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Ex situ upgrading of pyrolysis vapors over PtTiO₂: extraction of apparent kinetics via hierarchical transport modeling†

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Chemical reaction kinetics enable predictive scaling studies and process sensitivity analyses that can substantially accelerate commercial deployment of new catalytic transformation technologies. The absence of suitable kinetic parameters for catalytic fast pyrolysis (CFP) of biomass feedstocks has precluded such de-risking simulation activities. In this work we consider *ex situ* CFP using a Pt/TiO₂ catalyst in a packed bed vapor phase upgrading reactor (VPU) with co-fed H₂. We develop a multiscale simulation framework to de-couple apparent kinetics from both intraparticle and reactor-scale transport phenomena. The transport model is integrated with a kinetic scheme that predicts (1) lumped yields of product partially deoxygenated compounds, hydrocarbons, light gases, water, and coke, as well as (2) active site concentration and deactivation of the catalyst. We employ recent advancements in mathematical treatments of cascading reaction systems in the context of an axial-dispersion packed bed reactor model to achieve a rapidly-solving simulation framework that is amenable to iterative regression for kinetic parameter extraction. Results demonstrate accurate predictions of CFP yields within 5% for a variety of conditions, including different reaction times, Pt loadings, and variations in feedstock attributes.

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Introduction

Catalytic fast pyrolysis (CFP) in an *ex situ* configuration is a technology wherein lignocellulosic feedstocks like wood undergo fast pyrolysis and then the vapors are deoxygenated/upgraded to an oil product over a heterogeneous catalyst in a second reactor. Compared to non-catalytic fast pyrolysis, CFP produces a stabilized oil with reduced oxygen content. The CFP oil could then be hydroprocessed to a finished fuel blendstock,^{1,2} co-processed with refinery intermediates in an oil refinery,³ utilized as a heating fuel, or further transformed into bio-based chemicals or materials.^{4,5} Co-fed H₂ can be also introduced to promote hydrodeoxygenation (HDO) and direct deoxygenation (DDO)⁶ over a bifunctional metal-acid catalyst such as Pt/TiO₂ (ref. 1) or Mo/Al₂O₃.⁷ HDO and DDO remove oxygen as water and can improve carbon yields in the CFP oil product by reducing light gas formation and coke deposition on the catalyst.^{1,4,5,7–12} Recent microreactor⁶ and bench-scale work^{1,13} have shown particularly promising CFP

oil yields where pine was pyrolyzed in a fluidized bed reactor and the vapors were upgraded over a packed bed of a bifunctional metal-acid Pt/TiO₂ catalyst with co-fed H₂ at near atmospheric pressure. The PtTiO₂ catalyst is considered a class 1 bifunctional catalyst with a noble metal (Pt) dispersed on an acidic reducible/amphoteric oxide (*e.g.* TiO₂) that has demonstrated activity for direct deoxygenation of lignin-derived aromatic oxygenates like phenol in the literature.^{14–16} The high activity of this catalyst has been attributed to synergistic effects between the metallic phase and the reducible/amphoteric TiO₂ support, which promote C–O bond scission at or near the interface.⁶ Given the encouraging results of these bench-scale experiments, there is considerable interest in further technological development and process scale-up.¹³

As with any catalytic transformation process, an understanding of the chemical and physical mechanisms that govern the process performance is critical to developing predictive models that will guide and de-risk scale-up. Reasonably advanced predictive models exist for the fast pyrolysis step^{17–19} but robust predictive models do not exist for the upgrading step using the bifunctional metal-acid Pt/TiO₂ catalyst. Developing a predictive model of the upgrading step in *ex situ* CFP using a Pt/TiO₂ catalyst is the focus of this work. Obtaining kinetic rate parameters is essential to

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predictive model development. Apparent rate constants in this work are not collated with intraparticle diffusion or bulk transport effects and thus allow for parametric studies across different particle sizes, and support porosities, as well as reactor-scale dimensions and operating conditions. Previously, our group employed a protocol based on back-calculation of kinetic rate constants by iterative regression of a multi-scale model to experimental data.²⁰ Here we extend this methodology to obtain apparent kinetic rates for the *ex situ* CFP Pt/TiO₂ process. These are apparent rate constants rather than intrinsic because they combine adsorption, reaction, and desorption at an active site, an exercise approached with microkinetic modelling.²¹

Modeling packed bed reactors was one of the first tasks taken on by chemical reaction engineers, and differential equation models have been developed that can quickly describe 1-D, idealized plug-flow reactor systems.^{22–24} Several groups have used 2-D and 3-D computational fluid dynamics (CFD) modeling techniques to model realistic packings and geometries in packed-bed catalytic reactors, as reviewed by Jurtz *et al.*²⁵ It is evident that these more physically realistic techniques are critical for accurately representing the behaviour for reactors with small tube-to-particle diameter ratios, irregularly shaped particles, non-ideal packing, *etc.* While these models can indeed add a great deal of accuracy to simulations of packed bed reactors, their use is more computationally demanding and should be balanced with less computationally expensive alternatives such as plug-flow “axial dispersion” models.

In porous catalyst supports with dispersed active sites, surface reaction equations do not accurately describe the effective rate of reaction for reactants and products diffusing into and out of these porous particles. Intraparticle diffusion–reaction behavior has a major impact on the observed reaction rate. Jurtz *et al.*²⁵ surveyed a variety of tools for modeling intraparticle physics, but the fastest by far is the Thiele effectiveness factor, which is an analytical solution for spherical reaction–diffusion with homogeneous porosity and active site dispersion.²⁶ One of the limitations of the classic Thiele effectiveness factor is that it is only valid for single-step reactions. However, HDO processes using bi-functional metal–acid catalysts involve many reactions, occurring in parallel and in series, and are not accurately represented by single-step reaction approximations. In this work the upgrading process is referred to as the HDO step for simplicity because hydrogen is fed and we are deoxygenating the vapors, but it does not mean that the only reaction pathway occurring is formally hydrodeoxygenation. To expand the range of validity for the effectiveness factor approach, our group recently developed an analytical approach to deal with reactions in series called the multistep effectiveness vector (MEV).²⁷ In this present work, we aim to use an axial dispersion model with an MEV subgrid catalyst model to extract lumped apparent reaction rate constants and explore different experimental conditions that may be used to optimize the HDO step in *ex situ* catalytic fast pyrolysis using a Pt/TiO₂ catalyst (Fig. 1).

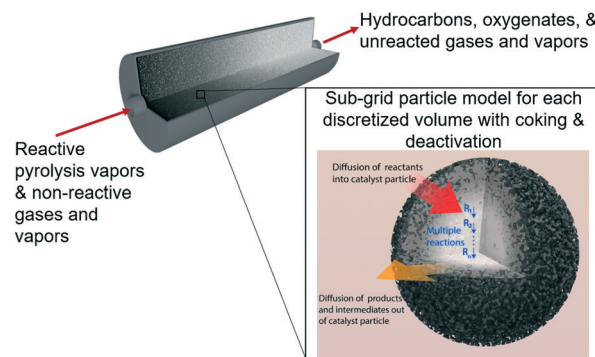


Fig. 1 Illustration of the packed bed model with discretized volumes for tracking sites and coke formation and the subgrid multistep effectiveness vector (MEV) model describing reaction–diffusion within catalyst particles.†

Experimental methods

Ex situ catalytic fast pyrolysis experiments

The experiments were performed in a bench scale *ex situ* catalytic fast pyrolysis reactor comprised of a 2" inner diameter fluidized bed fast pyrolysis reactor fed with mixtures of clean pine and forest residue (milled to 2 mm) coupled to a packed bed reactor loaded with Pt/TiO₂ catalyst. A detailed description of this system was provided in a previous report¹ and illustrated in Fig. 2 and feedstock characterization can be found in Klinger *et al.*⁴⁴ Briefly, the pyrolysis reactor was supplied with 85% H₂/15% N₂ at near atmospheric pressure. The fast pyrolysis vapors together with the gases leave the fluidized bed reactor and flow through a cyclone and then pleated stainless steel hot gas filter with nominal 2 μm openings. These vapors were piped through heat-traced lines into the packed bed VPU reactor, 3.18 cm i.d. by 14 cm long, containing ~100 g Pt/TiO₂ catalyst. The collected liquids spontaneously separate into an oil and aqueous phase and the top aqueous phase is removed by pipetting.

The composition of the vapor leaving the VPU was monitored with a mass spectrometer (residual gas analyzer, RGA). After the RGA, the vapors were condensed, and the liquids collected. The non-condensable gases were analyzed with a micro GC for light gases (LG, consisting of CO, CO₂, and C_{1–4} hydrocarbons) and an online GC-MS-FID for gases and vapors not detected by the micro GC (light condensables, consisting of light oxygenates and some C₅₊ hydrocarbons). The collected liquids were separated into an oil and aqueous phase, and the oil was analyzed for composition by a GC × GC-TOFMS. After each CFP experiment, the catalyst is regenerated by oxidation of coke and subsequent catalyst reduction. The ESI† contains a more detailed description of the reactor dimensions, flowrates, and analytical procedures.

† Fig. 1 is reprinted (adapted) from Chemical Engineering 380, Aaron M. Lattanzi, M. Brennan Pecha, Vivek S. Bharadwaj, Peter N. Ciesielski, Beyond the effectiveness factor: Multi-step reactions with intraparticle diffusion limitations, 1, Copyright (2020), with permission from Elsevier.



The experimental data used for fitting the rate constants was based on four replicate experimental runs, each with a cumulative biomass to catalyst mass ratio (B:C) of 12, a 7.1 h run time, with 0.5 wt% Pt/TiO₂ catalyst, a weight hourly space velocity of 1.5 g biomass per g catalyst h⁻¹, and a 50%/50% clean pine (CP)/forest residue (FR) wood blend as the biomass feed. The pyrolysis temperature was 500 °C and the average internal temperature of the VPU was 450 °C. The biomass ultimate analysis and further analytical details are included in the ESI.† Yields from the pyrolysis + HDO reactor were compared to results from pyrolysis alone. Further experimental data collected for the same catalytic conditions with B:C of 6 and 21 as well as a data set for a 1 wt% Pt on the same TiO₂ support and 100% clean pine as feed at B:C 3 were used to verify the model. The equations used to calculate lumped product yields across the VPU are detailed in the ESI.†

Catalyst preparation and characterization

The catalyst had been prepared by a strong electrostatic adsorption (SEA) method on a spherical TiO₂ support with an average particle diameter of 0.5 mm provided by Johnson Matthey. A detailed description of the catalyst preparation method can be found in Griffin *et al.*¹ In short, Pt was added to TiO₂ particles at pH 11.5 using 1.98 g Pt(NH₃)₄(NO₃)₃ and

soaked for >24 h in aqueous solution then dried >24 h at 60 °C. This catalyst was reduced in 5% H₂/N₂ at 450 °C and passivated under 1% O₂/N₂ at room temperature before loading into the packed bed. Pt particles had an average diameter of 2.7 nm. BET surface area was 44 m² g⁻¹. For the 0.5% Pt catalyst, Pt CO chemisorption suggested Pt dispersion of 33%. TPD acid site density was 220 μmol g⁻¹ and acid:metal site ratio is 4.9.

The catalyst was characterized in part with imaging by scanning electron microscopy (SEM) and transmission electron microscopy (TEM). For scanning electron microscopy, the whole catalyst particles were mounted on aluminum stubs with conductive carbon adhesive. Samples were imaged without coating using a FEI Quanta 400 FEG using an accelerating voltage of 25 kV. For transmission electron microscopy and tomography, catalyst particles were gently crushed and the powder was suspended in ethanol at ~0.5% wt/vol. A volume of 5 μl of the suspension was placed on carbon-coated copper TEM grids with a grid sized of 200 mesh (SPI Supplies, West Chester, PA). Samples were allowed to air dry prior to imaging. A FEI Tecnai G2 20 Twin 200 kV LaB6 TEM (FEI, Hillsboro, OR) instrument was used at an accelerating voltage of 200 kV; images were collected with a Gatan UltraScan 1000 camera (Gatan, Pleasanton, CA). Tomography was performed by first obtaining dual-axis ±60° tilt series of the region of interest at a pixel size of ~0.5 nm. Single axis tomograms were constructed from the tilt series using the *R*-weighted back projection algorithm and then combined to yield the final reconstruction using the IMOD software package.²⁸ Tomographic slices were visualized with IMOD and the volume reconstruction was visualized with PyMOL.²⁹ Particle porosity of 0.592 and an average pore diameter of 28 nm were measured by N₂ adsorption with BJH calculations.³⁰

The multiscale nature of this material is shown through light microscopy which shows the spherical shape (Fig. 3a) and scanning electron microscopy which shows the porous support structure formed by the agglomeration of TiO₂ nanoparticles (Fig. 3b). Transmission electron tomography was performed to investigate the three-dimensional morphology of intraparticle porosity to inform parameterization of the intraparticle transport model, as

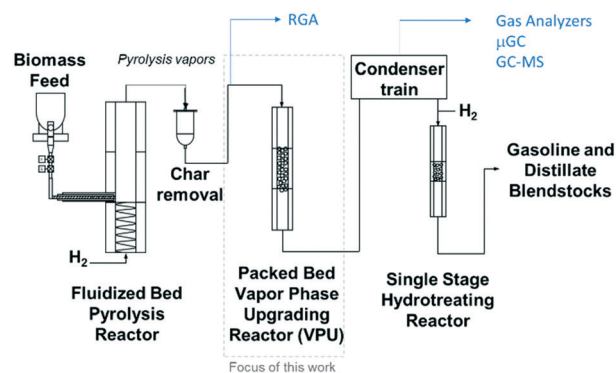


Fig. 2 Schematic of the bench scale *ex situ* catalytic fast pyrolysis system utilized in this work with a packed bed (fixed bed) of catalyst.

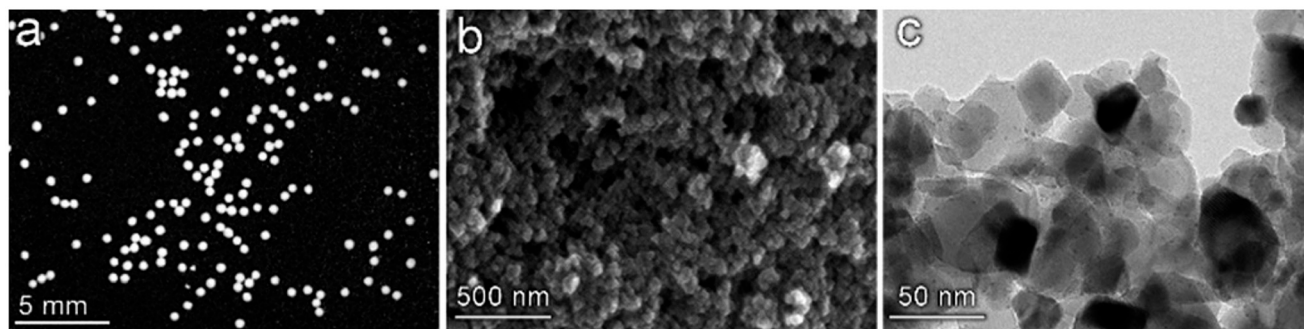


Fig. 3 Multiscale imaging of the Pt/TiO₂ catalyst particles. (a) Light microscopy of catalyst particles showing the spherical bulk geometry with narrow size distribution. (b) Scanning electron microscopy (SEM) of the particle surface reveals a porous support structure formed by the agglomeration of TiO₂ nanoparticles. (c) Transmission electron microscopy shows the presence of ~5 nm Pt particles visualized as dark spots on the surface of the larger TiO₂ support structure.



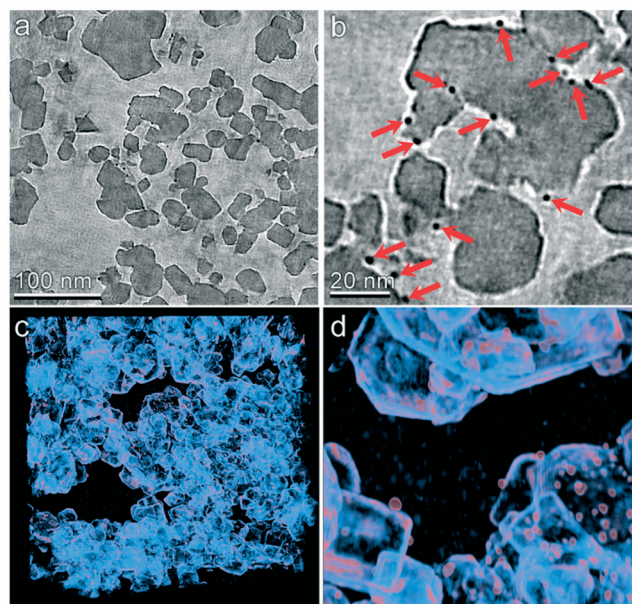


Fig. 4 TEM tomography of the TiO_2 catalyst particle mesostructure. (a and b) Slices through the tomographic volume are shown at two different magnifications. Pt particles are clearly identified by their higher electron density (indicated by red arrows in panel b). (c and d) 3D visualizations of the reconstructed volume are shown at two different magnifications. Density primarily corresponding to the TiO_2 support is shown as light blue and that roughly corresponding to the Pt particles is shown as orange.

shown in Fig. 3c. This technique involves acquiring a series of TEM images at different angles with respect to the electron beam which enables the reconstruction of the 3D sample volume *via* computational image processing. Slices through the 3D volume obtained using this method are shown at two different magnifications in Fig. 4a and b. Due to the large number of images (~ 160) that are averaged to produce the 3D density field, the contrast between the TiO_2 support and Pt nanoparticles is enhanced with respect to that observed in 2D TEM images. A 3D visualization of the reconstructed volume is shown in Fig. 4. This analysis reveals that Pt

clusters with diameters of roughly 5 nm are well-dispersed on the surface of the support material.

The tomographic reconstruction facilitates direct quantification of mesoporous dimensions including void space and distribution of pore dimensions. The void volume was determined by first calculating the average and standard deviation of the tomographic density of a clearly indefinable void region within the reconstructed volume. The void fraction of the entire volume was then determined by the number of voxels within the volume that displayed a density greater than two standard deviations above the mean density of the void volume. The entire reconstruction was divided into 64 sub-volumes and this calculation was performed within each to determine variability of the void space. By this method the average void fraction was determined to be 57% with a standard deviation of 6.8%. The pore size distribution was calculated using a combination of automated computational image transformations. The average pore radius was computed to be 7.82 nm with a standard deviation of 7.84 nm based on microscopy. However 28 nm is used as the pore diameter in the model because the adsorption experiments are more representative of the entire catalyst particle. The SEM and TEM analysis described in further detail in the ESI.†

Experimental results

Catalytic fast pyrolysis yields

A series of catalytic fast pyrolysis experiments were performed in the *ex situ* pyrolyzer followed by the hydrodeoxygenation (HDO) vapor phase upgrading (VPU) packed bed of porous Pt/ TiO_2 catalysts. Table 1 shows the mass yields on a dry biomass basis for the datasets used in this work. Table 2 shows the GC \times GC composition summary of the collected liquids with lumped species assignments that are used in this model. For the base case experiments used for kinetic parameter extraction (0.5% Pt, B:C 12), the pyrolysis reactor converted 89% of 50/50 CP/FR wood into volatile and gaseous species that pass into the HDO reactor (Table 1), with the remainder being

Table 1 Mass yields from pyrolysis and CFP, wt% dry biomass basis, normalized to 100% (mass closure >98%)

Description	Pyrolysis only	Pyrolysis only	Base case	Test case	Test case	Test case
B:C, kg/kg	—	—	0.5% Pt	1% Pt	0.5% Pt	0.5% Pt
Feed	CP/FR	CP	CP/FR	CP	CP/FR	CP/FR
# of runs	3	3	4	3	1	1
Organics in oil	58.5 \pm 0.5	56.3 \pm 0.1	24.7 \pm 0.1	20.9 \pm 0.1	23	26.6
Organics in aqueous phase	0.0 \pm 0.0	0.0 \pm 0.0	3.8 \pm 0.3	2.3 \pm 0.0	3.5	4.0
Water	13.3 \pm 0.2	17.5 \pm 0.4	21.7 \pm 0.0	22.9 \pm 0.3	21.9	20.1
Light condensables	2.3 \pm 0.1	0.5 \pm 0.2	10.0 \pm 0.2	8.0 \pm 0.1	8.6	9.9
Light gases	15.2 \pm 0.3	13.9 \pm 0.8	28.2 \pm 0.5	31.6 \pm 0.3	29.2	27.4
CH_4	1.3 \pm 0.0	0.5 \pm 0.1	3.2 \pm 0.1	3.2 \pm 0.0	3.2	3.0
CO	7.5 \pm 0.2	6.1 \pm 0.4	14.5 \pm 0.6	15.7 \pm 0.2	15.0	14.6
CO ₂	5.6 \pm 0.1	6.0 \pm 0.3	7.6 \pm 0.1	8.1 \pm 0.1	8.0	7.3
C ₂ –C ₄	0.9 \pm 0.0	0.7 \pm 0.0	2.8 \pm 0.1	4.0 \pm 0.0	3.0	2.5
Char	10.7 \pm 0.0	11.8 \pm 0.6	10.3 \pm 0.0	11.8 \pm 0.3	12.3	10.9
Coke	—	—	1.4 \pm 0.0	2.5 \pm 0.1	1.8	1.1



Table 2 GC × GC oil composition summary, wt% wet oil

	Run	Pyrolysis only	0.5% Pt	1% Pt	0.5% Pt	0.5% Pt
	B : C, kg/kg	—	12	3	6	21
Description	Feed	CP	CP/FR	CP	CP/FR	CP/FR
HC	Aromatic hydrocarbons	0.02	1.3 ± 0.2	2.8 ± 0.8	1.6	0.8
HC	Alkanes + alkenes	0.02	0.5 ± 0.1	0.6 ± 0.3	1.0	1.3
OX	Cyclopentene/anones	0.99	8.0 ± 0.2	9.2 ± 0.5	7.8	8.3
PV	Other ketones/aldehydes	4.15	5.4 ± 0.1	4.2 ± 0.1	6.6	3.4
OX	Phenol	0.19	4.6 ± 0.2	7.0 ± 0.7	5.2	4.6
OX	Methylphenols	0.18	1.7 ± 0.1	2.9 ± 0.2	2.0	1.6
OX	Other phenols	0.36	6.1 ± 0.6	11.5 ± 2.2	7.4	6.2
OX	Furans	2.29	4.1 ± 0.1	1.9 ± 0.1	2.6	7.2
PV	Acids	1.42	0.9 ± 0.1	1.3 ± 0.3	1.3	2.6
PV	Methoxyphenols	0.92	0.6 ± 0.1	1.2 ± 0.4	0.4	1.7
PV	Sugars	0.57	0.0 ± 0.0	0.0 ± 0.0	0.0	0.2
PV	Other oxygenates	2.54	1.4 ± 0.2	1.9 ± 0.4	0.7	1.9
	Total, wt% in oil	13.66	34.7	44.4	36.5	39.8

char. However, not all of the vapors are expected to participate in the catalytic upgrading reactions in the subsequent VPU reactor. Here, light gases (LG), light condensables (LC), and water vapor from pyrolysis were considered to pass through the VPU without being majorly transformed, as the yields of these

families increase after the HDO. They may couple but appear to remain in the same category.

LC compounds consist of primarily acetaldehyde, acetone, misc. C₅₊, butanone, furan, and 2-methyl furan so additional LC is considered to be in the intermediate OX family. In

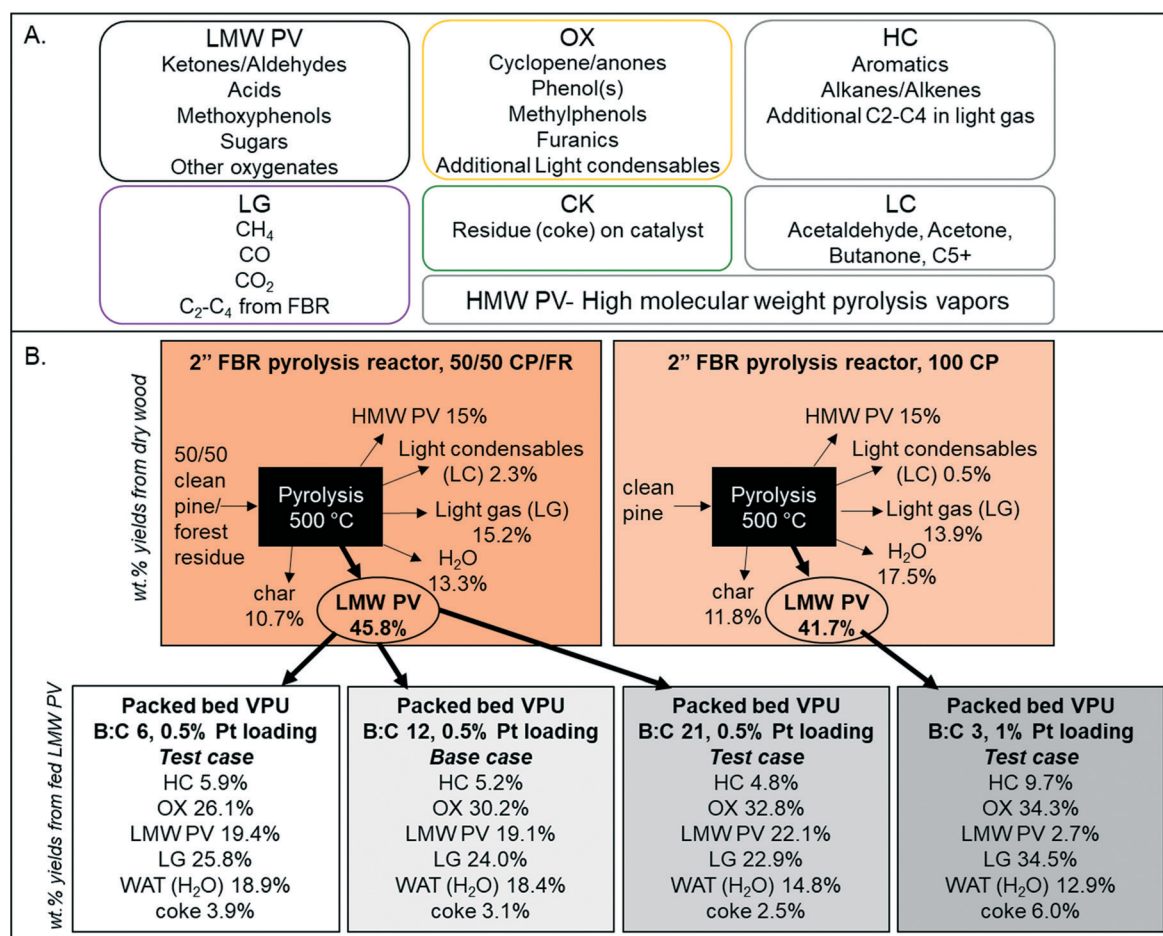


Fig. 5 (A) Lumped species definitions based on GC/MS quantification and (B) illustrative mass balances of the model development and three test cases with weight percent yields from dry wood for the pyrolysis reactor and weight percent yields from LMW PV for the vapor phase upgrading reactor.



addition, the fraction of CFP oil that was not identifiable by GC \times GC-TOFMS (Table 2), which consisted of high molecular weight pyrolysis vapor (HMW PV) that do not volatilize below the injector temperature of 300 °C and oven temperature of 255 °C. GC \times GC methods can be seen in the ESI.† The HMW PV component corresponded to 15% of the biomass. Previous research with similar fluidized beds and pine wood have shown that it is typical to have 15% high molecular weight “pyrolytic lignin”.^{31–33} For this work, HMW PV is considered generally unreactive, as there is a similar amount of unidentified high MW liquid in all catalytic experiments (~15%). With these assumptions, only 46 wt% of the biomass is considered “reactive pyrolysis vapors” and included in the kinetic HDO model. This consolidated mass balance is graphically illustrated for clarification in Fig. 5 for the four experimental conditions used in this work. Further oil characterization data can be found in the ESI.†

Real-time tracking of the vapor products leaving the reactor was accomplished through the RGA. However, due to the response factor variation and sensitivity, not all species were detected. Thus “ambassador compounds” were chosen to represent the chemical families of low molecular weight pyrolysis vapors (LMW PV), partially deoxygenated compounds (OX), and hydrocarbons (HC). For LMW PV, acetic acid/hydroxyacetaldehyde (m/z 60) was chosen; for OX, phenol (m/z 94), furans (m/z 68) and cyclopentenone/methyl furan (m/z 82); for HC, toluene (m/z 91). The lumping procedure is shown in Fig. 5. Data from 0.5% Pt B:C 12

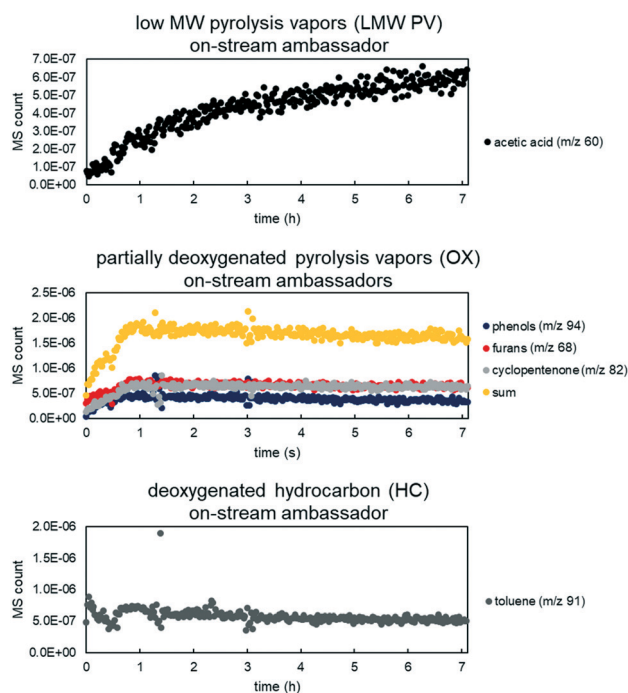


Fig. 6 MS counts vs. time for three experiments (overlaid) for ambassador compounds with high signal to noise ratio for LMW PV (top), OX (middle), and HC (bottom). Each data set was fit to a regression for model rate fitting. Data from 50/50 CP/FR, B:C 12, 0.5% Pt loading.

experiments were summed together, and for OX all tracked species were also summed together (Fig. 6). The component groups analyzed by GC \times GC were also divided into the same families (LMW PV, OX, HC), (Table 2), and the final yields for each family were calculated. The RGA data provides transient intensities representing these three primary vapor chemical families whose integrals are normalized to equal the yield of each lumped product. The light gases and light condensables were also monitored throughout the experimental runs; the additional light gases formed during VPU were tracked as a separate product family (LG); the additional condensables formed were divided into OX or HC depending on the compound. Thus the data was transformed to represent cumulative yield of products as a function of time based on total LMW PV fed during the experiment for the base case of biomass to catalyst (B:C) ratio of 12 with a 0.5 wt% Pt/TiO₂. In addition, the final yield of coke (CK) and additional water formed (WAT) was available. The final yields of each compound family based on the LMW PV from non-catalytic pyrolysis are shown in Fig. 5. The equations for determining the lumped yields are listed in the ESI.†

Model description

Packed bed model

The reaction scheme for this catalyst was developed based on prior experimental insights into the proposed mechanisms that this Pt/TiO₂ catalyst follows for conversion of pyrolysis vapors into partially oxygenated compounds (OX), hydrocarbons (HC), light gas (LG), water (WAT), and coke (CK). Pt/TiO₂ is a bifunctional metal-acid catalyst, and HDO is generally considered to progress in two stages: partial deoxygenation of pyrolysis vapors and nearly complete deoxygenation of pyrolysis vapors through a variety of mechanisms including decarbonylation, ketonization, aldol condensation, direct deoxygenation, and hydrogenation.¹ In

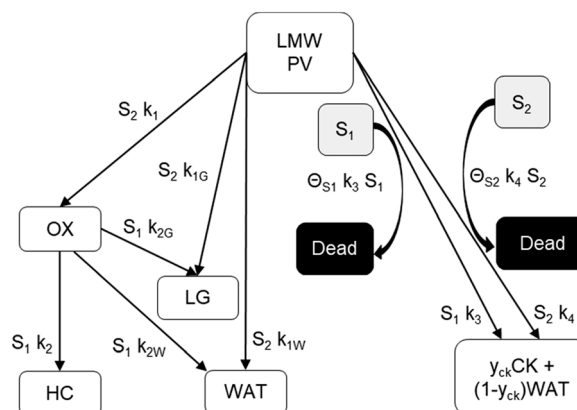


Fig. 7 Simplified reaction scheme for reactive pyrolysis vapors converting to partially deoxygenated compounds (OX), hydrocarbons (HC), light gas (LG), coke (CK), and water (WAT) over two different reactive catalyst sites (S₁ and S₂) that can deactivate to fully a dead site. y_{ck} is 0.67, representing a mass loss of 33% to remove oxygen in the form of H₂O from pyrolysis vapor to leave a nearly graphitic coke.



general, we could consider these as two sequential reaction families where $\text{LMW PV} \rightarrow \text{OX} + \text{LG} + \text{WAT} \rightarrow \text{HC} + \text{LG} + \text{WAT}$ where the first reaction is over catalytic active site S_2 and the second is over S_1 , as shown in Fig. 5. It could be that S_2 is an acid site and S_1 is a metal site, but the aim of this work was not to prove this hypothesis. There are also simultaneous, competitive coke formation and deactivation reactions for these two catalytic sites. The simplified mechanism used here is shown in Fig. 7. The following equations describe the rates for each species represented in this model. It should be noted that these reaction rates are attenuated by diffusion within the porous catalyst particles, as described later.

In the packed bed reactor, the vapor reactants and products experience advection, diffusion, and chemical reactions. The following equations describe change in concentration of the reacting species in the void fraction of the packed bed.

$$\frac{\partial \text{PV}_{\text{LMW}}}{\partial t} = -u \frac{\partial \text{PV}_{\text{LMW}}}{\partial x} + D_{\text{PV}} \frac{\partial^2 \text{PV}_{\text{LMW}}}{\partial x^2} - R_{\text{PV}_{\text{LMW}}, \text{eff}} (1 - \varepsilon_p) \quad (1)$$

$$\frac{\partial \text{OX}}{\partial t} = -u \frac{\partial \text{OX}}{\partial x} + D_{\text{OX}} \frac{\partial^2 \text{OX}}{\partial x^2} - R_{\text{OX}, \text{eff}} (1 - \varepsilon_p) \quad (2)$$

$$\frac{\partial \text{HC}}{\partial t} = -u \frac{\partial \text{HC}}{\partial x} + D_{\text{HC}} \frac{\partial^2 \text{HC}}{\partial x^2} - R_{\text{HC}, \text{eff}} (1 - \varepsilon_p) \quad (3)$$

$$\frac{\partial \text{LG}}{\partial t} = -u \frac{\partial \text{LG}}{\partial x} + D_{\text{LG}} \frac{\partial^2 \text{LG}}{\partial x^2} - R_{\text{LG}, \text{eff}} (1 - \varepsilon_p) \quad (4)$$

$$\frac{\partial \text{WAT}}{\partial t} = -u \frac{\partial \text{WAT}}{\partial x} + D_{\text{WAT}} \frac{\partial^2 \text{WAT}}{\partial x^2} - R_{\text{WAT}, \text{eff}} (1 - \varepsilon_p) \quad (5)$$

$$\frac{\partial S_1}{\partial t} = R_{S_1, \text{eff}} (1 - \varepsilon_p) \quad (6)$$

$$\frac{\partial S_2}{\partial t} = R_{S_2, \text{eff}} (1 - \varepsilon_p) \quad (7)$$

$$\frac{\partial \text{CK}}{\partial t} = R_{\text{CK}, \text{eff}} (1 - \varepsilon_p) \quad (8)$$

$$\text{PV}_{\text{LMW}} = \text{PV}_{\text{LMW}, 0}, \quad x = 0 \quad (9)$$

$$\text{HC} = \text{OX} = \text{LG} = \text{WAT} = 0, \quad x = 0 \quad (10)$$

$$\frac{d\text{PV}}{dt} = \frac{d\text{OX}}{dt} = \frac{d\text{HC}}{dt} = \frac{d\text{LG}}{dt} = \frac{d\text{WAT}}{dt} = 0, \quad x/L = 1 \quad (11)$$

where D_i is the effective Fick's diffusion coefficient, x is axial length of the reactor bed, ε_p is void fraction of the reactor bed, t is time, u is nominal gas velocity, eff indicates effective reaction rates attenuated by intraparticle diffusion, and $R_{i, \text{eff}}$ indicates reaction rate for the solid domain as calculated by the MEV approximation (*à la* a Thiele effective reaction rate) which is detailed thoroughly elsewhere.²⁷ Relevant parameters for the model were taken from the full reactor

description shown in the ESI† or calculated with basic mass balance equations and a summary of model-relevant parameters is shown in Table 3. These effective reaction rates are a function of each reactions respective local active site concentration, reactant concentration, and rate constant, as illustrated in Fig. 5. The eight individual apparent reaction rate equations are shown below:

$$R_1 = \text{PV}_{\text{LMW}} k_1 S_2 \quad (12)$$

$$R_{1G} = \text{PV}_{\text{LMW}} k_{1G} S_2 \quad (13)$$

$$R_{1W} = \text{PV}_{\text{LMW}} k_{1W} S_2 \quad (14)$$

$$R_2 = \text{OX} k_2 S_1 \quad (15)$$

$$R_{2G} = \text{OX} k_{2G} S_1 \quad (16)$$

$$R_{2W} = \text{OX} k_{2W} S_1 \quad (17)$$

$$R_3 = \text{PV}_{\text{LMW}} k_3 S_1 \quad (18)$$

$$R_4 = \text{PV}_{\text{LMW}} k_4 S_2 \quad (19)$$

As shown in Fig. 5, the relative mass yield of coke to water + coke, y_{ck} , is 0.67, representing a mass loss of 33% to remove

Table 3 Key parameters used in the packed bed reactor model

Parameter	Value	Description
L	0.14 m	Length of reactor bed
P_{tot}	1×10^5 Pa	Pressure in reactor, inlet
u_{inf}	0.947 m s^{-1}	Nominal gas velocity
U	2.167 m s^{-1}	Void fraction-corrected gas velocity
T_{in}	450 C	Temperature reactor
ρ_g	0.2247 kg m^{-3}	Density gas ([85% H_2 + 15% N_2] + PV + LG + H_2O), inlet
μ_{vap}	$1.97 \times 10^{-5} \text{ kg m}^{-1} \text{ s}^{-1}$	Estimated viscosity of gas + PV
AFR	$7.5 \times 10^{-4} \text{ m}^3 \text{ s}^{-1}$	Actual volumetric flowrate gas
mfPVin _{50/50}	0.458	Mass fraction LMW PV in for 50/50 CP/FR
mfPVin _{CP}	0.417	Mass fraction LMW PV in for CP
D_i	$4 \times 10^{-5} \text{ m}^2 \text{ s}^{-1}$	Bulk diffusion coefficient for all species
$D_{i, \text{eff}}$	$D_i \varepsilon_{\text{ps}}$	Effective bulk diffusion coefficient
ε_{ps}	0.437	Void fraction reactor bed
ε_{pp}	0.592	Void fraction catalyst particle
D_p	$0.5 \times 10^{-3} \text{ m}$	Diameter catalyst particle
D_{pore}	$2.8 \times 10^{-8} \text{ m}$	Pore diameter of catalyst particles
Re	12.4	Reynolds number with respect to particle
T	7	Tortuosity within catalyst particle
K_D	$48.5 D_{\text{pore}} \sqrt{T/MW}$	Knudsen diffusion coefficient calculation
Re	28	Reynolds number with respect to a single catalyst particle with inlet fluid properties
Sc	5	Schmidt number
Pe	141	Peclet number
Sh	152	Sherwood number for creeping flow around sphere ³⁹



oxygen in the form of H₂O from pyrolysis vapor to leave a nearly graphitic coke. The purpose of these ratios is to reduce the number of fitted parameters. Parameters Θ_{S_1} and Θ_{S_2} are fitted deactivation ratios that relate how quickly sites 1 and 2 deactivate relative to coke formation and are connected to rate constants 3 and 4, respectively.

Pressure drop is accounted for *via* the Ergun equation:³⁴

$$\frac{\Delta P}{\Delta L} = 150 \frac{\mu_f u_{\text{inf}}}{D_{\text{particle}}^2} \frac{(1 - \varepsilon_p)^2}{\varepsilon_p^3} + 1.75 \frac{u_{\text{inf}}^2 \rho_f}{D_{\text{particle}}} \frac{(1 - \varepsilon_p)}{\varepsilon_p^3} \quad (20)$$

such that the fluid density and fluid velocity can be determined using the ideal gas law based on inlet boundary conditions, which is generally considered to be valid at near-atmospheric pressure.

Deactivation is modeled by calculating the dynamic population of active sites (S_1 and S_2) and coke formation. These quantities are immobile which requires that the packed bed model be modeled with discretized volume fractions in the axial direction. Plug flow reactor assumptions are made to simplify the reactor model into a 1D simulation. The method of lines numerical approach was used to solve the system of partial differential equations that describe the reactor. A Matlab® script called MatMOL was used, which has been successfully demonstrated for similar problems.^{35–38}

Intraparticle model to decouple porous transport from chemical reactions

To represent intraparticle diffusion–reaction, the multistep effectiveness vector (MEV) approach was implemented. This MEV solution represents a pseudo-steady state analytical solution for diffusion and reaction of multistep reactions within catalyst particles, and is an extension of the Thiele modulus solution.²⁷ The solution represents a single catalyst particle, so it was multiplied by the total number of catalyst particle in each discrete axial subvolume to calculate the total rate of reaction in that subvolume. This approach is illustrated as shown in Fig. 1. The full derivation of the method for using a Thiele modulus with multiple reactions in series can be found in Lattanzi *et al.*²⁷ (a la the MEV). In short, the unsteady advection–diffusion model is used to describe a spherical porous catalyst particle,

$$\frac{dC_i}{dt} + u \cdot \Delta C_i = \Delta J_i - \sum_{j=1}^N R_{ij} + \sum_{m=1}^N R_{im} \quad (21)$$

where C_i is the molar concentration of vapor species i , u is velocity of gas, $\sum_{j=1}^N R_{ij}$ is the sum of reactions consuming i ,

and $\sum_{m=1}^N R_{im}$ is the sum of reactions forming i , and J_i is the flux of species i . Assuming no advection and using spherical coordinates the equation becomes

$$\frac{dC_i}{dt} = \frac{1}{r^2} \frac{\partial}{\partial r} \left(r^2 D_{i,\text{eff}} \frac{\partial C_i}{\partial r} \right) - \sum_{j=1}^N R_{ij} + \sum_{m=1}^N R_{im} \quad (22)$$

where r is the radial coordinate and $D_{i,\text{eff}}$ is the intraparticle diffusion coefficient. Next to non-dimensionalize the problem,

$$\hat{C}_i = \frac{C_i}{C_{i,\infty}} \quad (23)$$

where $C_{i,\infty}$ is the concentration of reactant i outside the particle,

$$\hat{r} = \frac{r}{r_p} \quad (24)$$

where r_p is the particle radius, and

$$\hat{t}_i = t \frac{D_{i,\text{eff}}}{r_p^2} \quad (25)$$

The Thiele modulus for consumption reactions must include all the reactions which consume i , and thus

$$\Phi_i = \sqrt{\frac{r_p^2 \sum_{j=1}^N k_{ij} S_l}{D_{i,\text{eff}}}} \quad (26)$$

where k is the rate constant for reaction j and S_l is the normalized active site concentration of site l remaining, from 1 to 0. Thus the remaining sites function as a deactivation factor. Similarly, for reactions that produce i from species m ,

$$\Phi_{im} = \sqrt{\frac{r_p^2 k_{im} S_l}{D_{i,\text{eff}}}} \quad (27)$$

If we then assume quasi steady state and apply boundary conditions we arrive at a dimensionless boundary value problem

$$\frac{d^2 \hat{C}_i}{d\hat{r}^2} + \frac{2}{\hat{r}} \frac{d\hat{C}_i}{d\hat{r}} - \phi_i^2 \hat{C}_i = - \sum_{m=1}^N \phi_{im}^2 \hat{C}_m \quad (28)$$

where at the center

$$\frac{d\hat{C}_i}{d\hat{r}} = 0 \quad (29)$$

and at the surface

$$\frac{d\hat{C}_i}{d\hat{r}} = \text{Bi}(1 - \hat{C}_i) \quad (30)$$

where Bi is the Biot number. Using matrix–vector form with the matrix of Thiele moduli for consumption and production, the whole system of reaction–diffusion can be described as:

$$\frac{d^2 \hat{C}}{d\hat{r}^2} + \frac{2}{\hat{r}} \frac{d\hat{C}}{d\hat{r}} - \bar{\phi}^2 \hat{C} = 0 \quad (31)$$

When the eigenvalues (λ) are real, the solution is a hyperbolic function:

$$\hat{U}_i = A_1 \sinh(\sqrt{\lambda_i} \hat{r}) + A_2 \cosh(\sqrt{\lambda_i} \hat{r}) \quad (32)$$



Converting back to concentration and applying the boundary conditions, where P is the eigenvector and D is the determinant, we find the concentration for all species at a given timestep

$$\hat{C} = \bar{P} \bar{D} \left(\frac{C_{Bi} \sinh(\sqrt{\lambda_i} \hat{r})}{\sinh(\sqrt{\lambda_i} \hat{r})} \right) \bar{P}^{-1} \hat{C}_{\text{Rat},\infty} \quad (33)$$

Thus the concentration for each species as a function of radius can be applied to the reaction rate and integrated over the whole particle:

$$\langle R_{ij} \rangle \equiv \frac{4\pi r_p^3 k_{ij} S_i C_{i,\infty}}{4/3\pi r_p^3} \int_0^1 \hat{C}_i \hat{r}^2 d\hat{r} = k_{ij} S_i C_{i,\infty} \eta_i \quad (34)$$

where η is the multistep effectiveness factor for that reaction

$$\eta_i = \bar{P} \bar{D} \left(\frac{3C_{Bi}}{\lambda} \left(\sqrt{\lambda} \coth(\sqrt{\lambda}) - 1 \right) \right) \bar{P}^{-1} \hat{C}_{\text{Rat},\infty} \quad (35)$$

and thus the effective rate for each species becomes

$$\langle R_{ij} \rangle \equiv C_{i,\infty} \left(\sum_m S_i k_{im} \eta_m - \sum_m S_i k_{ij} \eta_i \right) \quad (36)$$

In this particular reaction scheme, the effective reaction rate vector is the following:

$$\begin{aligned} \bar{R}_{\text{eff}} = & \begin{aligned} & -C_{\text{PV},\infty} (k_1 S_2 + k_{1G} S_2 + k_{1W} S_2 + k_3 S_1 + k_4 S_2) \eta_1 & \text{PV rate} \\ & C_{\text{PV},\infty} k_1 S_2 \eta_1 - C_{\text{OX},\infty} (k_2 S_1 + k_{2G} S_1 + k_{2W} S_1) \eta_2 & \text{OX rate} \\ & C_{\text{OX},\infty} k_2 S_1 \eta_2 & \text{HC rate} \\ & C_{\text{PV},\infty} k_{1G} S_2 \eta_1 + C_{\text{OX},\infty} k_{2G} S_1 \eta_2 & \text{LG rate} \\ & C_{\text{PV},\infty} (1 - y_{\text{CK}}) (k_3 S_1 + k_4 S_2) \eta_1 + C_{\text{PV},\infty} k_{1W} S_2 \eta_1 + C_{\text{OX},\infty} k_{2W} S_1 \eta_2 & \text{WAT rate} \\ & C_{\text{PV},\infty} y_{\text{CK}} (k_3 S_1 + k_4 S_2) \eta_1 & \text{CK rate} \\ & -C_{\text{PV},\infty} \theta_{S_1} k_3 S_1 \eta_1 & S_1 \text{ rate} \\ & -C_{\text{OX},\infty} \theta_{S_2} k_4 S_2 \eta_1 & S_2 \text{ rate} \end{aligned} \end{aligned}$$

The combined packed bed with sub-grid intraparticle MEV model was validated for simple reaction–diffusion following an example presented by Rawlings²³ as described in detail in the ESI†

Constant y_{ck} is 0.67, representing a mass loss of 33% to remove oxygen in the form of H_2O from pyrolysis vapor to leave a nearly graphitic coke. Initial values of S_1 and S_2 are 1 within the catalyst, and initial concentration of LMW PV is set to a smooth function along the length of the reactor for stability.

Other key model assumptions include:

- 1) Uniform coking and deactivation throughout the particle
- 2) Plug flow model with axial dispersion is valid for this packed bed reactor
- 3) Void fraction is homogeneous
- 4) The particles are spherical, particle size is uniform, and interparticle mass transfer is negligible
- 5) HMW PV, light gas, water, and light condensables are non-reactive
- 6) H_2 is present in excess.

After the model was parameterized and validated, it was used to fit apparent rate constants for the reactions that were

decoupled from the mass transport between the particles and within the particles. The simplex method⁴⁰ was used for numerical optimization of rate constants k_1 , k_2 , k_3 , k_4 , k_{1G} , k_{1W} , k_{2G} , k_{2W} , θ_1 , and θ_2 with the residual function representing the sum of the sum of least squares for the yield curves LMW PV, HC, and OX, and final yields of CK and LG, weighted by the inverse of each respective yield. Initial rate constant guesses were determined manually, and then refined by the simplex method using Matlab's `fminsearch` function. Optimization was run for 100 iterations with negative values prohibited. Then a mesh refinement study was performed to determine that ~ 100 nodes are required for accurate modeling, and the optimization was repeated, or until convergence tolerance $< 2 \times 10^{-4}$ was reached for sum of R^2 . The fitting was restarted with random numbers if convergence was not reached in 100 iterations.

Simulation results

The packed bed vapor phase upgrading (VPU) reactor model was used to iteratively fit apparent rate constants for our reaction scheme for catalytic upgrading of pyrolysis vapors over the deactivating Pt/TiO₂ catalyst up to biomass:catalyst ratio of 12. Note that these rate constants are for 450 °C and are independent of the catalyst porosity and particle size. Results from this optimization are shown in Table 4. A comparison of model predictions of lumped yields *vs.* experimental lumped yields *vs.* time is shown in Fig. 8, wherein the model and experiment match well and trend towards deactivation together.

It is notable that the model predicts that S_2 deactivates much faster than S_1 , as k_4 is much larger than k_3 . Under this model formulation S_2 deactivates faster than S_1 as indicated by k_4 and θ_{S_2} , parameters which are intrinsically coupled with respect to deactivation. If indeed this site does represent acid sites, which this paper does not prove *per se*, it is consistent with how quickly acid-dominant HZSM-5 deactivates in catalytic fast pyrolysis.⁴¹ Since the S_1 site corresponds to conversion of OX to HC, the first reaction step, PV to OX, occurs over the rapidly deactivating site. This matches the observation that OX does not dramatically dominate the product profile while the catalyst is deactivating. Deactivation factor θ_{S_2} is essentially irrelevant because k_3 , to which it is

Table 4 Reaction rate parameters fit by the optimizer coupled to the packed bed reactor

Rate constant	Fitted value
k_1 [s ⁻¹]	76
k_{1G} [s ⁻¹]	50.5
k_{1W} [s ⁻¹]	39
k_2 [s ⁻¹]	5.4
k_{2G} [s ⁻¹]	0.7
k_{2W} [s ⁻¹]	7.9×10^{-10}
k_3 [s ⁻¹]	7×10^{-14}
k_4 [s ⁻¹]	3.7×10^{-4}
θ_{S_1}	1.2×10^{-3}
θ_{S_2}	15.2



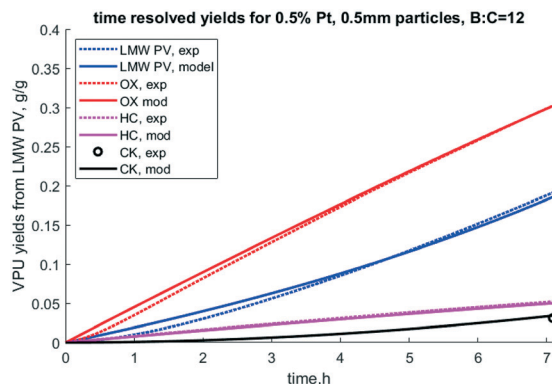


Fig. 8 Time resolved yields for 0.5% Pt, 0.5 mm particles, B:C = 12, 50/50 clean pine/forest residue. N.b. notation hydrocarbon HC, partially deoxygenated hydrocarbon (OX), coke (CK), and reactant low MW pyrolysis vapor (LMW PV), and experimental (exp).

intrinsically coupled, is so small. Further, k_1 is much larger than k_2 , which indicates higher reactivity of the S_2 site. Formation of light gas (LG) and water (WAT) are predicted to form at similar rates from PV, but the model fit seems to predict slower formation of these chemical families for the second stage of the reaction cascade.

To evaluate the accuracy of this model, cumulative yields from three other test case experiments were used: the same catalyst and biomass with B:C of 6 and 21 and an experiment with a catalyst with twice the platinum loading (1%), a different feed (100% clean pine vs. a 50/50 blend of forest residues and pine), and B:C 3. Changing the feedstock was accounted for by reviewing product characterization of the raw pyrolysis vapors. Doubling of the active sites was accounted for in the model by doubling S_1 and S_2 concentrations. In other tested simulations not presented here, doubling of a single catalytic site only resulted in behavior that did not match the experiments. This does not prove there are exactly two sites, but it does hint that that is the general physical phenomenon here.

Results shown in Fig. 9 show cumulative yields with the mass balance across VPU alone, based on LMW PV. For the base case at B:C 12 the model matches with experiments within the error, as expected. At B:C of 6 and 21, which had no replicates, the model predicts yields with an accuracy of 5% with the largest error being in the most abundant product OX. At a B:C of 21, the model predicts lower values for HC, OX and LG, but over predicts CK and PV. This could be because the experiment was done in two cycles. Cycle 1 was run until a B:C of 12 and cycle 2 was done the following day from B:C 12 to 21. The catalyst was left in N_2 flow overnight and was reduced in H_2 prior to running cycle 2, which reactivated some active sites. This may explain the discrepancies in the coke yields and shows the need for a thorough examination of coke formation and degradation over these catalysts.

However, most interesting is the case with twice the Pt, a quarter time on stream, and a slightly different biomass feed. Even with all those changes, all yields over the HDO reactor were within 5% with the exception of coke for which the

Yields from low MW pyrolysis vapors, VPU only

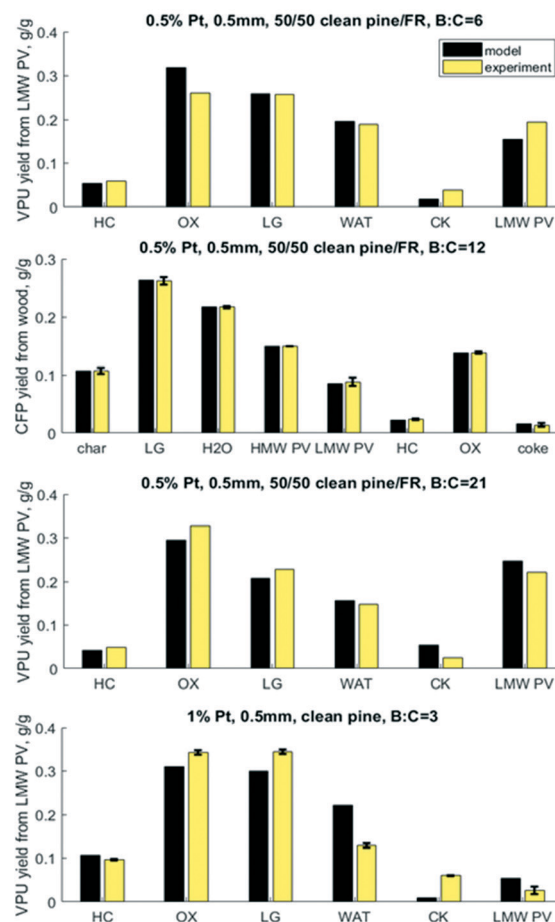


Fig. 9 Predicted vs. experimental yields of the six lumped species for the mass balance across the vapor phase upgrader (VPU) for three B:C ratios with 50/50 clean pine/forest residue and 0.5% Pt as well as one case with clean pine and 1% Pt. N.b. notation hydrocarbon HC, partially deoxygenated hydrocarbon OX, light gas (LG), reaction water (WAT), coke (CK), and reactant low MW pyrolysis vapor (LMW PV).

model underpredicted 1% yield vs. experimental 6% from LMW PV. It is possible that the different pine blend resulted in larger changes in the pyrolysis vapors that could not be accounted for in this lumped model, or the coke changes and degrades over time. Future work should explore these hypotheses around coking behavior and mechanisms.

When the cumulative yields are calculated over the mass balance of the whole system – wood to upgraded vapors – the relative variance decreases (Fig. 10). Again, the 0.5% Pt B:C 12 experiment with 50/50 CP/FR matches within the experimental error. More interestingly, the variance between experiment and model is reduced for the 1% Pt B:C 3 with clean pine. For light gas the model predicts 26% vs. experimental 28%; for water the model predicts 27% vs. experimental 23%; for breakthrough of reactant LMW PV the model predicts 2% vs. experimental 1%; for HC the model predicts 4.4% vs. experimental 4%; for OX the model predicts 13% vs. experimental 14%; and for coke the model predicts 0.4% vs. experimental 2.5%. For such a different experimental



Yields from dry wood for pyrolysis + VPU

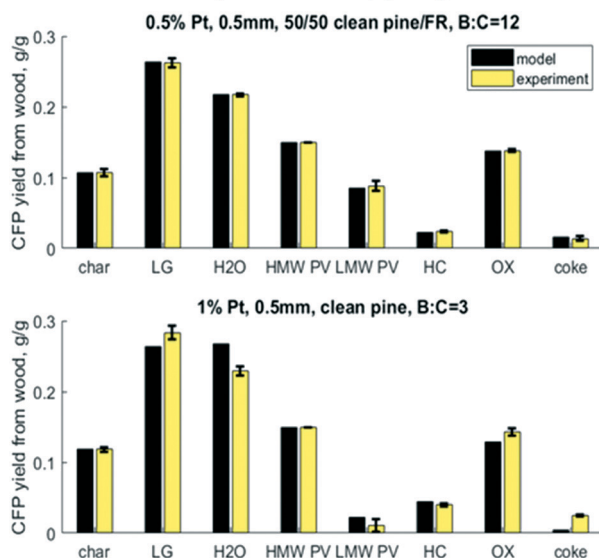


Fig. 10 Predicted vs. experimental yields of the six lumped species for the mass balance across the pyrolysis reactor and vapor phase upgrader (VPU) for 50/50 clean pine/forest residue, 0.5% Pt, B : C 12 as well as one case with clean pine, 1% Pt, B : C 3.

condition, these predictions exceeded expectations and show that while there is room for improvement, the existing model has valuable predictive capabilities. One of the limitations of this modeling approach is that the coke formation reaction rates were fit to final yields, not time-resolved yields. This could explain why the coke yields are not as accurate across different operating conditions than the other lumped chemical groups. Future modeling work exploring time-resolved coke yields and axial profiles would greatly improve the validity of this modeling approach. However, in general it is logical that if coke is formed from reactive pyrolysis vapors, then it is likely to accumulate near the entrance where there is a higher concentration of these compounds.

Nevertheless, this model can be used to perform sensitivity analysis to predict trends over a variety of operating conditions. An important output from the model is the prediction of accumulation of coke. The mass of coke, especially carbon content, directly impacts carbon yield in CFP oil as well as coke oxidation during catalyst regeneration. As shown in Fig. 11, more coke is predicted to accumulate near the entrance of the reactor as the reactive pyrolysis vapors are present in higher abundance there. As time progresses, more coke accumulates down the length of the reactor near the outlet, but the inlet has nearly twice the concentration compared to exit of the reactor.

Accurately predicting coke formation and localization is critical to de-risk this technology. If coke accumulates at the entrance, regeneration might be more effectively accomplished by feeding the oxidant through the outlet to reduce the risk of overheating the higher-coked inlet region. Over-heating through oxidative regeneration can induce sintering and even particle fragmentation. Further work should be done to

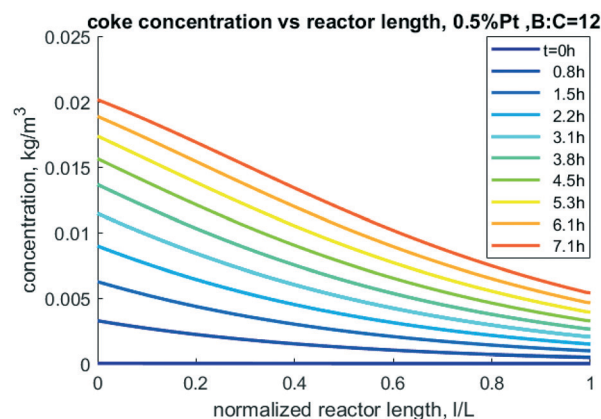


Fig. 11 Model-predicted coke density profiles through the reactor at different times for 0.5% Pt, 50/50 CP/FR example.

validate these predictions and use results to de-risk scaleup of the sensitive regeneration process. Similarly, incorporation of second order reactions with respect to consumption of hydrogen would also be relevant in future work, as the complex dynamics of consumption and generation of this gas has the potential to influence the rates of reaction.

To inform future experimental design, parametric sweeps were performed considering spherical particles from 0.25 mm to 5 mm catalyst diameters. This parameter is important because while it could be intuited that smaller particles will allow for full access to distributed active sites, it also increases the pressure drop in the bed which has an energy cost. Pressure drop from smaller particle sizes can be balanced by increasing the reactor size, to a degree. In these simulations all other parameters remained the same (bed void fraction, inlet conditions, particle porosity, *etc.*). The fluid density and velocity were back-calculated from the Ergun equation for each unique particle size simulation. Fig. 12 illustrates that reducing the particle diameter below 0.35 mm dramatically increases the pressure drop, and this should likely be avoided unless the reactor diameter is increased to 3.2 cm or larger for this size of catalyst bed (100 g). Going to larger D/L ratios requires expensive mixing distributors, so this may not be feasible. Interestingly, simulations to predict PV conversion over these particle sizes identified a notable regime for optimal performance. Nearly 100% conversion of reactive LMW PV to products is predicted at B:C 12 with 0.5% Pt on TiO_2 when the reactor diameter is less than 3.2 cm with particle sizes up to 5 mm, also shown in Fig. 10. This “sweet spot” would allow for longer operation before regeneration, less catalyst, and /or lower catalyst loading.

While the model development described in this work does warrant further examination and refinement, it does manage to achieve predictive capacity for lumped yields of products from catalytic fast pyrolysis. It is very rare to find any reports of rate constants for a lumped reaction scheme for the pyrolysis vapor phase HDO,^{42,43} let alone reaction kinetics that are decoupled from porous diffusion limitations. Lumped kinetics may be more applicable to this process than



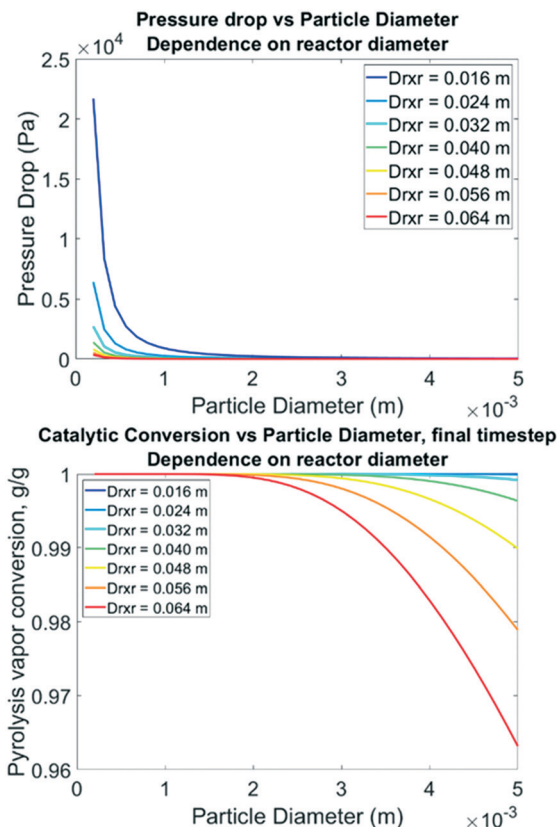


Fig. 12 Top: Model-predicted pressure drop from inlet to outlet vs. catalyst particle diameter following the Ergun equation with fixed inlet pressure & velocity. Bottom: Model-predicted LMW PV conversion vs. particle diameter for B : C 12, 0.5% Pt, 50/50 CP/FR.

rate constants for individual reactions when they are extracted from operational data because synergistic or competitive reactions may exist that are not observed in model compound studies. Further, this lumped reaction scheme approach allows for chemical engineers to design reactor systems and predict yields of products with this catalyst and vapor feed with some level of confidence, effectively reducing the risk of scaling up this technology for producing gasoline, diesel, and chemical precursors from renewable resources.

Conclusions

A simulation framework consisting of intraparticle diffusion-reaction for a cascading multistep reaction scheme coupled to an axial-dispersion packed bed reactor model was used to extract apparent reaction rate constants for a lumped catalytic fast pyrolysis scheme. The extracted rate constants were employed to simulate additional conditions beyond those from which they were obtained. The results provided reasonably accurate predictions of yields from a variety of different conditions in the same basic packed bed reactor system which was highly instrumented for near-100% mass closure. The model could be further improved by additional speciation of the pyrolysis vapor composition, as the current version is only valid for pine and some pine blends at 450 °C. Further

development should also be done to expand the mathematical underpinnings of the subgrid particle model to account for different particle shapes like cylinders. Nevertheless, this fast-solving fully-integrated multiscale modeling approach can be applied to a variety of catalytic chemical reaction processes with cascading reactions to de-risk critical emergent technologies in fuels and chemical production.

Abbreviations

Notation	Definition
B : C	Fed biomass to catalyst ratio, g : g
CFP	Catalytic fast pyrolysis
CK	Coke deposited on catalysts
CP	Clean pine
CP/FR	Clean pine/forest residue, 50/50
HC	Deoxygenated hydrocarbons, derived from OX
HDO	Hydrodeoxygenation; upgrading reactor
HMW PV	High molecular weight, assumed non-reactive pyrolysis vapors
LMW PV	Low molecular weight, reactive pyrolysis vapors
LG	Light gas
MEV	Multistep effectiveness vector (Thiele modulus-style intraparticle reaction-diffusion approximation)
OX	Partially deoxygenated hydrocarbons, precursor to HC
RGA	Residual gas analyzer
S_1	Catalytic reaction site 1
S_2	Catalytic reaction site 2
VPU	Packed (fixed) bed catalytic vapor phase upgrading reactor after the pyrolysis reactor
WAT	HDO reaction water

Conflicts of interest

There are no conflicts to declare. The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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References

- M. B. Griffin, K. Iisa, H. Wang, A. Dutta, K. A. Orton, R. J. French, D. M. Santosa, N. Wilson, E. Christensen, C. Nash, K. M. Van Allsburg, F. G. Baddour, D. A. Ruddy, E. C. D. Tan, H. Cai, C. Mukarakate and J. A. Schaidle, *Energy Environ. Sci.*, 2018, **11**, 2904–2918.
- K. Iisa, R. J. French, K. A. Orton, A. Dutta and J. A. Schaidle, *Fuel*, 2017, **207**, 413–422.
- S. D. Stefanidis, K. G. Kalogiannis and A. A. Lappas, *Wiley Interdiscip. Rev. Energy Environ.*, 2018, **7**, e281.
- A. N. Wilson, M. J. Price, C. Mukarakate, R. Katahira, M. B. Griffin, J. R. Dorgan, J. Olstad, K. A. Magrini and M. R. Nimlos, *ACS Sustainable Chem. Eng.*, 2017, **5**, 6615–6625.
- A. N. Wilson, A. Dutta, B. A. Black, C. Mukarakate, K. Magrini, J. A. Schaidle, W. E. Michener, G. T. Beckham and M. R. Nimlos, *Green Chem.*, 2019, **21**, 4217–4230.
- M. B. Griffin, G. A. Ferguson, D. A. Ruddy, M. J. Biddy, G. T. Beckham and J. A. Schaidle, *ACS Catal.*, 2016, **6**, 2715–2727.
- A. Eschenbacher, A. Saracian, B. H. Shanks, P. A. Jensen, C. Li, J. Ø. Duus, A. B. Hansen, U. V. Mentzel, U. B. Henriksen, J. Ahrenfeldt and A. D. Jensen, *Sustainable Energy Fuels*, 2020, **4**, 1991–2008.
- K. Wang, D. C. Dayton, J. E. Peters and O. D. Mante, *Green Chem.*, 2017, **19**, 3243–3251.
- S. Wan, T. Pham, S. Zhang, L. Lobban, D. Resasco and R. Mallinson, *AIChE J.*, 2013, **59**, 2275–2285.
- K. Murugappan, C. Mukarakate, S. Budhi, M. Shetty, M. R. Nimlos and Y. Román-Leshkov, *Green Chem.*, 2016, **18**, 5548–5557.
- M. A. Machado, S. He, T. E. Davies, K. Seshan and V. Teixeira da Silva, *Catal. Today*, 2018, **302**, 161–168.
- P. Cross, K. Wang, J. Weiner, E. Reid, J. Peters, O. Mante and D. C. Dayton, *Energy Fuels*, 2020, **34**, 4678–4684.
- A. Dutta, M. K. Iisa, M. Talmadge, C. Mukarakate, M. B. Griffin, E. C. Tan, N. Wilson, M. M. Yung, M. R. Nimlos and J. A. Schaidle, *Ex Situ Catalytic Fast Pyrolysis of Lignocellulosic Biomass to Hydrocarbon Fuels: 2019 State of Technology and Future Research*, National Renewable Energy Lab.(NREL), Golden, CO (United States), 2020.
- S. Boonyasuwat, T. Omotoso, D. E. Resasco and S. P. Crossley, *Catal. Lett.*, 2013, **143**, 783–791.
- T. O. Omotoso, B. Baek, L. C. Grabow and S. P. Crossley, *ChemCatChem*, 2017, **9**, 2612.
- P. M. de Souza, L. Nie, L. E. Borges, F. B. Noronha and D. E. Resasco, *Catal. Lett.*, 2014, **144**, 2005–2011.
- M. B. Pecha, E. Ramirez, G. M. Wiggins, D. Carpenter, B. Kappes, S. Daw and P. N. Ciesielski, *Energy Fuels*, 2018, **32**, 10683–10694.
- E. Ranzi, P. E. A. Debiagi and A. Frassoldati, *ACS Sustainable Chem. Eng.*, 2017, **5**, 2867–2881.
- E. Ranzi, P. E. A. Debiagi and A. Frassoldati, *ACS Sustainable Chem. Eng.*, 2017, **5**, 2882–2896.
- V. S. Bharadwaj, M. B. Pecha, L. Bu, V. L. Dagle, R. A. Dagle and P. N. Ciesielski, *Catal. Today*, 2019, **338**, 141–151.
- L. D. Dellon, C.-Y. Sung, D. J. Robichaud and L. J. Broadbelt, *Ind. Eng. Chem. Res.*, 2019, **58**, 15173–15189.
- O. Levenspiel, *Ind. Eng. Chem. Res.*, 1999, **38**, 4140–4143.
- J. B. Rawlings and J. G. Ekerdt, *Chemical reactor analysis and design fundamentals*, Nob Hill Pub., 2002.
- H. S. Fogler, *Elements of chemical reaction engineering*, Pearson PTG, London, England, UK, 6th edn, 2021.
- N. Jurtz, M. Kraume and G. D. Wehinger, *Rev. Chem. Eng.*, 2019, **35**, 139–190.
- E. W. Thiele, *Ind. Eng. Chem. Res.*, 1939, **31**, 916–920.
- A. M. Lattanzi, M. B. Pecha, V. S. Bharadwaj and P. N. Ciesielski, *Chem. Eng. J.*, 2020, **380**, 122507.
- J. R. Kremer, D. N. Mastronarde and J. R. McIntosh, *J. Struct. Biol.*, 1996, **116**, 71–76.
- PyMOL, *The PyMOL Molecular Graphics System*, 2017, Version 2.0.
- E. P. Barrett, L. G. Joyner and P. P. Halenda, *J. Am. Chem. Soc.*, 1951, **73**, 373–380.
- A. Oasmaa, Y. Solantausta, V. Arpiainen, E. Kuoppala and K. Sipilä, *Energy Fuels*, 2010, **24**, 1380–1388.
- A.-C. Johansson, H. Wiinikka, L. Sandström, M. Marklund, O. G. W. Öhrman and J. Narvesjö, *Fuel Process. Technol.*, 2016, **146**, 9–19.
- E. Hoekstra, R. J. M. Westerhof, W. Brilman, W. P. M. Van Swaaij, S. R. A. Kersten, K. J. A. Hogendoorn and M. Windt, *AIChE J.*, 2012, **58**, 2830–2842.
- S. Ergun, *Chem. Eng. Prog.*, 1952, **48**, 89–94.
- F. Logist, P. Saucez, J. Van Impe and A. V. Wouwer, *Chem. Eng. J.*, 2009, **155**, 603–616.
- A. Vande Wouwer, P. Saucez and W. Schiesser, *Ind. Eng. Chem. Res.*, 2004, **43**, 3469–3477.
- A. V. Wouwer, P. Saucez, W. Schiesser and S. Thompson, *J. Comput. Appl. Math.*, 2005, **183**, 245–258.
- P. Saucez, L. Some and A. V. Wouwer, *Appl. Math. Comput.*, 2009, **215**, 1821–1829.
- K. K. Sirkar, *Chem. Eng. Sci.*, 1974, **29**, 863–869.
- J. C. Lagarias, J. A. Reeds, M. H. Wright and P. E. Wright, *SIAM J. Control*, 1998, **9**, 112–147.
- C. Mukarakate, X. Zhang, A. R. Stanton, D. J. Robichaud, P. N. Ciesielski, K. Malhotra, B. S. Donohoe, E. Gjersing, R. J. Evans and D. S. Heroux, *Green Chem.*, 2014, **16**, 1444–1461.
- J. D. Adjaye and N. N. Bakhshi, *Biomass Bioenergy*, 1995, **8**, 131–149.
- J. D. Adjaye and N. N. Bakhshi, *Biomass Bioenergy*, 1995, **8**, 265–277.
- J. Klinger, D. L. Carpenter, V. S. Thompson, N. Yancey, R. M. Emerson, K. R. Gaston, K. Smith, M. Thorson, H. Wang, D. M. Santosa and I. Kutnyakov, *ACS Sustainable Chem. Eng.*, 2020, **8**(7), 2793–2805.

