Development of a simulation model for controlling and improving the productivity of batch reactors

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This paper describes the development of a dynamic simulator model for a jacketed batch reactor equipped with a mono-fluid heating/cooling system. The Mono-fluid flows at constant flow-rate through the jacketed reactor. The heating and cooling are assured respectively by electrical resistance and two plate heat exchangers. A detailed description of the equations leading to the development of simulation model is presented. The model is based on writing the equations of the mass balance and the heat balance for the reactor and the thermal loop in unsteady state. To validate the simulation model, we first studied the thermal behavior of the reaction mixture during heating and cooling, using water as the reaction mixture. We then considered the consecutive chemical reaction of the synthesis of cyclopentanediol from cyclopentadiene by studying the yield of this reaction. The results show that heating the reaction mixture increases significantly the yield of this synthesis reaction.

**Keywords:** batch chemical reactor, thermal control, dynamic simulator, mono-fluid, heating/cooling system.

**INTRODUCTION**

Recent researches shows that dynamic process simulation is increasingly used in the pharmaceutical industry and the fine chemicals industries. Typically, these industries use batch processes, where exothermic reactions are involved. In such scenarios, the thermal behavior of the system has a strong influence on the reaction selectivity and the safety of the process. Therefore, the correct control of the reactor temperature is fundamental. The dynamic simulation allows researchers to study alternative equipment systems of heat transfer and chemical reactions. Its use extends more and more in these industries of chemicals of high added value.

Process simulation is now a well-established technique and which is, frequently, used in many problems in process engineering. It makes it possible to deal with important issues such as risk reduction, improving the understanding of physical phenomena, sustainability knowledge, reducing costs by optimizing the performance or the reduction of time placing on the market of new products. In order for a simulator to be effective in all these areas, it is advisable that it uses models of knowledge (or phenomenological) constructed on the basis of the fundamental laws of physics and chemistry: the conservation equations (mass, energy, momentum), thermodynamics, kinetics, ... These equations have the advantage to provide a real prediction capacity in a domain of validity much more spread than the one for which data experiments are available.

However, the immediate advantage that can be expected to make an effective dynamic simulation in real processes is to verify the control strategy process. This is why the first industries that have incorporated dynamic simulation as a tool active were oil and refineries, where the control and preservation of production processes within a very small interval of operation provide a continuous stream of the product at a low added value type. A dynamic simulation requires the simultaneous solution of algebraic and differential equations, so the adequate mathematical algorithms need to be incorporated into software. Such software has already the rigorous implemented model and mathematical algorithms ready to be used, which requires the user to enter the information relevant to chemical processes to study. In this context, the dynamic simulation becomes easy to use.

This work, thus, joins within the framework of the thermal conduct of multi-batch reactors used in industry of fine chemicals and specialties. The main tool for the implementation of chemical syntheses in the field of fine chemicals and pharmaceutical remains the batch reactor. The studies led to the elaboration of methodologies allowing the development of modus operandi adapted to the functional constraints and safety of these devices such as thermal control of batch reactors or fed-batch. However, these reactors, even if they offer features flexibility and versatility to present a number of technological limitations. In particular, poor conditions of evacuation of the heat released by chemical reactions pose a serious security problem.

The study in this paper aims to contribute to the development of a dynamic simulator for thermal control of a one-liter batch reactor equipped with a mono-fluid heating/cooling system. This system uses only one fluid for heating and cooling of the reactor contents. The heating and cooling are provided by the implementation on the thermal circuit of an electrical resistance and two heat exchangers of different cooling capacity.

**PROCESS DESCRIPTION**

A schematic diagram of the pilot plant is depicted in Fig. 1. The experimental device consists of a one-liter jacketed glass reactor, fitted with a mono-fluid heating/cooling system. The mono-fluid used in this work is a mixture of ethylene glycol and water, in a ratio of 50% in weight, with a flow rate of 1000 l h⁻¹ and at a temperature which varies between ~35 and 110°C. The mono-fluid flow rate is measured by means of two flow-meters, one installed on the main thermal loop (Flow-meter 1) and the other on the secondary thermal loop (Flow-meter 2). The heating/cooling system uses a 2000 W electrical resistance and two plate heat exchangers (PHE). One PHE uses cold...
water as a utility fluid at a temperature around 15°C and a maximum flow rate of 1500 l h\(^{-1}\), while the other uses a mixture of ethylene glycol and water, in a ratio of 50% in weight, at a temperature around -10°C and a maximum flow rate of 1500 l h\(^{-1}\). The flow rates of the utility fluids are also measured. Three on-off valves allow the mono-fluid to be heated or cooled. Two other on-off valves are used to manipulate the utility fluids. A three-ways air-to-open valve ensures the division of the mono-fluid in two-parts during the cooling phases. A gearing pump ensures the circulation of the mono-fluid in the thermal loop at maximum flow rate of 1500 l h\(^{-1}\).

The reactor has the following physical specifications: an internal diameter of 82 mm, a wall thickness of 9 mm, an external jacket diameter of 125 mm, a jacket wall thickness of 5 mm, a maximal reactant mixture-reactor heat transfer area of 0.039 m\(^2\) and a jacket volume of 0.15 l. A propeller rotated at 260 rpm. The reactor is operated in batch and fed-batch modes. A piston pump allows the variation of the liquid reactant flow rate from 0 to 336 cm\(^3\) h\(^{-1}\).

**Figure 1.** Pilot-plant of the reactor and its thermal environment

**PROCESS MODELING**

**Modeling of the reactor**

For a system involving nc species (subscript i) and nr reactions (subscript j), the model has been established using the following assumptions:

- the reactor is perfectly mixed with homogeneous temperature in the reaction mixture;
- the physicochemical properties of each component of the reaction mixture are constant in the considered range of temperature, and the physical properties of the reaction mixture are revalued at each computation step as functions of temperature;
- feeding of the reactive is performed without volume concentration;
- the reactions are carried out in a pseudo-homogeneous liquid phase;
- the kinetic laws of the chemical reactions involved are assumed to be such that the reaction rate constant follows the Arrhenius law and the reaction rate is a function of the reactant concentrations.

For the reaction j, the reaction rate is given by:

\[
r_j = k_j^0 \exp\left(-\frac{E_j}{RT_j}\right)\prod_{i=1}^{nc} C_i^{\alpha_{ij}}
\]

(1)

For each reaction, the activation energy \(E_j\), the rate constant \(k_j^0\), the partial orders \(\alpha_{ij}\), and the stoichiometric coefficients \(C_i(i,j)\) (> 0 for products and < 0 for reactants) have to be given. The dynamic simulation model is obtained by writing mass and heat balances for the reaction mixture, the reactor wall, the mono-fluid flowing inside the jacket and the jacket wall, electrical resistance, plate heat exchangers.

**MASS AND HEAT BALANCE ON THE REACTION MIXTURE**

A molar balance on each of the nc components yields nc differential equations:

\[
\frac{dn_i}{dt} = f_c x_i + \sum_{j=1}^{nr} C_i(i,j) r_j \frac{m_r}{\rho_r} 10^3 \quad i = 1, \ldots, nc
\]

(2)

The mass of the reaction mixture is given by:

\[
m_r = \sum_{i=1}^{nc} n_i M_i
\]

(3)

The density of the reaction mixture (\(\rho_r\)) is estimated by assuming that the various components form a perfect mixture. The total volume \(m_r/\rho_r\) is equal to the sum of partial volumes, by:

\[
\frac{1}{\rho_r} = \sum_{i=1}^{nc} \frac{n_i M_i}{\rho_i}
\]

(4)

i.e.:

\[
\frac{1}{\rho_r} = \frac{1}{\rho_c} + \sum_{i=1}^{nc} \frac{n_i M_i}{\rho_i}
\]

(5)

Classical mixing laws give the physico-chemical properties of the mixture, according to:

\[
C_{pi} = \frac{1}{m_r} \sum_{i=1}^{nc} n_i M_i C_{pi}
\]

(6)

\[
\lambda_r = \frac{1}{m_r} \sum_{i=1}^{nc} n_i M_i \lambda_i
\]

(7)

\[
\mu_r = \frac{1}{m_r} \sum_{i=1}^{nc} n_i M_i \mu_i
\]

(8)

Values of \(C_{pi}\), \(f_c\), \(\lambda_i\), and \(\mu_i\) (properties of pure component i) are assumed to be constant over the range of considered temperatures, and are given as the data to the simulation model.

The heat balance on the reaction mixture gives the following differential equation:

\[
\frac{d(H_r(T_r))}{dt} = -U_{r,pr} A_{r,pr} (T_r - T_{pr}) + f_c M_c H_c(T_c)
\]

(9)

Let us take as the basis for the enthalpy computation, the enthalpy of pure liquid component at temperature \(T^*\). Then \(H_r(T_r)\) is expressed as:

\[
H_r(T_r) = \sum_{i=1}^{nc} n_i M_i H_i(T_r) = \sum_{i=1}^{nc} n_i M_i \left[ h_i(T^*) + C_p i (T_r - T^*) \right]
\]

(10)

After development, the final expression is obtained:

\[
\frac{m_r C_p r}{dt} = -\sum_{j=1}^{nr} r_j \frac{m_r}{\rho_r} \Delta H_{fr}(T_r) 10^3 - U_{r,pr} A_{r,pr} (T_r - T_{pr}) + \frac{f_c M_c}{\rho_c} C_p(T_c(T_r - T_c))
\]

(11)

With:

\[
\Delta H_{fr}(T_r) = \sum_{i,j} C_{pi}(T^*) C_{pj}(T_r - T^*)
\]

(12)
MODELLING OF THE JACKET

The jacket is modelled by means of a succession of N tanks supposed perfectly stirred (N being the number of tanks where the mono-fluid is in direct contact with the reactor wall and the jacket). The heat exchanges between the mono-fluid in the tank (i), the reactor wall, the wall of the jacket, the tank (i-1) and the tank (i+1) are schematized in Fig. 2.

Figure 2. Schematic of heat transfer in the tank (i) of the jacket

The heat balance on the mono-fluid in the tank (i) is given by the following equation:

\[
\frac{d[m_i \cdot Cp_i \cdot T_i]}{dt} = \left( F_i \cdot (p_i \cdot Cp_i \cdot T_i - p_{i-1} \cdot Cp_{i-1} \cdot T_{i-1}) + U_{i,i-1} \cdot \frac{A_{i,i-1}}{N} (T_i - T_{i-1}) - \sum_{j=1}^{N} U_{i,j} \cdot \frac{A_{i,j}}{N} (T_i - T_j) \right)
\]  

(13)

With:

\[
m_{i} = \rho_{i} \cdot \frac{V_{i}}{N}
\]  

(14)

HEAT BALANCE ON THE REACTOR WALL

The heat balance on this wall is defined by the following equation:

\[
\frac{d[U_{reactor} \cdot A_{reactor} \cdot (T_{wall} - T_{reactor}) + \sum_{j=1}^{N} U_{i,j} \cdot \frac{A_{i,j}}{N} (T_i - T_j)]}{dt} = F_i \cdot (p_i \cdot Cp_i \cdot T_i - p_{i-1} \cdot Cp_{i-1} \cdot T_{i-1}) + U_{i,i-1} \cdot \frac{A_{i,i-1}}{N} (T_i - T_{i-1}) + \sum_{j=1}^{N} U_{i,j} \cdot \frac{A_{i,j}}{N} (T_i - T_j)
\]  

(15)

HEAT BALANCE ON THE JACKET WALL

The jacket wall exchanges heat with mono-fluid, flowing inside the jacket and the ambient surroundings. A heat balance gives the following equation:

\[
\frac{d[m_{j} \cdot Cp_{j} \cdot T_{j}]}{dt} = U_{j} \cdot A_{j} \cdot (T_{j} - T_{ambient}) + \sum_{j=1}^{N} U_{i,j} \cdot \frac{A_{i,j}}{N} (T_i - T_j)
\]  

(16)

Modelling of the thermal loop

MODELLING OF HEATING PENCIL

The heating pencil is formed of a steel stem inside which is placed an electrical resistance, the mono-fluid flows in an annular space formed by the stem and the electric resistance as shown in Fig. 3.

To model the heating pencil, we have considered the annular space as a stirred block in which the mono-fluid exchanges heat with the wall of the stem and receives a heat flow resulting from the electrical resistance.

a / Heat balance on the wall of the electrical resistance

The heat balance on this wall, which exchanges heat with the mono-fluid and instantly receives electric power, is given by the following equation:

\[
\frac{dT_{wall}}{dt} = \frac{P_{electric} + U_{f,i} \cdot A_{f,i} \cdot (T_{wall} - T_{input})}{m_{f,i} \cdot Cp_{f,i}}
\]  

(17)

b / Heat balance on the mono-fluid

The balance of the mono-fluid involves the heat exchange with the walls of the electrical resistance and that of the stem and the amount of heat exchange between the input and the output of the tank is given by the following equation:

\[
\frac{dT_{mono}}{dt} = \left( \frac{U_{f,i} \cdot A_{f,i} \cdot (T_{input} - T_{wall}) + U_{f,i+1} \cdot A_{f,i+1} \cdot (T_{output} - T_{wall})}{m_{f,i} \cdot Cp_{f,i}} \right)
\]  

(18)

With: \( m_{f,i} = \rho_{f,i} \cdot V_{f,i} \)

The heat balance on this wall which exchanges heat with mono-fluid and the ambient surroundings is given by the following equation:

\[
\frac{dT_{wall}}{dt} = \left( \frac{U_{j} \cdot A_{j} \cdot (T_{wall} - T_{ambient}) + \sum_{j=1}^{N} U_{i,j} \cdot \frac{A_{i,j}}{N} (T_i - T_j)}{m_{j} \cdot Cp_{j}} \right)
\]  

(19)

MODELLING OF PLATE HEAT EXCHANGERS

Both heat exchangers used are plate heat exchangers (PHE). This type of exchanger is characterized especially for its good heat transfer. Seen their importance, several authors have contributed to the modelling of these exchangers\(^6, 7, 8\).

– Differential equations relative to the model of the heat exchanger

We consider a differential element of the plate heat exchanger which includes a volume element of the liquid and a volume element of the plate as shown in Fig. 4.
Starting from the fundamental principle of conservation of thermal energy, the following equations describe the heat transfer between the fluid (mono-fluid or utility cooling) flowing through a channel $j$ and two plates that cover:

$$V_j \rho_j C_p \left[ \frac{dT_j}{dt} + n_j u_j \frac{\partial T_j}{\partial z} \right] = U_{j,i} A_{j,i} (T_{p,j,i} - T_j) + U_j A_j (T_p - T_j) \quad (20)$$

For the plate $j$ located between the two fluids (mono-fluid and cooling utility):

$$V_{p} \rho_{p} C_{p} \frac{dT_{p}}{dt} = U_{j,i} A_{j,i} (T_{p,j,i} - T_{p}) + U_j A_j (T_p - T_j) + V_{p} \rho_{p} \frac{\partial T_{p}}{\partial z} \quad (21)$$

The main advantage of this model is that it allows to describe the mechanism of heat transfer inside the heat exchanger whatever the mode of arrangement chosen (in series, loop or complex)\(^3\). As in our case the type of arrangement of the two fluids flow inside the heat exchanger loop is against the current.

Both derivatives can be expressed by discretization according to $z$, what gives equations as the following ones:

$$V_j \rho_j C_p \left[ \frac{dT_{j,i}}{\Delta z} + n_j u_j \frac{T_{j,i} - T_{j,i-1}}{\Delta z} \right] = U_{j,i} A_{j,i} (T_{p,j,i} - T_{j,i}) + U_j A_j (T_p - T_j) \quad (22)$$

For the plate $j$:

$$V_{p} \rho_{p} C_{p} \left[ \frac{dT_{p,i}}{\Delta z} \right] = U_{j,i} A_{j,i} (T_{p,j,i} - T_{p,i}) + U_j A_j (T_{p,i} - T_{p}) + V_{p} \rho_{p} \frac{T_{p,i} - T_{p,i-1}}{\Delta z} \quad (23)$$

**MODELING PIPES**

a / Heat balance of the mono-fluid in each tank pipe $j$ ($j = 1, \ldots, 12$)

The heat balance translates the heat exchanges in the tank (i) of a pipe (j) between the mono-fluid and the wall of this pipe on one hand, and the tank (i) and the tank (i-1) on the other hand.

$$m_j' \rho_j C_p \frac{dT_{cond,i}}{dt} = U_{cond,i} A_{cond,i} \left[ \frac{T_{cond} - T_{i}}{n_i} \right] + D_j \rho_j C_p \frac{T_{cond} - T_{cond,i}}{n_i} \quad (24)$$

With: $m_j' = \rho_j V_{cond,i}/n_i$

b / Heat balance on the pipe wall $j$ ($j = 1, \ldots, 12$)

The heat balance on the wall of the pipe $j$, which exchanges heat with the mono-fluid and the ambient surroundings, is determined by the following equation:

$$m_{j,cond} C_p \frac{dT_{cond,i}}{dt} = \sum_{i=1}^{n_i} U_{cond,i} A_{cond,i} \left[ \frac{T_{cond} - T_{i}}{n_i} \right] + U_{j,cond} A_{j,cond} (T_{j} - T_{cond,i}) \quad (25)$$

**MODELLING OF THE THREE-WAY VALVE**

For its modelling, we consider that this valve is a mixture tank of two currents.

The heat balance is given by the following linear equation:

$$F_p \rho_p C_p \left[ T_{cond} - T_{cond} \right] = F_p \left( \cdot - \rho \right) + \int F_p \rho_p C_p \left[ T_{cond} - T_{cond} \right] \quad (26)$$

From this balance, and taking as reference temperature $T_{ref}$, the temperature at the outlet of the valve $T_{out}$, we obtain the following relation:

$$T_{out} = \left( 0 - \rho \right) + \int F_p \rho_p C_p \left[ T_{cond} - T_{cond} \right] \quad (27)$$

**STRATEGY FOR THE DIGITAL RESOLUTION**

Our model consists, on the one hand, of ordinary differential equations of the following form:

$$18 + nc + N + \sum_{i=1}^{n_i} (2. n_i)$$

differential variables (Mass and heat balances), and on the other, of algebraic equations (heat transfer coefficient, physical properties ...).

The strategy for solving such a problem, we chose the method of Gear. The principle of the Gear method is to transform the system of differential and algebraic equations into a system of nonlinear algebraic equation\(^9\),\(^10\).

**PROGRAMMING METHOD**

We used the Fortran language to simulate our model. The program is conversed in several modules or subroutines. The organizational dynamic simulation of thermal conduct of a batch reactor with jacket is given in Fig. 5.

**VALIDATION OF THERMAL MODEL**

To validate the simulation model, we first studied the thermal behavior of the reaction mixture during heating and cooling, using water as the reaction mixture. We register the evolution of the temperature of the reaction mixture the inlet and outlet of the jacket over time. Then we examine the consecutive chemical reaction of the synthesis of cyclopentanediol from cyclopentadiene by studying the yield of this reaction. We record the changing concentrations of the constituents of the reaction mixture and the temperature thereof over time.

**Validation of the simulator with pure water**

These operations consist in introducing 0.7 liters of water into the reactor. The heating and the cooling begin after 120 seconds. For heating, we take powers of 1000 W and 2000 W. For cooling, we take the same flow rates of mono-fluid in both heat exchangers with cold water and glycol water to 250 l/h and 750 l/h. The evolutions of the temperatures of the reaction mixture (Tr) as well as in the input (Tede) and in the output (Tsde) of the jacket are shown in Figs. 6, 7 by heating simulations and in Figs. 8, 9, by the simulation of cooling in the case of the exchanger with cold water and finally in Figs. 10, 11 for the simulation of cooling by using the heat exchanger with glycol water.
Temperature of the reaction mixture and the temperature of the jacket are very close. For example with heating by 1000 W the established regime is obtained at about 5000 seconds with a temperature of about 75°C, Fig. 6;

– The operation of heating or cooling makes even faster as we increase the electric power for the heating (one gains around 15°C for an increase of 500 W) and monofluid flow rate entering the heat exchanger.

– For cooling, the reaction mixture temperature decreases towards lower values with the glycol water heat exchanger than the heat exchanger with cold water because the temperature of the glycol water can be reduced to –10°C while the temperature water achieves 15°C.

– The difference in temperature between the reaction mixture and the inlet and outlet of the jacket increases...
We compare our simulation results with the experimental results of Bouhenchir\textsuperscript{11, 12}, when heated by the power of 1000 and 2000 W of 0.7 liters of pure water injected into the reactor during the cooling by the heat exchanger with cold water with a flow rate of mono-fluid entering exchanger of 250 and 750 l/h and during the cooling by the heat exchanger with glycol water with a flow rate of the mono-fluid of 250 and 750 l/h.

The comparison of simulation results and experiments for the three operations: heating, cooling by cold water and cooling by glycol water shows that our thermal model of simulation reproduces approximately the real dynamic behaviour of the reactor. However, we observe a slight difference especially at the beginning, which can be explained by the fact that the fluid temperature is experimentally stabilized after a while functioning time. In the results of Bouhenchir, one sees that the temperature changes even during the introduction of reagents we call time of casting. On the other hand, we cannot take this evolution by the simulator because we integrate after the entry of the cast. We notice a difference of 4°C during heating and 2°C during cooling between our results and the experimental results of Bouhenchir\textsuperscript{11, 12} for the temperatures of the reaction mixture (Tr), the entry (Tede) and output (Tsde) of the jacket. The differences observed between our simulations and the experimental results of Bouhenchir can be linked to calculation errors resulting from the transformation of a continuous problem to problem discontinuous. On the other hand, it seems to us that we used correlations and expressions for calculating the physical properties of fluids and heat transfer coefficients different to those used by Bouhenchir\textsuperscript{11, 12}.

**Validation of the simulator with a chemical synthesis**

Another example of application of this simulator is the chemical synthesis of cyclopentanediol from cyclopentadiene which can be illustrated as follows:

\[
C_2H_6 + H_2O \rightarrow C_2H_5OH
\]

\[
C_2H_5OH + H_2O \rightarrow C_2H_8(OH)_2
\]

\[
C_2H_6 \rightarrow 0.5C_{10}H_{12}
\]

The reactions are controlled by an Arrhenius law whose kinetic parameters are summarized in Table 1:

| Table 1. The kinetic parameters of the three synthesis reactions of cyclopentanediol\textsuperscript{13} |
| --- | --- | --- |
| Reaction (1) | Reaction (2) | Reaction (3) |
| Frequency factor \(k_0\) \([\text{mol}^{-1}\text{ s}^{-1}]\) | \(3.575 \times 10^6\) | \(3.575 \times 10^6\) | \(2511.94 \times 10^7\) |
| Activation energy \(E_a\) \([\text{J mol}^{-1}]\) | 81.189 | 81.189 | 71.219 |

The physical properties of the constituents involved in the three reactions are presented in Table 2.

To simulate well this reaction, we took two types of operations.

**SIMULATION WITH CASTING**

The principle of the first operation consists in introducing an initial load of 0.6 kg of water into the reactor in the ambient temperature then fed to the reactor by a casting of 0.2 kg of cyclopentadiene for 100 seconds at a temperature of 57°C.
The results obtained by the simulator in terms of evolution of the concentrations in the reaction mixture and the temperature during heating, during the cooling and without heating and cooling are presented in Figs. 12, 13, 14, 15.

From the three Figures 12, 13, 14 representing the evolution of the concentrations of constituents of the reaction mixture during the heating, cooling and without these two operations, we note that:

– The yield of cyclopentanol (the product of the first reaction) is of the order of 1.7 mol/l for cooling, Fig. 13, and 3.5 mol/l without heating and cooling as shown in Fig. 14 and achieved 3.7 mol/l for heating, Fig. 12.

– The yield cyclopentanediol (the desired product by conversion of the cyclopentanol) is of the order of 1.5 mol/l without heating and cooling Fig. 14, while it achieves 2.5 mol/l for heating Fig. 12.

– Appearance of the third reaction (produced unwanted), which occurs when the cyclopentadiene is in excess.

– The three reactions are faster during the heating that can be achieved after about 800 seconds in Fig. 12, while without heating and cooling exceed 2000 seconds Fig. 14, whereas for cooling over 2000 seconds.

So we can say that the synthesis of cyclopentanediol from cyclopentadiene is better and faster when heating the reaction mixture during the heating, cooling and without these two operations, we note that:

– The yield of cyclopentanol (the product of the first reaction) is of the order of 1.7 mol/l for cooling, Fig. 13, and 3.5 mol/l without heating and cooling as shown in Fig. 14 and achieved 3.7 mol/l for heating, Fig. 12.

– The yield cyclopentanediol (the desired product by conversion of the cyclopentanol) is of the order of 1.5 mol/l without heating and cooling Fig. 14, while it achieves 2.5 mol/l for heating Fig. 12.

– Appearance of the third reaction (produced unwanted), which occurs when the cyclopentadiene is in excess.

– The three reactions are faster during the heating that can be achieved after about 800 seconds in Fig. 12, while without heating and cooling exceed 2000 seconds Fig. 14, whereas for cooling over 2000 seconds.

So we can say that the synthesis of cyclopentanediol from cyclopentadiene is better and faster when heating the reaction mixture. One can see it clearly from Fig. 15 which shows a comparison between the evolution of the temperature of the reaction mixture during heating and cooling and without heating and cooling. We observe an increase of the temperature of the reaction in the three operations to reach 102°C during the heating. This point corresponds to a maximum conversion in the three reactions. This can be explained as follows: in the first reaction which is an endothermic reaction, the quantity of heat entrante with the cyclopentadiene introduced in 57°C is consumed to produce cyclopentanol. As two other reactions are exothermic and as they take place at the same time as the first reaction, we observe a slight increase in the reactor temperature which does not exceed the 57°C during cooling but which explains the low yield of these reactions when cooling. We have then a drop in the temperature of the reaction mixture even during heating because the temperature in the reactor exceeds the temperature of the mono-fluid in the jacket.

| Physical parameters of the various constituents$^{13}$ | Cyclopentadiene | Cyclopentanol | Cyclopentanediol | Dicyclopentadiene |
|-------------------------------------------------------|-----------------|---------------|-------------------|-------------------|
| Molar mass $M_i$ [kg mol$^{-1}$]                       | 0.066           | 0.084         | 0.102             | 0.132             |
| Density $\rho_i$ [kg m$^{-3}$]                         | 934.2           | 934.2         | 934.2             | 934.2             |
| Specific heat $C_p$ [kJ kg$^{-1}$ K$^{-1}$]            | 3.0096          | 3.0096        | 3.0096            | 3.0096            |
| Dynamic viscosity $\mu$ [kg m$^{-1}$ s$^{-1}$]        | 0.001           | 0.001         | 0.001             | 0.001             |
| Thermal conductivity $\lambda$ [kJ m$^{-1}$ s$^{-1}$ K$^{-1}$] | $6.688 \times 10^4$ | $6.688 \times 10^4$ | $6.688 \times 10^4$ | $4.18 \times 10^4$ |
| Molar enthalpy of formation at 25 °C h$^{-1}$ [kJ mol$^{-1}$] | $-92.378$ | $-425.599$ | $-412.691$ | $-285.79$ |

Figure 12. Simulation of heating the reaction mixture by 2000Watt

Figure 13. Simulation of cooling the reaction mixture by glycol water

Figure 14. Simulation of the reaction mixture without heating and cooling

Figure 15. Comparison of temperatures of the reaction mixture during heating, during cooling and without these two operations
by effect of the two exothermic reactions. Therefore, there will be a heat exchange between the reactor and the jacket to achieve thermal equilibrium.

SIMULATION WITHOUT CASTING

The operation without casting consists in making load of the reactor by two reagents without casting. We tested a load of 0.6 kg of water and 0.2 kg of cyclopentadiene at 57°C then at 77°C then a load of equimolar 0.4 kg of water and 0.4 kg of cyclopentadiene to 57°C. The simulations were performed with heating the reaction mixture by the power of 2000 watts. The results obtained by the simulator in terms of concentrations evolution in the reaction mixture and from the temperature during heating are presented in Figs. 16, 17, 18, 19, 20.

Figure 16. Simulation of heating the reaction mixture without casting reagents 57°C

Figure 17. Simulation of heating the reaction mixture without casting reagents 77°C

According to Figs. 16, 17 and 18 of evolution of the concentrations of the constituents of the reaction mixture without reagent casting at 57°C and 77°C and with a load of 0.6 kg of water and 0.2 kg of cyclopentadiene and in 57°C with an equimolar load (0.4 kg of water and 0.4 kg of cyclopentadiene) by heating the reaction mixture, we notice the total conversion of cyclopentadiene in cyclopentanol for the three cases. We also notice a total conversion of cyclopentanol to cyclopentanediol in the case of equimolar reagents which gives approximately 8 mol/l of cyclopentadiene and reagents for the case of non-equimolar in 77°C of the order of 4 mol/l. While in the case of non-equimolar reagents in 57°C we have a concentration of 3 mol/l cyclopentanediol and the rest of the cyclopentanol of 1 mol/l. We also notice faster reactions by the simulation of heating without casting from the casting simulation with heating of reagent. We also see the absence of the intermediate reaction which is of for the total conversion of cyclopentadiene in cyclopentanol in the three figures. The comparison of the temperatures of these three cases in Fig. 19 shows...
that the good yield accompanies the highest maximum temperature because of the large amount of heat released during the second reaction.

CONCLUSION

This paper presents the development of a dynamic simulation model of batch chemical reactor. This model is based on writing the equations of the mass balance and the heat balance for the reactor and the thermal loop in unsteady state.

The validation of the simulator is an important step to test the performance of this simulator. In this article, validation was performed by two examples:

– The study of the thermal behavior of the reaction mixture during the heating and cooling water as reaction mixture. The validation showed that the simulator faithfully reproduces the real behavior of the process (5% difference).

– The second validation is to introduce the chemical reactions of the synthesis of cyclopentanediol from cyclopentadiene in the reactor and study the yield of these reactions in simulations of heating and cooling the reaction mixture.

The results show that the heating of the reaction mixture significantly increases the yield of this synthesis reaction.

We can conclude that this synthesis will be better applied in a continuous reactor in a batch reactor, that due to the rapid reaction especially when heating the reaction mixture and reagents without casting. This conclusion can be considered as an advantage for our simulator to choose the suitable reactor for chemical synthesis. Chemical synthesis is a tool to validate our simulator and shows the importance of temperature control in a chemical reaction.

The simulator is a valuable tool that describes the dynamic evolution of the reactor and thus limits the number of experiments.

NOMENCLATURE

- $A_{fc, pde}$ heat transfer area between the mono-fluid and the jacket wall (m$^2$)
- $A_{fc, fc}$ heat transfer area between the mono-fluid and the reactor wall (m$^2$)
- $A_{fc, res}$ heat transfer area between the electrical resistance wall and the mono-fluid (m$^2$)
- $A_{fc, pch}$ heat transfer area between the mono-fluid and the stem wall (m$^2$)
- $A_{pch, ext}$ heat transfer area between the stem wall and the ambient surroundings (m$^2$)
- $A_{cond, pch}$ heat transfer area between the mono-fluid and the pipe wall j (m$^2$)
- $A_{ext, pch}$ heat transfer area between the pipe wall j and the ambient surroundings (m$^2$)
- $A_i$ and $A_{i+j}$ heat transfer area of the plate j and j+1 (m$^2$)
- $C_i$ concentration of component i (mol/l)
- $C_{(i, j)}$ stoichiometric coefficient of the component i
- $C_{pf}$ specific heat of the mono-fluid flowing inside the tank i (KJ Kg$^{-1}$ K$^{-1}$)
- $C_{pf}$ specific heat of the mono-fluid at the temperature $T_{ch}$ (KJ Kg$^{-1}$ K$^{-1}$)
- $C_{pf}$ average specific heat of the resistance wall (KJ Kg$^{-1}$ K$^{-1}$)
- $C_{pf}$ average specific heat of the wall of the stem (KJ Kg$^{-1}$ K$^{-1}$)
- $C_{pf}$ specific heat of the fluid flowing inside the channel j (KJ Kg$^{-1}$ K$^{-1}$)
- $C_{pf}$ specific heat of the plate wall j (KJ Kg$^{-1}$ K$^{-1}$)
- $D_{pf}$ molar flow rate of the mono-fluid (m$^3$ s$^{-1}$)
- $E$ index that represents the thermal state of mono-fluid at the inlet of the electrical resistance
- $E_{ch}$ activation energy for the reaction j (J mol$^{-1}$)
- $f_c$ liquid reactant feed molar flow rate (mole s$^{-1}$)
- $F_{jch}$ mono-fluid flow rate through the jacket (m$^3$ s$^{-1}$)
- $k$ rate constant for reaction j
- $M_i$ molar mass of component i (Kg mol$^{-1}$)
- $m_{jch}$ mass of mono-fluid present in the jacket of the electrical resistance (Kg)
- $m_{jch}$ mass of the mono-fluid present in the tank i of the pipe j (Kg)
- $m_{jch}$ mass of the jacket wall (Kg)
- $m_{jch}$ mass of the electrical resistance wall (Kg)
- $m_{jch}$ mass of the pipe wall j (Kg)
- $m_{jch}$ total mass of the stem wall (Kg)
- $N$ number of tanks of the jacket
- $n_c$ number of components present in the reaction mixture
- $n_i$ number of moles for component i
- $n_j$ index which indicates the direction of fluid flow in the channel j (plate heat exchangers)
- $n$ number of reactions
- $P_{elec}$ electrical power value (KJ s$^{-1}$)
- $R$ perfect gas constant (R = 8,32 J mol$^{-1}$ K$^{-1}$)
- $r_j$ rate of the reaction j (mol l$^{-1}$ s$^{-1}$)
- $T_{ch}$ reaction mixture temperature (°C)
- $T_{ch}$ temperature of the mono-fluid in the tank i of the jacket (°C)
- $T_{cond}$ temperature of mono-fluid in the tank i (°C)
- $T_{cond, n10}$ and $T_{cond, n10}$ mono-fluid temperature in the latter tanks of two pipes 10 and 11(°C)
- $T_{jch}$ pipe wall j temperature (°C)
- $T_j$ temperature of the fluid flowing in the channel j (°C)
- $T_{j-1}$ temperature of the fluid flowing in the channel j-1 (°C)
- $T_{j+1}$ plate j temperature (°C)
- $T_{j+1}$ plate j + 1 temperature (°C)
- $T_{jch}$ electrical resistance wall temperature (°C)
- $T_{jch}$ stem wall temperature (°C)
- $T_{jch}$ mixing temperature at the outlet of the three-way valve (°C)
- $T_{ref}$ reference temperature (°C)
- $U_{cond, pch}$ heat transfer coefficient between the mono-fluid and the pipe wall j in the tank i (KJ m$^2$ s$^{-1}$ °C$^{-1}$)
- $U_{ext, pch}$ heat transfer coefficient between the pipe wall j and the ambient surroundings (KJ m$^2$ s$^{-1}$ °C$^{-1}$)
heat transfer coefficient between the reactor wall and the mono-fluid in the tanks i (KJ m⁻² s⁻¹ °C⁻¹)

heat transfer coefficient between the jacket wall and the ambient surroundings (KJ m⁻² s⁻¹ °C⁻¹)

heat transfer coefficient between the mono-fluid and the electrical resistance wall (KJ m⁻² s⁻¹ °C⁻¹)

heat transfer coefficient between the mono-fluid and the stem wall (KJ m⁻² s⁻¹ °C⁻¹)

heat transfer coefficient between the stem wall and the ambient (KJ m⁻² s⁻¹ °C⁻¹)

heat transfer coefficient between the mono-fluid and the plate j (KJ m⁻² s⁻¹ °C⁻¹)

heat transfer coefficient between the mono-fluid and the plate j - 1(KJ m⁻² s⁻¹ °C⁻¹)

mono-fluid flow velocity (m s⁻¹)

volume of the tank occupied by the mono-fluid (m³)

volume of the pipe j (m³)

volume of the jacket (m³)

volume of the channel j (m³)

volume of the pipe wall j (m³)

volume of the electrical resistance wall (m³)

volume of the plate wall j (m³)

molar fraction of component i

partial order of component i for the reaction j

fraction of the mono-fluid flow rate which must pass through the heat exchange

thermal conductivity of the plate j (KJ m⁻¹ s⁻¹ K⁻¹)

density of the mono-fluid at the temperature Tch (Kg m⁻³)

density of the mono-fluid in the tank i (Kg m⁻³)

density of the mono-fluid at the mixture temperature (Kg m⁻³)

density of the fluid flowing in the channel j (Kg m⁻³)

density of the jacket wall (Kg m⁻³)

density of the plate wall j (Kg m⁻³)

density of the pipe wall (Kg m⁻³)

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