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Population balance models for polymer upcycling: signatures of the mechanism in the molecular weight evolution

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Chemical and catalytic upcycling processes could help realize a circular plastics economy, but current models for testing mechanistic hypotheses and designing catalysts remain primitive. This work shows how proposed catalytic mechanisms can be incorporated into population balance models to predict the time evolution of molecular weight distributions. We develop models for homogeneous and heterogeneous catalysts, including catalysts that cut at chain ends and catalysts that cut at random locations. For heterogeneous catalysts, we illustrate the effect of adsorption constants that depend on polymer chain length. We discuss ongoing efforts and challenges in measuring and modeling the time evolving molecular weight distributions in polymer upcycling processes.

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1. Introduction

Millions of tons of plastic are produced each year. Most is discarded in landfills, lost to the natural environment, or incinerated.¹ Polymer upcycling efforts aim to transform plastics into value-added products.² In its broadest definition, polymer upcycling includes a variety of strategies: producing novel composites,³ use of functionalization⁴ and compatibilizers,⁵ conversion to carbon materials (nanosheets, nanotubes, *etc.*),⁶ and selective degradation to fuels, lubricants, *etc.*⁷ For this work, we focus primarily on upcycling *via* selective catalytic degradation.

Before polymer upcycling technologies can be implemented, challenges in plastics collection and sorting, process design, and catalyst development must be addressed.^{6,8-11} Catalyst development for polymer upcycling is complicated by some particularly unique challenges.¹²⁻¹⁶ First, the starting reactants in polymer upcycling are a jumble of polymers with many different molecular weights. Second, the process involves thousands of intermediates all being consumed and generated en route to products. Third, if things go badly, the process may yield multiphase mixtures containing hundreds or thousands of different products.

Because so many species are involved, we cannot use the familiar "initial rates" or "integrated rate law" analyses for small molecule reactions.¹⁷ Moreover, experiments cannot monitor the rates at which each product is formed. They can monitor time-evolving molecular weight distributions (MWDs) by

sampling reactions at various time points, *e.g.* with chromatographic or spectrographic methods.^{13,18-20}

Many experimental studies have reported the number average and/or weight-average molecular weight. Across multiple studies, with entirely different catalysts, polymers, and reaction conditions, the results show a fast initial drop in molecular weight with slower and slower decreases in molecular weight at long times.^{7,12,17,21} These results could be due to catalyst deactivation or to length-dependent selectivity.²¹ Zhang *et al.* provided a quantitative model to help explain these results.¹² In brief, cleavage of long polymers at early times results in a massive reduction of molecular weight, but as depolymerization proceeds there are more chains to cut and each cut results in a smaller reduction of molecular weight. In the model of Zhang *et al.*, the rate of cutting was also affected by product inhibition.¹²

Here we further simplify the model from Zhang *et al.*¹² Let N_0 be the initial number of molecules and let r(t) be the rate of cleavage events per unit time. Each cleavage event increases the number of molecules by one, and so the total mass is gradually distributed across an increasing number of molecules. Using only the definitions of the average molecular weight M_N , with no further assumptions, it can be shown that the number average molecular weight declines with time (t) as²²

$$\frac{M_{\rm N}(t)}{M_{\rm N}(0)} = \left(1 + \frac{1}{N_0} \int_0^t r \, \mathrm{d}t\right)^{-1} \tag{1}$$

Eqn (1) holds regardless of how, which, or where polymers are cleaved. The result is always an $M_{\rm N}(t)$ vs. t curve with approximately the same shape, whether chains are cut by hydrogenolysis^{12,21,23,24} or tandem metathesis/chain isomerization,^{7,18} longest chains first or shortest chains first,²⁵ at the

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chain ends^{26,27} or at random locations along the chain,^{28–30} *etc.* It also applies to linear or branched polymers. In this sense, eqn (1) is a universal feature of depolymerization. Eqn (1) also provides a simple way to extract a cleavage rate (cuts per time) from data, and (given the amount of catalyst) to obtain a quantitative catalyst activity, for example from r(t) divided by the number of catalyst sites. Because of its generality, it enables activity comparisons across different catalysts and different polymer chemistries.

Despite the utility of eqn (1), the fact that all catalysts, mechanisms, and polymers lead to a similar M_N vs. t curve limits its value as a tool for identifying the underlying mechanism. Additional information, like weight-average molecular weight vs. time data can be used to construct the polydispersity as a function of time, but this has a similarly mechanism-agnostic shape for similar reasons. Moreover, M_W primarily reflects the longest chains and thus provides limited information about small reaction products. Mechanistic studies require more detailed models that can predict both the product distribution and the entire MWD as a function of time.

The widely-used bottom-up strategy for modeling catalytic processes starts from molecular mechanistic hypotheses, *ab initio* calculations, microkinetic modeling, and ultimately predicts the product formation kinetics, reaction orders, activation parameters, *etc.*³¹⁻³³ Bottom-up models have successfully predicted kinetics and MWDs in some polymerizations, and the bottom-up strategy may also be successful for polymer upcycling.³⁴⁻³⁶ However, polymer upcycling by a heterogeneous catalyst involves polymer adsorption at sites, surfaces, or pores, in conjunction with multistep reactions involving many rate constants. With so many parameters to compute, errors in the quantum chemistry, force fields, or adsorption models may impair the predictions even if the hypothesized upcycling mechanisms are correct.

An alternative approach is to construct phenomenological kinetic models based on specific mechanistic hypotheses and test them against experimental data.^{37–40} This can be done with deterministic rate equations or kinetic Monte Carlo,⁴¹ *cf.* studies of long-chain *vs.* short-chain selectivity and product distributions from selectivity for different cleavage locations.^{21,42–45} Note that, even at the lab scale, experiments begin with upwards of billions of chains. Therefore the stochastic simulation results (if converged) should match the predictions of the corresponding deterministic rate laws⁴⁶ and (when they can be solved) the deterministic models are more easily fitted to quantitatively extract rate parameters from experimental data.⁴⁷

What types of deterministic kinetic models can predict MWDs for depolymerization? Kinetic lumping models^{48,49} predict the rates at which groups or "lumps" are consumed and generated, *e.g.* the reactants and products may be lumped into gases, liquids, and wax fractions. Recent work by Wu *et al.*²⁵ takes this direction.

Population balance equations (PBEs) go a step further, predicting the entire evolving MWD.³⁹ This work shows how, without specifying the mechanism at the most detailed level of elementary steps, we can already categorize upcycling mechanisms into a few "motifs" and construct the appropriate PBE. For constructing population balance models, most catalytic depolymerization processes will fit into one of sixteen categories.

• A catalyst may be homogeneous or heterogeneous.^{21,50}

It may operate processively or non-processively.^{51,52}

• It may cleave chains near their ends or at random locations along the chain.^{13,21}

• The polymers themselves may be dissolved in a solvent or a liquid melt with no solvent.^{44,45}

Each category gives a different population balance model, different solutions for the molecular weight evolution in time, and a different interpretation for the kinetic parameters. Each upcycling strategy implies a specific type of PBE with solutions that predict characteristic "signatures" of the mechanistic motif. Our goal is to discover easily identified signatures that can help to identify the underlying mechanism.

In constructing the population balance model, note that polymer fillers and even chemically inert particles in a composite may add new scission pathways. For example, they may change the importance of mass transfer and/or increase levels of shear induced mechanical chain scission. In some cases, the observed behavior is a combination of multiple mechanisms. Some processes employ "tandem" catalysts to combine advantages of the individual catalyst mechanism categories.^{7,18,45,53} We restrict ourselves here to ideal cases with single mechanisms.

In the following sections, we illustrate the predicted molecular weight evolution for the proposed mechanisms in several upcycling strategies. Each calculation starts from a lognormal MWD with $M_{\rm N}(0) = 3.0$ kDa and $M_{\rm W}(0) = 3.3$ kDa. First, in Section 2, we define a dimensionless time scale that helps to place the molecular weight evolution on a commensurate time scale regardless of the mechanism, the rate parameters, and their absolute sizes.

2. Activity comparisons

The mechanisms we discuss in this work span a range of conditions, catalyst concentrations, and reaction orders. Accordingly, the absolute time t is often inconvenient for comparing them. As previously noted by Helfferich, there is no universal way to nondimensionalize the rate equations that emerge from different mechanisms.⁵⁴ To allow comparison between mechanisms after a similar number of cleavage events, we invoke a dimensionless "cut time" τ . τ is the equivalent of the integral in eqn (1), and may be alternatively written based on the number of newly created molecules in the population:

$$\tau(t) = \frac{1}{N_0} \int_0^t r \, dt$$

$$= \frac{1}{N_0} \int_1^\infty (\rho(n, t) - \rho(n, 0)) dn$$
(2)

where $\rho(n,t)$ is the continuous concentration of polymers of length *n*, *i.e.* the count per unit volume of chains with length between *n* and *n* + d*n*. All models considered in this work exclude recombination, so each cleavage reaction creates one new chain.

3. Homogeneous mechanisms

We begin our discussion of mechanistic motifs for homogeneous reactions (catalytic or not) of the form

$$C_n \xrightarrow{k} C_{n-m} + C_m \tag{3}$$

where C_n is an arbitrary polymer with *n* repeat units and *k* is the pseudo-elementary rate constant. Models for the kinetics of these reactions have been used to describe radical depolymerization,⁵⁵ thermal degradation,^{40,56} and radiation-induced degradation,⁵⁷ among others.^{25,58,59}

3.1. Random cleavage

Random cleavage usually refers to any process in which all bonds are equally likely to be cut, although nonuniform cases have also been studied.⁶⁰ An schematic of random cleavage for a single starting chain is shown in Fig. 1. Examples of polymer upcycling strategies based on this method include an aminolysis process for polyesters⁶¹ and homogeneous cross alkane metathesis schemes based on work of Goldman and Brookhart.⁶²

A kinetic model may be constructed by considering polymer length as a discrete or continuous quantity. Discrete models are (in principle) more accurate because the number of monomers is countable. However, continuous models tend to be more mathematically convenient for large chains, so we focus on models with a continuous chain length here.⁶³ We borrow the continuous formulation of Staggs to account for cleavage steps:⁶⁴

$$\frac{\mathrm{d}\rho(n,t)}{\mathrm{d}t} = -kn\rho(n,t) + 2k \int_{n}^{\infty} \rho(m,t)\mathrm{d}m \tag{4}$$

Eqn (4) describes the change in $\rho(n,t)$, *i.e.* the population of chains of length *n* at time *t*, as *n*-mers are cut to form smaller species and as larger chains are cut to form *n*-mers. As an *n*-mer possesses approximately *n* bonds that can break independently, the first term is weighted by that length. The integrand is likewise weighted by *m*, but the probability of cutting an *m*-mer to an *n*-mer is $2m^{-1}$ d*m*. The overall timescale of the process is determined by *k*, which in this case is an effective first order rate constant with units of cuts per time per bond.



Fig. 1 Illustrating homogeneous random cleavage. Circles represent monomers in a large polymer. Long chains are repeatedly broken into smaller chains, with each cleavage site marked with a red X.



Fig. 2 Time evolution of the MWD for homogeneous random cleavage. Each line denotes the MWD at a point in rescaled time τ , with $\tau = 0$ (purple) the initial distribution. Lines are evenly spaced in τ , with $\Delta \tau = 0.3$.

The simplicity of this model makes it amenable to analytic and numerical solutions, and we direct the reader to the existing literature for a more detailed discussion of those solutions.^{29,30,65,66} The time evolution for homogeneous random cleavage is shown below in Fig. 2.

Homogeneous random cleavage manifests as a rising plateau, particularly among small species that are not present in the initial distribution. As depolymerization proceeds, the plateau grows narrower due to the preferential cutting of the longest chains, and taller, as further cuts increase the number of small fragments.

Homogeneous random cleavage quickly generates all possible chain lengths, from monomer to the largest initial chain. This rapid accumulation of small chain lengths can cause a rapid initial increase in the dispersity of the population, as shown in Fig. 3.

As random cleavage quickly generates lengths that were not part of the initial MWD, dispersity will rapidly increase. At later stages, long chains will be depleted, and the dispersity will



Fig. 3 Plot of number average chain length ($M_{N'}$, solid blue), weight average chain length ($M_{W'}$, dashed blue), and dispersity ($D = M_{W'}/M_{N'}$, dotted red) for homogeneous random cleavage as shown in Fig. 2.

eventually decrease again. In principle, at infinite time, all bonds will be cleaved, and all polymers will be transformed to monomer, with $M_{\rm N} = M_{\rm W} = D = 1$. An analysis of the moments of the MWD for homogeneous random cleavage processes has been given by McCoy and Madras.⁶⁰

3.2. Non-processive chain-end scission

These mechanisms remove a fixed-size oligomer or monomer from an end of a polymer chain. Such behavior is common in biological systems, *e.g.* β -amylase depolymerizes large, polymeric starches (blood sugar) into glucose by hydrolyzing glucosidic linkages.⁶⁷ An example upcycling process based on this mechanism is the tandem chain migration + olefin metathesis scheme proposed by Guironnet and Peters.¹⁸ An illustration of this mechanistic motif is shown in Fig. 4.

The discrete nature of the cuts make fully continuous formulations difficult, and in most cases the smallest species must be explicitly modeled.^{63,68} Zeman and Amundson developed tools to model chain length as a continuous variable, with a simple pseudo-elementary rate constant k, and with the monomer concentration separately determined by conservation of mass.^{18,69} The balance equation at the continuum population level for chain-end cleavage in homogeneous solution is:²⁶

$$\frac{\partial \rho(n,t)}{\partial t} = k \sum_{i=1}^{\infty} \frac{1}{i!} \frac{\partial^i \rho(n,t)}{\partial n^i}$$
(5)

The required number of terms for the continuous treatment to be valid depends on the smoothness and broadness of the polymer distribution. Typically, the second derivative term (i = 2) is sufficient.²⁶ Because cuts are always made at the end of a chain, the rate constant k is first order in chains rather than bonds. Guironnet and Peters have shown how the appropriate pseudo-elementary rate constant k can be derived from more detailed kinetic schemes.¹⁸ The monomers and long chain populations must both be considered when computing the number and weight averages. The time evolution for a chainend scission process is shown in Fig. 5.

Chain-end scission results in a gradual broadening and translation of the MWD towards lower chain lengths. The rate of this broadening and translation depends on the initial width of the MWD and the size of the fragment removed. Systems that



Fig. 4 Illustration of a chain-end cleavage process. Circles represent monomers in a large polymer. With each cleavage reaction, marked by a red X, a monomer is removed from the end of the chain.



Fig. 5 Time evolution the MWD by homogeneous chain-end cleavage. Monomer concentration is not shown to prevent distortion of the vertical axis. Lines are evenly spaced in τ , with $\Delta \tau = 200$.

cleave variable length oligomers will give faster broadening than systems that cleave the same fragment each time.¹⁸ When viewed on a log *n* scale, this broadening does result in a rising plateau like that of homogeneous random scission (Fig. 2). However, the initial MWD gradually shifts leftward from chainend scission, while the initial MWD is decimated after orders of magnitude fewer random cleavage steps. In this sense, random cleavage is more efficient at rapidly altering the MWD than chain-end cleavage.

Unlike random cleavage, there is typically a clear demarcation between volatile and nonvolatile products in chain-end cleavage, particularly when the initial distribution is primarily long polymer chains.⁷⁰ An easy separation between products and long polymers may have practical reaction engineering advantages at the process design stage. However, it can create difficulties for data analysis in bench-scale experiments. Discarding the small molecule products when characterizing the MWD will affect the results.⁷⁰ Procedures that include *vs.* exclude the very small fragments produced by chain-end scission can give very different trends in $M_{\rm N}$, $M_{\rm W}$, and D, as shown in Fig. 6.

With the simplest cases of random and chain-end scission discussed, it is important to note that many processes are in fact combinations of the two.^{30,38,56,68,71-73} For example, in the polysaccharide/amylase system, there are three classes of enzymes that work together to depolymerize starches: α -amylase catalyzes random cleavage, while β - and γ -amylase catalyze chain-end cleavage.⁷⁴⁻⁷⁶ The relative concentration and activity of these amylases can dramatically alter the evolution of the MWD.^{77,78} A single catalyst that targets bonds at random may also have a significant chain-end scission preference due to different chemistries at chain ends.⁴⁰

Heterogeneous mechanisms

We now focus on models for polymer upcycling by heterogeneous catalysts. Heterogeneous catalytic mechanisms introduce additional theoretical difficulties. In particular, one must



Fig. 6 Plot of number average chain length (M_N , solid blue), weight average chain length (M_W , dashed blue), and dispersity ($D = M_W/M_N$, dotted red) for the population evolution in Fig. 2. Note the large difference in dispersity scale between plots. (a) Small molecule products are not counted, and (b) small molecule products are counted when calculating averages and dispersity.

consider the bulk MWD and the distribution of molecular weights for adsorbed polymers.⁴⁷ Long chains and short chains compete for adsorption sites on the surface, with preferential adsorption determined by the loss of entropy upon adsorption and enthalpic interactions that (in some cases) favor adsorption.^{79,80} We address two regimes in this work: a polymer melt in which a polymer in the bulk is surrounded by like polymers, and a dilute solution in which the polymer in the bulk primarily interacts with a solvent. In both cases, we utilize quasiequilibrium adsorption models. The models will be inaccurate if the reaction is too fast for polymer conformations in the interfacial layer to relax to a local conformational equilibrium, or if boundary layer transport (where applicable) is too slow to maintain an equilibrium with the bulk.

4.1. Random cleavage at melt-catalyst interface

In this section we consider a polymer melt in contact with a heterogeneous catalyst that cleaves chains at random locations. A schematic for this case is shown in Fig. 7. Examples of catalysts that likely work *via* this mechanism include hydrogenolysis by Ru/C⁴³ and Ru/TiO₂,⁴² tandem hydrogenolysis and



Fig. 7 For a polymer melt in contact with a heterogeneous catalyst, only bonds within the layer near the catalyst (red dashes) can be cleaved. Enthalpy is largely irrelevant for adsorption because all surface sites will be in contact with portions of some chemically similar chain. Preferential adsorption of small chains may occur because the conformational entropy loss upon adsorption is smaller for small chains.

aromatization on $Pt/\gamma\text{-}Al_2O_3,^{12}$ and melt hydroconversion by Pt/ WO_3/ZrO_2.^{81}

In a melt, the entire catalyst surface is in contact with polymers, but the contacts may be segments from chains with different lengths. Therefore, we define a surface coverage such that the fraction of all catalytic sites occupied by segments of an *n*-mer is given by $\theta(n)$. Because the surface must be covered entirely, $\int \theta(n) dn = 1$. The MWD of the adsorbed chains may deviate from the MWD of chains in the bulk. We may account for this nonideality by writing

$$\theta(n) = \theta^{\text{ideal}}(n) + \theta^{\text{excess}}(n) \theta^{\text{ideal}}(n) = \phi^{\text{bulk}}(n)$$
(6)

where the ideal surface coverage of a species is given by the bulk volume fraction of that species ϕ^{bulk} . A model for the surface excess was given by Van der Gucht *et al.* as

$$\theta^{\text{excess}}(n)/\phi^{\text{bulk}}(n) = A(1 - n/M_{\text{W}})$$
(7)

where *A* is a constant related to the enthalpic and entropic differences experienced by chain ends relative to an internal monomer.⁸² *A* is determined jointly by the surface, polymer end, and polymer backbone chemistries. A > 0 indicates a surface that is attractive to small chains, either due to favorable enthalpic effects of the chain-end chemistry being proportionally larger for smaller chains, or due to the lesser entropic penalty for confining small chains at the surface.

If we assume quasi-equilibrated coverages, then the fraction of the surface covered by *n*-mers is

$$\theta(n,t) = \frac{n\rho(n,t)}{\int_{1}^{\infty} m\rho(m,t) dm} (1 + A(1 - n/M_{\rm W}))$$
(8)

In formulating eqn (8) we have ignored excess mixing volumes by assuming that, for each *n*, $n\rho(n,t)$ is proportional to

the bulk volume fraction $\phi^{\text{bulk}}(n)$. Now the bulk MWD changes in response to the equilibrium adsorption and kinetics of scission as

$$\frac{\mathrm{d}\rho(n,t)}{\mathrm{d}t} = -r\theta(n,t) + 2r\int_{n}^{\infty}\frac{1}{m}\theta(m,t)\mathrm{d}m \tag{9}$$

where r is the effective rate of depolymerization, per eqn (1).

Eqn (9) parallels the result for homogeneous random scission, eqn (4), except the surface coverage θ allows the dependence on chain length to be more complex. In homogeneous random scission, the polymer reactivity is proportional to the number of bonds in the chain (eqn (4)). Non-zero values of parameter *A* lead to length-dependent adsorption and to a length-dependent cleavage selectivity that deviates from the bulk volume fractions. For example, when A < 0, long chains are favored to adsorb and cleave. When A > 0, short chains are favored. The preferential adsorption relative to the bulk populations is shown in Fig. 8 for a series of different *A* values.

If products of an intermediate molecular weight are desired, *e.g.* in hydrogenolysis of polyethylene, the catalyst should be designed such that A < 0 if possible. If A > 0 instead, then long chains will be excluded from the surface, causing the short chains to be repeatedly cut and resulting in a mixture with large fractions of over-hydrogenolysis products (like methane) and uncut chains. Fig. 9 shows that, according to eqn (8) and (9), the MWD evolves in a manner like that for homogeneous random scission. Fig. 9 shows the dependence on parameter *A* by plotting solutions for A = -1.0 and A = +1.0.

As in Section 3, we may also consider the dispersity of the resulting polymer. This is demonstrated in Fig. 10. Surfaces that favor the adsorption of longer species (A < 0) result in a less disperse polymer product relative to the homogeneous case shown in Fig. 3. Short chain favoring surfaces (A > 0) do exactly the opposite, with an increase in dispersity.

In the melt, where enthalpic driving forces largely cancel with those of other chemically similar chains, the value of A will be largely determined by entropic factors. Long chains lose



Fig. 9 Time evolution of the MWD for random-cleavage at a meltcatalyst interface. Black line denotes initial polymer population. Red denotes a long chain favoring surface (A = -1), dashed blue denotes a short chain favoring surface (A = +1), per Fig. 8. Lines are evenly spaced in τ , with $\Delta \tau = 0.5$.

more conformational entropy upon adsorption to a flat surface than short chains, so catalysts that present a flat interface to the melt may lead to the situation with $A > 0.^{42-45,83}$ It may be possible to tune the value of *A* (and the product distribution) by altering the chemistry of the chain ends to disfavor the surface or to design special pore geometries that favor adsorption of long chains as done by Tennakoon *et al.* and Wu *et al.*^{13,25}

Note that the surface segregation model of van der Gucht *et al.* breaks down for very large polymers in highly polydisperse melts. For positive *A*, *i.e.* a surface that preferentially adsorbs smaller polymers, there may even be chains for which $1 + A(1 - n/M_W)$ is negative. According to this linear relationship, these extremely long chains are entirely unreactive, kept away from the surface by the abundance of shorter chains. These chains will not become reactive with the passage of time, as depolymerization can only decrease the average chain length. The possibility of a nonreactive set of very large chains must be considered when using these



Fig. 8 Surface coverage (eqn (8)) for varying surface segregation parameter *A*, for the initial distribution shown in Fig. 2. Red denotes a long-chain favored surface; blue denotes a short-chain favored surface. The bulk mass fraction for all cases is equal to the surface coverage for the non-interacting surface (black).



Fig. 10 Dispersity for melt-surface random scission. Dashed black line denotes the homogeneous case (A = 0, Fig. 3). Red denotes a long chain favoring surface (A < 0), blue denotes a short chain favoring surface (A > 0), per Fig. 8. Lines are evenly spaced in A, with $\Delta A = 0.2$.

equations with very large maximum chain lengths n_{maxo} or very small average chain lengths M_{W} . When generating Fig. 9 and 10, we replaced any negative values of $\theta(n,t) < 0$ with $\theta(n,t) = 0$. Further development of preferential adsorption models would be a useful direction. Note that the ideal and excess coverages in eqn (6) are additive, in contrast to activity models where non-ideality is included *via* a multiplicative activity coefficient. It would be useful in future work to develop models which automatically satisfy the proper asymptotes in the long chain and dilute limits, like regular solution models⁸⁴ and Margules models.⁸⁵

4.2. Solute-surface random scission

Now we consider heterogeneous catalytic cleavage of polymers that are dissolved in solution. In this case, the surface is no longer guaranteed contact with polymers, and this necessitates a more complicated handling of the surface-polymer interactions and adsorption.^{86–89} We assume that polymer chains adsorb to a reactive surface from solution and proceed to either desorb back to the bulk or react, as shown in Fig. 11. As an examples of this type of system, Ellis *et al.* considered a SnPt/ γ -Al₂O₃ and Re₂O₇/ γ -Al₂O₃ system with *n*-pentane solvent.⁴⁵

Again, we assume well-mixed conditions where coverages are quasi-equilibrated with the bulk polymer concentrations. Note that this assumption may be lifted using standard techniques for treating exterior transport limitations. Namely, one would replace the bulk populations in the models below with unknown populations near the catalyst surface and equate the resulting rates to the rate of chain transport from the bulk through a boundary layer to the surface. External mass transport models are currently under development.

To build the PBE, we again consider the fractional surface coverage $\theta(n)$ representing the fraction of sites in contact with segments from an *n*-mer. The bulk MWD evolves as in the melt-surface case, by the same PBE as that in eqn (9). The difference



Fig. 11 Illustration of solute-surface random scission. (a) Long chains adsorb to the surface from the bulk. (b) The adsorbed chain is irreversibly cut into two or more smaller fragments. (c) Smaller fragments desorb back to the bulk or remain on the surface for further scission. Adsorption and desorption steps (a) and (c) are assumed to be reversible and guasi-equilibrated.

lies in the isotherms for the bulk-surface coverage relationship. To determine $\theta(n)$ in this case, we invoke a common multi-site generalization of the Langmuir isotherm:⁹⁰

$$s_n K_n \rho(n,t) = \theta(n,t) / \theta_0(t)^{s_n}$$
(10)

Here s_n is the average number of catalytic sites occupied by an *n*-mer, K_n is a length dependent equilibrium constant, and $\theta_0(t)$ is the fraction of unoccupied sites, *i.e.* $\theta_0(t) = 1 - \int \theta(n,t) dn$ where the integration bounds are from n = 1 to ∞ . We split the free energy within the equilibrium constant into an entropic component (assumed constant) and a per-adsorbed site enthalpic component

$$K_n = k_e \exp[-\Delta H_{ads} s_n + T \Delta S_{ads}]$$
(11)

where k_e is a constant prefactor with units of inverse concentration, like a standard reference volume. We assume here that the entropic and enthalpic terms are independent of the changing composition of the solution. Given another layer of theory that connects the composition to the adsorption energies, a time- or composition-dependent K_n may be incorporated into the model. We leave this development for future work.

Multiplying both sides of eqn (11) by $\theta_0(t)s^n$, and integrating both sides from n = 1 to ∞ yields a single equation for the fraction of empty sites:

$$\theta_0(t) = 1 - \int_1^\infty s_n K_n \rho(n, t) \theta_0(t)^{s_n} \, \mathrm{d}n$$
 (12)

At the initial time step, the fraction of empty sites may be solved to a tight error tolerance for the initial coverages of adsorbed polymers, and then included in the system of differential equations to compute changes in coverage over the integration period. Differentiation of eqn (12) and rearrangement yields

$$\frac{\mathrm{d}\theta_0}{\mathrm{d}t} = \frac{\int_1^\infty s_n K_n \theta_0(t)^{-s_n} \frac{\mathrm{d}\rho}{\mathrm{d}t} \,\mathrm{d}n}{1 - \int_1^\infty s_n^2 K_n \theta_0(t)^{s_n - 1} \rho(n, t) \mathrm{d}n} \tag{13}$$

Unlike previous models discussed in this work, eqn (10)–(13) depend on the total concentration N_0 . As the total concentration of chains rises, the surface becomes more occupied. While this effect does impact the evolving MWD, the contribution is minor relative to that of the adsorption energetics. We focus here on the convenient case where $k_e N_0 = 1$.

When integrating forward in time, only one nonlinear solution needs to be computed at the first step. From the computed fraction of empty sites, the individual species coverages may be estimated and interpolated by eqn (10). The change in the population for the current timestep follows from eqn (9), and (13) then also predicts the revised fraction of empty sites.

Solute-surface adsorption behavior can cause drastic departures from the prior cases. The form of eqn (10) reveals that the primary factor controlling a species surface coverage is the number of sites it demands. Polymer adsorption theories



Fig. 12 Relative preferential adsorption for quasi-equilibrated bulk polymer ρ (log normal MWD with $M_{\rm N}(0) = 3.0$ kDa and $M_{\rm W}(0) = 3.3$ kDa, $k_{\rm e}N_0 = 1$) and surface θ . Positive numbers (red) denote long chains are preferred for adsorption. Negative numbers (blue) denote short chains are preferred for adsorption.

suggest that the preferred number of contacts between a polymer and a surface scales roughly as $n^{1/2}$.⁹¹⁻⁹⁴ Based on this, and the fact that an adsorbed monomer should have one contact with the surface, $(s_1 = 1)$ we perform calculations for such that a monomer occupies one site. Then, the adsorption energetics may be varied. We initially consider the surface coverage in quasi-equilibrium with the initial polymer population, relative to the bulk population, by analogy to Fig. 8, in Fig. 12.

$$s_n = \sqrt{n} \tag{14}$$

Fig. 12 reveals that this model, like the melt-surface case, may prefer either long or short chains depending on the energetics of adsorption. This preference is primarily driven by the constant entropic penalty for adsorption. When this penalty is strong, large chains are preferred, as they adsorb to multiple sites and thus have a greater enthalpy of adsorption. When this penalty is weak, small chains are preferred, as they demand fewer sites each to adsorb and do not require a large enthalpy to offset the entropic penalty.

Preferential adsorption strongly impacts the evolving MWD as cleavage proceeds. Minor differences in the behavior of the initial distribution may be magnified as polymers are cleaved and the MWD evolves. We demonstrate this in Fig. 13, as small adjustments to the adsorption enthalpy drive significant changes in the product distribution. According to the model, solute-surface random scission creates significant amounts of small products (n = 1-100). Therefore, we show the massweighted MWD, $n\rho(n,t)$, to highlight the differences between distributions.

As the system evolves, it develops a new peak in the massweighted MWD that represents the primary product size by mass. The size of the primary product is associated with both the site-dependent ΔH and site-independent $T\Delta S$ parameters. At fixed conversion, smaller values of ΔH and/or $T\Delta S$ cause the primary product size to decrease, approximately correlated with





Fig. 13 Time evolution of the mass-weighted MWD by solute-surface random cleavage for varying per-site enthalpy of adsorption. ΔH and $T\Delta S$ in $k_{\rm b}T$ units, $k_{\rm e}N_0 = 1$. Lines are evenly spaced in τ , with $\Delta \tau = 0.3$. Arrows denote the length of the product species with the greatest mass fraction at $\Delta \tau = 3$.

the initial preference shown in Fig. 12. Increased conversion will also decrease the primary product size.

When adsorption strongly favors small species a bimodal distribution may arise, with the primary product coexisting with the initial distribution, even at high conversion (Fig. 13, bottom). After the first few catalytic turnovers, the products of cleaving the initial, large polymers preferentially adsorb. The products of their cleavage in turn are preferentially adsorbed, and so on, creating an abundance of small molecular weight products.

5. Conclusions

Population balance models have long been a powerful tool to predict product distributions for both polymerization and depolymerization processes, but most research has focused on the simplest of problems for which analytical solutions exist. Here we have demonstrated the beginnings of a framework to translate specific mechanistic proposals into population balance models. The population balance models then yield specific predictions about the molecular weight evolution. We have developed models for a variety of processes and mechanisms. Each mechanism and its corresponding population balance model predicts "fingerprints" in the molecular weight evolution by which an observed behavior may be matched to an appropriate underlying mechanism. Efforts are underway to fit these models to experimental data as a quantitative way of mechanism identification.

Conflicts of interest

There are no conflicts to declare for this work.

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