

Emulsion Polymerization of Butyl Acrylate: Safe Optimization Using Topological Criteria

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ABSTRACT: Fast and strongly exothermic emulsion polymerization processes are particularly difficult to optimize from both safety and productivity points of view because of the occurrence of a number of undesired side reactions (e.g., propagation of tertiary radicals, chain transfer to monomer, backbiting, termination by disproportionation, etc.) and the triggering of boiling phenomena with consequent stable foam formation under atmospheric pressure. Therefore, it would be useful to develop a suitable combined theoretical and experimental procedure able to detect both the optimum process dosing time and initial reactor temperature. In this work, it is discussed how an extended version of the topological criterion theory, originally developed for isoperibolic semibatch reactors, can be used to safely optimize an indirectly cooled isothermal semibatch reactor. Moreover, such a methodology is applied to a case study represented by the synthesis of poly(butyl acrylate) through the radical emulsion polymerization of butyl acrylate.

1. INTRODUCTION

Runaway phenomena are a very common problem in pharmaceutical and fine chemical industries. However, despite only a small number of these accidents injuring workers or the inhabitants of the neighborhood of the damaged factory, when strong runaways occur, the consequences can be really serious. This has motivated the great amount of work that has been done on runaway phenomena for batch (B) and semibatch (SB) reactors during the last 30 years.^{1–11} Particularly, a general criterion able to detect the so-called “runaway boundary” in multiple reaction systems and different reactor typologies is the generalized parametric sensitivity criterion.⁴ This criterion identifies the runaway boundary looking for the maximum of a suitable objective sensitivity coefficient, the normalized sensitivity coefficient of the maximum reactor temperature with respect to any operating parameter. A different general criterion for determination of the runaway boundary is the divergence criterion.^{5–7} This criterion states that if the system of ordinary differential equations (ODEs) that describes the analyzed process exhibits positive divergence, the synthesis is operating under runaway conditions. The criterion shows the relevant advantage of being usable for both online and offline evaluations.

However, from a practical point of view, the optimization of all (or some) operating parameters (e.g., coolant temperature and dosing time) is the relevant aspect. Knowing the runaway boundaries allows one to avoid the selection of potentially hazardous operating conditions, but it does not allow one to optimize the analyzed process. The first combined “safety-optimization” approach for isoperibolic semibatch reactors (SBRs) was developed by Hugo and Steinbach.¹ Their work, based on homogeneous systems, introduced the

accumulation criterion for analysis of SBR thermal behavior: coreactant accumulation into the system, which arises from a nonnegligible characteristic time of the chemical reaction with respect to that of the coreactant supply, must be kept at sufficiently low values to avoid the reactor thermal loss of control (runaway). According to this criterion, operating conditions characterized by a sufficiently low coreactant accumulation are considered not only safe but also productive. Steensma and Westerterp² extended the results obtained by Hugo and Steinbach¹ to heterogeneous (liquid–liquid) SBRs, introducing the concept of target temperature and producing the so-called boundary diagrams (BDs) for single reactions of (1,1) reaction-order kinetics. Particularly, such diagrams are generated in a suitable dimensionless space that allows end users to easily discriminate between safe (“quick onset, fair conversion, and smooth temperature profile”, QFS) and excessive accumulation operating conditions. According to the BD method, the QFS region is considered the most safe and productive for an exothermic SBR. However, it can be noted that the definition of such a region is somehow arbitrary.³ The problem of having an arbitrary definition of the QFS conditions was discussed by Alós et al.,³ who proposed, for reacting systems involving a single reaction, a definition of QFS using a suitable objective function.

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In particular, the QFS of an isoperibolic SBR is individuated, for a given value of the dosing time, by the coolant temperature in correspondence with which the objective function “time at which the maximum reactor temperature occurs” exhibits a local minimum. The intrinsic nature of this criterion can be proven by using parametric sensitivity. Recently, a more general optimization criterion, based on topological tools and able to consider different reactor typologies and temperature control modes, has also been developed.^{12,13}

Polymerization processes are one of the industrial processes most frequently involved in thermal runaway accidents¹⁴ (that is, accidents involving an exothermic phenomenon caused by a loss of reactor temperature control that occurs whenever the heat evolution rate is higher than the removal one provided by a dedicated cooling system) for three main reasons: (1) polymerizations can exhibit high reaction enthalpies¹⁵ (up to 110 kJ/mol); (2) huge free-radical accumulation, due to both a termination reaction rate decrease and a propagation reaction rate increase (Trommsdorff or gel effect^{16,17}), causes an autoacceleration of heat evolution; (3) under atmospheric pressure, boiling phenomena followed by stable foam formation can be triggered above a threshold temperature [in the following, referred to as MAT, the “maximum allowable temperature”¹⁸ (note that the MAT value could also be lower than such a value when reaction mixture decompositions or unwanted side reactions, such as the cross-linking ones) is triggered at lower temperatures], which is characteristic of the analyzed reacting system. Therefore, it is difficult to find a scale-independent optimization procedure able to identify the optimum values of the relevant operating variables (e.g., for a SBR, the dosing time and initial reactor temperature values) in order to maximize both reactor safety and productivity.

Among all polymerization processes, emulsion polymerization is widely diffused around the world to produce waterborne resins with a great variety of colloidal and physicochemical properties. Particularly, colloidal properties of latex products are of great importance from both academic and industrial viewpoints.¹⁹ A generic heterogeneous free-radical polymerization involves emulsification of a relatively hydrophobic monomer in water by an oil-in-water emulsifier [e.g., sodium lauryl sulfate (SLS)], followed by the initiation reaction with either a water-soluble initiator [e.g., sodium persulfate or potassium persulfate (KPS)] or an oil-soluble initiator [e.g., 2,2'-azobis(isobutyronitrile) (AIBN)]. It is a complex process because nucleation, growth, and stabilization of polymer particles are controlled by the free-radical polymerization mechanisms in combination with various colloidal phenomena. The most striking feature seems to be the free-radical segregation among monomer-swollen polymer particles: this phenomenon greatly reduces the bimolecular termination probability and, therefore, it results in a faster polymerization rate and higher polymer molecular weight.¹⁹ This advantageous characteristic of emulsion polymerization cannot be achieved simultaneously in bulk or solution polymerization.

Industrially, because of the exothermic nature of free-radical polymerizations and the limited heat-transfer capacity in large-scale reactors, most commercial latex products are manufactured by semibatch or continuous reaction systems. One of the main differences among the polymerization processes cited above is the growing particle residence time distribution within the reactor. Particularly, residence time distribution broadness is, in decreasing order, continuous > semibatch. As a consequence, the resultant particle size distribution broadness is, in decreasing order, continuous > semibatch. On the contrary, the polymerization rate generally follows the trend semibatch > continuous. Furthermore, at the full-plant scale, reproducibility of emulsion

polymerization processes within a narrow range is desirable; this means that the final solids content should be constant within $\pm 1\%$, the particle size, emulsion viscosity, and polymer average molecular mass should vary little, and any residual monomer should be maintained within minimum possible narrow limits (<0.5%). In order to achieve these conditions, the formulation of the emulsion polymerization (namely, the recipe) should not be subject to variations such as minor changes of either raw materials or operative conditions.²⁰ Anyway, a certain degree of operational flexibility is usually permitted, and both semibatch and continuous emulsion polymerization processes can produce latex products with controlled polymer composition and particle morphology even if subjected to the minor changes cited above.

Moreover, other restrictions apply on an emulsion polymerization process in industrial practice: the process should be completed in the shortest possible time, and the latex concentration must be as high as possible. With regard to time, if the process is operated discontinuously, an overall time including loading of the chemicals, polymerizing, cooling, and unloading should be within 8–12 h to allow two or three shifts per day; on the contrary, if the process is operated continuously, the reactor residence time should be as short as possible, once taking into account a suitable reaction time. With regard to the latex concentration, it is desirable to prepare latex at the highest concentration possible to save time in production, unlike most theoretical works in laboratories.²⁰ The upper limit of the maximum solids content is generally the high viscosity of the latex product, which prevents proper stirring and heat transfer, and the reactor temperature operating mode (e.g., the solids content for isoperibolic processes should be under 35%, while for isothermal ones, it can be up to 55%).

Poly(butyl acrylate) (PBA) lattices are an interesting case study to investigate in detail the safe optimization of a generic emulsion polymerization process because its synthesis involves lots of substance loadings, heating and cooling steps, a quite high propagation reaction enthalpy, and a huge tendency to accumulate free radicals.

In this work, the topological criterion theory^{12,13} originally developed for isoperibolic SBRs has been extended to SBRs operated in the isothermal temperature control mode and, then, used to safely optimize the free-radical emulsion homopolymerization of butyl acrylate (BA) thermally initiated by KPS. Particularly, an indirectly cooled SBR (RC1, 1 L, Mettler Toledo) operated in the isothermal control mode (which is safer than the isoperibolic one for fast and exothermic polymerizations) has been chosen for the laboratory tests aimed to validate the developed optimization procedure. Because such an optimization procedure is scale-independent, it can be easily used to identify the optimum operating conditions even for a full-plant reactor.

2. EXTENDED TOPOLOGICAL CRITERION AND OPTIMIZATION PROCEDURE

The topological criterion theory^{12,13} states that, for a semibatch process carried out under isoperibolic temperature control mode, the boundary between “runaway” (RW) and QFS^{21–25} conditions with respect to a desired product X (QFS_X) is identified by an inversion of the topological curve (TC) showing a concavity toward the right. Furthermore, as discussed in detail elsewhere,^{12,13} this curve shows all of the possible thermal behavior regions of an isoperibolic SBR obtainable by varying one system constitutive parameter (e.g., the dosing time) or initial condition (e.g., the initial reactor or coolant temperature) in a suitable range. The TC can be drawn by solving material and reactor

energy balance equations describing the analyzed system for each investigated value of the generating parameter and then reporting onto a 2D diagram the ratio of the obtained reactor temperature maxima to the initial reactor temperature, $\psi_{\max, \text{theo}} = T_{\max, \text{theo}}/T_0$, versus the conversion with respect to the desired product in correspondence with such maxima, $\zeta_X(\psi_{\max, \text{theo}})$. In the presence of only one exothermic reaction (in the polymerization cases, the “global” propagation reaction), such a curve can be generated also experimentally²⁵ by measuring, through a set of isoperibolic tests carried out in a laboratory calorimeter, the maximum reactor temperature, $T_{\max, \text{exp}}$ and the enthalpic conversion in correspondence with such a maximum, $\zeta_{\text{cal, max}} = \int_0^{t_{\max}} \dot{Q}_{\text{rxn}}(t) dt / \Delta \hat{H}_{\text{rxn}} m$.

Moreover, it can be shown that, whenever a TC whose generating parameter is the dosing time exhibits an inversion with concavity toward the left, one of the following system thermal behavior boundaries is encountered: transition (the system thermal loss of control shifts its occurrence from times lower/larger than the dosing period to times larger/lower) or starving (the system state shifts from QFS conditions to low-productivity operating conditions characterized by a squared pseudostationary reactor temperature profile and a forced linear conversion during the dosing period).

It has been also demonstrated²⁶ that the topological criterion theory for QFS detection does not depend on the particular system of ODEs used to describe the process dynamics provided that only two dependent variables contribute effectively to characterize the system thermal behavior: temperature and conversion with respect to the desired product X. Such variables appear in all possible systems of equations writable to describe a generic-complex-controlled process: they can be referred to as “core constants”. Accounting for this feature, dosing stream and jacket temperature control equations, mixing rules for volume determination, global material, and jacket energy balance equations can be added to the system of ODEs describing an isoperibolic SBR without invalidating the topological criterion; this leads to an “extended form” of the topological criterion.

Accordingly, in this paper the topological criterion has been extended to optimize semibatch processes operated in the isothermal temperature control mode. Along the same lines as those used for developing the original topological criterion, the developed optimization procedure is based on analysis of the TC: the QFS inversion is considered to be a boundary beyond which the optimum operating conditions (that is, the dosing time and initial reactor temperature) can be searched, accounting for both the MAT value and all of the desired productivity and quality constraints.

Moreover, because we are dealing with controlled isothermal processes, we expect that before the QFS inversion, a “loss of control” (LoC) happens involving a more general behavior than that identified for isoperibolic SBRs, where the loss of thermal control is always associated with a sharp temperature peak. In controlled isothermal SBRs, a loss of thermal control can lead either to a sharp temperature peak or simply to a loss of the temperature set point. The latter situation is obviously less dangerous than the first one, where a real thermal explosion can arise if the venting system is not able to deal with the reactor pressurization, but it cannot be considered suitable for operating the reactor from neither a product quality nor a safety point of view.

Because no real control systems can maintain the reactor temperature value exactly equal to the set point, it is necessary to define a range around the set point beyond which a LoC is assumed. In this work, a maximum reactor temperature fluctuation of 2.5 °C has been assumed as the threshold value that, if

overcome, evidences a LoC situation independently on the temperature versus time profiles.

The combined theoretical–experimental optimization procedure proposed for a controlled isothermal SBR involves six main steps, and it is summarized in the synoptic of Figure 1.

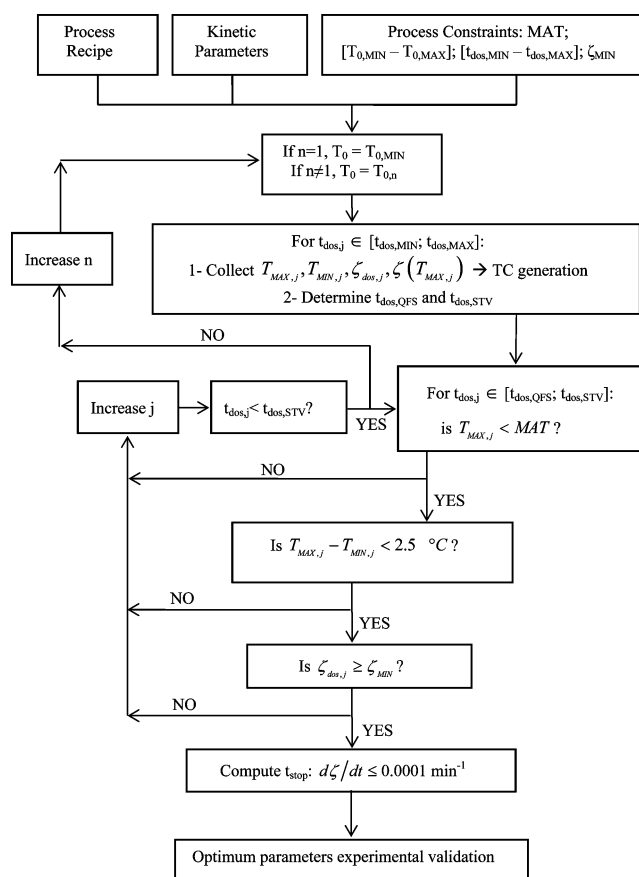


Figure 1. Isothermal optimization procedure scheme.

Step 1. Definition of the recipe. Provided that the reactor topology (semibatch) and the temperature control mode (isothermal) are given, the operational list (that is, the recipe) has to be defined together with the successive computation of the cooling system characteristics (e.g., jacket volume, coolant flow rate, coolant saturation temperatures, etc.). This information will be used for both theoretical simulations and laboratory tests.

Step 2. Execution of calorimetric screening tests. This step allows for both characterization of all substances thermal behavior in the selected temperature range and MAT determination.

In particular, dynamic and isothermal differential scanning calorimetry (DSC) experiments must be performed on reactants and additives to detect if any thermal decomposition or oxidation can be triggered. Moreover, a set of batch isoperibolic syntheses at different coolant temperatures should be carried out to detect the minimum temperature in correspondence with which the reaction mixture, under atmospheric pressure, starts to boil, vigorously forming foams. Batch conditions have been selected because it is known that they are the most dangerous from the thermal stability point of view.

Step 3. Estimation of reaction rate parameters and mass-transfer coefficients. This step can be performed either experimentally (through a data fitting of adiabatic, isoperibolic, or

isothermal tests) or theoretically (looking for the required parameters in the literature).

With regard to the experiments, a single isoperibolic laboratory test (RW and starving conditions must be obviously avoided) must be carried out with RC1 equipment. However, if the isoperibolic test does not cover the full temperature range of interest, either PHI-TEC II or ARC adiabatic tests can be used for to obtain the data needed for kinetic parameter fitting in the entire range of normal and upset operating temperatures. However, if the process is critical from the thermal stability point of view and neither adiabatic nor isoperibolic tests can be carried out safely, also isothermal tests at different initial reactor temperatures can be used for data fitting.

Step 4. Determination of the optimum initial reactor temperature—dosing time couple at the laboratory scale. This is the core of the optimization procedure. It must be carried out through the simulation model until convergence is reached, and it is based on the following substeps.

Substep a. Definition of suitable initial reactor temperature ($T_{0,\min}$ and $T_{0,\max}$) and dosing time ($T_{\text{dos},\min}$ and $T_{\text{dos},\max}$) ranges and selection of the iterating parameter in the optimization procedure and of the generating parameter of the TC. It should be noted that because polymerization reactions require a high product quality (e.g., peaked molecular weight distribution, high average molecular weight, little branching, etc.), which can be obtained only by limiting as much as possible temperature fluctuations, the initial reactor temperatures are forced to vary into a narrow range. For this reason, the initial reactor temperature is not a suitable parameter for generating a TC because the generating parameter must be varied in a sufficiently wide range (conversely, the TC would be incomplete and, consequently, useless). Therefore, the dosing time, which can be varied in a wide range, has been chosen as the generating parameter of the TC, and the initial reactor temperature has been selected as the iterating parameter in the optimization procedure.

Moreover, a minimum threshold value for the conversion of species X at the end of the dosing time, $\zeta_{X,\min}$, has to be defined.

Substep b. Identification of the dosing time values for QFS_X ($t_{\text{dos},\text{QFS}_X}$) and for starving ($t_{\text{dos},\text{STV}}$) conditions in the TC. Through the procedure discussed elsewhere^{12,13} and using a given temperature value (starting from the lower one allowed), a TC has to be generated to bind the QFS operating region (namely, the one whose dosing time ranges between $t_{\text{dos},\text{QFS}_X}$ and $t_{\text{dos},\text{STV}}$), where the optimum dosing time at the current reactor temperature has to be searched for.

Substep c. Verification of the MAT constraint. For increasing dosing times into the QFS operating region, the computed maximum reactor temperatures, $T_{\text{max},j}$, have to be checked for fulfillment of the constraint:

$$T_{\text{max},j} < \text{MAT} \quad (1)$$

If it is found that, for all of the allowed dosing times, $T_{\text{max},j} > \text{MAT}$, the initial reactor temperature should be increased and substep b repeated with this new value.

Substep d. Verification of the LoC constraint. In this substep, it is necessary to check that the following constraint is fulfilled, where $T_{\text{max},j}$ is the computed maximum reactor temperature and $T_{\text{max},j}$ the computed minimum reactor temperature:

$$\Delta T = T_{\text{max},j} - T_{\text{min},j} \leq 2.5^\circ\text{C} \quad (2)$$

If it is found that this constraint is not fulfilled, the dosing time has to be increased and substep c repeated with this new value.

Substep e. Verification of the $\zeta_{X,\min}$ constraint. As a final check, conversion of the target species X at the end of the dosing time, $\zeta_{X,\text{dos},j}$, is required to be larger than a given threshold value, $\zeta_{X,\min}$:

$$\zeta_{X,\text{dos},j} \geq \zeta_{X,\min} \quad (3)$$

If this is not true, the dosing time has to be increased and substep c repeated. If all $\zeta_{X,\text{dos},j} < \zeta_{X,\min}$, the initial reactor temperature should be increased and substep b repeated.

Step 5. Stop time determination. Once the optimum initial reactor temperature and dosing time have been determined, the maximum time at which the process has to be stopped must be identified. Such a time can be defined as that corresponding to conversion variations lower than 0.0001 min^{-1} , that is, to a negligible production rate. Experimentally, such a time can be identified when there are no more reactor temperature fluctuations (that is, temperature fluctuations are within normal temperature background noise). From a practical point of view, in these conditions, the process may be meaningless and the real stop time (lower than both of these two values) should be determined by a cost/benefit analysis.

Step 6. Experimental validation at laboratory scale. Finally, the optimum laboratory initial reactor temperature and dosing time must be experimentally validated through an isothermal RC1 test carried out at these operating conditions.

Once the procedure has been validated for the recipe of interest, it is possible to take advantage of its scale insensitivity to repeat steps 1, 4, and 5 for identifying the optimum operating parameters for the real-scale SBR.

3. RESULTS

As mentioned before, the BA emulsion polymerization case study has been investigated to assess the practical feasibility of the optimization procedure proposed in section 2 and to validate its results.

3.1. Experimental Setup. In order to perform a PBA laboratory synthesis as close as possible to that usually carried out at an industrial scale, RC1 equipment (MP06, 1 L, Mettler Toledo), indirectly cooled by means of an external jacket, has been used with the following operational list: (1) distilled water (W, continuous medium), SLS (surfactant), and sodium carbonate (Na_2CO_3 , buffer) loaded into the reactor and then heated to a reactor set-point temperature (70, 75, or 78 °C) in 35–40 min, activating an isothermal temperature control mode; (2) reaction mixture kept at the set-point temperature for 20–30 min in order to facilitate SLS micelles formation; (3) KPS loaded in one shot into the reactor, providing a waiting time of 20 min to allow for the reactor temperature reequilibration at its set-point value; (4) BA dosed through a pump.

Reactants and reactor characteristics are summarized in Table 1.

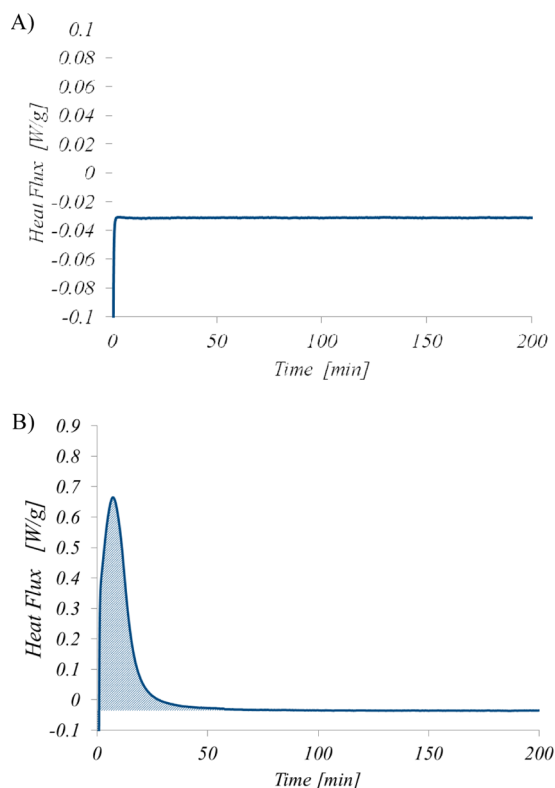
3.2. Calorimetric Screening Tests. In order to characterize the thermal behavior of all of the substances involved in the process and to identify the MAT value, DSC tests and a set of 50 mL scale isoperibolic syntheses at different coolant temperatures have been carried out.

In particular, dynamic DSC tests have been performed on KPS and SLS (as discussed elsewhere²⁷), while a set of isothermal DSC tests have been carried out on BA (Figure 2) using a Mettler Toledo DSC 823 apparatus and stainless steel, medium-pressure Viton/120 μL crucibles. All DSC tests have been performed under a nitrogen atmosphere to avoid eventual oxygen contributions.

It has been found that KPS exhibits an exothermic effect in correspondence with 188 °C. Such a phenomenon can be ascribed

Table 1. Process Recipe and Reactor Characteristics for the Free-Radical Emulsion Homopolymerization of BA Thermally Initiated by KPS

		Initial Load	
283 g	water	0.5 g	Na ₂ CO ₃
3.0 g	SLS	1.2 g	KPS
		Dosed Stream	
200 g	BA		
Reactor and Cooling System			
reactor nominal volume	1.0 × 10 ⁻³ m ³	jacket nominal volume	5.0 × 10 ⁻⁴ m ³
cooling	external jacket	coolant flow rate	5.0 × 10 ⁻⁴ m ³ /s
coolant	silicon oil	UA ₀	2.33–2.43 W/K

**Figure 2.** Isothermal DSC characterization of BA in a nitrogen atmosphere: (A) without initiator, sample amount = 4.98 mg, and investigated temperature = 70 °C; (B) with AIBN as the initiator, sample amount = 2.5 mg, and investigated temperature = 70 °C.

to thermal decomposition of the crystalline solid. On the contrary, SLS exhibits two endothermic effects (the former at about 47 °C and the latter at about 92 °C). Such phenomena correspond to a solid-phase transition between two different crystalline phases and a melting phase, respectively.²⁷

With regard to BA, it can be observed from the isothermal DSC test reported in Figure 2A that no self-initiated polymerization reactions occur at the selected temperature (70 °C) for all of the investigated time; it is necessary to add an organic-soluble initiator (in this case AIBN) in order to observe a polymerization reaction (see Figure 2B). The same behavior is observed at 75 and 78 °C.

With particular reference to the MAT parameter determination, a set of three isoperibolic 50-mL-scale tests at different coolant temperatures, respectively 70, 75, and 78 °C, have been

carried out. Particularly, a 100 mL flask (supplied with a thermocouple connected with an automatic temperature data recorder and a magnetic stirrer) has been inserted into a thermostatic water bath, whose temperature has been maintained at the desired set-point temperature (to realize the isoperibolic mode), and loaded with 24.25 g of a water/SLS emulsion previously prepared. When water bath and flask thermocouple equilibria are reached, 17 g of monomer is loaded into the flask, and a new thermocouple equilibration is expected. Then 0.1 g of KPS are added in one shot.

The main experimental conditions and results are summarized in Table 2, which lists the recipe, set point of the coolant

Table 2. Process Recipe and Coolant and Reaction Mixture Boiling Temperatures for the Investigated 50-mL-Scale BA Emulsion Homopolymerization Synthesis

Recipe			
24.0 g	water	0.10 g	KPS
0.25 g	SLS	17.0 g	BA
0.07 g	Na ₂ CO ₃		
Set of 50-mL-Scale Synthesis			
run	T _{cooling} (°C)	T _{boiling} (°C)	
1	70	85.2	
2	75	86.2	
3	78	85.9	

temperatures for each synthesis, and temperature values at which boiling phenomena (if any) have been observed. It is possible to see that the minimum temperature at which the reaction mixture

Table 3. Constitutive Model and Kinetic Parameters for the Free-Radical Emulsion Homopolymerization of BA Thermally Initiated by KPS^a

A _i	s ⁻¹	8.00 × 10 ¹⁵	28
E _i	kJ/kmol	1.62 × 10 ⁵	28
f		0.60	28
A _p ^s	³ /(kmol s)	2.21 × 10 ⁷	29
E _p ^s	kJ/kmol	1.79 × 10 ⁴	29
A _p ^t	m ³ /(kmol s)	1.52 × 10 ⁶	30
E _p ^t	kJ/kmol	2.89 × 10 ⁴	30
A _{bb}	m ³ /(kmol s)	4.84 × 10 ⁷	30
E _{bb}	kJ/kmol	3.17 × 10 ⁴	30
A _t ^{ss}	m ³ /(kmol s)	1.34 × 10 ⁹	31
E _t ^{ss}	kJ/kmol	5.60 × 10 ³	31
A _t st	m ³ /(kmol s)	2.74 × 10 ⁸	30
E _t st	kJ/kmol	5.60 × 10 ³	30
A _t ^{tt}	m ³ /(kmol s)	1.80 × 10 ⁷	30
E _t ^{tt}	kJ/kmol	5.60 × 10 ³	30
k _{nucl,omo}		6.01 × 10 ⁻³	this work
f _{nucl,mic}		1.00 × 10 ⁻⁵	this work
k _{coal}	m ³ /s	2.80 × 10 ⁻²⁸	this work
D _{diff,wp}	m ² /s	3.91 × 10 ⁻¹⁰	this work
D _{diff,pw}	m ² /s	3.10 × 10 ⁻¹⁰	this work
ΔH _{rxn}	kJ/mol	64.6	this work
ε		0.6964	this work
R _H		0.5613	this work
UA _{ext}		0.4200	this work
α		0.1530	this work
r _{mic}	M	2.50 × 10 ⁻⁹	32
E _{w,sat}	kmol/m ³	2.43 × 10 ⁻⁹	32

^aIn the last column, "this work" means that the value has been estimated by fitting a set of isothermal RC1 data.

starts to boil vigorously is equal to about 85 °C. Such a temperature can be considered, as a first approximation, to be the MAT parameter value because no other side reactions, such as cross-linking reactions, have been observed (these reactions start to occur at temperatures much higher than 85–90 °C and, therefore, they can be disregarded without a loss of reliability of the kinetic model).

3.3. Kinetics Determination. Because the adiabatic temperature rise for this BA emulsion polymerization is equal to about 95 °C, it is not possible to perform an adiabatic RC1 test to determine the reaction kinetics because boiling phenomena would start before the reaction completion. The same risk can be encountered even with an isoperibolic RC1 experiment (see section 3.2 with regard to isoperibolic tests at 50 mL scale); therefore, a series of three isothermal tests at different initial reactor temperatures and dosing times (that is, 70 °C and 60 min, 75 °C and 40 min, and 78 °C and 30 min) have been carried out. Using these RC1 tests and the mathematical model reported in Appendix A, a data fitting of the constitutive model parameters summarized in Table 3 has been performed. It is worthwhile to note that the mathematical model used in this work (see equations in Appendix A) is able to fully represent the polymerization process in an isothermal SBR, including dosing policy, temperature control loop, volumetric mixing rules, and polymer particle radius evolution.

3.4. Process Optimization. Using the model parameters summarized in Table 3, a series of TCs have been generated (see Figure 3) by solving the equations summarized in Appendix

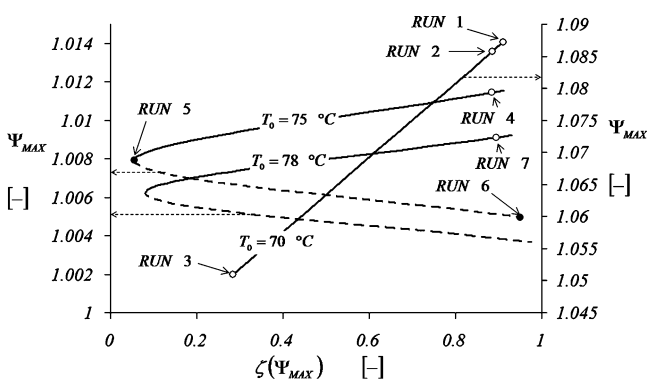


Figure 3. TCs (—, LoC region; ---, QFS region) at different initial reactor temperatures for the free-radical emulsion homopolymerization of BA thermally initiated by KPS. $T_0 = 70, 75,$ and 78 °C, $t_{\text{dos}} = 5\text{--}60$ min, and model parameters as in Table 3. Experimental runs are reported according to the following criterion: (○) LoC run; (●) QFS run.

A for different initial reactor temperatures (that is, 70, 75, and 78 °C, respectively) and using dosing times as TC generating

parameters (from $t_{\text{dos,min}} = 5$ min up to $t_{\text{dos,max}} = 60$ min, with steps of 150 s).

We can see that for $T_0 = 70$ °C neither transition, QFS, nor starving inversions occur in the investigated dosing time range. Such a thermal behavior indicates that the reaction is always carried out under high accumulation conditions because the set point of the reactor temperature is too low and, consequently, LoC operating conditions always dominate and no optimum dosing time can be detected. However, for TC generated at $T_0 = 75$ and 78 °C, there is a clear QFS inversion, in correspondence with which there is a transition between LoC and QFS conditions. No starving inversion has been detected in the investigated dosing time range for analyzed initial reactor temperatures.

After this preliminary evaluation aimed at characterizing the analyzed system thermal behavior, it is possible to perform the optimization procedure previously presented. Therefore, an optimum dosing time has been iteratively searched for, starting from $T_0 = 70$ and 78 °C, by checking constraints (1)–(3) from the dosing time at which the QFS inversion occurs, $t_{\text{dos,QFS}}$ (QFS point in Figure 3) to $t_{\text{dos,max}}$ (because no starvation inversion has been detected). For $T_0 = 70$ °C, no QFS inversion can be detected, and therefore the optimization algorithm continues to increase the initial reactor temperature. For $T_0 = 75$ °C, the QFS boundary is detected for a dosing time equal to 40 min. In correspondence with this point, a local minimum (because it refers to the investigated dosing time range) of the conversion with respect to the desired polymer species has been observed. The minimum dosing time, able to satisfy constraints (1)–(3), has been found to be equal to 60 min. This dosing time can be considered optimized from both safety ($T_{\text{max,theo}} = 76$ °C < MAT = 85 °C) and productivity ($\zeta_{\text{dos,theo}} = 0.937 > \zeta_{\text{min}} = 0.930$) points of view. Moreover, the run can be considered to be controlled (that is, no LoC is evidenced) because $\Delta T \sim 1.5$ °C ≤ 2.5 °C.

3.5. Stop-Time Determination. The time at which the batch period (which follows the dosing period) can be terminated is referred to as the stop time, and it is defined as that corresponding to a conversion variation lower than 0.0001 min^{-1} . In this case, it is equal to about 70 min. Using the aforementioned experimental criterion related to the temperature fluctuations, the stop time has been found to be equal to about 75 min, in fair agreement with the value predicted by the model. Of course, as previously mentioned, the real stop time can be lower than these values, and it must be determined by a cost/benefit analysis.

3.6. Experimental Validation. Both the theoretical TC and optimum ($T_{0,\text{opt}}$ and $t_{\text{dos,opt}}$) couple have been validated through a set of isothermal RC1 runs, whose initial reactor temperature and dosing time values cover the range used to generate the TC.

In particular, seven isothermal experiments, summarized in Table 4, have been carried out. The corresponding experimental

Table 4. Comparison between Model Predictions and Experimental Results for the Free-Radical Emulsion Homopolymerization of BA Thermally Initiated by KPS^a

run	T_0 (°C)	t_{dos} (min)	ΔT (°C)	$\zeta_{\text{max,cal}}$	$\zeta_{\text{dos,exp}}$	exp class	model class	notes
1	70	10	30.2	0.908	0.005	LoC	LoC	strong boiling and foam formation
2	70	20	30.1	0.883	0.000	LoC	LoC	strong boiling and foam formation
3	70	60	16.7	0.286	0.937	LoC	LoC	moderate boiling and foam
4	75	20	4.2	0.913	0.840	LoC	LoC	no boiling, no foam
5	75	40	2.9	0.056	0.934	LoC	LoC/QFS	no boiling, no foam
6	75	60	1.8	0.955	0.936	QFS	QFS	no boiling, no foam
7	78	30	3.3	0.931	0.884	LoC	LoC	incipient boiling, no foam

^aThe consequences of each run have been reported by specifying the level of risk associated with the specific operating conditions.

temperature and enthalpic conversion versus time profiles are shown in Figure 4, while Table 4 also reports, for each experiment,

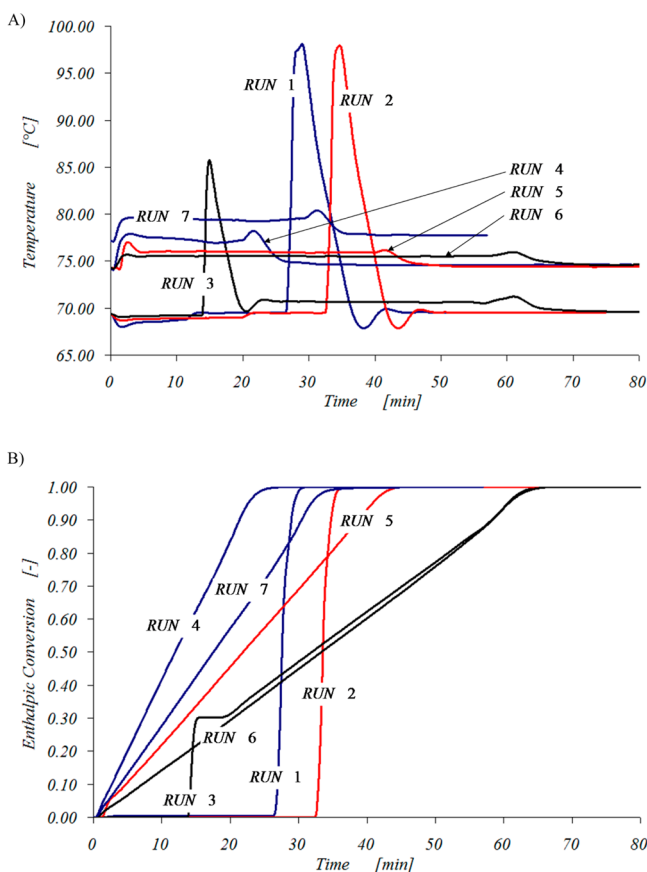


Figure 4. Experimental temperature (A) and enthalpic conversion (B) versus time profiles for the free-radical emulsion homopolymerization of BA thermally initiated by KPS (experimental details as in Tables 3 and 4).

the difference between the maximum and minimum reactor temperatures ($\Delta T = T_{\max, \text{exp}} - T_{\min, \text{exp}}$) and the enthalpic conversion corresponding to both the time at maximum temperature ($\zeta_{\text{cal, max}}$) and the end of the dosing period ($\zeta_{\text{dos, exp}}$). Moreover, the same table also compares the theoretical and experimental classification of each run in terms of thermal behavior (LoC or QFS). Each ($T_{\max, \text{exp}}$ and $\zeta_{\text{cal, max}}$) couple has been located on the diagram reported in Figure 3. From these results, we can see that $T_{0, \text{opt}} = 75 \text{ }^\circ\text{C}$ and $t_{\text{dos, opt}} = 60 \text{ min}$, which are the optimum operating parameter values calculated from the optimization procedure and are identified as optimal also from the experiments because all thermal, productivity, and control definition constraints are fulfilled (see Table 4). Moreover, the boundary between the LoC and QFS thermal behavior is the same from both model and experimental analysis, as can be seen in Figure 3. These cross-validations strongly support the reliability of the obtained results.

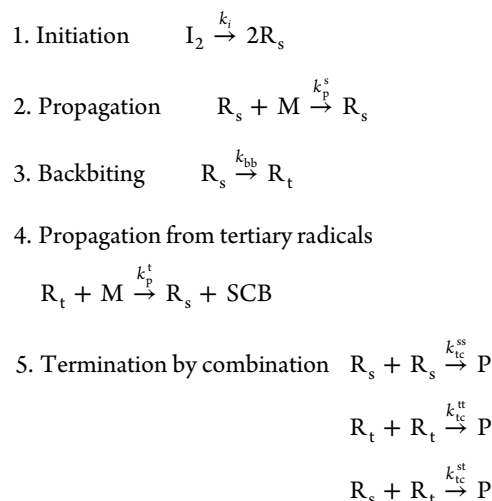
4. CONCLUSIONS

In this work, it has been shown that the topological criterion theory, originally developed for isoperibolic SBRs, can be successfully extended to isothermal reacting systems when providing all of the necessary equations (e.g., the temperature control mode) needed to simulate the whole process. Moreover, a suitable combined experimental–theoretical optimization procedure has been developed and validated using several RC1 tests at different initial reactor temperatures and dosing times. Particularly, the case study

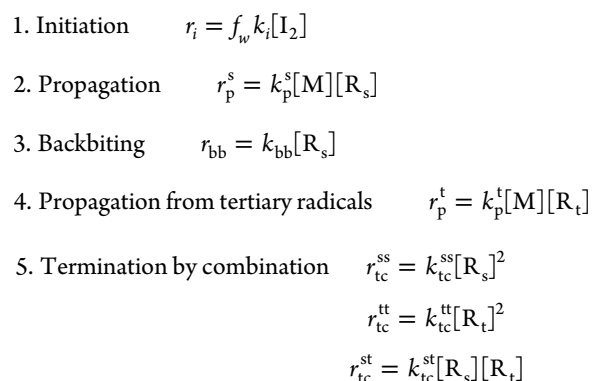
of BA emulsion homopolymerization thermally initiated by KPS has been analyzed showing a good agreement between the theoretical predictions (namely, the shape of the TC and its thermal behavior regions) and the experimental results.

■ APPENDIX A: MATHEMATICAL MODEL

The following kinetic scheme summarizes the main reactions considered in the PBA synthesis:



where I represents the initiator species (in this case, KPS), $R_s = \sum_{n=1}^{\infty} R_{s,n}$ is a pseudoradical that represents all radical species of type s (secondary radicals) independent of their chain length (as usual assumed in the frame of the so-called “terminal kinetic model”, TMK), $R_t = \sum_{n=1}^{\infty} R_{t,n}$ is a pseudoradical that represents all radical species of type t (tertiary radicals) independent of their chain length, M is the monomer (in this case, BA), and P represents the dead polymer chain (neither branching nor molecular weight distribution analysis has been considered according to the TMK theory). For the sake of simplicity, it is possible to consider all reaction rates corresponding to the above reactions independent by the phase (aqueous or polymer) where they occur. According to such a hypothesis, reaction rates can be expressed as follows:



The reactive steps (dosing and curing) can be modeled through the following system of differential–algebraic equations (DAEs; see the Nomenclature section for the meaning of the different symbols): dosing policy (first equation), isothermal temperature proportional–integral–derivative control loop (second equation), energy balances on both the jacket (third equation) and reactor (fourth equation), global material balance (fifth equation), mixing rule (sixth equation), polymer particle concentration (seventh equation), radius evolution (eighth equation), micelle conversion (ninth equation), and all component [initiator (10th equation), radicals (11th through 14th

equations), and monomer (15th equation)] material balances (for each required phase, aqueous and polymeric).

In the time range $0 \leq \vartheta < 1$:

$$\frac{dv_{\text{dos}}}{d\vartheta} = 1$$

$$\frac{d\tau_{\text{cool,in}}}{d\vartheta} = -K_{\text{PT}} \left[\frac{d\tau}{d\vartheta} - \frac{t_{\text{dos}}}{K_{\text{IT}}} (\tau_{\text{set}} - \tau) + \frac{K_{\text{dT}}}{t_{\text{dos}}} \frac{d^2\tau}{d\vartheta^2} \right]$$

$$\frac{d\tau_{\text{cool}}}{d\vartheta} = \theta_{\text{cool}} (\tau_{\text{cool,in}} - \tau_{\text{cool}}) + \text{NTU}\nu (\tau - \tau_{\text{cool}})$$

$$\rho\nu \frac{d\tau}{d\vartheta} = \Delta\tau_{\text{ad}} \frac{d\zeta_{\text{M}}}{d\vartheta} + \frac{dv_{\text{dos}}}{d\vartheta} \varepsilon R_{\text{H}} (\tau_{\text{dos}} - \tau) - St \times \nu (\tau - \tau_{\text{cool}}) - St_{\text{ext}} (\tau - \tau_{\text{ext}})$$

$$\frac{d\nu}{d\vartheta} = \frac{\varepsilon}{\rho} \frac{dv_{\text{dos}}}{d\vartheta} - \frac{\nu}{\rho} \frac{d\rho}{d\vartheta}$$

$$\frac{d\rho}{d\vartheta} = \frac{\varepsilon \frac{dv_{\text{dos}}}{d\vartheta} \left(1 - \frac{(1 - \alpha\zeta_{\text{M}})}{\rho_{\text{dos}}} \right) + \alpha \frac{\varepsilon}{\rho_{\text{dos}}} v_{\text{dos}} \frac{d\zeta_{\text{M}}}{d\vartheta} (1 + \varepsilon\nu_{\text{dos}})}{\left(1 + \frac{\varepsilon}{\rho_{\text{dos}}} v_{\text{dos}} (1 - \alpha\zeta_{\text{M}}) \right)^2}$$

$$\frac{d\eta}{d\vartheta} = Da_{\text{nucl,mic}}^s \zeta_{\text{R}_s} (1 - \zeta_{\text{mic}}) + Da_{\text{nucl,mic}}^t \zeta_{\text{R}_t} (1 - \zeta_{\text{mic}}) + Da_{\text{nucl,omo,bis}}^s \kappa_{\text{p}}^s \zeta_{\text{R}_s} + Da_{\text{nucl,omo,bis}}^t \kappa_{\text{p}}^t \zeta_{\text{R}_t} - Da_{\text{coal}} \eta^2 \sqrt{1 - \zeta_1}$$

$$\frac{d\delta}{d\vartheta} = C_{\delta,s} \kappa_{\text{p}}^s n_s \delta^{-2} + C_{\delta,t} \kappa_{\text{p}}^t n_t \delta^{-2}$$

$$\zeta_{\text{mic}} = \delta\eta$$

$$\frac{d\zeta_1}{d\vartheta} = Da_i \kappa_i (1 - \zeta_1)$$

$$\begin{aligned} \frac{d\zeta_{\text{R}_s}^w}{d\vartheta} = & 2Da_i \kappa_i (1 - \zeta_1) + Da_{\text{p,w}}^t \kappa_{\text{p}}^t \zeta_{\text{R}_t} - Da_{\text{bb,w}} \kappa_{\text{bb}} \zeta_{\text{R}_s} \\ & - 2Da_{\text{tc,w}}^{ss} \kappa_{\text{tc}}^{ss} \zeta_{\text{R}_s}^2 - Da_{\text{tc,w}}^{st} \kappa_{\text{tc}}^{st} \zeta_{\text{R}_s} \zeta_{\text{R}_t} - Da_{\text{diff,wp}}^s \zeta_{\text{R}_s} \\ & \times \eta\delta - Da_{\text{nucl,mic}}^s \zeta_{\text{R}_s} (1 - \zeta_{\text{mic}}) - Da_{\text{nucl,omo}}^s \kappa_{\text{p}}^s \zeta_{\text{R}_s} \\ & + Da_{\text{diff,pw}}^s n_s \eta^2 \delta \end{aligned}$$

$$\begin{aligned} \frac{d\zeta_{\text{R}_t}}{d\vartheta} = & -Da_{\text{p,w}}^t \kappa_{\text{p}}^t \zeta_{\text{R}_t} + Da_{\text{bb,w}} \kappa_{\text{bb}} \zeta_{\text{R}_s} - 2Da_{\text{tc,w}}^{tt} \kappa_{\text{tc}}^{tt} \zeta_{\text{R}_t}^2 \\ & - Da_{\text{tc,w}}^{st} \kappa_{\text{tc}}^{st} \zeta_{\text{R}_s} \zeta_{\text{R}_t} - Da_{\text{diff,wp}}^t \zeta_{\text{R}_t} \eta\delta - Da_{\text{nucl,mic}}^t \\ & \times \zeta_{\text{R}_t} (1 - \zeta_{\text{mic}}) - Da_{\text{nucl,omo}}^t \kappa_{\text{p}}^t \zeta_{\text{R}_t} + Da_{\text{diff,pw}}^t n_t \eta^2 \delta \end{aligned}$$

$$\begin{aligned} \frac{dn_s}{d\vartheta} = & Da_{\text{p,p}}^t \kappa_{\text{p}}^t n_t - Da_{\text{tc,p}}^{ss} \kappa_{\text{tc}}^{ss} \frac{n_s (n_s - 1)}{\delta^3} - Da_{\text{tc,p}}^{st} \kappa_{\text{tc}}^{st} \frac{n_s n_t}{\delta^3} \\ & - Da_{\text{bb,p}} \kappa_{\text{bb}} n_s + C_{\text{diff,wp}}^s \zeta_{\text{R}_s} \delta - C_{\text{diff,pw}}^s n_s \delta \end{aligned}$$

$$\begin{aligned} \frac{dn_t}{d\vartheta} = & -Da_{\text{p,p}}^t \kappa_{\text{p}}^t n_t - Da_{\text{tc,p}}^{tt} \kappa_{\text{tc}}^{tt} \frac{n_t (n_t - 1)}{\delta^3} - Da_{\text{tc,p}}^{st} \kappa_{\text{tc}}^{st} \frac{n_s n_t}{\delta^3} \\ & + Da_{\text{bb,p}} \kappa_{\text{bb}} n_s + C_{\text{diff,wp}}^t \zeta_{\text{R}_t} \delta - C_{\text{diff,pw}}^t n_t \delta \end{aligned}$$

$$\begin{aligned} \frac{d\zeta_{\text{M}}}{d\vartheta} = & Da_{\text{pM}}^s \kappa_{\text{p}}^s \left[[I]_0 \zeta_{\text{R}_s} + \frac{[N_{\text{mic}}]_{\text{w},0}}{N_{\text{A}}} n_s \eta \right] \left[\frac{(v_{\text{dos}} - \zeta_{\text{M}})}{\nu} \right] \\ & + Da_{\text{pM}}^t \kappa_{\text{p}}^t \left[[I]_0 \zeta_{\text{R}_t} + \frac{[N_{\text{mic}}]_{\text{w},0}}{N_{\text{A}}} n_t \eta \right] \left[\frac{(v_{\text{dos}} - \zeta_{\text{M}})}{\nu} \right] \end{aligned}$$

This DAE system has to be solved together with the following initial conditions for $\vartheta = 0$:

$$v_{\text{dos}} = 0, \tau_{\text{cool,in}} = \tau_{\text{cool,in},0}, \tau_{\text{cool}} = \tau_{\text{cool},0}, \tau = \tau_0, \nu = 1,$$

$$\rho = 1, \eta = 0, \delta = 1, \zeta_1 = 0, \zeta_{\text{R}_s} = 0, \zeta_{\text{R}_t} = 0,$$

$$n_s = 0, n_t = 0, \zeta_{\text{M}} = 0$$

Some of the equations above change for $\vartheta \geq 1$ as follows:

$$\frac{dv_{\text{dos}}}{d\vartheta} = 0$$

$$\frac{d\nu}{d\vartheta} = -\frac{\nu}{\rho} \frac{d\rho}{d\vartheta}$$

$$\frac{d\rho}{d\vartheta} = \frac{\alpha \frac{(1 + \varepsilon)\varepsilon}{\rho_{\text{dos}}} \frac{d\zeta_{\text{M}}}{d\vartheta}}{\left(1 + \frac{\varepsilon}{\rho_{\text{dos}}} (1 - \alpha\zeta_{\text{M}}) \right)^2}$$

$$\rho\nu \frac{d\tau}{d\vartheta} = \Delta\tau_{\text{ad}} \frac{d\zeta_{\text{M}}}{d\vartheta} - St \times \nu (\tau - \tau_{\text{cool}}) - St_{\text{ext}} (\tau - \tau_{\text{ext}})$$

Note that radical diffusion (inside and outside the micelle/polymer particles) and homogeneous (in water) and micelle nucleation with polymer particle coagulation phenomena have also been considered.³² Moreover, in the energy balance equation, the following heat contributions have been considered: heat accumulation (left-hand side) responsible for the temperature increase during the process, heat evolution (first term on the right-hand side) accounting for the heat developed by all exothermic reactions (in this case, only the propagation reaction), heat removal due to the dosing stream action (second term on the right-hand side, which disappears for times larger than the dosing one), heat removal due to the cooling jacket action (third term on the right-hand side), and heat removal due to both ambient dispersions and evaporation losses (fourth term on the right-hand side).

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Notes

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NOMENCLATURE

A	preexponential factor [s^{-1} or $m^3 \cdot s / kmol$] or area [m^2]
$C_{\text{diff,pw}}$	$k_{\text{diff,wp},0} \times 4\pi r_{\text{mic}}^2 N_{\text{A}} [I]_0 t_{\text{dos}}$, global constant for radical diffusion from polymer particles to water
$C_{\text{diff,wp}}$	$k_{\text{diff,wp},0} \times 4\pi r_{\text{mic}}^2 N_{\text{A}} [\text{mic}]_0 t_{\text{dos}}$, global constant for radical diffusion from water to polymer particles
C_{δ}	$k_{\text{p,rit}} t_{\text{dos}} [\varepsilon_{\text{mp}} / (1 - \varepsilon_{\text{mp}})] (\rho_{\text{M}} / \rho_{\text{P}}) (1 / N_{\text{A}}) (1 / 4\pi v_{\text{mic}})$, volume increase constant for a polymer particle
\hat{c}_{p}	specific heat capacity [$J / (kg \cdot K)$]
D_{diff}	diffusion coefficient [m^2 / s]

Da_{bb}	$k_{bb,rif}t_{dos}$ Damköhler number for the backbiting reaction
Da_{coal}	$k_{coal}([I]_0V_0N_A)^{1/2}[\text{mic}]_0V_0t_{dos}$ Damköhler number for coalescence of the polymer particles
$Da_{diff,pw}$	$k_{diff,pw}[\text{mic}]_0^2 \times 4\pi r_{mic}^2 t_{dos}/[I]_0N_A$ Damköhler number for radical diffusion from polymer particles to water
$Da_{diff,wp}$	$k_{diff,wp} \times 4\pi r_{mic}^2 [\text{mic}]_0 t_{dos}$ Damköhler number for radical diffusion from water to polymer particles
Da_i	$f_w k_{i,rif} t_{dos}$ Damköhler number for the initiation reaction (in water)
$Da_{nucl,mic}$	$4\pi D_{diff,wp} r_{mic} f_{nucl,mic} [\text{mic}]_0 t_{dos}$ Damköhler number for micellar nucleation
$Da_{nucl,omo}$	$k_{nucl,omo} k_{p,rif} [M]_{w,sat} t_{dos}$ Damköhler number for homogeneous nucleation
$Da_{nucl,omo,bis}$	$k_{nucl,omo} k_{p,rif} [M]_{w,sat} ([I]_0 N_A / [\text{mic}]_0) t_{dos}$ Damköhler number for homogeneous nucleation (variation)
Da_{pM}	$k_{p,rif} t_{dos}$ Damköhler number for the global propagation reaction
$Da_{p,p}$	$k_{p,rif} [M]_{p,sat} t_{dos}$ Damköhler number for the propagation reaction in the polymeric phase
$Da_{p,w}$	$k_{p,rif} [M]_{w,sat} t_{dos}$ Damköhler number for the propagation reaction in the aqueous phase
$Da_{tc,p}$	$(k_{tc,rif}/v_{mic} N_A) t_{dos}$ Damköhler number for the termination reaction in the polymer phase
$Da_{tc,w}$	$k_{tc,rif} [I]_0 t_{dos}$ Damköhler number for the termination reaction in the aqueous phase
E_{att}	activation energy [kJ/kmol]
$f_{nucl,mic}$	micellar nucleation efficiency
f_w	initiator thermal decomposition efficiency (in water)
$\Delta \tilde{H}_{rxn}$	j th reaction enthalpy [J/kmol]
K_{dT}	derivative time [s]
K_{iT}	60, reset time [s]
K_{pT}	6, proportional gain
k	$A \exp(-E_{att}/RT)$, kinetic constant [s^{-1} or $m^3 \cdot s / \text{kmol}$]
k_{coal}	kinetic constant for polymer particle coalescence
k_{diff}	kinetic constant for radical diffusion [m/s]
$k_{nucl,omo}$	constant for homogeneous nucleation
I	initiator species
m	total reacting mass [kg]
M	monomer species
MAT	maximum allowable temperature [$^{\circ}\text{C}$ or K]
MW	molecular weight [kg/kmol]
n	run index, number of monomer units in a radical chain, number of moles [kmol], or number of species units
N_A	6.022×10^{26} , Avogadro's number [kmol $^{-1}$]
NTU	$UA_0 t_{dos} / \hat{p}_{cool} \hat{c}_{p,cool} V_{cool}$, number of transfer units
P	dead polymer chain
R	gas constant = 8314 [J/(kmol·K)] or generic pseudoradical species
R_H	$\hat{c}_{p,dos} / \hat{c}_{p,mix}$ heat capacity ratio
RW	runaway operating conditions
r	particle radius [m]
STV	starving conditions
St	$UA_0 t_{dos} / \hat{p}_0 \hat{c}_{p,mix} V_0$ or Stanton number
St_{ext}	$UA_{ext} t_{dos} / \hat{p}_0 \hat{c}_{p,mix} V_0$ or Stanton number referred to ambient dispersion and evaporation losses
t	time [s]
T	temperature [K]
U	overall heat-transfer coefficient [W/(m 2 ·K)]

V	volume [m 3]
\dot{V}	volumetric flow rate [m 3 /s]
ν	V/V_0 , dimensionless reactor volume
ν_{dos}	$V_{dos}/V_{TOT,dos}$, dimensionless dosing volume

Subscripts and Superscripts

att	activation energy
ad	adiabatic
cool	coolant or external jacket
dos	dosing stream or dosing time
ext	external heat losses
in	inlet stream
i	initiation reaction
j	index of the dosing times
max	maximum value of a quantity or at the maximum value of a quantity
min	minimum value of a quantity or at the minimum value of a quantity
mic	micelles
mix	reaction mixture
p	propagation reaction or polymer particles
QFS	“quick onset, fair conversion, and smooth temperature profile”
rif	reference conditions
S, s	radical species of type s
sat	saturation value
set	desired set point
T, t	radical species of type t
TOT	total amount
w	continuous phase (water)
0	start of the dosing period

Greek Symbols

α	$1 - \hat{p}_M / \hat{p}_P$, volume contraction when monomer is converted in a dead chain
$\Delta \tau_{ad}$	$-\Delta \tilde{H}_{rxn} [M]_{dos} / \hat{p}_0 \hat{c}_{p,mix} T_{rif}$ dimensionless adiabatic temperature rise
δ	r/r_{mic} dimensionless polymer particles radius
ϵ_{mp}	$[M]_{p,sat} MW_M / \hat{p}_M$ ratio between the monomer volume and polymer particle volume when the monomeric phase still exists
γ	E_{att} / RT_{rif} dimensionless activation energy
ϵ	$m_{TOT,dos} / m_0$, relative mass increase at the end of the semibatch period
ζ	dimensionless concentration or conversion of a generic species
η	$[P]_w / [\text{mic}]_0$, dimensionless polymer particle concentration (in water)
κ	$\exp[\gamma(1 - 1/\tau)]$, dimensionless kinetic constant
ρ	\hat{p} / \hat{p}_0 , dimensionless reaction mixture density
\hat{p}	reaction mixture density [kg/m 3]
θ	$\dot{V}_{cool} t_{dos} / V_{cool}$ dimensionless coolant contact frequency into the jacket
ϑ	t/t_{dos} dimensionless time
τ	T/T_{rif} dimensionless temperature
Ψ	T/T_0 , dimensionless temperature with respect to the initial reactor temperature

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