Analysing the effect of screw configuration using a stochastic twin-screw granulation model

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Abstract

In this work we present a framework for modelling twin-screw granulation processes with variable screw configurations using a high-dimensional stochastic population balance method. A modular compartmental approach is presented and a method for estimating residence times for model compartments based on screw element geometry is introduced. The model includes particle mechanisms for nucleation, primary particle layering, coalescence, breakage, and consolidation. We introduce a new twin-screw breakage model which takes into account the differing breakage dynamics between two types of screw element. The resulting model is used to simulate a twin-screw system with a number of different screw configurations and the predictive power of the model is assessed through comparison with an existing experimental data set in the literature. The model demonstrates the ability to qualitatively capture experimental trends, including the reduction in fines associated with an increase in the number of kneading elements in the screw configuration. Based on model results, a number of key areas for future model development are identified and discussed.

Highlights

- Existing model extended to include novel layering mechanism
- Novel compartment residence time model developed
- New breakage model reflects screw element geometry effects
- Model captures the liquid distributing/growth promoting effects of kneading elements
1 Introduction

Twin-screw wet granulation (TSG) is a relatively new method of continuous granule man-
ufacture. TSG devices, such as the one depicted in Figure 1, consist of two co-rotating
screws enclosed within a barrel. The screws are constructed from an inter-changeable se-
quence of elements with various geometries. In the case of wet granulation, a solid blend
of excipient/ active-pharmaceutical ingredient (API) and liquid binding agent are sepa-
rately and continuously fed to the system. The solid and liquid phases interact in a high-
shear environment of the screw barrel to create a potential mixture of granules and often
some ungranulated powder mass, depending on the operating condition [27, 43, 48, 62].
The resulting granules find application as tabletting feed stock within the pharmaceutical
industry, the fertiliser industry and foodstuffs [30].

Figure 1: Variation zone of the a twin-screw granulator

TSG has a number of benefits over traditional batch granulation, namely, reduced plant
footprint, ease of scale-up [69], the ability to create granules with high drug loading [65]
and a highly configurable set-up, primarily due to the variable screw configuration. How-
ever, the configurability of the device also creates a very large operating space for process
optimisation during formulation, which can be problematic when only small quantities
of API may be available. For this reason, a number of experimental studies [43, 68]
have tried to investigate the relationship between the screw configuration, in terms of the
number/position of certain types of element, and key properties of the product granules
(such as particle size distribution (PSD), porosity and liquid distribution). Some studies
have also focused on defining the role of specific types of screw element by using them
in isolation [57, 62]. Experimental investigations into particle breakage in twin-screw
devices [22, 32, 43, 57, 68] have highlighted the role of screw element geometry on the
breakage dynamics along the barrel. For example, several studies [43, 57] have shown
that that the large agglomerates in conveying elements undergo size reduction through
cutting or edge chipping, where small fragments are continually taken from the edge of
the agglomerates. The particle size distribution transformation induced by distributive
mixing elements (DME) in Pradhan et al. [57] suggests that breakage in these elements
occurs through a combination of crushing and chipping [57].
Because of the combinatorial nature of the twin-screw operating space, it is desirable to develop a process model where the screw sections can be treated in a modular manner, such that the performance of new screw configurations may be quickly assessed without the usage of excipient/API or the need to set-up the device etc. This has generally been attempted through the use of compartmental population balance models (PBM) [58]. Several examples of compartmental twin-screw PBMs exist in the literature [2, 3, 5, 31, 35]. In these examples, the screw barrel domain is modelled as a number of connected compartments that permit process conditions and thus particle morphology to vary along the length of the simulation domain. These examples have used a sectional solution approach [36–38] which allows the compartmental PBM to be approximated and solved as a system of ordinary differential equations. This numerical approach generally limits the particle representation to taken on three dimensions at most. The Stochastic particle method [20, 54, 56, 59, 71–74] is alternative approach that has been employed to solve PBMs for batch granulation systems [9–14, 23, 39, 40], silica [50] and TiO₂ [7] nano-particle synthesis, soot formation [15, 52], and more recently twin-screw granulation [45, 46]. Unlike sectional methods, stochastic particle methods permit much more complex particle representations, which can then be leveraged within the process model description, whilst still yielding a numerical problem that can be solved with acceptable levels of computational effort.

The main aims of this paper are:

1. Improve the stochastic TSG model in McGuire et al. [45, 46] based on the areas identified for improvement.
2. Construct a modelling framework that allows for the compartmental representation of arbitrary screw configurations and incorporates an element rate constant library. This library should contain optimised, re-usable, model rate parameters for different types of screw element.
3. Use the modelling framework to optimise a model parameter library against existing experimental data.
4. Assess the predictive ability of the complete model framework/parameter library using experimental data associated with a screw configuration not used in the optimisation of the library.

The remainder of the paper is structured as follows: firstly we present the TSG PBM and a discussion of the compartmentalisation methodology. The stochastic weighted particle method used to simulate the PBM is then presented in detail in Section 3. The experimental systems used for the optimisation step and the simulation conditions are described in Section 4. This is followed by a discussion of the model results in Section 5 and concluding remarks in Section 6.
2 Twin-screw population balance model

2.1 Particle type-space and population balance equation

In this TSG model, particles are described using the type-space $X$. In $X$, each element is characterised using a four-dimensional vector $x = (s_o, l_e, l_i, p) \in X$ where: $s_o$ is the volume of original solid, $l_e$ is the volume of external liquid, $l_i$ is the volume of internal liquid and $p$ is the pore volume. The key derived particle properties are summarised in Table 1.

<table>
<thead>
<tr>
<th>Property (Nomenclature)</th>
<th>Expression</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volume ($v$)</td>
<td>$s_o + l_e + p$</td>
<td>m$^3$</td>
</tr>
<tr>
<td>Diameter ($d$)</td>
<td>$(6v/\pi)^{1/3}$</td>
<td>m</td>
</tr>
<tr>
<td>Mass ($m$)</td>
<td>$\rho_s s_o + \rho_l (l_i + l_e)$</td>
<td>kg</td>
</tr>
<tr>
<td>Porosity ($\varepsilon$)</td>
<td>$p/v$</td>
<td>-</td>
</tr>
<tr>
<td>External surface area ($a_{surf}$)</td>
<td>$\pi^{1/3}(6v)^{2/3}$</td>
<td>m$^2$</td>
</tr>
</tbody>
</table>

‘Free’ primary particles (i.e. those that are not part of an agglomerate) are comprised purely of original solid. Since the width of the primary particle size distribution is significantly smaller than that of the aggregate distribution of interest, these particles are modelled as a mono-disperse phase with representative particle diameter $d_{pp}$ and volume $v_{pp}$. Furthermore, since the set of all primary particles $X_{pp} \subset X$ is only permitted to occupy a very limited region of the type-space ($X_{pp} = \{x_{pp}\}$, $x_{pp} = (v_{pp}, 0, 0, 0)$) it is sufficient to characterise this phase by the number concentration of primary particles $c_{pp} \in \mathbb{R}^+$ or, equivalently, the number of primary particles $N_{pp} \in \mathbb{R}^+$ that exist within a given volume of the system to be modelled. The aggregate type space may then be defined as $X_{agg} = X \setminus X_{pp}$.

Elements of $X_{agg}$ take positions in a bounded domain of compartments. Similarly, each compartment has an associated number of primary particles $N_{pp}$. Each compartment is denoted by its index $z \in \mathbb{L}$. Particles (both primaries and aggregates) are permitted to move between compartments according to the connections defined by the compartmentalisation of the system (discussed further in Section 2.4). Only particles within the same compartment are permitted to interact with each other.

In this work, we make use of the idea of deferred processes as defined by the Linear Process Deferment Algorithm [55]. This algorithm is utilised to defer the applications of linear process operators that are particularly computationally intensive, such as the layering of primary particles onto the surface of large aggregates.

Since the population balance model is to be solved using the stochastic particle method, it is constructed in weak form. That is to say, each of the terms is integrated against some suitable test function. Let

1. $\lambda(z, t, dx)$ be a concentration measure on $X_{agg}$ at time $t$ in compartment $z$,
2. addition and subtraction on \( X \) correspond to particle coagulation and breakage, respectively,

3. \( \varphi(z, x) : X \mapsto \mathbb{R} \) be a suitable test function which is smooth with compact support,

4. \( D_t : ([0, \infty), X_{agg}, \mathbb{R}^+) \mapsto X_{agg} \) be the aggregate deferment function that carries out the deferred aggregate processes. For any particle \( x \), compartment \( z \) and primary particle concentration \( c_{pp} \), \( D_t(z, x, c_{pp}) \) is distributed as the value at time \( t \) of the Markov chain defined by the deferred jump processes and their associated rates. In this way, \( \mathbb{P}(D_t(z, x, c_{pp}) = dz) \) defines the probability that particle \( D_t(z, x, c_{pp}) \) will lie within the type-space \( [\xi, \xi + d\xi] \).

The weak form of the aggregate PBE to be solved is then

\[
\begin{align*}
\frac{d}{dt} \int_{x \in X_{agg}} \varphi(z, x) \lambda(z, t, dx) &= \int_{x \in X_{agg}} \varphi(z, x) I_{nuc}(z, t, c_{pp}, dx) \\
&+ \frac{1}{2} \int_{x, y, \xi, \zeta \in X_{agg}} [\varphi(z, \xi + \zeta) - \varphi(z, x) - \varphi(z, y)] K_{coag}(z, \xi, \zeta) \\
&\quad \mathbb{P}(D_t(z, x, c_{pp}) = d\xi) \mathbb{P}(D_t(z, y, c_{pp}) = d\zeta) \\
&+ \frac{1}{2} \int_{x, y, \xi, \zeta \in X_{agg}} [\varphi(z, \xi) + \varphi(z, \zeta - y) - \varphi(z, x)] F(z, \xi, dy) \\
&\quad \mathbb{P}(D_t(z, x, c_{pp}) = d\xi) \lambda(z, t, dx) \\
&+ \int_{x, \xi \in X_{agg}} \varphi(z, \xi) r_{inflow}(z) \mathbb{P}(D_t(z - 1, x, c_{pp}) = d\xi) \lambda(z - 1, t, dx) \\
&- \int_{x, \xi \in X_{agg}} \varphi(z, \xi) \frac{1}{\tau(z)} \mathbb{P}(D_t(z, x, c_{pp}) = d\xi) \lambda(z, t, dx) \quad \forall z \in \mathbb{L},
\end{align*}
\]

and corresponding primary particle PBE can be written as

\[
\frac{d}{dt} c_{pp}(z, t) = I_{trans, pp}(z, t) - I_{nuc, pp}(z, t) - \int_{x \in X_{agg}} r_{layer}(z, t, x, c_{pp}) \lambda(z, t, dx). \tag{2}
\]

In this form, each integral on the RHS of Equation (1) represents an aggregate particle processes within the model. In order of appearance, these are: nucleation, collision (which may lead to coagulation), breakage, inflow and outflow (illustrated in Figure 2). The \( \varphi(\cdot) \) component of each integrand represents the particle transformation associated with the related mechanism. The remainder of the integrand defines the rate at which this process occurs. In the primary particle PBE (Equation (2)) the terms, in order of their appearance on the RHS, represent inflow/outflow processes, nucleation and layering (onto aggregates). Each of the terms in Equation (1) and (2) will be fully defined for the TSG model in the section to follow.
2.2 Aggregate particle processes

In the next section we describe the form of the various kernels and operators in the aggregate PBE (1) in the context of the twin-screw model. Details of the implementation of these mechanisms including the jump transforms are presented later in Section 3.

2.2.1 Nucleation

The first term in Equation (1) represents the nucleation (formation) of aggregate particles. The nucleation process involves the addition of a liquid droplet to the first compartment and rapid addition of primary particles to the droplet, producing a nucleus particle with form $x_{nuc} \in X_{agg}$.

As in [45], the nucleation model is formulated to resemble the process of a large liquid droplet (relative to the primary particle size) penetrating into a porous powder bed (immersion nucleation). Using this model, a nucleus incepted into compartment $z$ takes the form

$$x_{nuc}(z, c_{pp}, t) = \begin{cases} 
\left( \frac{v_{drop}}{\phi_{max}}, 0, \frac{v_{drop}}{s^*} \right), & \text{if } \frac{v_{drop}}{\phi_{max}} \leq c_{pp}(z) V_{real}(z) v_{pp}, \\
\left( c_{pp}(z) V_{real}(z) v_{pp}, v_{drop} - c_{pp}(z) V_{real}(z) v_{pp} \phi_{max}, c_{pp}(z) V_{real}(z) v_{pp} \phi_{max} \right), & \text{otherwise}.
\end{cases}$$

Here, $N_{pp}(z)$ is the number of primary particles in compartment $z$, $s^*$ is the pore liquid saturation limit, $\phi_{max}$ is maximum liquid saturation of solid material during nucleation and $V_{real}$ is the physical volume of the compartment occupied by the granular mass (including the bed voidage), which sets the scale of the ‘physical’ system. $v_{drop}$ is the number average volume of a droplet, which, in this study, takes the volume of a sphere with the
same diameter as the liquid addition nozzle $d_{\text{nozzle}}$. As in McGuire et al. [45], the liquid saturation limit is given as

$$\phi_{\text{max}} = \frac{(1 - \varepsilon_{\text{bed}})s^*}{\varepsilon_{\text{bed}}}.$$  \hspace{1cm} (3)

The two cases in (3) cover the situations in which:

i) there is sufficient primary particle mass in the droplet zone to permit formation of a complete nucleus particle

ii) there is insufficient primary particle mass in the droplet zone and a partially formed nucleus is created, which has a non-zero amount of external liquid

In the model, a single droplet creates a single nucleus particle, hence the nucleation rate $R_{\text{nuc}}(z, t)$ is equal to the rate of droplet addition $R_{\text{drop}}(z, t)$, which is defined as

$$R_{\text{drop}}(z, t) = \begin{cases} \frac{(\text{LSR})m_{\text{feed}}}{v_{\text{drop}}p}, & \text{if } z = 1, \\ 0, & \text{otherwise.} \end{cases}$$  \hspace{1cm} (4)

Here, LSR is the operating liquid-solid mass feed ratio to the twin-screw device and $m_{\text{feed}}$ is the operating mass feed rate.

In the context of the aggregate PBE (1), we may define

$$I_{\text{nuc}}(z, t, c_{\text{pp}}, dx) = \frac{R_{\text{nuc}}(z, t)\delta_{\text{nuc}}(z, c_{\text{pp}}, t)(x)dx}{V_{\text{real}}(z)},$$  \hspace{1cm} (5)

where $\delta_{\text{nuc}}(x)$ is to be understood as the Dirac delta function, centred on $x_i$.

### 2.2.2 Collision/coagulation

The second term in the PBE (1) represents binary collisions/coagulation between particles. The collisions fall into the following categories:

1. **Coagulating collision**: collision pair forms a new aggregate particle

   $$x_i, x_j \mapsto T_{\text{coag}}(x_i, x_j),$$  \hspace{1cm} (6)

   where, following the approach of Braumann et al. [10], the vector valued function $T_{\text{coag}}(x_i, x_j)$ is characterised as

   $$T_{\text{coag}}(x_i, x_j) = (T_{\text{coag}}(x_i, x_j)_{s}, T_{\text{coag}}(x_i, x_j)_{l}, T_{\text{coag}}(x_i, x_j)_{l'}, T_{\text{coag}}(x_i, x_j)_{p})^T$$  \hspace{1cm} (7)
with coordinate functions

\[ T_{\text{coag}}(x_i, x_j)_s = s_o(x_i) + s_o(x_j) \]  
\[ T_{\text{coag}}(x_i, x_j)_l = l_e(x_i) + l_e(x_j) - l_{e\to i}(x_i, x_j) \]  
\[ T_{\text{coag}}(x_i, x_j)_l = l_i(x_i) + l_i(x_j) + l_{e\to i}(x_i, x_j) \]  
\[ T_{\text{coag}}(x_i, x_j)_p = \frac{a_{\text{surf}}^+(x_i, x_j)^{3/2}}{6\pi^{1/2}} \]  
\[ - [s_o(x_i) + s_o(x_j) - l_e(x_i) - l_e(x_j) + l_{e\to i}(x_i, x_j)]. \]  

Here \( l_{e\to i}(x_i, x_j) \) represent the amount of surface liquid that is internalised due to the contact area between the colliding particle pair. This is computed as [10]

\[ l_{e\to i}(x_i, x_j) = \left\{ \begin{array}{l} l_e(x_i)l_e(x_j) \left[ 1 - \sqrt{\frac{1 - \left( \sqrt{v(x_i)} - l_e(x_i) \right)^2}{\sqrt{v(x_i)} + \sqrt{v(x_j)}}} \right] \\
\times \left[ 1 - \sqrt{\frac{1 - \left( \sqrt{v(x_j)} - l_e(x_j) \right)^2}{\sqrt{v(x_i)} + \sqrt{v(x_j)}}} \right] \right\}^{1/2}, \]  

and

\[ a_{\text{surf}}^+(x_i, x_j) = (1 - e_{\text{coag}}) \left[ a_{\text{surf}}(x_i)^{3/2} + a_{\text{surf}}(x_j)^{3/2} \right]^{2/3} + e_{\text{coag}} \left[ a_{\text{surf}}(x_i) + a_{\text{surf}}(x_j) \right], \]  

and \( e_{\text{coag}} \) is the coefficient of restitution of the granular material.

2. Non-coagulating collision: colliding pairs do not coalescence and the collision pair remain unchanged

\[ x_i, x_j \mapsto x_i, x_j. \]  

Upon collision of two aggregates, the collision is deemed to be successful (i.e. coagulation takes place) provided that the colliding particles meet the Stokes criterion as detailed in Braumann et al. [10], otherwise the collision is non-coagulating and the colliding particles remain unchanged. The Stokes criterion takes into account the material coefficient of restitution \( e_{\text{coag}} \) and the amount of surface liquid present, relative to the presumed height of asperities on the surface of the aggregates \( h_a \).

Particles collide according to the size-dependent collision kernel \( K_{\text{col}} \), which takes the form

\[ K_{\text{col}}(z, x_i, x_j) = k_{\text{col}}(z)n_{\text{screw}}C(d(x_i), d(x_j)) \]
Here, \( k_{\text{col}}(z) \) is the collision rate constant in compartment \( z \), \( n_{\text{screw}} \) is the screw speed, and \( C(d_1, d_2) \) is the collision rate function which defines the collision frequency between particles with diameter \( d_1 \) and \( d_2 \), respectively.

In this work we use the Equi-partition of kinetic energy collision rate function \([66]\), which takes the form

\[
C(d_i, d_j) = (d_i + d_j)^2 \sqrt{\frac{1}{d_i^3} + \frac{1}{d_j^3}}. \tag{17}
\]

Studies using the Discrete Element Method (DEM) \([17]\) have shown that the form of the collision function in (17) adequately describes the collision dynamics within batch granulation systems \([41]\). Furthermore, preliminary twin-screw simulations with this form of collision rate function showed that it promoted collisions between large agglomerates and primary particles over those between particles which were similar in size. Thus, in the absence of an existing twin-screw specific collision kernel, the kernel in Equation (17) was deemed acceptable for the purposes of aggregate collisions and layering (to be introduced in Section 2.2.5).

The coagulation kernel \( K_{\text{coag}} \) of PBE (1) is then

\[
K_{\text{coag}}(z, x_i, x_j) = \mathbb{I}_{\{x_i, x_j | \text{St}(x_i, x_j) \leq \text{St}^*_{\text{v}}(x_i, x_j)\}}(x_i, x_j), \tag{18}
\]

where \( \text{St}_v(x_i, x_j) \leq \text{St}^*_v(x_i, x_j) \) indicates that the Stokes criterion has been met and \( \mathbb{I}_A \) is to be understood as the indicator function of set \( A \).

### 2.2.3 Breakage

The third term in PBE (1) represents the aggregate breakage process. In this work, aggregates may undergo binary breakage according to the transform

\[
x \mapsto x - y, y \tag{19}
\]

at rate

\[
g_{\text{break}}(z, x) = \begin{cases} k_{\text{att}}(z)n_{\text{screw}}^2 \left( \frac{v(x)}{\hat{v}_{\text{break}}} \right)^{\omega_{\text{att}}(z)}, & \text{if } v(x) \geq v_{\text{min}}^\text{parent} \text{ and } l_e(x) + l_i(x) + p(x) \neq 0, \\ 0, & \text{otherwise}, \end{cases} \tag{20}
\]

where \( k_{\text{att}} \) is the attrition rate constant, \( v_{\text{min}}^\text{parent} \) is the minimum agglomerate size that can undergo breakage, \( \omega_{\text{att}} \) is the breakage rate exponent and \( \hat{v}_{\text{break}} \) is a normalisation parameter. In this work we set \( v_{\text{min}}^\text{parent} = v_{pp} \). The breakage of primary particles is not permitted in the current model.

In the context of the aggregate PBE (1), the breakage kernel takes the form \([40]\)

\[
F(z, x, dy) = \mathbb{I}_{\{x, y | m(y) < m(x)\}}(x)g_{\text{break}}(z, x)B_{\text{frag}}(z, x, dy), \tag{21}
\]

11
where $B_{\text{frag}}(z, x, dy)$ is the probability that the first particle formed from the breakage of particle $x$ (according to (20)) lies within the space $[y, y + dy]$. $B_{\text{frag}}(z, x, dy)$ is characterised by first considering the probability distribution of the volume of daughter particle $y$ (denoted $v_y$). In this work, we define

$$v_y(z, x, d\chi_{\text{frag}}) = v_{\text{parent}}^\text{min} + d\chi_{\text{frag}} \left[v(x) - v_{\text{parent}}^\text{min}\right],$$  \hspace{1cm} (22)

where, following Braumann et al. [10], $\chi_{\text{frag}}$ is a random measure with beta distribution $f(z, \chi_{\text{frag}})$ with skewness parameters $\alpha_{\text{daughter}}(z)$ and $\beta_{\text{daughter}}(z)$.

As in previous granulation works [10, 39], we assume that the composition of the abraded particles $y$ and $x - y$ are the same as $x$. Hence, $y$ can be defined as

$$dy = \frac{v_y(z, x, d\chi_{\text{frag}})}{v(x)} dx.$$  \hspace{1cm} (23)

As mentioned in Section 1, conveying elements have been observed to break particles through cutting/edge chipping. No daughter distribution breakage data is currently available for kneading elements. However, based on their somewhat similar geometry, it is expected that the primary breakage mechanism in these elements will be similar to that in so called distributive mixing elements (DME) [62]. In this work, it is hypothesised that the primary breakage mechanisms in kneading elements will be the crushing behaviour observed in DME, with less emphasis on the chipping mechanism, due to the absence of the pronounced blades that are present in DME. To incorporate this information into the model, breakage exponent and daughter distribution parameters differ between different types of compartment (or, equivalently, screw element). The probability density distribution for $\chi_{\text{frag}}$ used for each element in this study are illustrated in Figure 3.

![Figure 3: Element specific probability density functions for agglomerate breakage. In compartments that represent pure conveying zones $\alpha_{\text{daughter}} = \beta_{\text{daughter}} = 0.5$ and those representing pure kneading zones $\alpha_{\text{daughter}} = \beta_{\text{daughter}} = 2.$](image-url)
2.2.4 Aggregate transport

The fourth and fifth terms in the PBE (1) represent aggregate inflow and outflow processes on compartment \( z \), respectively. Agglomerates and primary particles are permitted to flow uni-directionally along the network of compartments from the feed zone to the exit zone. Each compartment is modelled as a perfectly stirred tank such that each particle flows out of compartment \( z \) with rate \( 1/\tau(z) \), where \( \tau(z) \) is the compartment residence time. Similarly, aggregate particles flow into compartment \( z \) from compartment \( z - 1 \) at rate \( r_{\text{inflow}} \) where

\[
r_{\text{inflow}}(z) = \begin{cases} 
1/\tau(z-1) & \text{if } z > 1 \\
0 & \text{otherwise.}
\end{cases}
\]  

(24)

2.2.5 Continuous/deferred aggregate processes

A number of particle mechanisms are modelled as continuous processes that act on the agglomerates (and indirectly on the primary particles) in the system according to the deferment function \( D_t : (L, X_{\text{agg}}, R^+) \rightarrow X_{\text{agg}} \) introduced in Section 2.1.

These processes are:

1. **Layering**
   
   Layering is the processes by which primary particles attach to the surface of the agglomerates. This process is modelled as a collision between agglomerates and primary particles using the same size-dependent collision kernel used for aggregate coagulation (Equation (15)).

   Layering is only permitted on agglomerates which have a volume of external liquid with height \( h_l \) exceeding the fixed height of asperities of the agglomerate particles \( h_a \). The height of the external liquid on particle \( x \) is defined as

   \[
h_l(x) = \frac{1}{2} \sqrt{\frac{6}{\pi}} \left[ \sqrt{v(x)} - \sqrt{v(x) - l_e(x)} \right].
\]  

(25)

In the current model, a successful layering event is supposed to ‘dry out’ the surface of the agglomerate particle. This drying is captured by the transformation of external liquid to internal liquid. For the addition of a single primary particle onto the surface of an agglomerate, the amount of liquid moved from the exterior to the interior is modelled as

\[
l_{e \rightarrow i}(x) = \min(v_{pp}, \pi d_{pp}^2 h_l(x)).
\]  

(26)

The form of (26) was constructed to cover the situations where the agglomerate surface is liquid rich and liquid poor. In the liquid rich case, a small primary may become fully immersed in the thick binder layer. In liquid poor situation, the primary particle is more likely to simply stick to the surface, hence the amount of internalisation is hypothesised to be controlled by the projected area of the primary.
It follows from the definitions above that the agglomerate particle $x$ undergoes layering with rate

$$ r_{layer}(z, t, c_{pp}, x) = \begin{cases} c_{pp}(z, t) K_{col}(z, d(x), d_{pp}), & \text{if } h_t(x) > h_a, \\ 0, & \text{otherwise}, \end{cases} \quad (27) $$

and the rates of change of each particle property induced by the layering process are

$$ \frac{d\varepsilon_0(x)}{dr}\bigg|_{layer} = r_{layer}(z, x, c_{pp}, t) v_{pp}, \quad (28) $$

$$ \frac{dl_e(x)}{dr}\bigg|_{layer} = -r_{layer}(z, x, c_{pp}, t) l_{e \rightarrow i}(x), \quad (29) $$

$$ \frac{dl_i(x)}{dr}\bigg|_{layer} = r_{layer}(z, x, c_{pp}, t) l_{e \rightarrow i}(x), \quad (30) $$

$$ \frac{dp(x)}{dr}\bigg|_{layer} = r_{layer}(z, x, c_{pp}, t) l_{e \rightarrow i}(x). \quad (31) $$

2. Consolidation

Consolidation of particles within the twin-screw system is presumed to occur primarily due to impacts between the particles and screw/walls of the barrel. The consolidation processes is modelled as the reduction in particle porosity and subsequent squeezing of the internal liquid to the surface. It is presumed that the rate of consolidation is dependent on the screw speed and the geometry of the screw element in which the process is taking place. Since the effect of screw speed is not being investigated in this study, a simple consolidation model is employed. This takes the form:

$$ \frac{d\varepsilon(x)}{dr}\bigg|_{consol} = -k_{comp}(z) n_{screw}[\varepsilon(x) - \varepsilon_{min}], \quad (32) $$

where $k_{comp}(z)$ is the compaction rate constant in compartment $z$ and $\varepsilon_{min}$ is the minimum porosity permitted for agglomerate particles.

The associated changes in the tracked particle properties due to consolidation are

$$ \frac{d\varepsilon_0(x)}{dr}\bigg|_{consol} = 0, \quad (33) $$

$$ \frac{dl_e(x)}{dr}\bigg|_{consol} = -l_i(x) \frac{dp(x)}{dr} / p(x), \quad (34) $$

$$ \frac{dl_i(x)}{dr}\bigg|_{consol} = -\frac{dl_e(x)}{dr}, \quad (35) $$

$$ \frac{dp(x)}{dr}\bigg|_{consol} = v(x) \frac{d\varepsilon(x)}{dr}. \quad (36) $$

Note that this has no effect on the primary particles since they have an $\varepsilon = 0$. 

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The aggregate deferment function $D_t$ maps a particle $x$ and primary particle concentration measure $c_{pp}(z)$ in $z$ at time $t_p$ to a particle $D_t(z, x, c_{pp})$ with time $t$ (where $t_p$ is the current time of particle $x$, $t_p \leq t$). This is done by evolving $x$ in time according to the rates

$$\frac{ds_o(x)}{dt}_{\text{defer}} = \frac{ds_o(x)}{dt}_{\text{layer}} + \frac{ds_o(x)}{dt}_{\text{consol}},$$

$$\frac{dl_c(x)}{dt}_{\text{defer}} = \frac{dl_c(x)}{dt}_{\text{layer}} + \frac{dl_c(x)}{dt}_{\text{consol}},$$

$$\frac{dl_i(x)}{dt}_{\text{defer}} = \frac{dl_i(x)}{dt}_{\text{layer}} + \frac{dl_i(x)}{dt}_{\text{consol}},$$

$$\frac{dp(x)}{dt}_{\text{defer}} = \frac{dp(x)}{dt}_{\text{layer}} + \frac{dp(x)}{dt}_{\text{consol}}.$$  

Hence, if

$$\frac{dx}{dt}_{\text{defer}} = \left( \frac{ds_o(x)}{dt}_{\text{defer}}, \frac{dl_c(x)}{dt}_{\text{defer}}, \frac{dl_i(x)}{dt}_{\text{defer}}, \frac{dp(x)}{dt}_{\text{defer}} \right)^T,$$

then

$$D_t(z, x, c_{pp}) = x + \int_{t_p}^{t} \frac{dx}{dt}_{\text{defer}} dt.$$  

### 2.3 Primary particle processes

In this section we describe each of the operators in the primary particle PBE (Equation (2)).

The first term in Equation (2) represents the transport operator (inflow and outflow) acting on the primary particles phase in compartment $z$. This is characterised by

$$I_{\text{trans,pp}}(z, t) = \begin{cases} \frac{m_{\text{feed}}}{v_{pp}\rho_s} \frac{N_{pp}(z)}{\tau(z)} \frac{1}{V_{\text{real}}(z)}, & \text{if } z = 1, \\ \left[ \frac{N_{pp}(z-1)}{\tau(z-1)} - N_{pp}(z) \right] \frac{1}{V_{\text{real}}(z)} & \text{otherwise.} \end{cases}$$

The second term in Equation (2) represents the nucleation sink term, which complements the aggregate nucleation model by taking the form

$$I_{\text{nuc,pp}}(t, z) = \begin{cases} \frac{m_{\text{feed}}\text{LSR}}{v_{pp}\rho_s\phi_{\text{max}} V_{\text{real}}(z)}, & \text{if } z = 1, \\ 0, & \text{otherwise.} \end{cases}$$

The final term in (2) accounts for the primary particles depletion through the deferred aggregate layering process (Equations (27)-(31)).
2.4 Compartmentalisation

In this paper, each compartment represents exactly one type of screw element. However, a screw element may be represented by more or less than one compartment (as demonstrated in Figure 4). As such, the total number of compartments employed may vary depending on the screw configuration being modelled. This allows increased resolution in areas where the changes in particle characteristics along the screw length are expected to be significant, such as the region around the liquid inception port. This is also the case in regions where the material flow is expected to more closely resemble a plug-flow, with limited back mixing, such as kneading elements.

As in existing twin-screw modelling efforts, we assume that the material undergoes very little change prior to the point of liquid inception (termed the “metering” zone), hence the screw configuration is modelled from the liquid inlet port on-ward (termed the “variation” zone). This liquid inception zone (droplet zone) in modelled as a conveying compartment with length 0.33D (where D is the diameter of the screw) which serves as the first compartment in all screw configurations investigated. An example of the compartmentalisation of a screw configuration used in this study is presented in Figure 4.

![Figure 4: Example screw configuration (variation zone only) and the associated compartmental representation. The number in the centre of each compartment represents the compartment length (normalised by the screw diameter). Compartments representing conveying zones are in blue and compartments representing kneading zones are shown in orange.](image)

2.5 Residence time estimation

The compartment residence times $\tau(z)$ that control the aggregate and primary particle flow rates are dependent on the compartmentalisation of the screw and the operating conditions. In previous TSG modelling studies, compartmental residence times have been estimated through the use of DEM [1] and chemical imaging techniques [5, 33, 34]. In the case where DEM has been used, the PBM and DEM solvers are coupled, allowing collision statistics [4] and residence time estimation to be made, however, the significant computational cost of the DEM step negatively impacts the overall time required to solve the model. Hence, it is advantageous to have the ability to estimate the residence time of individual compartments simply from the screw configuration and device operation (mass and liquid flow rates). It is worth noting that, in studies of alternative granulation devices,
such as Barrasso et al. [6], the costly DEM step has been replaced by an Artificial Neural Net (ANN) with promising results, though a sizeable amount of DEM data must be gathered to train such networks.

In order to create a framework for the prediction of residence times, information from a number of existing twin-screw material flow studies must be considered. A number of experimental studies have examined the mass distribution over the twin-screw system, through the use of Positron Emission Particle Tracking (PEPT) [42, 64]. Recently, it has also been proposed [26] that barrel fill level be used as way to define the design space in TSG. This has been motivated by the observation that very similar PSDs can be obtained for very different screw speed and mass feed rates, provided the fill level is the same across these experiments. From a screw element perspective, it is noted that, as a result of their geometry, kneading elements do not have any significant conveying capacity and thus they will generally be completely/almost completely filled with material at steady state. In contrast, conveying elements generally have a much lower fill level that is dependent on the operating speed of the screw [64] and the feed material [61]. This non-uniformity in the mass distribution ultimately leads to varying residence times across different sections of the screw, which must be captured within a model, if model rates are to be applied to various screw configurations. Attempts have been made to predict such mass distribution in the context of twin-screw extrusion systems [21]. Though the resulting models showed good performance, the flowing material consisted of a polymer melt, which has quite different flow characteristics compared to the partially wet powder mass in the TSG system. This limits its applicability to TSG.

In order to be able to produce reasonable estimates of particle residence times of screw sections/model compartments for arbitrary screw configurations, a novel but simplistic approach was developed for use in this study. The approach involves two key stages: firstly the mean residence time of the complete device $\tau_{\text{screw}}$ is linearly interpolated from existing experimental data based on the screw configuration, screw speed and mass feed rate. In the second stage, this total device residence time is distributed over the compartmental network by estimating the mass distribution profile across the compartments. In this work we use data from the residence time analysis experiments performed by Kumar et al. [33]. In that study, the authors used a chemical imaging approach where a dye-impulse was introduced into the powder feed zone in order to extract residence time distributions. These residence time estimates consider the full screw (i.e. both metering and variation zones) and the metering zone consisted purely of conveying elements.

The total mean screw residence time is split into the contribution from the metering zone and the variation zone as

$$\tau_{\text{screw}} \approx \tau_{\text{metering}} + \tau_{\text{variation}}. \quad (45)$$

Assuming that each section behaves as perfect plug-flow with no back-mixing then,

$$\tau_{\text{screw}} \approx \frac{M_{\text{metering}}}{m_{\text{feed}}} + \frac{M_{\text{variation}}}{m_{\text{feed}}(1 + \text{LSR})}, \quad (46)$$

where $M_{\text{metering}}$ and $M_{\text{variation}}$ are dynamic mass hold-up (i.e. the mass of material that is not ‘stuck’ to the barrel wall/screws) within the metering and variation zones, respectively.
These are given as

\[ M_{\text{metering}} = \sum_{i=1}^{N_{\text{m,convey}}} L_{\text{m,convey}}(i) v_{\text{convey}} e_{\text{bed}} \rho_s f_{\text{m,convey}}(i), \]  

(47)

\[ M_{\text{variation}} = \sum_{i=1}^{N_{\text{v,convey}}} L_{\text{v,convey}}(i) v_{\text{convey}} e_{\text{bed}} \rho_{\text{eff}} f_{\text{v,convey}}(i) \]

\[ + \sum_{i=1}^{N_{\text{knead}}} L_{\text{v,knead}}(i) v_{\text{knead}} e_{\text{bed}} \]  

(48)

where the subscripts \(m\) and \(v\) refer to the metering and variation zones respectively and

- \(N_{j,\text{convey}}\) is the number of conveying elements in zone \(j\);
- \(N_{j,\text{knead}}\) is the number of kneading elements in zone \(j\);
- \(L_{j,e}(i)\) is the length of the \(i\)th screw section consisting of element type \(e\) in zone \(j\) (measured in screw diameters \(D\));
- \(v_e\) is the specific volume available in a screw section with element type \(e\) (measured in unit volume/length in screw diameters \(D\));
- \(f_{j,e}(i)\) is the volumetric fill fraction of the \(i\)th screw section consisting of element type \(e\) in zone \(j\);
- \(\rho_{\text{eff}}\) is the effective density of the solid material being held up in the variation zone;

For simplicity we assume \(\rho_{\text{eff}}\) is constant along the length of the variation zone. The effective density is computed as the weighted averaged density of the solid and liquid feed such that:

\[ \rho_{\text{eff}} = \frac{(LSR + 1)\rho_s \rho_l}{\rho_l + \rho_s LSR}. \]  

(49)

Again, \(\rho_s\) is the density of solid feed material and \(\rho_l\) is the density of the liquid binder.

We impose that all kneading elements are filled to capacity with material (i.e. \(f_{v,knead}(i) = f_{v,knead} = 1 \forall i \in \{1, \ldots, N_{v,knead}\}\)). Though PEPT studies [64] have shown that conveying elements that precedes a kneading element will generally have a higher fill fraction, for the purposes of the residence time estimation in this paper we assume that each compartment of element type \(e\) has the same fill fraction, such that \(f_{m,\text{convey}}(i) = f_{v,\text{convey}}(k) = f_{\text{convey}} \forall i \in \{1, \ldots, N_{m,\text{convey}}\}, k \in \{1, \ldots, N_{v,\text{convey}}\}\). This assumption permits \(f_{\text{convey}}\) to be solved by substitution of (47) and (48) into (46) and re-arranging to yield

\[ f_{\text{convey}} = \frac{\tau_{\text{screw}} h_{\text{feed}} (1 + LSR) - f_{v,knead} e_{\text{bed}} v_{\text{knead}} \sum_{i=1}^{N_{v,knead}} L_{v,knead}(i)}{e_{\text{bed}} v_{\text{convey}} \left[ \rho_l (1 + LSR) \sum_{i=1}^{N_{m,\text{convey}}} L_{m,\text{convey}}(i) + \rho_{\text{eff}} \sum_{i=1}^{N_{v,\text{convey}}} L_{v,\text{convey}}(i) \right]} \]  

(50)

Assuming that kneading elements are filled to capacity, knowledge of \(f_{\text{convey}}\) specifies the mass distribution across the full screw.
Given that compartment $z$ represents a section of the screw composed of elements of type $e$, with length $L(z)$ and steady state mass hold-up $M(z)$ then the compartment residence time is approximated as

$$\tau(z) := \frac{M(z)}{\dot{m}_{\text{feed}}} = \frac{L(z)v_e \epsilon_{\text{bed}} \rho_{\text{eff}} f_{v,e}}{\dot{m}_{\text{feed}}}.$$ \hspace{1cm} (51)

Since no residence time data was presented for pure conveying screws in Kumar et al. [33], residence times for such screw configurations was estimated using the derived steady state fill fraction predicted using the above flow model applied to a configuration with two kneading blocks (13.3%).

Derived fill fractions for conveying elements ranged from 10 - 13.3%, and overall derived fill levels between 13.3 - 24.1% with $\epsilon_{\text{bed}} = 0.3$, depending on the screw configuration. This gives the fraction of screw bed occupied by material (i.e. discounting the packing fraction) as 4-7.23%. These values are lower than the ranges observed in alternative twin-screw devices (10-30%) [47], however, the method presented in this paper does not account for non-dynamic mass (i.e. material that is stuck to the walls), which could be significant as demonstrated in other experimental studies [61].

Given a fill fraction for each element type, the real volume of material (including the bed voidage) modelled in each compartment $V_{\text{real}}(z)$ is computed as

$$V_{\text{real}}(z) = L(z)v_e f_{v,e}.$$ \hspace{1cm} (53)

### 3 Numerical treatment

#### 3.1 Stochastic particle methods for twin-screw granulation

The aggregate phase of the twin-screw population balance model is solved using the stochastic weighted algorithm (SWA) [54, 56, 59, 60]. The SWA has been successfully employed to solve population balance problems in the fields of granulation [41], soot formation, aerosol and nano-particle synthesis [7, 49, 50] and more general coagulation/fragmentation processes [73]. In the SWA, each compartment $z$ is simulated with a discrete list of computational particles

$$(z, x_i, w_i), \quad i = 1, \ldots, N_{\text{agg}}(z, t),$$ \hspace{1cm} (54)

which describes the population dynamics in (1). In (54), $x \in X_{\text{agg}}, w_i \in (0, w_{\text{max}}]$ is the statistical weight of the particle with index $i$ and $N_{\text{agg}}(z, t)$ is the total number of aggregate stochastic particles in compartment $z$ at time $t$. $w$ can be thought of as indicator of the number of physical particles that are represented by computational particle $(z, x, w)$. Each simulated compartment has an associated scaling parameter or sample volume denoted $V_{\text{sample}}(z, t)$ and the measure valued solution $\lambda$ to the population balance equation (1) is
approximated by the stochastic particle systems such that, for all $\phi$ ins a large class of test functions

$$
\frac{1}{V_{\text{samp}}(z,t)} \sum_{i=1}^{N_{\text{agg}}(z,t)} w_i \phi(z,x_i) \xrightarrow{\text{V_{samp}(z,0) \rightarrow \infty}} \int_{X_{\text{agg}}} \phi(z,x) \lambda(t,z, dx)
$$

and the initial sample volume (i.e. that at $t = 0$) is characterised by

$$
\frac{1}{V_{\text{samp}}(z,0)} \sum_{i=1}^{N_{\text{agg}}(z,0)} w_i \approx \int_{X_{\text{agg}}} \lambda(t,z, dx). \quad (56)
$$

The system of stochastic particles is evolved in time through a Markov jump process. This process is characterised by the possible jumps and their associated rates. Each available jump and the associated rate is a function of the state of the system at that point in time. At each $t$ there exists a list of possible jumps which have independent, exponentially distributed waiting times. The waiting time between any two jumps $\Delta t_{\text{wait}}$ is described by the distribution \cite{24}:

$$
P(t, \Delta t_{\text{wait}} \geq \theta) = \exp(-R_{\text{SWA total}}(t) \theta), \quad \theta \geq 0, \quad (57)
$$

where $R_{\text{SWA total}}$ is the total jump rate, which has the form

$$
R_{\text{SWA total}}(z,t) = R_{\text{SWA nuc}}(z,t) + R_{\text{SWA break}}(z,t) + R_{\text{SWA trans}}(z,t) + R_{\text{SWA coag}}(z,t). \quad (58)
$$

Here, $R_{\text{SWA nuc}}(z,t)$, $R_{\text{SWA break}}(z,t)$, $R_{\text{SWA trans}}(z,t)$ and $R_{\text{SWA coag}}(z,t)$ are the individual jump rates for nucleation, breakage, transport and coagulation, respectively, which are detailed in Section 3.3.

Upon selection of the waiting time, jump process $p$ is carried out with probability

$$
\frac{R_p(z,t)}{R_{\text{SWA total}}(z,t)} \quad (59)
$$

and the system moves forward in time. This process is continued until the stopping condition $t \geq t_{\text{stop}}$ is satisfied.

### 3.2 Splitting scheme

Due to the small size of primary particles relative to large agglomerates within twin-screw devices, the number concentration of these species may differ by several orders of magnitude. In such situations it is unfeasible to solve the primary particle part of the twin-screw population balance problem using a stochastic particle method (unless the primary particle size is significant relative to the mean aggregate size, or in cases where the physical collision rates to be simulated are relatively low). This is due to the fact that the collision jump rates required for significant transfer of mass between the two species becomes too computationally intensive to simulate within reasonable time-scales. For this reason, the primary particle population balance equation (2) is solved using an implicit ODE solution technique (discussed further in Section 3.4). The use of both stochastic particle and ODE
methods to solve different parts of the coupled population balance equations (1) and (2) closely follows the operator splitting technique presented by Celnik et al. [16]. The nature and implementation of the splitting are discussed further in the following sections.

Due to the interaction between individual stochastic particles and the primary particle phase, it is often more convenient to work in terms of the absolute number of primary particles within a given sample volume (as opposed to the number concentration.) Hence, for the remainder of the paper we will work in terms of primary particle number and 

\[ N_{pp}(z,t) \]

is to be understood as the number of primary particles that exist within the sample volume \( V_{samp}(z,t) \) corresponding to physical compartment \( z \) at time \( t \).

### 3.3 Jump processes

The stochastic jump processes associated with the twin-screw population balance model are presented in this section. Since all of the weight transfer functions employed within these jump processes have been shown to converge to the appropriate forms in the un-weighted aggregate PBE (2), the associated weighted population balance equation is not presented here. For details of the various convergence proofs the reader is directed to works [40, 41, 53, 56].

#### 3.3.1 Nucleation

In this work, the nucleation jump involves the inception of particles of the form

\[ (z,x_nuc,w_{nuc}) \]

at rate

\[
R_{nuc}^{SWA}(z,t) = \begin{cases} 
R_{drop}(z,t)V_{samp}(z,t) & \text{if } z = 1, \\
0 & \text{otherwise},
\end{cases}
\]

(61)

where \( w_{nuc} \) is the statistical weight of the nucleus particle to be added. Since the nucleation is the only jump process in this work which increases the number of computational particles in the droplet zone (\( z = 1 \)) and transport is the only process which reduces the number of stochastic particle in this zone, the number of particle in \( z = 1 \) can be held approximately constant by enforcing

\[
R_{nuc}^{SWA}(z,t) \approx R_{trans}^{SWA}(z,t),
\]

(62)

where \( R_{trans}^{SWA}(z,t) \) is the transport jump rate in compartment \( z \) (detailed in Section 3.3.4). From (4), (61) and (62) it follows that

\[
w_{nuc}(z,t) = \frac{(LSR)\dot{m}_{\text{feed}}V_{samp}(z,t)}{v_{drop}\rho_{l}V_{real}(z)R_{trans}^{SWA}(z,t)}.
\]

(63)
### 3.3.2 Coagulation

Aggregate collision jumps take the following number-conserving form

**Coagulating collision:**

\[
(z, x_i, w_i), (z, x_j, w_j) \mapsto (z, T_{\text{coag}}(x_i, x_j), \gamma_{\text{coag}}(x_i, w_i, x_j, w_j), (z, x_j, w_j)), \tag{64}
\]

**Non-coagulating collision:**

\[
(z, x_i, w_i), (z, x_j, w_j) \mapsto (z, x_i, w_i), (z, x_j, w_j), \tag{65}
\]

where \(\gamma_{\text{coag}}\) is the coagulation weight transfer function. In this work we employ a mass-conserving form of \(\gamma_{\text{coag}}\) whose convergence has been demonstrated in [53, 56] and has been utilized in previous SWA studies [40, 41]. This has the form

\[
\gamma_{\text{coag}}(x_i, w_i, x_j, w_j) = w_i \left( \frac{m(x_i)}{m(x_i) + m(x_j)} \right). \tag{66}
\]

The SWA collision kernel associated with the weight transfer function in (66) is [56]

\[
K_{\text{col}}^{\text{SWA}}(z, x_i, w_i, x_j, w_j) = K_{\text{col}}(z, x_i, x_j) w_j \tag{67}
\]

\[
= k_{\text{col}}(z) n_{\text{screw}}(d_i + d_j) \sqrt{\frac{1}{d_i^3} + \frac{1}{d_j^3}} w_j, \tag{68}
\]

and the SWA coagulation kernel is

\[
K_{\text{coag}}^{\text{SWA}}(z, x_i, w_i, x_j, w_j) = K_{\text{col}}^{\text{SWA}}(z, x_i, w_i, x_j, w_j) \mathbb{1}_{\{x_i, x_j | St(v(x_i), x_j) \leq St(v(x_i), x_j)\}}(x_i, x_j). \tag{69}
\]

Due to the complex form of (68), the repeated evaluation of the associated total collision rate in each compartment is very computationally intensive, since it requires looping through each pair of aggregates within the ensemble. For this reason, a majorant kernel is employed. An in-depth treatment of majorant techniques and their application to the solution of population balance problems can be found in [19, 25, 28, 49, 56]. The majorant form of (68) used is

\[
\hat{K}_{\text{col}}^{\text{SWA}}(z, x_i, w_i, x_j, w_j) = k_{\text{maj}} k_{\text{col}}(z) n_{\text{screw}}(d_i^2 + d_j^2) \left( \frac{1}{d_i^{1.5}} + \frac{1}{d_j^{1.5}} \right) w_j, \tag{70}
\]

where \(k_{\text{maj}}\) is the majorant scaling factor. As in Lee et al. [41], \(k_{\text{maj}}\) was set to 1.42 in order to satisfy the inequality \(K_{\text{col}}^{\text{SWA}} < \hat{K}_{\text{col}}^{\text{SWA}}\) throughout the simulation.

The majorant collision jump rate in compartment \(z\) at time \(t\) is (adapted to the current
model from [41])

\[
R_{\text{SWA, col}}(z, t) = \frac{1}{V_{\text{samp}}(z, t)} \sum_{i \neq j} N_{\text{agg}}(z, t) \left[ \sum_{i=1}^{N_{\text{agg}}(z, t)} d_i^{0.5} \sum_{j=1}^{N_{\text{agg}}(z, t)} w_j - \sum_{i=1}^{N_{\text{agg}}(z, t)} d_i^{0.5} w_i \right]
+ \left[ \sum_{i=1}^{N_{\text{agg}}(z, t)} d_i 1 - \sum_{i=1}^{N_{\text{agg}}(z, t)} d_i^{0.5} w_i \right]
+ \left[ \sum_{i=1}^{N_{\text{agg}}(z, t)} d_i^{1.5} \sum_{j=1}^{N_{\text{agg}}(z, t)} d_j^{0.5} w_j - \sum_{i=1}^{N_{\text{agg}}(z, t)} d_i^{0.5} w_i \right]
+ \left[ \sum_{i=1}^{N_{\text{agg}}(z, t)} 1 - \sum_{j=1}^{N_{\text{agg}}(z, t)} d_j^{0.5} w_j - \sum_{i=1}^{N_{\text{agg}}(z, t)} d_i^{0.5} w_i \right]
\]
\(\text{(71)}\)

Using the majorant rate expression in (71), jumps are accepted with probability

\[
\frac{K_{\text{SWA, col}}(z, x_i, w_i, x_j, w_j)}{\hat{K}_{\text{SWA, col}}(z, x_i, w_i, x_j, w_j)},
\]
\(\text{(72)}\)

otherwise the jump is fictitious, in which case the ensemble remains unchanged and time moves forward according to (57). Again, the collision pair must satisfy the Stokes criterion in order for the jump to proceed, otherwise the collision is non-coagulating and the system remains unchanged, since the jump (65) has no effect on the system state.

For a full derivation of the majorant rate expression in (71) and the associated particle selection measures, the reader is referred to Lee et al. [41].

### 3.3.3 Breakage

In this paper, aggregate particles undergo breakage according to the jump [40]

\[
(z, x_i, w_i) \mapsto (z, y, \gamma_{\text{frag}}(x_i, w_i, y)),
\]
\(\text{(73)}\)

which occurs at rate

\[
g_{\text{break}}(z, x).
\]
\(\text{(74)}\)

Here, \(\gamma_{\text{frag}}\) is the breakage weight transfer function which takes the form [40]

\[
\gamma_{\text{frag}}(x_i, w_i, y) = w_i \frac{m(x_i)}{m(y)}.
\]
\(\text{(75)}\)

As in [40, 41], convergence is achieved by selecting \(y\) according to the following probability measures:

\[
P(y = x_j) = \frac{m(x_j)}{m(x_i)},
\]
\(\text{(76)}\)

\[
P(y = x_i - x_j) = 1 - \frac{m(x_j)}{m(x_i)}.
\]
\(\text{(77)}\)
The volume of $x_j$ is selected according to (22) and its composition is that of particle $x_i$.

The total breakage jump rate in compartment $z$ is

$$R_{\text{break}}^{\text{SWA}}(z,t) = \sum_{i=1}^{N_{\text{agg}}(z,t)} g_{\text{break}}(z,x_i).$$

(78)

### 3.3.4 Transport

Aggregate particle inflow and outflow processes are grouped and carried out under an overarching ‘transport’ jump process. The transport jump takes the following spatially-dependent forms

1. **Droplet zone** ($z = 1$)
   
   Due to the absence of aggregate inflow in to the first reactor, the transport jump only captures outflow through the deletion of a stochastic particle as
   
   $$(z,x,w) \mapsto \text{deleted}.$$  
   (79)

2. **Non-droplet zone** ($z > 1$)
   
   $$(z,x_i,w_i) \mapsto \text{deleted},$$  
   (80)
   $$(z-1,x_j,w_j) \mapsto (z,x_j,F_c(z,t)w_j).$$  
   (81)

Here, $F_c$ is the transport scaling factor required to maintain continuity, which takes the form

$$F_c(z,t) = \frac{V_{\text{samp}}(z,t)N_{\text{agg}}(z-1,t)}{V_{\text{samp}}(z-1,t)N_{\text{agg}}(z,t)}.$$  
(82)

A complete derivation of (82) is provided in Appendix A.

In all zones, the transport jump is carried out with total rate

$$R_{\text{trans}}^{\text{SWA}}(z,t) = \frac{N_{\text{agg}}(z,t)}{\tau(z)}.$$  
(83)

### 3.4 Continuous Processes

Primary particle transport, primary particle depletion through nucleation, layering of primary particles onto stochastic particles (aggregates) and aggregate consolidation are carried out as continuous processes using the Linear Process Deferment Algorithm (LPDA) [55]. LPDA can be thought of as an operator splitting technique with a “just in time” feature. LPDA has been employed in the simulation of stochastic population balance equations for granulation [40], silica nanoparticle synthesis [50] and soot particle formation [55, 56]. As part of the LPDA, each computational particle is tagged with the time $t_p$ that it has been simulated to. In the LPDA, deferred processes are carried out at two specific stages during the simulation:
1. **Local Application**

Here, stochastic particles that are selected to take part in one of the jump processes described in Section 3.3 are brought up to the current simulation time $t$ by applying the aggregate deferment function $D$ to each of the selected particles in turn (see Section 2.2.5). This is done just before the jump transforms are applied (i.e. “just in time”).

2. **Global Application**

In these instances, the continuous processes are periodically applied to the full ensemble of stochastic particles between jump processes. This global deferment is carried out with rate $1/\Delta_{\text{defer}}$, where $\Delta_{\text{defer}}$ is the deferment time step. The global deferment procedure is described in detail in Algorithm 1. Primary particle transport processes are also carried out after each global deferment step (this is discussed further in 3.4.2). The global deferment step ensures that the time over which a particle is integrated is small, such that any linearisations have minimal effect on the solution. This also ensures that the computed jump rates remain close to that of the ‘true’ undeferred system.

### 3.4.1 Layering as a deferred process

Since layering is not strictly a linear process (the layering rate for each stochastic aggregate is a function of primary particle concentration), application of the LPDA to the layering processes described in (28)-(31) requires linearisation of the these equations. This is done by assuming that $N_{\text{pp}}$ is approximately constant over the deferment time step. In the context of local deferment, this requires one to hold $N_{\text{pp}}(z)$ constant while all selected particles are integrated to the target time. In the context of global deferment steps, the assumption requires one to hold the $N_{\text{pp}}(z)$ constant across each integration interval $\Delta_{\text{defer}}$ (i.e. all particles are integrated using the same value of $N_{\text{pp}}(z)$ for each global deferment step and $N_{\text{pp}}(z)$ is updated between deferment steps). Such assumptions have been employed using LPDA to describe the surface growth of stochastic particles within the context of soot models [16]. Preliminary simulations with the twin-screw model confirmed that this assumption was appropriate, provided that a small enough deferment time step $\Delta_{\text{defer}}$ was enforced.

### 3.4.2 Primary particle update

As mentioned in the previous section, $N_{\text{pp}}$ is assumed constant over the course of the deferment step and only updated at the end of each deferment step. These updates take different forms depending on the context in which the deferment is being applied. Let us denote the time at the end of the deferment step as $t_{\text{target}}$, then the possible forms of primary particle update are
1. Local Application

\[ N_{pp}(z, t_{\text{target}}) \leftarrow N_{pp}(z, t) - \frac{1}{v_{pp}} \sum_{i=1}^{N_{\text{jump}}} \left[ s_o(x_i)_{l_{\text{target}}} - s_o(x_i)_{l} \right] w_i, \]  

(84)

where \( N_{p,\text{jump}} \) is the number of particle involved in the selected jump process and the second term on the RHS represents the linearised SW A form of the final (layering) term in (2), taken over only the particle(s) involved in the local deferment. The derivation of this term is provided in Appendix B.

2. Global Application

\[ N_{pp}(z, t_{\text{target}}) \leftarrow N_{pp}(z, t) - \frac{1}{v_{pp}} \sum_{i=1}^{N_{\text{agg}}(z)} \left[ s_o(x_i)_{l_{\text{target}}} - s_o(x_i)_{l} \right] w_i + (t - t_{\text{target}}) \left[ \frac{\sum_{i=1}^{N_{pp}(z,t)}}{v_{pp}} \sum_{i=1}^{N_{\text{agg}}(z)} \left[ s_o(x_i)_{l_{\text{target}}} - s_o(x_i)_{l} \right] w_i \right] + (t - t_{\text{target}}) \left[ \sum_{i=1}^{N_{pp}(z,t)} \frac{\sum_{i=1}^{N_{\text{agg}}(z)} \left[ s_o(x_i)_{l_{\text{target}}} - s_o(x_i)_{l} \right] w_i}{v_{pp}} \right] \]  

\[ + (t - t_{\text{target}}) \left[ \frac{\sum_{i=1}^{N_{pp}(z,t)}}{v_{pp}} \sum_{i=1}^{N_{\text{agg}}(z)} \left[ s_o(x_i)_{l_{\text{target}}} - s_o(x_i)_{l} \right] w_i \right] \]  

(85)

The second term on the RHS of (85) is the same as the local case (84), except that the layering term has been integrated across the full ensemble of stochastic particles in compartment \( z \). The third term on the RHS of (85) represents the integral of the linearised primary particle transport rate (Equation (43)), which, applied to the network of sample volumes, takes the form

\[ \frac{dN_{pp}(z,t)}{dt} \bigg|_{\text{transport}} = \begin{cases} \frac{\bar{m}_{\text{feed}}V_{\text{sample}}(z,t)}{V_{\text{sample}}(z)\rho_{pp}v_{pp}} - \frac{N_{pp}(z,t)}{\tau(z)}, & \text{if } z = 1, \\ \frac{V_{\text{real}}(z-1)}{V_{\text{real}}(z)} \frac{V_{\text{sample}}(z)}{V_{\text{sample}}(z-1)} \frac{N_{pp}(z-1,t)}{\tau(z-1)} - \frac{N_{pp}(z,t)}{\tau(z)}, & \text{otherwise.} \end{cases} \]  

(86)

A full derivation of (86) is provided in Appendix C.

Finally, the last term in (85) is the integral of the nucleation depletion term (43), scaled according to the sample volume.

3.5 Solver implementation

In this paper, we are primarily interested in the steady state solution of the twin-screw population balance problem described by Equations (1)-(2). As such, this permits us to solve the compartment network in a sequential manner, starting from \( z = 1 \) and moving along the network in a linear fashion until the final reactor with \( z = \max L \) is reached. This dramatically reduces the computational effort required to achieve converged solutions, to the extent that large-scale parameter estimation procedures become possible. Since the
networks studied here are strictly linear (i.e. no back-flow or recycle loops) a single pass of
the network is sufficient to fully converge the compartmental network. Algorithm 2
is a simplified account of the sequential solver algorithm used to solve the twin-screw
population balance problem in this study. Preliminary twin-screw simulations using the
model presented in this paper and previous twin-screw SWA works [45, 46] showed that, if

Algorithm 1: The global deferment algorithm used to carry out continuous pro-
cesses, applied to compartment $z$.

\begin{algorithm}
\STATE START
\STATE \textbf{if} $t == 0$ \textbf{then}
\STATE \hspace{1em} Set $t_{\text{next-defer}} \leftarrow \Delta t_{\text{defer}}$.
\STATE \textbf{while} $t_{\text{next-defer}} < t$ \textbf{do}
\STATE \hspace{2em} \textit{Integrate all aggregate particles to required time in stages.}
\STATE \hspace{2em} Set $t_{\text{target}} \leftarrow \min \{t_{\text{next-defer}}, t\}$. 
\STATE \hspace{2em} \textbf{for} $i = 1, N_{\text{agg}}(z,t)$ \textbf{do}
\STATE \hspace{3em} Numerically integrate particle $(x_i, w_i)$ to $t_{\text{target}}$ according to Equation (42).
\STATE \hspace{2em} Update $N_{\text{pp}}(z)$ to $t_{\text{target}}$ according to (85).
\STATE \hspace{2em} Set $t_{\text{next-defer}} \leftarrow t_{\text{next-defer}} + \Delta t_{\text{defer}}$.
\STATE STOP
\end{algorithm}

Algorithm 2: The SWA sequential solver algorithm for a single realisation.

\begin{algorithm}
\STATE START
\STATE Set $z = 1$.
\STATE \textbf{while} $z \leq \max \mathbb{L}$ \textbf{do}
\STATE \hspace{1em} Set $t \leftarrow 0$.
\STATE \hspace{2em} Compute $t_{\text{stop}}(z)$ from (87).
\STATE \hspace{2em} Initialise particle ensemble and $N_{\text{pp}}(z,0)$.
\STATE \hspace{2em} \textbf{while} $t < t_{\text{stop}}(z)$ \textbf{do}
\STATE \hspace{3em} Apply global deferment Algorithm (1).
\STATE \hspace{3em} Compute $R_{\text{SWA}}^{\text{total}}(z,t)$ according to (58).
\STATE \hspace{3em} Compute $\Delta t_{\text{wait}}$ according to (57).
\STATE \hspace{3em} Set $t \leftarrow t + \Delta t_{\text{wait}}$.
\STATE \hspace{3em} Select jump process according to (59).
\STATE \hspace{3em} Apply local deferment transformation $D_t$ to the selected
\STATE \hspace{3em} particle(s).
\STATE \hspace{3em} Update the particle ensemble according to the selected jump
\STATE \hspace{3em} process.
\STATE \hspace{2em} Set $z \leftarrow z + 1$.
\STATE STOP
\end{algorithm}
the initial state of each compartment (i.e. at $t = 0$) has an overall particle concentration that is sufficiently close to the solution of the population balance problem, then the simulation time required to reach steady state is determined by the residence time of the compartment in question. In this paper we define the steady state solution to the population balance problem as the state of the ensemble which does not exhibit any dynamic drift in the key measures (particle moments, overall mass density, mean porosity, liquid content, etc.). Thus each compartment is solved to time

$$t_{\text{stop}}(z) = n_\tau \tau(z),$$

where $n_\tau \in \mathbb{R}^+$. The choice $n_\tau = 8$ was observed to yield simulation stop times that were sufficiently long enough to reach steady state. Since the reactors are solved in sequence, the transport terms that have a dependence on the previous reactor (such as $N_{\text{agg}}(z-1, t)$) should be replaced with the equivalent steady state value (in this example $N_{\text{agg}}(z-1, t_{\text{stop}}(z-1))$).

Algorithm 2 is applied numerous times (each time with a different seed to the random number generator) in order to generate multiple independent measures of the steady state solution to the twin-screw population balance problem. From here on, each repetition of the algorithm for a fixed set of operating conditions will be referred to as a realisation.

The first compartment is initialised such that 50% of the steady state mass hold-up is allocated to aggregate phase in the form of particles with $x = (v_{\text{drop}}, 0, 0, 0), w = 1$. The remaining mass specifies $N_{pp}(1, 0)$. In each realisation, the first compartment is initialised with $N_{pp}(1, 0) = 0.75N_{\text{agg}}^{\text{max}}$, where $N_{\text{agg}}^{\text{max}}$ is the maximum number of computational particles permitted (per compartment). In all subsequent compartments, the particle ensemble and $N_{pp}(z, 0)$ are initialised with the final state of particle ensemble and $N_{pp}$ in the previous compartment ($z-1$). The particle doubling and reduction procedures described in Lee et al. [40] are employed to control the number of computational particles in each compartment. Note that this is only important in the case of $z = 1$, since it is the only compartment where inflow and outflow processes are not directly coupled (the coagulation, breakage and non-droplet zone transport jump processes presented in Section 3.3 are all constant number).

4 Application

4.1 Twin-screw operating conditions

The predictive abilities of the twin-screw model are assessed using a two-step process. Firstly, unknown model parameters are estimated using data from selected experiments carried out by Vercruysse et al. [68]. In the second stage, these parameter estimates are used in the simulation of additional experiments by Vercruysse et al. [68], that were not featured in the estimation stage. In the experimental work used, the authors investigated the effect of the screw configuration on the product particle size distribution (PSD) using a ConsiGma\textsuperscript{TM} twin-screw granulator (length-to-diameter ratio of 20:1) with $\alpha$-lactose mono-hydrate as the feed material and distilled water as the binding liquid. The authors
tested a large number of screw configurations with slight permutations in the number of specific element types, number of blocks (i.e. groups of the same element type) of certain element types, with all other controllable operating conditions fixed. In order to better understand the role of the most common element types (conveying and kneading), only screw configurations that consisted of a combination of kneading and conveying elements (and for which product PSD were presented) were simulated here. The simulated screw configurations and their corresponding compartmental representations are outlined in Figure 5. Details of the model parameters corresponding to the experimental conditions in Vercruysse et al. [68] are presented in Table 2.
Figure 5: Screw configurations modelled in this work and their respective compartmental representations. The number in the centre of each compartment represents the compartment length (normalised by the screw diameter). Compartments representing conveying zones are in blue and compartments representing kneading zones are shown in orange. These correspond to configurations A, B, D and E in Vercruysse et al. [68]
Table 2: Un-optimised model inputs.

<table>
<thead>
<tr>
<th>Parameter (symbol)</th>
<th>Type</th>
<th>Value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liquid:solid mass feed ratio (LSR)</td>
<td>Operating parameter</td>
<td>0.09</td>
<td>-</td>
</tr>
<tr>
<td>Mass feed rate ($\dot{m}_{\text{feed}}$)</td>
<td>Operating parameter</td>
<td>20.0</td>
<td>kg hr$^{-1}$</td>
</tr>
<tr>
<td>Screw speed ($n_{\text{screw}}$)</td>
<td>Operating parameter</td>
<td>13.33</td>
<td>rev s$^{-1}$</td>
</tr>
<tr>
<td>Screw residence time ($\tau_{\text{screw}}$) (CE,2KE,6KE,2x6KE) [33]</td>
<td>Operating parameter (interpolated)</td>
<td>-, 3.02, 3.09, 4.52</td>
<td>s</td>
</tr>
<tr>
<td>Liquid addition nozzle diameter ($d_{\text{nozzle}}$)</td>
<td>Equipment geometry</td>
<td>1.6</td>
<td>mm</td>
</tr>
<tr>
<td>Screw diameter (D)</td>
<td>Equipment geometry</td>
<td>25</td>
<td>mm</td>
</tr>
<tr>
<td>Specific available volume CE ($\nu_{\text{convey}}$)</td>
<td>Equipment geometry</td>
<td>1.218×10$^{-5}$</td>
<td>m$^3$/D</td>
</tr>
<tr>
<td>Specific available volume KE ($\nu_{\text{knead}}$)</td>
<td>Equipment geometry</td>
<td>1.284×10$^{-5}$</td>
<td>m$^3$/D</td>
</tr>
<tr>
<td>Coefficient of restitution ($e_{\text{coag}}$) [44]</td>
<td>Material property</td>
<td>0.2</td>
<td>-</td>
</tr>
<tr>
<td>Liquid binder viscosity ($\mu_{\text{binder}}$)</td>
<td>Material property</td>
<td>10$^{-3}$</td>
<td>Pa s</td>
</tr>
<tr>
<td>Liquid binder density ($\rho_{l}$)</td>
<td>Material property</td>
<td>998</td>
<td>kg m$^{-3}$</td>
</tr>
<tr>
<td>Solid original density ($\rho_{s}$)[70]</td>
<td>Material property</td>
<td>1545</td>
<td>kg m$^{-3}$</td>
</tr>
<tr>
<td>Height of surface asperities ($h_{a}$)</td>
<td>Material property (estimated)</td>
<td>5×10$^{-6}$</td>
<td>m</td>
</tr>
<tr>
<td>Representative volume mean primary particle diameter ($d_{pp}$)</td>
<td>Material property</td>
<td>27.3</td>
<td>$\mu$m</td>
</tr>
<tr>
<td>Droplet diameter ($d_{\text{drop}}$)</td>
<td>Model parameter</td>
<td>1.6</td>
<td>mm</td>
</tr>
<tr>
<td>Minimum particle size for breakage ($v_{\text{parent}}$)</td>
<td>Model parameter</td>
<td>27.3</td>
<td>$\mu$m</td>
</tr>
<tr>
<td>Daughter distribution parameter 1 ($\alpha_{\text{daughter}}$) (CE,KE)</td>
<td>Model parameter</td>
<td>0.5, 2.0</td>
<td>-</td>
</tr>
<tr>
<td>Daughter distribution parameter 2 ($\beta_{\text{daughter}}$) (CE,KE)</td>
<td>Model parameter</td>
<td>0.5, 2.0</td>
<td>-</td>
</tr>
<tr>
<td>Minimum particle porosity ($\epsilon_{\text{min}}$)</td>
<td>Model parameter</td>
<td>0.3</td>
<td>-</td>
</tr>
<tr>
<td>Particle bed packing fraction ($\epsilon_{\text{bed}}$)</td>
<td>Model parameter</td>
<td>0.3</td>
<td>-</td>
</tr>
<tr>
<td>Pore saturation limit for nucleation ($s^*$)</td>
<td>Model parameter</td>
<td>0.12</td>
<td>-</td>
</tr>
</tbody>
</table>
4.2 Simulation conditions

All simulations were carried out using the numerical inputs given in Table 3 on a single core of an Intel® Sandy Bridge™ E5-2670 3.30GHz Processor with 4GB of RAM per core. Temporal functionals $M(t)$ from the simulations are reported as averages taken over all realisations as

$$\eta(t) = \frac{1}{n_{\text{realisations}}} \sum_{i=1}^{n_{\text{realisations}}} M_i(t),$$

(88)

where the functional may be a particle ensemble property such as the mass fraction of particles in a particular sieve class.

The half-width of the confidence intervals are

$$c(t) = 1.64 \sqrt{\frac{\sum_{i=1}^{n_{\text{realisations}}} (M_i(t) - \eta(t))^2}{n_{\text{realisations}}^2}},$$

(89)

which corresponds to a confidence interval $P = 0.9$ [40].

4.3 Parameter estimation

In this work, compartments with the same element type (conveying, kneading) are assumed to have the same process rate constants (4 for each element type, 8 in total), regardless of position. In this way, a central screw element library can be used to store element specific rate constants. This library can then be accessed to retrieve/define the rate constants for each compartment in an representation of arbitrary screw configurations. Using this framework, the model can be calibrated against experimental data through the optimisation of the relevant variables in the screw element library. The procedure for optimising each of the variables within the screw element library follows that described in McGuire et al. [45] and is only described briefly here for clarity. The rate constants are fitted against experimental data [68] corresponding to screw configurations CE, 2KE and 2x6KE (see Figure 5). Screw configuration 6KE is then used to measure the predictive power of the model. An initial parameter scan is carried out using 10000 Sobol points [8] across the parameter ranges outlined in Table 4. A Hooke Jeeves optimisation [29] is then carried out using the best 4 Sobol points from the first optimisation step as the starting
positions. Both stages are carried out using the Model Development Suite (MoDS) [51]. The objective function used in the fitting takes the form

$$OF = \sum_{i=1}^{N_{\text{exp}}} \sum_{j=1}^{N_{\text{response}}} \left( \frac{y_{i,j}^{\text{model}} - y_{i,j}^{\text{exp}}}{\sigma_j} \right)^2.$$  \hspace{1cm} (90)

Here, $y_{i,j}^{\text{model}}$ is the $j$th model response for the $i$th screw configuration modelled and $y_{i,j}^{\text{exp}}$ is the associated experimental response. $N_{\text{exp}}$ is the number of different screw configurations fitted and $N_{\text{response}}$ model/experimental responses from each configuration. Mass based percentiles diameters $d_{25}, d_{50}, d_{75}$ and $d_{95}$ of the granular product in [68] are used as the model/experimental responses. These are weighted, respectively, using weighting factors $\sigma$ of 25$\mu$m, 50$\mu$m, 75$\mu$m and 95$\mu$m.

### Table 4: Bounds used for optimisation of model parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Conveying Lower</th>
<th>Conveying Upper</th>
<th>Kneading Lower</th>
<th>Kneading Upper</th>
<th>Scaling</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k_{\text{col}}$</td>
<td>$10^{-10}$</td>
<td>$5 \times 10^{-8}$</td>
<td>$10^{-10}$</td>
<td>$2 \times 10^{-8}$</td>
<td>Logarithmic</td>
<td>m$^3$</td>
</tr>
<tr>
<td>$k_{\text{comp}}$</td>
<td>$10^{-2}$</td>
<td>2.0</td>
<td>$10^{-2}$</td>
<td>2.0</td>
<td>Logarithmic</td>
<td>-</td>
</tr>
<tr>
<td>$k_{\text{att}}$</td>
<td>$1.68 \times 10^{-2}$</td>
<td>8.4</td>
<td>$1.68 \times 10^{-3}$</td>
<td>0.0168</td>
<td>Logarithmic</td>
<td>s</td>
</tr>
<tr>
<td>$\omega_{\text{att}}$</td>
<td>$10^{-2}$</td>
<td>0.2</td>
<td>$5 \times 10^{-2}$</td>
<td>0.6</td>
<td>Linear</td>
<td>-</td>
</tr>
</tbody>
</table>

### 5 Model results

### Table 5: Optimised model rate constants.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Conveying</th>
<th>Kneading</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k_{\text{col}}$</td>
<td>$10^{-10}$</td>
<td>$2.0 \times 10^{-8}$</td>
<td>m$^3$</td>
</tr>
<tr>
<td>$k_{\text{comp}}$</td>
<td>0.01</td>
<td>0.475</td>
<td>-</td>
</tr>
<tr>
<td>$k_{\text{att}}$</td>
<td>0.0168</td>
<td>0.0557</td>
<td>s</td>
</tr>
<tr>
<td>$\omega_{\text{att}}$</td>
<td>0.1095</td>
<td>0.387</td>
<td>-</td>
</tr>
</tbody>
</table>

The set of model parameters with the lowest objective function value (found through the optimisation procedure) are presented in Table 5. The associated mass fraction distributions for this parameter set are displayed, firstly, for screw configurations used in the optimisation procedure in Figure 6, and secondly, for the ‘new’ 6KE screw configuration (depicted in Figure 5(c)) in Figure 7.

Note that the model has qualitatively captured the experimental trend in which the amount of fines (defined here as particles in the smallest sieve class) reduces with an increasing number of kneading elements. This is expected since the kneading elements serve to distribute the binding liquid across the body of solid material passing through the element,
Figure 6: A comparison of the optimised model particle size distributions in the final compartment against the experimental results from Vercruysse et al. [68] for different screw configurations.

thereby promoting particle growth. Though the model has captured this trend, the degree of primary particle consumption is under-predicted for all screw configuration tested - thought the disparity in minimal in the 2x6KE case. This is to be expected since a very simplistic layering mechanism was employed in this work. Within the model, the degree of layering is strongly controlled by the amount of surface liquid present on agglomerates, however, the amount of surface liquid that should be consumed by layering is unclear and hence a very basic model has therefore been used. Implementation of more complex models in the future would require experimental data on the layering dynamics in isolation from other twin-screw processes.

In terms of breakage processes, the model has under-predicted the production of particles in the size range 200-1000µm in screw elements with a high number of conveying elements. This is an indication that the breakage parameters that generate this distribution (daughter distribution shape parameters $\alpha_{\text{daughter}}$, $\beta_{\text{daughter}}$ and breakage exponent $\omega_{\text{att}}$) may need to be revisited. As previously discussed, the daughter distribution parameters used for conveying elements in this work were selected to qualitatively capture the cutting/edge
chipping effects that were experimentally observed in such elements. Since these parameters were not fitted within this study, it is likely that these parameters would benefit from further investigation. This would merit a modelling investigation in its own right and is not pursued further here. It is noted that the compaction, breakage and coagulation processes have all been pushed to their lower bound as a result of the optimisation process. This indicates that the majority of the model particle processes occurred in the kneading elements. The bounds of these conveying parameters were not reduced further, since the effect on the results was deemed to be minimal. The fact that conveying element particle processes have been minimised indicates the need to incorporate mid-barrel particle size information in the optimisation process. At the moment, only the final distribution is in this process, which can lead to un-physical particle evolutions or concentration of particle processes in distinct sections, as observed here.

The evolution of the particle size distribution along the compartmental networks is shown for the 2x6KE screw configuration in Figure 8. The nucleation zone consists of a mixture of over-sized agglomerates and primary particle mass. Upon entering the first kneading compartment \((z = 2)\) these large agglomerates are broken down and consolidated. The consolidation process has resulted in the squeezing of internal liquid to the particle surface which has promoted consumption of fines through layering. This trend continues through the second and third kneading compartment. It is clear from the results there is little change in the granular mass as it passes through the central conveying section \((z = \{6, 7\})\). Only a minimal degree of breakage is observed in this zone, and, as discussed in the previous paragraph, the absence of the expected increase in intermediate size classes (200-400µm) for the 2x6KE simulation indicates that the daughter distribution of these elements requires further investigation. Almost all of the remaining fines are consumed within the second kneading section of the compartment network.

In order to further assess the quality of the model, the evolution of the agglomerate particle composition along the screw barrel for each simulated screw configuration is presented in Figure 9. The associated mass fraction of fines at each of the these barrel positions, in each screw configuration, is shown in Figure 10. From Figure 9 it is observed that, for
the conveying elements, a slight degree of particle compaction has reduced the porosity of large agglomerate particles, which resulted in the complementary increase in solid volume fraction. Though the solid volume fraction increase could be attributed to particle growth through layering/coagulation, the minimal fines consumption for the CE configuration in Figure 10 and the abundance of surface liquid in Figure 9 (which should promote aggregate coagulation if the collision rate is sufficiently high) demonstrates that this is almost purely the result of aggregate compaction. In all configurations with a kneading zone, it is observed that, for the first kneading zone, aggregate particles acquire solid mass though the consolidation of highly porous agglomerates. This consolidation has replenished surface liquid levels, which promoted layering of primary particles and coalescence of surface wet agglomerates. This decrease in surface liquid observed in Figure 9 is a consequence of the internalisation of liquid present in both the aggregate consolidation and primary particle layering mechanisms of the model. This raises an interesting point - though several experimental studies [18, 22, 63, 67, 68] have concluded that kneading elements squeeze liquid to the surface large agglomerates, promoting further growth, it is generally not understood if this growth takes place within the kneading element, or further down the barrel in conveying zones. From Figure 10, the model in this work would seem to indicate that the majority of this growth/fines reduction occurs within the kneading blocks.

An interesting feature that is observed in particle evolution of the 2x6KE screw configuration is the increase in gas volume fraction within the second kneading block. This occurred as the result of aggregate coalescence - which, according to the coalescence model of Braumann et al. [10] used in this work, results in the generation of trapped pore volume as aggregates coalesce. Though trapping of such ‘new’ pore volume may occur, it

Figure 8: Particle size distribution evolution along the compartment network for the 2x6KE screw configuration. Element types by compartment index: conveying $z = \{1, 5, 6, 10, 11\}$, kneading $z = \{2, 3, 4, 7, 8, 9\}$. 
is expected that a net reduction in porosity would be observed across this kneading block (as is the case in the first kneading block of all screw configurations tested). Thus, the competition between the consolidation forces and pore volume creation through coalescence would be an area of investigation in future modelling efforts.
Figure 9: Mass-averaged aggregate particle composition evolutions (by volume) along the barrel length for each screw configuration simulated. $s_o$ represents original solid, $l_e$ represents external liquid, $l_i$ represents internal liquid and $g$ is gas. Kneading zones are highlighted in orange. Data points are placed according to the centre of the associated compartment. Barrel positions correspond to the variation zone and are expressed as a normalised length.
Figure 10: Simulated evolution of the mass fraction of fines ($d < 75\mu m$) along the length of the barrel for all screw configurations tested. Data points are placed according to the centre of the associated compartment. Barrel positions correspond to the variation zone and are expressed as a normalised length. Kneading zones are highlighted in orange.
6  Conclusions

In this work we have adapted and extended the twin-screw granulation model of [45, 46] to include a primary particle layering mechanism. In order to estimate compartmental residence times for use within this model, a novel procedure was developed to infer this parameter based on the type of screw element associated with each compartment. A stochastic particle framework for the simulation of the aggregate particle phase of the granulation system was presented and coupled to an ODE solver in order to simulate the dynamics of the primary particle population and carry out continuous aggregate particle processes. Model parameters specific to different types of screw element were calibrated through simulation of systems with a number of different screw configurations and comparison with an existing experimental data set. The model was observed to qualitatively capture the reduction in the mass fraction of fines in the granular product as the number of kneading elements was increased. The element specific breakage model and the drying dynamics of the layering process were identified as key areas for future model refinement.

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Abbreviations

CE  Conveying element
DEM  Discrete element method
DME  Distributive mixing element
KE  Kneading element
LPDA  Linear process deferment algorithm
PBE  Population balance equation
PBM  Population balance model
PEPT  Positron emission particle tracking
MC-PBM  Monte Carlo population balance model
SWA  Stochastic weighted algorithm
TSG  Twin-screw granulation
Nomenclature

Roman symbols

- $a_{\text{surf}}$: particle external surface area \( \text{m}^2 \)
- $B_{\text{frag}}$: breakage fragment probability distribution
- $c_{\text{pp}}$: primary particle concentration measure \( \text{m}^{-3} \)
- $C$: collision rate function \( \text{m}^2 \)
- $d$: particle diameter \( \text{m} \)
- $d_{\text{nozzle}}$: liquid injection nozzle diameter \( \text{m} \)
- $d_{\text{pp}}$: primary particle diameter \( \text{m} \)
- $D_i$: deferment function
- $e_{\text{coag}}$: material coefficient of restitution
- $f_{\text{frag}}$: breakage fragment size parameter
- $f_j,e$: volumetric fill fraction of screw section with element type $e$ in zone $j$
- $F$: aggregate breakage kernel
- $F_c$: transport scaling factor
- $g_{\text{break}}$: particle breakage frequency \( \text{s}^{-1} \)
- $h_a$: height of surface asperities \( \text{m} \)
- $h_l$: height of external liquid layer \( \text{m} \)
- $I_{\text{nuc.pp}}$: primary particle depletion sink term (nucleation) \( \text{s}^{-1} \text{m}^{-3} \)
- $I_{\text{trans.pp}}$: primary particle depletion source term (transport) \( \text{s}^{-1} \text{m}^{-3} \)
- $k_{\text{att}}$: attrition rate constant \( \text{s} \)
- $k_{\text{col}}$: collision rate constant \( \text{m}^3 \text{s}^{-2} \)
- $k_{\text{comp}}$: compaction rate constant
- $k_{\text{maj}}$: majorant scaling factor
- $K_{\text{coag}}$: coagulation kernel \( \text{m}^3 \text{s}^{-1} \)
- $K_{\text{SWA}}$: SWA coagulation kernel \( \text{m}^3 \text{s}^{-1} \)
- $K_{\text{col}}$: collision kernel \( \text{m}^3 \text{s}^{-1} \)
- $K_{\text{col}}^\prime$: collision kernel term \( \text{m}^3 \text{s}^{-1} \)
- $K_{\text{SWA}}^\prime$: SWA collision kernel \( \text{m}^3 \text{s}^{-1} \)
- $K_{\text{SWA}}^\prime$: SWA majorant collision kernel \( \text{m}^3 \text{s}^{-1} \)
- $l_e$: external liquid volume \( \text{m}^3 \)
- $l_{e\rightarrow i}$: amount of surface liquid internalised \( \text{m}^3 \)
- $l_i$: internal liquid volume \( \text{m}^3 \)
- $L_{j,e}$: length of screws section consisting of element type $e$ in zone $j$ \( \text{D} \)
- $L$: set of all compartment indices
- $LSR$: operating liquid solid mass flowrate ratio
- $m$: particle mass \( \text{kg} \)
- $\dot{m}_{\text{feed}}$: operating mass feed rate \( \text{kg s}^{-1} \)
- $M_{\text{metering}}$: dynamics mass hold up in screw metering zone \( \text{kg} \)
- $M_{\text{variation}}$: dynamics mass hold up in screw variation zone \( \text{kg} \)
\( n_{\text{realisations}} \) number of simulation realisations
\( n_{\text{screw}} \) screw speed \( \text{rev s}^{-1} \)
\( n_{\tau} \) simulation stop time parameter -
\( N_{\text{agg}} \) number of aggregate/computational particles -
\( N_{\text{agg}}^{\text{max}} \) maximum number of aggregate/computational particles -
\( N_{\text{pp}} \) Number of primary particles -
\( OF \) objective function -
\( p \) pore volume \( m^3 \)
\( r_{\text{inflow}} \) single particle inflow rate \( s^{-1} \)
\( r_{\text{layer}} \) rate of primary particle layering onto agglomerate particle \( s^{-1} \)
\( R_{\text{drop}} \) model droplet addition rate \( s^{-1} \)
\( R_{\text{nuc}} \) model nucleation rate \( s^{-1} \)
\( R_{\text{outflow,agg}} \) model aggregate outflow rate \( s^{-1} \)
\( R_{\text{SWA,\text{break}}} \) SWA breakage jump rate \( s^{-1} \)
\( R_{\text{SWA,\text{coag}}} \) SWA coagulation jump rate \( s^{-1} \)
\( R_{\text{SWA,\text{nuc}}} \) SWA nucleation jump rate \( s^{-1} \)
\( R_{\text{SWA,\text{total}}} \) SWA total jump rate \( s^{-1} \)
\( R_{\text{SWA,\text{trans}}} \) SWA transport jump rate \( s^{-1} \)
\( s_{o} \) original solid volume \( m^3 \)
\( s^{*} \) pore saturation limit -
\( t \) time \( s \)
\( t_{p} \) particle current time \( s \)
\( t_{\text{stop}} \) simulation stop time \( s \)
\( t_{\text{target}} \) deferment stop time \( s \)
\( \Delta t_{\text{wait}} \) jump waiting time \( s \)
\( \Delta t_{\text{defer}} \) deferment time step \( s \)
\( T_{\text{coag}} \) coagulation type transformation -
\( v \) particle volume \( m^3 \)
\( \hat{v}_{\text{break}} \) breakage normalisation parameter \( m^3 \)
\( v_{\text{parent}} \) minimum volume for breakage \( m^3 \)
\( v_{\text{pp}} \) primary particle volume \( m^3 \)
\( v_{y} \) breakage daughter volume function \( m^3 \)
\( V_{\text{real}} \) physical volume of compartment occupied by mass \( m^3 \)
\( V_{\text{samp}} \) sample volume \( m^3 \)
\( w \) particle statistical weight -
\( w_{\text{max}} \) maximum particle statistical weight -
\( w_{\text{nuc}} \) nuclei particle statistical weight -
\( x \) particle vector \( m^3 \)
\( x_{\text{nuc}} \) nuclei particle vector -
\( x_{\text{pp}} \) primary particle vector -
\( X \) particle type-space -
\( X_{\text{agg}} \) aggregate type-space -
<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\mathbf{X}_{pp}$</td>
<td>primary particle type space</td>
<td>-</td>
</tr>
<tr>
<td>$y$</td>
<td>breakage fragment particle vector</td>
<td>m³</td>
</tr>
<tr>
<td>$z$</td>
<td>compartment index</td>
<td>-</td>
</tr>
</tbody>
</table>

**Greek symbols**

- $\alpha_{\text{daughter}}$: breakage daughter distribution shape factor
- $\beta_{\text{daughter}}$: breakage daughter distribution shape factor
- $\gamma_{\text{coag}}$: coagulation weight transfer function
- $\gamma_{\text{frag}}$: breakage weight transfer function
- $\epsilon$: particle porosity
- $\epsilon_{\text{bed}}$: particle bed packing fraction
- $\lambda$: aggregate particle concentration measure
- $v_{\text{convey}}$: specific volume available in conveying elements
- $v_{\text{knead}}$: specific volume available in kneading elements
- $v_{\text{convey}}$: specific volume available in conveying elements
- $\rho_{\text{eff}}$: effective density of the solid material in the variation zone
- $\rho_l$: binder density
- $\rho_s$: solid density
- $\phi_{\text{max}}$: maximum liquid saturation
- $\sigma$: objective function weighting factor
- $\tau$: compartment residence time
- $\tau_{\text{metering}}$: metering zone screw residence time
- $\tau_{\text{screw}}$: total mean screw residence time
- $\tau_{\text{variation}}$: variation zone screw residence time
- $\chi_{\text{frag}}$: breakage fragment size parameter
- $\varphi$: test function
- $\omega_{\text{att}}$: breakage rate exponent
A Appendix: derivation of transport scaling factor

We wish to show that the weight scaling factor applied to inflow particles within the transport jump process (81) has the form:

\[ F_c(z,t) = \left[ \frac{V_{\text{samp}}(z,t)}{V_{\text{samp}}(z-1,t)} \right] \left[ \frac{N_{\text{agg}}(z-1,t)}{N_{\text{agg}}(z,t)} \right] \]  

(91)
in the context of a non-droplet zone compartment \((z \neq 1)\) for a twin-screw model system.

Firstly, we start with the assertion that each compartment in the network is one of constant mass and volume (i.e. \(M_{\text{real}}(z), V_{\text{real}}(z) \neq f(t)\) and thus, to provide continuity, the mass flow rate through each compartment \(\dot{m}_{\text{feed}}\) is the same). It follows that concentration of mass flow entering/leaving the sample volume (denoted by the superscript \(SV\)) for compartment \(z\) must be the same as that entering the physical system i.e.

\[ \frac{\dot{m}_{\text{inflow}}^{SV}(z,t)}{V_{\text{samp}}(z)} = \frac{\dot{m}_{\text{outflow}}^{SV}(z,t)}{V_{\text{samp}}(z,t)} = \frac{\dot{m}_{\text{feed}} V_{\text{real}}(z)}{V_{\text{real}}(z)} \]  

(92)

Thus we can express the mass flow of aggregate particles entering the systems based on the mass fraction of aggregates (in a mixture of aggregates and primary particles) in the preceding compartment (at steady state) as

\[ \dot{m}_{\text{agg,inflow}}^{SV}(z,t) \in L, z > 1 = \frac{\dot{m}_{\text{inflow}}^{SV}(z,t) M_{\text{agg}}^{SV}(z-1,t)}{M_{\text{total}}^{SV}(z-1,t)} \]  

\[ = \frac{\dot{m}_{\text{feed}} M_{\text{agg}}^{SV}(z-1,t)V_{\text{samp}}(z,t)}{V_{\text{real}}(z) M_{\text{total}}^{SV}(z-1,t)}, \]  

(93)

(94)

where \(M_{\text{agg}}^{SV}(z,t)\) and \(M_{\text{total}}^{SV}(z,t)\) are the mass of aggregates and the total mass (primary particles and aggregates) in the sample volume, respectively.

Let the number average mass of computational particles (aggregates) in the sample volume associated with compartment \(z\) be defined as

\[ \langle mw \rangle_{z,t} := \frac{M_{\text{agg}}^{SV}(z,t)}{N_{\text{agg}}(z,t)}. \]  

(95)

It is required that the inflow process to compartment \(z\) sample the particle distribution from compartment \(z - 1\). Thus, we are free to scale the weights of particles sampled from the preceding reactor by some constant scaling factor \(F_c\) prior to their inception into the current reactor. This scaling has no effect on the physical particle distribution in compartment \(z - 1\). Incorporating this scaling factor, the average mass of a particle incepted into compartment \(z\) from compartment \(z - 1\) during an inflow event is

\[ F_c \langle mw \rangle_{z-1,t}. \]  

(96)

We may write the mass inflow rate into compartment \(z\) in an alternative form which is based on the rate of inflow events \(R_{\text{inflow}}^{SWA}(z,t)\) as

\[ \dot{m}_{\text{agg,inflow}}^{SV}(z,t) \in L, z > 1 = R_{\text{inflow}}^{SWA}(z,t) F_c \langle mw \rangle_{z-1,t}. \]  

(97)
Equating (94), (97) and substituting (95), we may write

$$R_{\text{inflow}}^{\text{SWA}}(z, t) = \frac{\dot{m}_{\text{feed}} V_{\text{samp}}(z, t) N_{\text{agg}}(z-1, t)}{V_{\text{real}}(z) M_{\text{total}}^{\text{SV}}(z-1, t) F_c}. \quad (98)$$

Using the residence time definition

$$\dot{m}_{\text{feed}} := \frac{M_{\text{real}}(z-1)}{\tau(z-1)} = \frac{M_{\text{total}}^{\text{SV}}(z-1, t) V_{\text{real}}(z-1)}{\tau(z-1) V_{\text{samp}}(z-1, t)}, \quad (99)$$

we may write (98) as

$$R_{\text{inflow}}^{\text{SWA}}(z, t) = \left[ \frac{V_{\text{real}}(z-1)}{V_{\text{real}}(z)} \right] \left[ \frac{V_{\text{samp}}(z, t)}{V_{\text{samp}}(z-1, t)} \right] \left[ \frac{N_{\text{agg}}(z-1, t)}{F_c \tau(z-1)} \right]. \quad (100)$$

Equation (101) can be further simplified by acknowledging that a series of compartments with constant mass and constant volume has the property

$$\frac{V_{\text{real}}(z-1)}{V_{\text{real}}(z)} = \frac{M_{\text{real}}(z-1)}{M_{\text{real}}(z)}, \quad (102)$$

and again applying the residence time definition to each compartment in (102) we see that

$$\frac{V_{\text{real}}(z-1)}{V_{\text{real}}(z)} = \frac{\tau(z-1)}{\tau(z)}. \quad (103)$$

Substitution of (103) into (101) yields

$$R_{\text{inflow}}^{\text{SWA}}(z, t) = \left[ \frac{V_{\text{samp}}(z, t)}{V_{\text{samp}}(z-1, t)} \right] \left[ \frac{N_{\text{agg}}(z-1, t)}{F_c \tau(z)} \right]. \quad (104)$$

Now, if the inflow jump is to be coupled with the outflow jump process for any non-droplet zone ($z \neq 1$) then we require

$$R_{\text{inflow}}^{\text{SWA}}(z, t) = R_{\text{outflow}}^{\text{SWA}}(z, t) = \frac{N_{\text{agg}}(z)}{\tau(z)}. \quad (105)$$

Finally, equating (104) and (105) we arrive at an expression for the inflow weight scaling factor

$$F_c(z, t) = \left[ \frac{V_{\text{samp}}(z, t)}{V_{\text{samp}}(z-1, t)} \right] \left[ \frac{N_{\text{agg}}(z-1, t)}{N_{\text{agg}}(z, t)} \right]. \quad (106)$$

45
Appendix: layering rate equation

The layering source term in the primary particle PBE (2) has the form

\[
\frac{dc_{\text{pp}}}{dr}\bigg|_{\text{layer}} = - \int_{X_{\text{agg}}} r_{\text{layer}}(z, t, x, c_{\text{pp}}) \lambda(z, t, dx) \quad (107)
\]
\[
\approx - \frac{1}{V_{\text{samp}}(z, t)} \sum_{i=1}^{N_{\text{agg}(t)}} r_{\text{layer}}(z_i, t, x_i, c_{\text{pp}}) w_i. \quad (108)
\]

Hence, in number form

\[
\frac{dN_{\text{pp}}}{dr}\bigg|_{\text{layer}} \approx - N_{\text{agg}}(z, t) \sum_{i=1}^{N_{\text{agg}(t)}} r_{\text{layer}}(z_i, t, x_i, c_{\text{pp}}) w_i, \quad (109)
\]

and the change in the number of primaries over the global deferment step is then characterised as

\[
N_{\text{pp}}(z, t_{\text{target}}) \leftarrow N_{\text{pp}}(z, t) - N_{\text{agg}(t)} \int_{t_p}^{t_{\text{target}}} r_{\text{layer}}(z, t, x_i, c_{\text{pp}}) w_i \, dt. \quad (110)
\]

Re-arranging (28) we have

\[
r_{\text{layer}}(z, t, x, c_{\text{pp}}) = \frac{1}{v_{\text{pp}}} \left. \frac{ds_o(x)}{dr} \right|_{\text{layer}} \quad (111)
\]

and since

\[
\left. \frac{ds_o(x)}{dr} \right|_{\text{consol}} = 0, \quad (112)
\]

then, by way of (37),

\[
r_{\text{layer}}(z, t, x, c_{\text{pp}}) = \frac{1}{v_{\text{pp}}} \left. \frac{ds_o(x)}{dr} \right|_{\text{defer}}. \quad (113)
\]

Substitution of (113) into (110) and assuming that changes in \(N_{\text{pp}}\) over the deferment step are small yields

\[
N_{\text{pp}}(z, t_{\text{target}}) \leftarrow N_{\text{pp}}(z, t_{\text{target}}) - \frac{1}{v_{\text{pp}}} \sum_{i=1}^{N_{\text{agg}(t)}} \left[ s_o(x_i)_{t_{\text{target}}} - s_o(x_i)_{t_p} \right] w_i, \quad (114)
\]

which completes the derivation.

The argument above can be applied in the context of local deferment steps by summing only over the sequence of particles which are involved in the local deferment step (and hence the subsequent jump process).
C Appendix: derivation of primary particle transport rate

We wish to show that the primary particle flow rate for a linear network of constant mass, constant volume compartments has the form

\[
\frac{dN_{pp}(z)}{dt} \bigg|_{\text{transport}} = \begin{cases} 
\dot{m}_{\text{feed}} V_{\text{samp}}(z,t) - N_{pp}(z,t) \frac{1}{\tau(z)}, & \text{if } z = 1, \\
\frac{V_{\text{real}}(z-1)}{V_{\text{real}}(z)} \frac{V_{\text{samp}}(z)}{V_{\text{samp}}(z-1)} \frac{N_{pp}(z-1,t)}{\tau(z-1)} - \frac{N_{pp}(z,t)}{\tau(z)}, & \text{otherwise.}
\end{cases}
\]

(115)

Firstly, we start with a component balance on the primary particles based on the transport processes (inflow and outflow)

\[
\frac{dN_{pp}(z)}{dt} \bigg|_{\text{transport}} = R_{\text{inflow,pp}}(z,t) - R_{\text{outflow,pp}}(z,t),
\]

(116)

where \( R_{\text{inflow,pp}}(z,t) \) and \( R_{\text{outflow,pp}}(z,t) \) are the rates of primary particle inflow and outflow (number based) to/from the sample volume for compartment \( z \), respectively. We wish to define the forms of both terms of the RHS of (116) \( \forall z \in \mathbb{L} \).

Let us first consider the outflow term in (116). Using the mass-based definition of residence time we can write the mass outflow rate of primary particles from the sample volume of compartment \( z \) \( \forall z \in \mathbb{L} \) as

\[
m_{\text{pp,outflow}}(z) = M_{\text{pp,SV}}(z,t) \frac{1}{\tau(z)},
\]

(117)

where \( M_{\text{pp}}(z,t) \) is the total mass hold-up of primary particles in the sample volume for compartment \( z \).

Hence the rate of primary particle outflow from the sample volume of compartment \( z \) is

\[
R_{\text{outflow,pp}}(z,t) = \frac{m_{\text{pp,outflow}}(z)}{\rho_s v_{pp}} = \frac{M_{\text{pp,SV}}(z,t)}{\rho_s v_{pp} \tau(z)},
\]

(119)

The total mass hold-up of primary particles can be written as

\[
M_{\text{pp}}(z,t) = N_{pp}(z,t) \rho_s v_{pp}.
\]

(120)

It follows from (119) and (120) that

\[
R_{\text{outflow,pp}}(z,t) = \frac{N_{pp}(z,t)}{\tau(z)} \quad \forall z \in \mathbb{L}.
\]

(121)
Moving on to the inflow term in the RHS or (116), let us consider the first reactor in the network i.e. \( z = 1 \). Since the mass feed to this compartment consists purely of primary particles, the mass feed rate to the physical first compartment is simply the operating mass feed rate i.e.

\[
\dot{m}_{\text{pp,inflow}}(1) = \dot{m}_{\text{feed}}.
\]  

(122)

Thus, the mass feed rate of primary particles into the first compartment with sample volume \( V_{\text{samp}}(z,t) \) is

\[
\dot{m}^{SV}_{\text{pp,inflow}}(1,t) = \dot{m}_{\text{pp,inflow}}(1) \frac{V_{\text{samp}}(1,t)}{V_{\text{real}}(1)} = \dot{m}_{\text{feed}} \frac{V_{\text{samp}}(1,t)}{V_{\text{real}}(1)}.
\]  

(123)

(124)

The rate of primary particle inflow (number based) into the sample volume of the first compartment is then

\[
R_{\text{inflow,pp}}(1,t) = \frac{\dot{m}^{SV}_{\text{pp,inflow}}(1,t)}{\rho_{s,v_{\text{pp}}}} = \frac{\dot{m}_{\text{feed}} V_{\text{samp}}(1,t)}{V_{\text{real}}(1) \rho_{s,v_{\text{pp}}}}.
\]  

(125)

(126)

Together, (116), (121) and (126) prove (115) for the case \( z = 1 \).

In order to formulate an expression for \( R_{\text{inflow,pp}}(z,t) \) \( \forall z \neq 1 \), we first enforce continuity (in terms of the primary particle phase) across the boundary between two compartments. This requires that the equality

\[
R_{\text{real,inflow,pp}}(z,t) = R_{\text{real,outflow,pp}}(z-1,t) \quad \forall z \neq 1
\]  

(127)

be satisfied \( \forall z \in \mathbb{L}, z > 1 \), where \( R_{\text{real,inflow,pp}}(z,t) \) and \( R_{\text{real,outflow,pp}}(z,t) \) are the number flowrate of primary particles into and out of the physical compartment with indices \( z \) and \( z - 1 \), respectively.

Let us suppose that the physical compartment \( z \) can be constructed from \( n(z,t) \) identical sample volumes of size \( V_{\text{samp}}(z,t) \). Applying the same logic to \( z - 1 \) we may write (127) as

\[
n(z,t) R_{\text{inflow,pp}}(z,t) = n(z-1,t) R_{\text{outflow,pp}}(z-1,t) \quad \forall z \neq 1.
\]  

(128)

Since all sample volumes are identical, we may also write

\[
n(z,t) = \frac{V_{\text{real}}(z)}{V_{\text{samp}}(z,t)}.
\]  

(129)

Substitution of (129) into (128) and re-arranging for \( R_{\text{inflow,pp}}(z,t) \) yields

\[
R_{\text{inflow,pp}}(z,t) = \frac{V_{\text{real}}(z)}{V_{\text{real}}(z-1) V_{\text{samp}}(z-1)} R_{\text{outflow,pp}}(z-1,t).
\]  

(130)
Using the expression for $R_{\text{outflow,pp}}$ in (121) in (130) gives

$$R_{\text{inflow,pp}}(z,t) = \frac{V_{\text{real}}(z-1)}{V_{\text{real}}(z)} \frac{V_{\text{samp}}(z)}{V_{\text{samp}}(z-1)} \frac{N_{pp}(z-1,t)}{\tau(z-1)},$$

which, combined with (116) and (121), completes the derivation.
References


