaction time), precursor density outside the porous material (n_P) influences the penetration depth. Fig. 9 presents the effect of varying precursor density (precursor pressure) introduced to the ALD chamber. With low precursor partial pressure, the precursor

molecules are reacting with a given sticking probability (β_0) on the available sites (σ_P) and depleting faster compared to higher partial pressures.

The deposition temperature has an effect on both penetration depth and achieved growth-per-cycle (GPC) in the TMA/ H₂O ALD process. Deeper penetration can be expected in an increased deposition temperature due to enhanced diffusion $(T_P \text{ at the eqn (6)})$ and a decreased surface OH group availability. 28,38 Having less OH groups for TMA precursor adsorption, the precursor can travel deeper into the catalyst structure (see Fig. 10). In addition to the penetration depth, the achieved layer thickness will change due to OH group availability. Fig. 11 presents thickness profiles given by eqn (11):

$$d_{\text{Al},O_2} = [1 - \Theta(t, z)] \times \text{GPC} \times \text{ALP cycles}$$
 (11)

where the deposition temperature has an effect on both the degree of surface coverage and the GPC. Increasing the

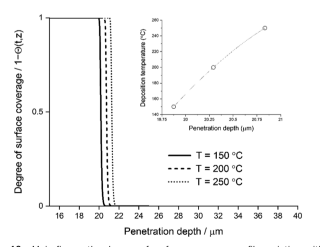


Fig. 10 Main figure, the degree of surface coverage profile variation with deposition temperature 150, 200 and 250 °C. Merged figure, the achieved penetration depth with varying deposition temperature.

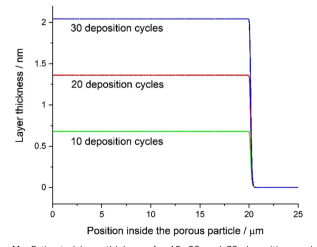


Fig. 11 Estimated layer thickness for 10, 20 and 30 deposition cycles. Layer thickness determined with eqn (11) having a growth-per-cycle (GPC) of 0.0681 nm per cycle from the TEM microscopy studies.

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deposition temperature will decrease the achieved layer thickness through decreased GPC.

Maximum density of particles adhering to the surface (σ_P) is dictated by the availability of reactive sites on the catalyst surface and precursor dimensions (see Fig. 12). Decreasing precursor size will allow for an increased surface saturation and less steric hindrance from neighbouring molecules. Altering the precursor structure would affect the σ_P through a different ligand arrangement and the bonding configuration.8 Similar to our work, Keuter et al.23 present a study with a tetrakis(ethylmethylamino)zirconium (TEMAZ) precursor on ZrO₂ support, where the small number of surface adsorption sites and relatively large TEMAZ molecular dimensions results in film growth restrictions due to steric hindrance. In their study, the porous material surface has open reactive sites; however, most of these sites have been blocked by alreadyadsorbed precursor molecules. With a TMA precursor, the molecular dimensions are smaller, and less restrictions are present for adsorption reactions.

3.3.2 Sensitivity related to catalyst structural parameters.

Catalyst porosity (ε) and tortuosity (τ) are present in the eqn (6). Dividing porosity per tortuosity, gives a simplified average of the catalyst structure. Typical high-range tortuosity values range between 6-10,39,40 therefore, the catalyst support used in our experiments had remarkably high tortuosity value (τ = 32.1). Fig. 13 presents the effect of varying tortuosity value to achieved penetration depth. In addition to catalyst tortuosity, porosity influences the achieved penetration depth. Fig. 14 presents the effect of varying values of porosity to the achieved penetration depth.

Fig. 15 presents the effect of porosity and tortuosity on the effective diffusion coefficient (D_{ei}) . With high tortuosity values, the porosity effect on the resulting D_{ei} decreases. Although D_{ei} is not greatly affected at the high tortuosity values, the slight change in the $D_{\rm ei}$ (4.56 imes 10⁻⁸ cm² s⁻¹ and 9.93 imes $10^{-8} \text{ cm}^2 \text{ s}^{-1}$ at $\varepsilon = 0.2$ and $\varepsilon = 0.5$ respectively with fixed τ = 32.1) has a rather high impact on the achieved penetration depth in Fig. 14.

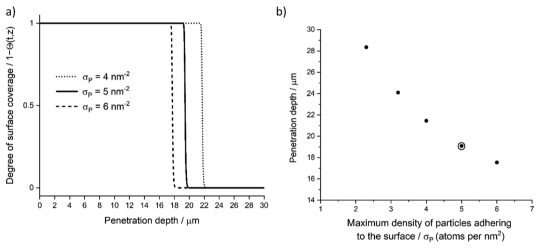


Fig. 12 Parameter sensitivity analysis, (a) degree of surface coverage profile variation with three σ_P values with 4, 5, and 6 atoms per nm², (b) achieved penetration depth (deepest penetration depth at $\Theta = 1$) with varying σ_P . Used model parameter marked with surrounding circle.

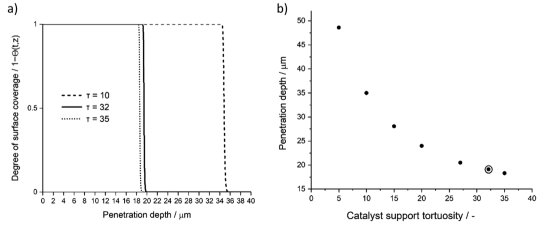


Fig. 13 Parameter sensitivity analysis, (a) degree of surface coverage profile variation with three catalyst tortuosity (τ) values, (b) achieved penetration depth (deepest penetration depth at $\Theta = 1$) with varying catalyst tortuosities (τ). Used model parameter marked with surrounding circle.

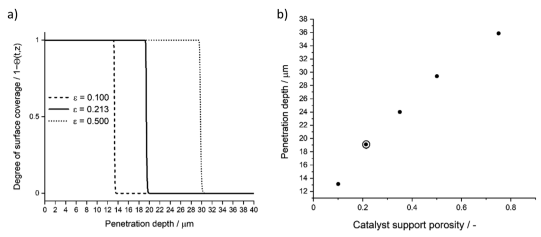


Fig. 14 Parameter sensitivity analysis, (a) degree of surface coverage profile variation with three catalyst porosity (ε) values, (b) achieved penetration depth (deepest penetration depth at $\Theta = 1$) with varying catalyst porosities (ϵ). Used model parameter marked with surrounding circle.

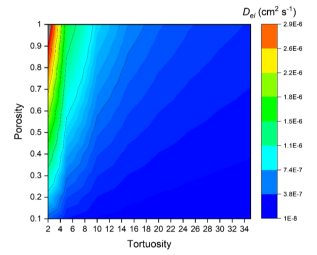


Fig. 15 The effect of porosity and tortuosity on the Knudsen diffusion coefficient (D_{ki}). Knudsen diffusion determined with fixed values for the mean pore radius (r), temperature (T_P) , and precursor mass (m_P) .

Catalyst characterisation

The catalyst nitrogen sorption measurement results are presented in Table 4. BET surface area for non-overcoated Catalyst A is 38.3 m 2 g_{cat} $^{-1}$, whereas ALD overcoated catalyst samples present the surface area between 41.3-41.6 m² g_{cat}⁻¹. The increased surface area is related to overcoating temperature

treatment, resulting in porous Al₂O₃ structure. Pore volumes remain closely unchanged with ALD overcoated samples, which could be expected, as overcoating thickness was between 0.68-2.04 nm; the resulting volume increase cannot be measured with the selected analysis method. However, the pore size measurement was able to give comparison data between catalyst samples. Comparing to the non-overcoated Catalyst A, the pore size was decreasing between 2.6-3.1 nm with overcoated samples. By assuming a cylindrical pore, the pore size decreasing was due to overcoating formation on the inner surface of the cylindrical hole. Therefore, dividing the pore size decrease by two would estimate layer thicknesses for overcoated samples, having a similar range as TEM and modelling results presented in Table 3.

The diffusion-reaction model utilised the Knudsen diffusion coefficient (D_{ki}) to estimate precursor transport inside the catalyst particle. Pore size distribution analysis was relevant to address whether or not Knudsen diffusion coefficient should be position dependent. If the overcoating layer-by-layer growth has a significant effect on the resulting pore size, the Knudsen diffusion coefficient should take this growth into account.²³ In order to determine which form of D_{ki} is suitable for our catalyst selection and target overcoating thicknesses, the pore size distribution from the BJH adsorption measurement presented useful information, with an average pore size ranging between 24.6-27.7 nm. From these results, it was concluded that the targeted maximum deposition thickness (2-3 nm) had no

Table 4 Nitrogen adsorption/desorption measurement results for the BET surface area, pore volume and pore size. Overcoated samples measured after temperature treatment

Catalyst	BET surface area (m ² g ⁻¹)	Pore volume (mL g^{-1})	Pore size (nm)
TiO ₂ support (Saint-Gobain Norpro, ACCU [®] SPHERE ST32244)	52	0.37	28.9
Catalyst A	38	0.27	27.7
Catalyst A + 10c ALD	42	0.26	25.1
Catalyst A + 20c ALD	41	0.25	24.6
Catalyst A + 30c ALD	42	0.26	24.8

Experimental error $(\pm 2\sigma)$ for surface area was ± 1 m² g⁻¹, pore volume ± 0.01 mL g⁻¹ and pore size ± 0.1 nm.

significant effect on the modelled results through layer-by-layer overcoating formation. Although a fraction of small pores (<6 nm) was present, the filling of these pores and coverage of cobalt particles < 6 nm had no significant effect on the Fischer-Tropsch reaction. In the FT reaction, the Co particle size should exceed 10 nm to achieve high turnover frequency. 41,42 For these reasons, the Knudsen diffusion coefficient was calculated with eqn (6), not considering layer-by-layer growth.

4 Conclusions

The Co-Pt/TiO2 Fischer-Tropsch porous catalyst was overcoated with an ALD (TMA/H2O) process. A diffusion-reaction differential equation model was prepared to address overcoating thickness and penetration to the porous catalyst sample. The model results were compared against SEM and TEM microscopy measurements, indicating wide distribution of penetration 0-18 μm, with an average penetration depth of 9.6 μm (corresponding to $\sim 5-10\%$ depth from throughout particle penetration). This variation is significantly different to ideal test surfaces and non-porous materials, where ALD overcoating can produce highly conformal and uniform inorganic layers. In addition to penetration depth distribution, the overcoating thickness had an intrinsic variation within the catalyst particle from 1.29 \pm 0.16 nm and 2.15 \pm 0.29 nm for 20- and 30-cycle ALD overcoated samples, respectively. The presented model was also used to estimate these thicknesses, and the results were corresponding well with 1.36 nm and 2.04 nm for 20- and 30-cycle ALD overcoated samples, respectively. The presented model can be applied with little effort to other catalyst support structures and different precursor compounds. The diffusionreaction model can be utilised to plan the ALD process on given catalyst structures.

Symbols and abbreviations

A_{O}	BET surface area (m ² g _{cat} ⁻¹)
β_0	Sticking coefficient of TMA
$D_{ m ki}$	Knudsen diffusion coefficient (m ² s ⁻¹)
3	Porosity
GPC	Growth-per-cycle
$k_{ m B}$	Boltzmann constant (m ² kg s ⁻² K ⁻¹)
$m_{ m P}$	Precursor mass (kg)
$n_{ m P}$	Precursor density outside the porous material (m ⁻³)
r	Mean pore radius (m)
\bar{s}	Ratio of surface area $(A_{\rm O})$ per pore volume $(V_{\rm P})$
s_0	Average surface area of adsorption sites (m ²)
$\sigma_{ m P}$	Maximum density of particles adhering to the
	surface (m^{-2})
τ	Tortuosity
T	Temperature (K)
$T_{ m P}$	Precursor temperature (K)
Θ	Surface coverage $(0 \le \Theta \le 1)$

trimethylaluminium, Al(CH₃)₃ Pore volume (m³ g_{cat}⁻¹)

Author contributions

Conceptualization (N. H., J. L. & M. R.), data curation (N. H., J. L., S. S. & Y. G.), formal analysis (N. H., J. L., L. K., J. Y., S. S. & Y. G.), investigation (N. H., S. S. & Y. G.), methodology (N. H. & J. L.), writing - original draft (N. H.), writing - review & editing (all authors).

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Conflicts of interest

The authors declare no conflict of interest.

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