Abstract
Pharmaceutical production remains one of the last industries that predominantly uses batch processes, which are inefficient and can cause drug shortages due to the long lead times or quality defects. Consequently, pharmaceutical companies are transitioning away from outdated batch lines, in large part motivated by the many advantages of continuous manufacturing (e.g., low cost, quality assurance, shortened lead time). As chemical reactions are fundamental to any drug production process, the selection of reactor and its design are critical to enhanced performance such as improved selectivity and yield. In this article, relevant theories, and models, as well as their required input data are summarized to assist the reader in these tasks, focusing on continuous reactions. Selected examples that describe the application of plug flow reactors (PFRs) and continuous-stirred tank reactors (CSTRs)-in-series within the pharmaceutical industry are provided. Process analytical technologies (PATs), which are important tools that provide real-time in-line continuous monitoring of reactions, are recommended to be considered during the reactor design process (e.g., port design for the PAT probe). Finally, other important points, such as density change caused by thermal expansion or solid precipitation, clogging/fouling, and scaling-up, are discussed.

Keywords
Flow chemistry · Continuous manufacturing · PFR · CSTR · PAT · Residence time distribution

Introduction
Continuous manufacturing, or continuous processing, is defined as “the material(s) and product are continuously charged into and discharged from the system respectively, throughout the duration of the process” [1]. Industries such as food, petrochemicals and automotive, have long since adopted automated and continuous manufacturing, whereas pharmaceutical production remains one of the last industrial processes that mainly use a non-continuous (i.e., “batch”) approach [2, 3]. This is because of the following differences that those other industries did not have to consider: structural complexity, quality and regulatory, and quantity requirements (i.e., the trend towards lower dose drugs) [4]. This inefficient batch process can cause drug shortages due to the long lead times (up to 12 months) [2] or quality defects [1]. The current pharmaceutical industry operates at approximately 2–3 sigma quality (~6.7–30.9% defects, i.e., failed / rejected products), thus, much improvement is required to achieve 6 sigma quality (~0.0003% defects) [5]. Motivated by the benefits shown in Fig. 1 [6–10], the pharmaceutical industry is transitioning to continuous processes, including end-to-end integrated continuous manufacturing (ICM) approaches [1, 6, 8, 10–18]. A first-of-its-kind research demonstration of an end-to-end ICM line was unveiled by MIT in 2011 [13]. The model drug was aliskiren hemifumarate, and the throughput of the process was 45 g/h, with a residence time of 47 h. Subsequently, the first commercial-setting end-to-end ICM pilot plant was reported by CONTINUUS Pharmaceuticals in 2019 [12]. The throughput of the process was 4800 tablets per hour, or 40.3 × 10⁶ tablets per year, with a total residence time of <30 h. There are examples of continuously manufactured drug products that have been approved in the US (e.g., Orkambi, Symdeko and Trikafta by Vertex, Prezista by Johnson & Johnson, Verzenio
by Eli Lilly, Daurismo by Pfizer), in the EU (e.g., Orkambi and Symkevi by Vertex, Prezista by Johnson & Johnson, Verzenios by Eli Lilly), and in Japan (e.g., Tramacet by Johnson & Johnson, Verzenio by Eli Lilly), as well as drugs that are under development [12, 16, 19].

The reaction is one of the necessary steps in the continuous manufacturing of pharmaceuticals. There are many types of reactions (e.g., imidazole cyclization, hydroformylation, reductive amination, thermal deprotection) that could be performed in continuous reactors. In addition, some can occur simultaneously with crystallization (i.e., reactive crystallization), when the solubility of the active pharmaceutical ingredient (API) is low but the reaction rate is high. Thus, appropriate reactor design and selection for different reactions are essential to ensure desirable performance of the manufacturing process (e.g., high selectivity and yield). There are three main categories of ideal reactors: batch reactors, plug flow reactors (PFR, also known as tubular reactors), and continuous-stirred tank reactors (CSTRs, also known as backmixing reactors) [20, 21]; the last two are continuous. In batch reactions, raw materials and solvents are charged into the reactor at the beginning, and the product is collectively discharged at the end [1]. The composition within the batch reactor changes with time, the residence time is uniform, and there is no flow through the process. These reactors are usually used for small-scale production and for testing new processes, especially those with complex chemistry.

In continuous reactions, raw materials and solvents are continuously charged into the system, and the product is continuously discharged from the system throughout the duration of the process [1]. An ideal PFR has no axial mixing, but perfect radial mixing. In a PFR, all materials passing through experience the same concentration and temperature profiles along its length and have the same residence time. Conversely, in a CSTR, although all the materials have uniform temperature, pressure, and concentration, the residence time is characterized by a distribution (residence time distribution, RTD). This is because some components that enter the reactor can leave immediately, while others remain for longer time periods. Usually one CSTR cannot achieve complete conversion, and CSTRs-in-series are used to approximate a PFR. PFRs and CSTRs-in-series are popular continuous reactors that are widely used in the pharmaceutical industry. Figure 2 and Table 1 compare the differences between a PFR and CSTRs-in-series [24–26]. This article summarizes theories for the design and selection of continuous reactors, and highlights some examples of PFRs and CSTRs-in-series within the pharmaceutical industry.

**Selected data collection**

The design and selection of a continuous reactor depends on many parameters including reaction kinetics, enthalpy of reaction, heat and mass transfer, etc. A more effective design can be made when a large amount of input data is available; however, this is not always the case. Reaction kinetics and enthalpy of reaction are parameters of paramount interest, since they dictate the amount of reaction time needed, reaction conditions (e.g., pressure, temperature, catalyst) and the heating management of the reactor, respectively.

**Reaction kinetics**

Reaction kinetics is the study of the reaction rate and mechanism by which the reactants are transformed into products [27]. In most cases the reaction rate constants and rate equations need be approximated, as they cannot be predicted from first principles [28]. Accordingly, the main process reactions are usually approximated as first-order or second-order, and over a narrow range of conditions (e.g., concentration, temperature, pressure) [29]. The reactor design and selection discussed in this article is based on non-zero-order reactions.

For a first-order reaction (i.e., a reaction that proceeds at a rate that depends linearly on the concentration of one reactant [20, 21]),

\[ A \rightarrow P \]  

The reaction rate equation for this first-order reaction is [20, 21].

\[ r = \frac{dc_A}{dt} = \frac{dc_P}{dt} = k_1 C_A \]  

(1)
Here \( r \) is reaction rate, and \( C_A \) and \( C_P \) are the reactant and product concentrations, respectively. \( k_1 \) is the rate constant for first-order reaction. Integrating eq. (1), we obtain eq. (2),

\[
k_1 = \frac{1}{t} \ln \frac{C_{A0}}{C_A}
\]

(2)

\( C_{A0} \) is the original reactant concentration, and the unit for \( k_1 \) is \( \text{time}^{-1} \).

For a second-order reaction, the sum of the exponents in the rate law is equal to two.

\[A(\text{reactant}) + B(\text{reactant}) \rightarrow P(\text{product})\]

The reaction rate equation for this second-order reaction is [20, 21].

\[
r = \frac{-dC_A}{dt} = \frac{-dC_B}{dt} = \frac{dC_P}{dt} = k_2 C_A C_B
\]

(3)

Here \( C_B \) is the concentration of reactant B, and \( k_2 \) is the rate constant for second-order reaction. Integrating eq. (3), we obtain eqs. (4) and (5),

\[
k_2 = \frac{1}{t} \left( \frac{1}{C_A} - \frac{1}{C_{A0}} \right), \text{ (if} \ C_{A0} = C_{B0})
\]

(4)

\[
k_2 = \frac{1}{t(C_{A0} - C_{B0})} \ln \frac{C_{B0} C_A}{C_{A0} C_B}, \text{ (if} \ C_{A0} \neq C_{B0})
\]

(5)

\( C_{B0} \) is the original concentration of reactant B, and the unit of \( k_2 \) is \( \text{(concentration)}^{-1} \cdot \text{(time)}^{-1} \).

The Arrhenius equation is an expression for the temperature dependence of reaction rates (eq. (6)) [20, 21].

\[
k = A \exp \left( - \frac{E}{RT} \right)
\]

(6)

where \( k \) is the rate constant at temperature \( T \), \( R \) is the molar gas constant, \( A \) is the pre-exponential factor, and \( E \) is the apparent activation energy.

**Enthalpy of reaction**

Enthalpy of a reaction, \( \Delta H_r \), is defined as the heat energy change that takes place when reactants convert into products. If the sum of the enthalpies of the products is greater than that of the reactants, the reaction will be endothermic. Conversely, if the reactants have larger enthalpies, the reaction will be exothermic.

Under standard condition (i.e., temperature of 25 °C and pressure of 1 atm), the standard enthalpy of reaction, \( \Delta H_r^\circ \), is calculated below,

\[
\Delta H_r^\circ = \Delta H_f^\circ(\text{products}) - \Delta H_f^\circ(\text{reactants})
\]

(7)

where \( \Delta H_f^\circ \) is the standard enthalpy of formation (i.e., the enthalpy change during the formation of 1 mol of...
the substance from its constituent elements under standard state.

Under process conditions of temperature $T$ and pressure $P$, the enthalpy of reaction $(\Delta H_r, P, T)$ is expressed as eq. (8) [29].

$$
\Delta H_r^{Q_2} = \Delta H_r^{Q_1} + \int_1^P \left[ \left( \frac{\partial H_{\text{prod}}}{\partial P} \right)_T - \left( \frac{\partial H_{\text{react}}}{\partial P} \right)_T \right] dP + \int_{298.15}^T \left[ \left( \frac{\partial H_{\text{prod}}}{\partial T} \right)_P - \left( \frac{\partial H_{\text{react}}}{\partial T} \right)_P \right] dT \tag{8}
$$

If temperature is the only parameter that needs to be accounted for, the equation is simplified [29],

$$
\Delta H_r^{Q_2} = \Delta H_r^{Q_1} + \Delta H_{\text{prod}} + \Delta H_{\text{react}} \tag{9}
$$

where $\Delta H_{\text{prod}}$ and $\Delta H_{\text{react}}$ are the enthalpy change of the products and reactants, respectively, when their temperature changes from 298.15 K to $T$.

Other parameters, such as equilibrium constant and Gibbs free energy, are also useful for reactor design. The heat and mass transfer properties are especially important for multiphase reactors. These parameters will not be discussed here.

### RTD theory, dispersion model and CSTRs-in-series model

#### RTD theory

Residence time describes the length of time that a molecule or reaction material spends in a reactor. It is an important characteristic of any reactor. Material in an ideal batch reactor or an ideal PFR has a single residence time. However, for all other reactor types, multiple residence times exist, and they are expressed as a function of time by the residence time distribution (RTD) [20, 21, 30].

For the pulse injection of the tracer, the mean residence time, $\bar{t}$, and variance, $\sigma^2$, can be obtained by [20, 21].

$$
\bar{t} = \frac{\int_0^\infty t C(t) dt}{\int_0^\infty C(t) dt} \tag{10}
$$

and

$$
\sigma^2 = \frac{\int_0^\infty t^2 C(t) dt}{\int_0^\infty C(t) dt} - \bar{t}^2 \tag{11}
$$

The RTD function of $E(t)$ is calculated from the pulse tracer experiments by [20, 21].

$$
E(t) = \frac{C(t)}{\int_0^\infty C(t) dt} \tag{12}
$$

The dimensionless function, $E(\theta)$, which offers a direct comparison of experimental results for different conditions (e.g., different flowrates, different reactor sizes), is calculated by [20, 21].

$$
E(\theta) = \frac{C(\theta)}{\int_0^\infty C(\theta) d\theta} \tag{13}
$$

where the dimensionless time, $\theta$, is calculated from the ratio of the real time, $t$, to the mean residence time, $\bar{t}$, [20, 21].

$$
\theta = \frac{t}{\bar{t}} \tag{14}
$$

#### Axial dispersion model

The axial dispersion model is usually used to describe non-ideal PFRs. The vessel dispersion number, $D^*/uL$, is the parameter that determines axial dispersion. When $D^*/uL < 0.01$, the system emulates plug flow, and $E(\theta)$ is expressed as [21].

$$
E(\theta) = \frac{1}{\sqrt{4\pi (D^*/uL)}} \exp \left[ - \frac{(1-\theta)^2}{4(D^*/uL)} \right] \tag{15}
$$
where \( u \) is the fluid velocity, \( L \) is the reactor length, and \( D^* \) is the axial dispersion coefficient, which is defined by [21].

\[
D^* = D + \frac{u^2 d^3}{192 D}
\]

(16)

where \( D \) is the diffusion coefficient and \( d \) is the tube diameter. The variance, \( \sigma^2_\theta \), is calculated by [21].

\[
\sigma^2_\theta = \frac{2 D^*}{u L}
\]

(17)

Accordingly, when \( D^*/uL > 0.01 \), the system is open and far from plug flow, and \( E(\theta) \) and its variance are expressed as [21].

\[
E(\theta) = \frac{1}{\sqrt{4\pi D^* / uL} \theta} \exp \left[ -\frac{(1-\theta)^2}{4 \theta (D^* / uL)} \right]
\]

(18)

and

\[
\sigma^2_\theta = \frac{2 D^*}{u L} + 8 \left( \frac{D^*}{uL} \right)^2
\]

(19)

The axial dispersion coefficient \( D^* \) can be estimated from the maximum peak heights of either \( E(\theta) \) curves.

### CSTRs-in-series model

The CSTRs-in-series model can be used when there is less deviation from plug flow. This model is simple, can be used with any kinetics, and can be extended to any arrangement of compartments (with or without recycle) [21].

For the \( n \)th CSTR, the RTD function of \( E(t) \) and its variance are expressed as [21, 31].

\[
E(t) = \frac{t^{n-1} n^n}{(n-1)!} e^{-tn/t}
\]

(20)

and

\[
\sigma^2 = \frac{2}{n}
\]

(21)

The dimensionless function \( E(\theta) \) and its variance are expressed as [21, 31].

\[
E(\theta) = n \frac{(n\theta)^{n-1}}{(n-1)!} e^{-n\theta}
\]

(22)

and

\[
\sigma^2_\theta = \frac{1}{n}
\]

(23)

\[PFR\]

Ideal PFRs and batch reactors are characterized by a single uniform residence time. For non-ideal reactors, a PFR’s RTD is broader than that of a batch reactor, but narrower than that of a CSTR. In the production of APIs, batch reactors and PFRs have been widely used, and complete conversion is achievable [30]. For some reactions such as organic azide or tetrazole formations, a PFR is preferable because it operates 100% liquid-filled, while a batch reactor or CSTR typically has a headspace where hydrazoic acid would partition [24, 32]. Compared with CSTRs-in-series, PFR is usually lower in cost (e.g., less materials required for research and development, and a PFR is cheaper than CSTRs-in-series) and in complexity. To manufacture the APIs continuously, a PFR is preferred, unless it is not practical (e.g., clogging issues). PFRs can be used for gas-, liquid-, gas-liquid, and liquid-liquid phases reactions, and some solid-related reactions (e.g., hydroformylation reaction [23]), although it is challenging to handle solids.

In a PFR, we assumed that the concentration varies continuously in the axial direction along the reactor length, but no change with time for a given location [20, 24]. The required volume \( (V) \) to achieve the specified conversion \( X \) is [20],

\[
V = F_{A0} \int_{X_0}^X \frac{dX}{-r_A}
\]

(24)

where \( F_{A0} \) is the molar flowrate of reactant A, and \( -r_A \) is the reaction rate of reactant A. Under experimental conditions, PFRs are not ideal because of the axial dispersion (Fig. 3). In pharmaceutical applications, the flow is usually laminar (i.e., \( Re < 2300 \)) because of the low throughput and small characteristic dimension. Turbulent flow is achieved when \( Re > 4000 \).

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**Fig. 3** Comparison of ideal plug flow and axial dispersed plug flow
The general form of correlation for $D^*/uL$ is described by equation [33, 34],

$$\frac{D^*}{uL} = (\text{intensity of dispersion})(\text{geometric factor}) \quad (25)$$

where intensity of dispersion is defined as $D^*/u_d$, which correlates to Reynolds number, $Re$, and Schmidt number, $Sc$, [21].

$$Re = \frac{\rho u d_i}{\mu} \quad (26)$$

$$Sc = \frac{\mu}{\rho D} \quad (27)$$

Where $\rho$ is the density, and $\mu$ is the dynamic viscosity. For straight tubing, the geometric factor is $d/L$ (i.e., the reverse of the common used form $L/d$). Typically, higher $L/d$ leads to lower $D^*/uL$. It is interesting to note that for smaller tubes, there was less axial dispersion at lower flow rates, but for larger tubes, there was more axial dispersion at lower flow rates [33]. For coiled tubes, the geometric factor is more complex, and the coil diameter $d_c$ matters. As reported, under certain assumptions the logarithm of $D^*$ decreased linearly with the logarithm of the dimensionless parameter $De^2Sc^{1.14}$ [35], where $De$ is the Dean number [36], a key parameter for coiled flows, [21].

$$De = \left( \frac{d_i}{d_c} \right)^{\frac{1}{2}} Re \quad (28)$$

When coil diameter decrease, $De$ and $De^2Sc^{1.14}$ will increase. Therefore, the axial dispersion reduced.

Furthermore, it is helpful for better understanding of the reaction system if we can estimate the conversion along the length of the PFR. The conversion is mainly influenced by residence time, reaction order, reaction rate and axial dispersion. Consider a PFR with length $L$, within which chemicals are mixing axially with a dispersion number of $D^*$. For a $n$th-order reaction with reactant A, the reaction conversion in this nonideal PFR can be estimated with simulation software (e.g., MATLAB) by the following equation [21, 37].

$$\frac{D^*}{uL} \frac{d^2X}{dz^2} - \frac{dX}{dz} + kT_C^{n-1}(1-X)^n = 0 \quad (29)$$

where $z = l/L$, and $l$ is the length from the entrance. At the exit, $l = L$. From this equation, we observe that the fraction conversion of reactant A through the PFR is controlled by $D^*/uL$ and $\tau KC_{A_0}^{-1}$ [21]. From Fig. 4a, if a first-order reaction requires 90 min to reach 99.9% conversion in an ideal PFR (i.e., $D^*/uL=0$), the mean residence time for $D^*/uL$ of 0.002, 0.025 and 0.2 are 91.2, 105 and 196 min, respectively (assuming the same reaction conditions) [33]. Figure 4b and Table 2 provide an example of the simulated conversion results as a function of $z$ for three different temperatures for a second-order reaction.

It is important to note that the conversion profile will not be as smooth as shown in Fig. 4, especially when there exists a “hot spot” (i.e., an area or point within a reaction system at which the temperature is appreciably higher than in the bulk of the reactor), for exothermic reactions. The existence of a “hot spot” depends mainly on a reactor’s specific area (i.e., surface area per reactor volume). Specific areas greater than $10^4 \text{ m}^2/\text{m}^3$ can only be achieved in microreactors. Usually the specific area in a PFR is less than $2000 \text{ m}^2/\text{m}^3$. Scaling up the inner diameter (i.d.) from 2 mm to 10 mm, decreases the specific area from 2000 to 400 $\text{ m}^2/\text{m}^3$ (Table 3).

Although it is not feasible to measure the temperature at all points along the length of a PFR, numerical modeling is an alternative method to estimate the temperature profile throughout the PFR. Fogler [20] derived the energy balance for a non-isothermal PFR, as expressed in eq. (30), which provides the temperature profile along the length of the PFR.

$$\frac{dT}{dV} = \frac{r_A(T)\Delta H_{rxn}(T) - U(a(T-T_a))}{\sum F_iC_{P_i}} \quad (30)$$

Where $\frac{dT}{dV}$ is the temperature change of the process fluids in a segment of the reactor, $\Delta H_{rxn}(T)$ is the heat of reaction at temperature $T$, $r_A(T)$ is the reaction rate of species $A$ at temperature $T$, $U$ is the overall heat-transfer coefficient, $a$ is the PFR heat-exchange area per volume of reactor, $T$ is the temperature of the process fluids, $T_a$ is the temperature of the heating or cooling media, $F_i$ is the mole flow rate of species $i$, and $C_{P_i}$ is the mean heat capacity of species $i$. Based on eq. (30), Johnson et al. [25] reported a steady-state temperature profile along a PFR for a homogeneous cryogenic lithiation reaction. The simulation result showed the existence of a hot spot, although it is not physically measurable.

Table 2  Comparison of the experimental and simulated conversion results in a PFR (Adapted from Ref. [16] with permission from the Royal Society of Chemistry)

| # | Temperature (°C) | Experimental X (%) | Simulated X (%) | Error | Error% |
|---|----------------|--------------------|-----------------|-------|-------|
| 1 | T-5            | 31.8               | 25.1            | -6.7  | -21.1 |
| 2 | T              | 36.2               | 32.2            | -4.0  | -11.3 |
| 3 | T+5            | 40.6               | 39.7            | -0.9  | -2.22 |
The first PFR was used to optimize a hydrazine condensation reaction in flow (step 1 in Table 4a). For reference, in batch mode, a large molar excess of hazardous hydrazine is required, and the transformation proceeds slowly. Conversely, in flow mode, stainless steel (SS) tubing with inner diameter (i.d.) of 4.57 mm, length of 91 m, and volume of 1.4 L was used. The throughput was 3.4 kg/day, the residence time was 60 min, and the reaction was operated under 130 °C and 34.5 bar, conditions not feasible with batch reactors. Minor impurities were observed at the end of the reaction. The second PFR was used for a nucleophilic aromatic substitution reaction coupling a pyrazole and pyrazine (step 2 in Table 4a). The flow reaction was conducted in low pressure PFA tubing with inner diameter (i.d.) of 6.35 mm, length of 91 m, and volume of 2.8 L. The liquid flowed from the bottom to the top of the horizontally coiled tube (immersed in a heated bath) to ensure liquid filled. The throughput was 2.9 kg/day, the residence time was 180 min, and the reaction temperature and pressure were 70 °C, and atmospheric pressure, respectively. The third PFR was used for deprotection with simultaneous gas and liquid handling (step 3 in Table 4a). This reaction exhibited excellent robustness between 20 and 40 °C, with a residence time between 2 and 6 h. The PFA coiled tubing (15.9 mm i.d.) was vertically oriented, and operated 50% filled with liquid and 50% filled with N2 carrier gas. Less impurities were produced when the generated CO2 and iso-butylene were removed in situ. The characterization of this PFR revealed plug flow operation with a $D^* / \mu L$ of 0.003, which approximates the RTD of 140 equal-volume CSTRs-in-series.

Mascia and Patrick et al. [13, 43] developed a multi-step synthesis and workup sequence for aliskiren hemifumarate, and PFRs were used in two steps (Table 4b). A melted chemical intermediate was pumped into the first PFR (i.d. of 11.7 mm, length of 25 m, volume of 2.7 L) at 100 °C, where it was mixed with 10 equiv. amine and 1 equiv. acid catalyst. This single-phase reaction was much faster when run neat, compared to the batch reaction (3–4 h vs. 72 h). The diffusion coefficient was $1 \times 10^{-9}$ m$^2$/s when the nominal flow rate of 675 mL/h was used. The second PFR (i.d. of 4 mm, length of 4.9 m, volume of 0.062 L) was used for the acid-catalyzed removal of the Boc protecting group. The reaction formed CO2 gas that created irregular flow patterns within the 4 mm i.d. tubing. It was observed that orientating the reactor coil horizontally resulted in 1–2% higher yield, with a peak yield being obtained at 30 °C [43].

Johnson and May et al. [25, 33] investigated imidazole cyclization (Table 4c) for GMP production of 29 kg of an advanced intermediate. They utilized a 7.1 L coiled stainless steel tube thermal PFR (151 m long, 7.75 mm i.d.) with a specific area of 516 m$^2$/m$^3$, and concluded that this specific area was sufficient to achieve adequate heat-up and cool-

![Fig. 4](image-url)

**Fig. 4** (a) Influence of axial dispersion on $\tau$ for a first-order reaction that requires 90 min to reach conversion of 99.9% in an ideal PFR (Fig. 4a is reproduced with permission from Ref. [33] Table 6. Copyright 2012 American Chemical Society), and (b) simulated conversion results as a function of $z$ for three different temperatures for a second order reaction. (Fig. 4b is adapted from Ref. [16] with permission from the Royal Society of Chemistry)

### Table 3 A comparison of the specific area for typical reactors

| Reactor type                      | Length/Diameter ratio ($L/d_t$) | Specific area ($m^2/m^3$) |
|----------------------------------|---------------------------------|---------------------------|
| 10 L batch reactor               | –                               | ~20                       |
| 1 L batch reactor                | –                               | ~50                       |
| 1 L PFR with inner diameter of 10 mm | $1.3 \times 10^3$               | 400                       |
| 1 L PFR with inner diameter of 2 mm | $1.6 \times 10^5$               | 2000                      |
| 140 µL microreactor with channel of 400 × 400 µm [38] | $2.0 \times 10^6$               | $10^4$                    |
Table 4  Selected examples of reactions performed in a PFR.

| #  | Reaction Scheme | PFR Characteristics | Conditions and Notes | Ref. |
|----|-----------------|---------------------|----------------------|------|
| a  | ![Reaction Scheme](image) | SS; 4.57 mm i.d.; 91 m long; 1.4 L; specific area 875 m²/m³ | T=130 °C (PFR in oven); P=34.5 bar; t=60 min | [39] |
|    |                 | PFA; 6.35 mm i.d.; 91 m long; 2.8 L; specific area 630 m²/m³ | T=70–100 °C; t=1–3 h; | |
|    |                 | PFA; 15.9 mm i.d.; $D^*/u_L=0.003$; specific area <250 m²/m³ | T=20–40 °C; t=2–6 h (liquid t=4 h, gas t=80 min); CO₂ formation | |
| b  | ![Reaction Scheme](image) | SS; 11.7 mm i.d.; 25 m long; 2.7 L; specific area 342 m²/m³ | T=100 °C (PFR in oven); t=4 h | [13,40] |
|    |                 | PFA; 4 mm i.d.; 4.9 m long; 0.062 L; specific area <1000 m²/m³ | T=30 °C (higher T resulting in increased degradation); CO₂ formation | |
| c  | ![Reaction Scheme](image) | SS; 7.75 mm i.d.; 151 m long; 7.1 L; specific area 516 m²/m³; coil diameter 0.28–0.43 m; coil height 0.43 m | T=140 °C (PFR in oven); P=69 bar (pressure drop ΔP=0.34 bar); t=90 min | [23,33] |
|    | Hydroformylation | SS; 16.5 mm i.d.; 152 m long; 32 L; specific area < 242 m²/m³; coil diameter 0.36–0.56 m; coil height 0.53 m | T=55 °C (PFR in heating bath); P=68 bar; t=24 h | [23] |
| d  | ![Reaction Scheme](image) | 45 SS vertical pipes in series; each 3.7 m tall and 53 mm i.d.; 360 L | T=20 °C; P=50 bar (pressure drop ΔP=7 bar); t=12 h; throughput was 100 kg per day | [23,45] |
| e  | ![Reaction Scheme](image) | SS; 2.5 L (plant-scale) | T= -30 °C; plant-scale throughput was 2 kg/h | [43] |
| f  | ![Reaction Scheme](image) | SS; 164 mL | T= -65 °C; t=0.48 min | [44] |
| g  | ![Reaction Scheme](image) | SS; 476 mL | T= -65 °C; t=0.78 min | |
| h  | ![Reaction Scheme](image) | PFA; 3.19 L | T= 50–60 °C; t=3.96 | |

[a] The tubing was operated 50% filled with liquid and 50% with N₂ gas; CO₂ gas was formed in the reaction; [b] the real specific area should be less than 1000 m²/m³ as CO₂ gas was formed in the reaction; [c] vapor and liquid mixture;
down times [25]. The reaction was carried out under extreme conditions (i.e., 140 °C, 69 bar), which exceed typical limitations of batch reactors. However, these conditions are feasible with SS PFRs without any scale-up issues. Under these reaction conditions, the pressure drop (∆P) from inlet to outlet is about 0.34 bar. Pressure drop along the length of the PFR can be estimated by the Hagen-Poiseuille equation [33, 48],

$$
\Delta P = \frac{8 \mu LQ}{\pi R^4} = \frac{8 \pi \mu LQ}{A^2}
$$

where $Q$ is the volumetric flow rate, $R$ is the pipe radius, and $A$ is the cross section of the pipe.

The scheme in Table 4d shows a hydroformylation reaction [23], where solid precipitated from the reaction mixture. A 32 L SS pulsating coiled tube reactor (with 16.5 mm i.d., 152 m length, see Fig. 5a-b) was operated under 68 bar and 55 °C. The syngas was composed of 1:1 CO and H₂, and the substrate to RhCOH[PPh₃]₃ catalyst ratio was 1000. It was not practical to increase the catalyst loading because: 1) the cost of Rh is high, and 2) the catalyst could precipitate from the reaction mixture, causing clogging and fouling issues. The reactor had pulsating flows in the forward and backward direction to prevent clogging. The residence time was 24 h, and decreasing the reaction time by increasing the temperature was not practical, as more linear aldehyde (undesired product) was generated at higher temperatures.

May et al. [44] investigated the GMP scale-up of a continuous Ir-catalyzed homogeneous reductive amination reaction (two phase gas-liquid reaction) between a secondary amine and a trans-aldehyde to produce a tertiary amine (Table 4e). A vertical pipes-in-series bubble flow reactor was constructed from 45 SS pipes (3.7 m tall, 53 mm i.d., $D^*/uL = 0.001$), connected by 4.6 mm i.d. down-jumper tubes (Fig. 5c) [25, 44]. The reaction was operated under 20 °C, 50 bar, a 12 h residence time, and with dissolved [Ir(Cod)Cl]₂ catalyst. The throughput was 100 kg/day and the plant was run continuously for 24 days. Compared with the batch process, this continuous reactor was safer because: 1) the amount of H₂ vapor space was low (~2% of the reactor volume), and 2) the reactor was located outside of the building, which was not feasible for batch reactors that required frequent reagent and catalyst charging. In addition, compared to the coiled tubing reactor, this pipes-in-series reactor was scalable to larger volumes.

Vieira et al. [40] reported a large-scale cyanation process to the synthesis of Remdesivir (Fig. 5d-e and Table 4f). The investigated step was exothermic and these exotherms could have an adiabatic temperature rise of 4–9 °C, which was easily controlled by the batch cooling as the reactor material was SS. When scaling up the input from 100 g to 2.75 kg, incomplete conversion of the starting material and much lower diastereoselectivity was obtained. However, the addition of 1.0 M equiv. of TFA solved this problem. The operation
temperature was −30 °C instead of −40 °C because TFA has limited solubility in the TMSOTf/DCM mixture, and clogged the line gradually. After applying the optimal conditions, high solution purity and diastereoselectivity were observed.

Cole et al. [41] built an equipment set for the production scale of continuous cryogenic lithium-halogen exchange (Fig. 5f and Table 4 g). Briefly, n-BuLi and bromide were precooled and mixed in a T-mixer and then entered the 164 mL SS lithiation reactor, then the solution mixed with precooled imine and entered the 476 mL SS addition reactor. The aqueous HCl in MeOH stream was precooled and met the process stream first, then entered the 3.19 L PFA chiral auxiliary cleavage reactor. Clogging happened during the 16 days campaign, however, disassembly was not required as the clogging could be resolved by warming the reactor while flow continued.

Table 4 h shows the scheme of thermal ethoxyethyl deprotection [25, 45]. Three PFR reactors were designed, and all of them were constructed from 7.75 mm i.d. Hastelloy. The lengths of these three reactors were 255, 53, and 153 m, with volumes of 12, 2.5, and 7.2 L, respectively. Accordingly, the reaction conditions were different, as described in Table 4 h [45–47]. It is important to note that the second PFR’s coil diameter was smaller (< 10 cm) than the other two, and that it fit inside a 10 cm inner-diameter /1.2 m tall steam heating pipe. Therefore, the heat transfer to and from the PFR was better than that of a forced convective oven.

In addition, the smaller diameter of the coil reduced the axial dispersion number \((D^* / uL = 0.001)\) in the second reactor, compared to the PFRs with larger coil diameters [36].

There are other publications that report on reactions performed in PFRs [23, 49–63], which are not discussed in detail here.

**CSTRs-in-series**

As mentioned above, a CSTR has a broader RTD than a batch reactor or PFR. Integrating more CSTRs (i.e., CSTRs-in-series) could narrow the RTD, but it is usually still broader than a PFR because of the limited reactor numbers. Figure 6a-b provide an example of the RTD measurements of a 5-stage CSTRs-in-series [16]. The designed residence time was 3.5 h in each stage, however, tracer molecules could spread for approximately 20 h in the 1st stage and for more than 40 h in the 5th stage (Fig. 6a). As more CSTRs were integrated, the RTD narrowed (Fig. 6b). The 5-stage CSTRs-in-series system described in Fig. 6a-b was designed for a reactive-crystallization process, and the yield obtained was 89.6% in the 5th stage [16]. To further increase the reaction yield, the authors integrated a PFR before the 1st stage CSTR. Figure 6c-d show the RTD measurements for the PFR-CSTRs-in-series system. Slightly narrower RTDs were observed, compared with the 5-stage CSTRs-in-series system, and a 91.3% yield was
obtained. As well known, there is an induction period (from the start of reaction to crystallization) for reactive crystallization processes. Therefore, it could be economically beneficial to integrate a PFR before the 1st stage CSTR (i.e., PFR was used before the onset of the crystallization).

A CSTR or CSTRs-in-series could be used for liquid, liquid-liquid, and liquid-solid phase reactions. Compared with a PFR, CSTRs-in-series have much higher solid-handling capacity [64, 65], and could buffer out the fluctuations of reagent feeds. In a CSTR, we assume the mixture within the reactor is perfectly mixed, and no time or position dependence of the temperature, concentration, or reaction rate inside the reactor at steady state [20]. The volume \( V \) necessary to achieve conversion \( (X) \) is [20].

\[
V = \frac{F_{A0}X}{-r_A} \tag{32}
\]

Compared with a PFR, a larger size CSTR is necessary to achieve the same conversion (except zero-order reactions), as the reaction occurs in a CSTR with the lowest reactant concentration. Consider a second-order reactive crystallization with a conversion of 97% [8], the volume of the corresponding PFR is 1.2 L, and 12.4 L for a single-stage CSTR. By approximating a PFR with a four-stage CSTRs-in-series, the volume is reduced from 12.4 L to 2 L [8].

The Damköhler number, \( Da \), which is defined as the ratio of the reaction rate of \( A \) to the convective transport rate of \( A \) at the entrance of the reactor, is often used to estimate the degree of conversion achieved in continuous reactors [20].

\[
Da = \frac{-r_{A0}V}{F_{A0}} \tag{33}
\]

For a first-order reaction, [20],

\[
Da = \tau k_1 \tag{34}
\]

where \( \tau \) is the space time, which can be obtained by dividing the reactor volume by the volumetric flow rate \( (v_0) \) entering the reactor,

\[
\tau = \frac{V}{v_0} \tag{35}
\]

Conversion \( X \) of a first-order liquid-phase reaction can be described in terms of \( Da \), [20].

\[
X = 1 - \frac{1}{(1 + Da)^n} \quad (\text{reactor number } n = 1, 2, 3, \ldots) \tag{36}
\]

For a single stage reactor, \( n = 1 \), and eq. (36) will simplify to

\[
X = \frac{Da}{1 + Da} \tag{37}
\]

Therefore, a conversion of greater than 90% is expected when \( Da \) is greater than 10; conversely, a conversion of less than 10% is obtained when \( Da \) is less than 0.1. Based on Fig. 7a-b, the number of CSTRs required to achieve a conversion of 90% were 1, 6, 13, 25 and 48 for \( Da \) of 10, 0.5, 0.2, 0.1 and 0.05, respectively.

For a second-order reaction, [20].

\[
Da = \tau k_2 C_{A0} \tag{38}
\]

Accordingly, conversion \( X \) of a second-order reaction can be expressed as, [20],

\[
X = \frac{(1 + 2Da)-\sqrt{(1 + 4Da)}}{2Da} \tag{39}
\]

From Fig. 7c, a \( Da \) value of 90 is necessary to achieve a conversion of 90%, while a \( Da \) value of 2 achieves 50% conversion. This means that a 45-fold increase in \( Da \) (this can be attained by increasing either the reaction temperature or the CSTR volume) will increase the conversion less than two times for a single CSTR. Thus, to obtain a relatively high conversion while minimizing reactor volume and reaction temperature, it is necessary to use CSTRs-in-series to approximate a PFR. From eq. (38), we observe that \( Da \) decreases as the reactant concentration decreases for a second-order reaction. Therefore, for a 4-stage CSTRs-in-series system, \( Da \) decreases stage by stage due to the decreasing concentration of reactants (Fig. 7d). In addition, the conversions for stages 1–4 are 71.6, 87.5, 93.0 and 97.0%, respectively [8]. This translates to conversion changes in each vessel are 71.6, 15.9, 5.5, and 4%, respectively. As the later stages provide less conversion, their addition should be dependent on the balance between simplicity and cost (fewer vessels), and a narrower RTD and higher conversion (more vessels). Typically, the number of stages in CSTRs-in-series is 3–5.

Figure 8 and Table 5 show several examples of reactions carried out in CSTRs-in-series. White et al. [66] investigated a Schotten-Baumann reaction that formed a cytotoxic API using a CSTRs-in-series system (three 12 L vessels). For this reaction, either a PFR or CSTRs-in-series would work. The reasons the authors selected the CSTRs-in-series are: 1) the evolution of CO\(_2\) gas during the reaction would easily partition into the head space of a CSTR, but result in a third phase in a PFR, 2) compared to a PFR, CSTRs could provide the benefit of dampening out some temporary inaccuracies without large fluctuation in the effective stoichiometry, and 3) impurity levels are high when mixing is poor. Figure 8a and Table 5a provide the process flow diagram and the scheme, respectively. The three feeds (i.e., 5-Bromothiophene-2-sulfonamide in MeTHF/iPrOAc, 2,4-dichlorobenzoyl chloride in Toluene, Na\(_2\)CO\(_3\) in water) were pumped into the first CSTR continuously, and the resulting two-phase mixture was then transferred stage to stage, maintaining a constant level. The
operating temperature was 65 °C, the $t$ was 1 h in each CSTR, and the throughput was 5 kg/day. As the equipment set was portable, dedicated, and disposable, the potential for cross-contamination was eliminated.

Braden et al. [67] reported a Barbier Grignard formation and coupling, along with quench and neutralization. The authors used three CSTRs-in-series (see Fig. 8b, the third CSTR from the left was not used) to produce 5 kg/day product in fume hoods. Table 5b provides the relevant scheme. The re-actor ran 90 min in the Grignard CSTR, 30 min in the quench, and 30 min in the carbonate wash. Compared with the batch process, this continuous process could reduce racemization of the unstable tetrahedral intermediate. This is because the product is not stable from racemization until after the carbonate wash, and the batch process takes longer between the Grignard coupling reaction and the quench reaction, which allowed more time for racemization. In addition, the continuous process is safer because: 1) the Grignard formation reaction has runaway potential in a 100 L batch reactor (operating in batch mode in a 100 L vessel provides the same overall throughput as the 2 L continuous reactors), and 2) much less excess Mg was required quenching at the end, and less hydrogen gas is generated. The authors selected CSTRs-in-series instead of a PFR because of the solid Mg reagent, which was sequestered in the CSTR by an internal settling pipe. During the entire 75 h continuous run, a high enantiomeric excess (ee) was obtained, which averaged >99%.

Hu et al. [8, 16] developed an automated multi-stage continuous reactive crystallization system with in-line PATs for a high viscosity process. A solitary PFR was not considered because of the high solid concentration; rather, a hybrid approach was taken. Figure 8c shows the process flow diagram of the PFR-CSTRs system. The reactive crystallization consists of a second order C-N bond-forming reaction between a nitrile species and a secondary amine hydrochloride, in which the resulting API adduct (N,N-dialkyl guanine) precipitates out as a hydrochloride salt (Table 5c). The integration of a PFR before the first CSTR increased the yield by 1.7%, as discussed above. In addition, for this highly exothermic reaction, the temperature was observed to be quite stable for the continuous process. This temperature stability could have contributed to lower impurity levels and higher yields. Conversely, the temperature of the corresponding batch process fluctuated over a wide range, as heat was quickly released during a very short time interval.

Duan et al. [68] investigated the flow epoxidation using catalytic methyltrioxorhenium (MTO) and aqueous H$_2$O$_2$ (Table 5d and Fig. 8d). The authors used a PFR initially, and low conversion was obtained due to the formation of a biphasic layer and insufficient mixing. CSTRs provided better mixing and yielded similar results to those in the batch
reaction; however, the heat dissipation capacity was low. As better mixing and a longer residence time facilitate the reaction conversion, a combination of a PFR-3 CSTRs-in-series system was used. The PFR coil, with its more efficient heat exchange capability, managed most of the reactive heat, while the CSTRs-in-series, which provides better mixing and a longer residence time, drove the epoxidation reaction to completion.

Susanne et al. [26] reported the synthesis of benzoxazole from N-pivaloyl 3,4-dichloro aniline. In this case, two steps utilized CSTRs-in-series systems (Fig. 8e and Table 5e). PFRs was not selected because of the clogging issues. For these two reaction steps, the cooling medium was set to $-30^\circ\text{C}$, and process temperatures above $-10^\circ\text{C}$ in the vessels were avoided to prevent the generation of a by-product. For the first reaction step, a 5-stage SS CSTRs-in-series system was used. In order to limit the maximal hot spot temperature, $n$-butyl lithium was added in both the first CSTR and the third CSTR (i.e., two-point dosing strategy), and the total residence time set to 1.4 min. Figure 9a shows the simulation results for the heat profiles in each vessel. Stages 1 and 3 exhibited the highest temperatures after the system stabilized because of the addition of $n$-butyl lithium. For the second reaction step, borosilicate glass was used because SS is not compatible with the high acidity of the reaction medium. Sulfur dioxide was added in the first vessel, causing it to have the highest temperature after the system stabilized. Figure 9b shows the simulation results for the heat profiles in each vessel.

Tom et al. [69] developed the methylation of octacycle to AMG 397 (Fig. 8f and Table 5f). Initially a PFR-CSTRs-in-series system was used; however, clogging occurred when the process was operated over a long period because of accumulation of the major by-product KI. A CSTR was used to replace the PFR to ensure the solid transfer during the deprotonation step. Thus 4-CSTRs-in-series was used for deprotonation, methylation, aging, and quench vessel, respectively. A pressure cascade was implemented to transfer the reaction mixture between the 300 mL CSTRs consistently under an inert atmosphere. The throughput was 30 g/h of starting material, and 115 g was successfully processed over 4 h. More specifically, no precipitation in the first CSTR, no transfer issues, and no clogging were observed. The reaction system

![Fig. 8](image-url)
achieved a state of control after approximately two residence times and complete conversion (> 95% AMG 397) was obtained.

There are other publications that report on reactions performed in CSTRs-in-series [71–77], which are not discussed in detail here.

### Selected PATs in continuous reactors

Process analytical technology (PAT) is defined by the U.S. Food and Drug Administration (FDA) as “a system for designing, analysing, and controlling manufacturing through timely measurements (i.e., during processing) of critical quality and performance attributes of raw and in-process materials and processes, with the goal of ensuring final product quality” [78]. Approximately 70% of PAT applications are developed for pharmaceutical manufacturing because of regulatory requirements [79]. Batch-produced APIs are analyzed by off-line instruments (e.g., HPLC, GC) to verify their quality attributes; however, in-line/on-line continuous monitoring largely based on PATs is necessary for the continuous manufacturing. These PATs could activate feed-forward and feedback control actions to ensure the quality of the APIs, as well as the drug product. During the reactor design process, it is important to consider the PATs that will be incorporated into the system (e.g., design ports for suitable PATs). Although there are many commonly used PATs [79–90], herein we only briefly discuss the ReactIR and focused beam reflectance measurement (FBRM), which could be used in continuous reactors.

ReactIR is a real-time, in situ mid-infrared based system that enables tracking of soluble reactants, intermediates, products and by-products during the reaction process. This provides useful information about initiation, conversion, intermediates and endpoints of a reaction [8, 16, 91–93]. A ReactIR probe could be located in a batch reactor, a CSTR, or any stage of a CSTRs-in-series, while a ReactIR flow cell can be placed at the exit of a PFR [94]. For example, Wernik et al. [75] used

| # | Reaction Scheme | CSTRs-in-series Characteristics | Conditions and Notes | Ref. |
|---|---|---|---|---|
| a | ![Chemical Structure](attachment://a.png) | 3 CSTRs, 12 L each | T=65 °C; t=1 h in each CSTR (3 h in total); throughput 5 kg/day | [23,66] |
| b | ![Chemical Structure](attachment://b.png) | 3 CSTRs, one is 2L, the other two are 1 L | T=55 °C initiation, and T=35 °C for the continuous reaction; t= 90 min (Grignard)+30 min (quench) + 30 min (carbonate wash) | [23,67, 70] |
| c | ![Chemical Structure](attachment://c.png) | 5 CSTRs (4 for Reaction + 1 for Crystallization) | High solid concentration (> 50 wt%); a PFR was integrated before the first CSTR | [8,16] |
| d | ![Chemical Structure](attachment://d.png) | 3 CSTRs, 12 L each | T=5–15 °C; t= ~50 min; a PFR was integrated before the first CSTR | [68] |
| e | ![Chemical Structure](attachment://e.png) | 5 SS CSTRs, 10 mL each | T=30–10 °C; t= 1.4 min | [22] |
| f | ![Chemical Structure](attachment://f.png) | 4 CSTRs; 300 mL each; (deprotonation, methylation, aging, quench) | T=20 °C; t= 5 min (20 min in total) | [69] |
ReactIR in the second CSTR to monitor the formation of diazoketone at 2107 cm$^{-1}$. The concentration of CH$_2$N$_2$ cannot be monitored because the overlap of the characteristic stretch between CH$_2$N$_2$ (2097 cm$^{-1}$) and diazoketone; Hu et al. [8, 16] used ReactIR in the last stage of CSTRs-in-series to monitor the concentration of reactant in the mother liquor; Beaver et al. [74] developed ReactIR method to monitor both the consumption of 2-bromopropene and the generation of product to reduce the safety risk relevant to the accumulation of the Grignard precursor.

FBRM is a probe-based instrument that tracks changing particle size and counts in real-time, and has been widely used in: 1) developing and optimizing crystallization process [95–99], 2) tracking and troubleshooting crystallizer systems [100–105], and 3) even monitoring polymorphic forms [103, 106]. The FBRM probe could be located in a batch crystallizer, or in CSTRs-in-series systems for reactive crystallization processes [8, 16]. For example, Hu et al. [8, 16] used FBRM in the last stage of CSTRs-in-series to track changing chord length in real time.

**Selected points for attention**

**Density change**

Johnson et al. [25] discussed thermal expansion in a PFR. For exothermic or endothermic reactions, the temperature profile along the length of a PFR is not uniform, and this will influence the reaction rate as well as the residence time [25]. When the reaction mixture is heated, it will generally expand, resulting in higher volumetric flow rate. Accordingly, the residence time will decrease, possibly causing incomplete conversion of the reactants (the conversion also depends on the reaction rate change). Conversely, when the reaction mixture is cooled, the residence time will increase, possibly resulting in over-reacted material (if the influence of the reaction rate change is less than that of the residence time change). Thus, it is useful to understand the thermal expansion phenomenon within a reactor, as described in eq. (40) [25, 107].

\[
\frac{\delta V_{\text{run mix}}}{\delta T} \bigg|_p \frac{1}{V_{\text{run mix}}} = \beta
\]
where $\frac{\partial V_{\text{rxn mix}}}{\partial T} \bigg|_P$ is the volume change of the reaction mixture for different temperatures at constant pressure, $V_{\text{rxn mix}}$ is the volume of the reaction mixture, and $\beta$ is the volumetric thermal expansion coefficient.

In the CSTRs-in-series systems for the described reactive crystallization process [8, 16], the precipitation of the product from the reaction mixture could result in the density increase, and accordingly influence the residence time. If the reaction system sensitive to the reaction time, measures should be taken to mitigate this impact.

### Clogging and fouling

Clogging and fouling are issues that need to be considered, as they can lead to process failure. Hu et al. [8] reported a high temperature and high viscosity continuous reactive crystallization system in CSTRs-in-series, and revealed that the transfer tubing between stages easily clogged, restricting continuous pumping of the slurry. A “forward-backward” burst pumping strategy was developed to enable transfer of the high viscosity slurry from one CSTR to the next. More specifically, the transfer tubing inserted into the bottom of the previous vessel, and on the top of the next vessel. In this way, the slurry could only be transferred in the forward direction. The feed pump had a constant flow rate of 3.3 mL/min, while the transfer pumps ran forward at 150 mL/min for 25 s, after which they ran backward at 150 mL/min for approximately 15 min. During the latter, the transfer tubing was emptied (i.e., made ready for the next cycle). In a hydroformylation reaction [23] where solids precipitate from the reaction mixture, potentially clogging the reactor tubing, a strategy of “pulsating flow” was used. The flow was forced back and forth by approximately 1 m, preventing solids from accumulating in the tube.

![Fig. 10 Heat transfer between hot reaction mixture, reactor wall and cooling medium](image)

Fouling, or encrustation, is a phenomenon in which solids precipitate and deposit on the internal surfaces of equipment, such as impellers, PAT probes, and the inner walls of vessels. Fouling on the vessel walls will reduce the heat transfer, which could influence the reaction rate, resulting in decreased yield. Many strategies [99, 108–114] have been developed to avoid fouling in continuous crystallization, which can also be used in reactive crystallization processes, including: (1) mechanical devices (e.g., addition of baffles, scrapes, rotating shafts), (2) surface energy or roughness modification via coating (e.g., gold), (3) operating conditions (e.g., mixing, supersaturation level, flow rate, seeding), (4) process dynamics (e.g., use of ultrasonic vibrations, temperature cycling), (5) polymeric excipients (e.g., hydroxypropyl methylcellulose (HPMC)), (6) model-based strategies, and (7) PAT-based strategies.

### Scaling-up

The ultimate goal of research and development (small scale, typically on the order of mg to 1 kg) is production (large scale, typically on the order of 10 kg and higher). There are several options for scaling a continuous process from lab to plant [7, 115]: 1) running for a longer period of time; 2) scaling-out (numbering up/parallelization); 3) scaling-up (increasing size). A combination of scaling-out and scaling-up is not uncommon for commercial-scale production, because the middle-sized reactor can still benefit from enhanced heat transfer and safety, without any further re-optimization [7]. However, the most common strategy to achieve target throughput is scaling-up. The technical challenges of this approach include the changes of reactor
performance (e.g., heat transfer, safety) and reaction outcome (e.g., yield, impurity), which largely caused by reduced specific area of the reactor.

As described by Berton et al. [115], a major concern during the scaling-up process is insufficient heat transfer, especially for fast reaction types such as nitration, organolithium, Grignard and reduction. The limitations of heat transfer within the continuous reactors are (see Fig. 10): 1) transferring heat from reaction medium to inside of reactor wall; 2) transferring heat across the reactor wall; 3) removing the heat from outside of reactor wall. Point 1) is the most challenging problem, which can be mitigated by inducing better mixing inside the reaction zone (e.g., static mixers, packing or porous media, baffled walls, Coriolis flow path) and applying multipoint-dosing strategy as mentioned in Fig. 9a. It is important to notice the space between the dotted line in the hot reaction mixture and the reactor wall (i.e., film thickness $\delta_1$), where the heat transfer rate is significantly lower than in the reaction mixture. This poor heat transfer is due to the laminar flow that is usually obtained near the wall. Better mixing could reduce film thickness $\delta_1$. Point 2) can be addressed by actively pumping the cooling media across the reactor surface instead of being immersed in the cooling media. Increasing the pumping rate could help to reduce film thickness $\delta_2$. Point 3) can be addressed simply by minimizing the wall thickness and choosing appropriate materials such as metal and glass.

Conclusions

In this article, several relevant theories and models (i.e., RTD theory, axial dispersion model, CSTRs-in-series model) are summarized to facilitate effective continuous reactor design and selection. Before the design and selection of continuous reactors can be made, input data such as reaction kinetics, and enthalpy of reaction are required. Several examples in the pharmaceutical industry that use PFRs and CSTRs-in-series are provided. PATs are important tools that provide real-time in-line continuous monitoring of reactions, and should be considered in the reactor design process. There are other points that require special attention, including density change caused by thermal expansion or solid precipitation, clogging/fouling, and scaling-up.

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Declarations

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