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Open loop thermal control of exothermal chemical reactions in multifunctional heat exchangers

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This paper presents an experimental study which demonstrates the potentiality of multifunctional heat exchangers (MHE) to carry out exothermal chemical reactions by the local control of the reactive environment temperature. Two highly exothermal chemical reactions with different kinetics (instantaneous and fast) have been investigated. The MHE is found to be very efficient in extracting the heat released by the chemical reactions. Due to its large heat transfer capacity, this process allows to increase considerably inlet reactant concentrations without thermal runaway and thus to enhance both chemical reaction conversion ratio and yield. This study also shows the limitations of the multifunctional heat exchanger for exothermal and instantaneous chemical reactions.

Keywords: Multifunctional compact heat exchanger; Heat exchanger; Chemical reactor; Heat exchanger–reactor; Exothermal chemical reaction; Micro-mixing; Offset strip fins; Thermal control of chemical reactions

1. Introduction

Process intensification (PI) is a new concept which aims at fundamental improvements in process engineering. On the one hand, it aims at the reduction of the physical size of the equipment, and on the other hand, it focuses on the achievement of a given product with specified properties and quality. More recent ideas of process intensification require a substantial decrease in the three following process properties [1]: rate of equipment size to production capacity (compactness), energy consumption (moderate production), and waste (clean production).

In this sense PI encompasses all methods of conducting a process and the usage of appropriate apparatus which allow to achieve the above goals. Therefore, improvements in materials such as catalysts and chemical path are not considered as PI. Integration of unit operations such as mixing, reaction and heat transfer in a single device is a paramount example of process intensification. If a heat exchanger is used to accomplish mixing and reaction as well as heat transfer, then it is called a “multifunctional heat exchanger” (MHE). It is clear that in this case process intensification is no longer based on pure unit operations, but involves multi-operations. The impact of this process intensification on equipment size reduction is even more dramatic, since some types of pure processes and their associate apparatus will disappear from the future plants. The result would be safer, cheaper, compact, sustainable (environmental friendly) and energy efficient equipments in the process industry.

Heat transfer and fluid mechanics of the process play a crucial role in the process intensification. Some authors [2] consider the process intensification as “a design philosophy whereby the fluid dynamics in a process is matched to its chemical, biological and/or physical requirements”. Following this line of reasoning a chemical process becomes
ideal if the following dynamic features of the process are matched with the physico-chemical requirements of the reaction [2]: matching reaction rate with mixing rate, matching mechanism with flow pattern, matching reaction time with residence time, matching exothermicity with heat transfer.

The multifunctional heat exchanger discussed here is designed based on compact heat exchanger technology. Compact heat exchangers appear to have a potential application as a reactor in the process industry. Indeed, the high mixing level and the high heat-transfer performance of turbulence promoters provided by a compact heat exchanger makes this device also potentially useful as a chemical reactor: it is then called a multifunctional heat exchanger (MHE). This technology offers many advantages such as better reaction control (from the thermal point of view), improved selectivity (through intensified mixing, more iso-thermal operation and shorter residence time, and sharper residence time distribution), by-products reduction and better safety.

Indeed, many traditional designs such as stirred tanks already incorporate heat transfer in the process, but in these devices there is a significant distance between the heat transfer site and the site of the chemical reaction [3–6]. The aim of the MHE is to annihilate this distance by supplying or removing the heat almost as rapidly as it is absorbed or generated by the reaction. Comparison of the heat transfer performances of different heat exchanger reactors, presented in Table 1, shows the major advantage of MHE [7]. Indeed, thanks to its considerable heat transfer performance, this process should be able to carry out strong exothermic reactions without thermal runaway.

In a previous paper Ferrouillat et al. [8] studied the micro-mixing time in three generic heat exchanger geometries by using a physico-chemical technique based on a parallel-competing reaction scheme and an incorporation model. They compared micro-mixing level achieved in a plain Poiseuille flow, in a Poiseuille flow equipped with vortex generators of offset strip fins (OSF) type and finally a Poiseuille flow filled with different types of metallic foam. Micro-mixing here refers to mixing on a scale near and below the Kolmogorov turbulence micro-scale, while macro-mixing refers to bulk scale blending. Ferrouillat et al. [8] showed that for a given flow rate, a simple channel generates a reasonable micro-mixing level, but the Reynolds number, i.e. the velocity must be sufficiently high. Hence, the residence time is not sufficient for the chemical reaction to be completed and it appears that, in order to avoid extremely long channels, inserts in the channel are necessary to generate a more intense mixing for the same flowrate. Metallic foams have been proved to be the geometry that generates the best micro-mixing level. On the other hand,
for a given flow rate, offset strip fins (OSF) need less pumping power but generate less micro-mixing than metallic foams, and have been retained for the present study.

In this paper, it is shown that the heat extraction capacity of the MHE is suitable for temperature limitation in the case of two exothermal chemical reactions. The reactor used in this study is composed of three independent and superimposed channels of rectangular cross-section, where reactants circulate in the central channel and coolant fluids in both side channels as is shown in Fig. 1. Inside the reactive section, mixing promoters can be inserted in order to intensify mass and heat transfer. In the present study, OSF inserts were used, and their performance were compared with those of an empty duct channel (as the reference case) to highlight the gain attributable to inserts.

In order to have a modular device, the OSF inserts have not been brazed to the heat transfer surface. Indeed, with this prototype, global heat transfer coefficients reach roughly at most 2000 W m$^{-2}$ K$^{-1}$ instead of at least 5000 W m$^{-2}$ K$^{-1}$ with a thermal optimized design. The thermal optimization is not addressed in this study but has been widely studied by many authors [9–11]. The available literature provides a large amount of thermal-hydraulic data on compact heat exchangers equipped or not with offset strip fins [10–13]. A short review of the two technologies selected here is given in this paper.

Two exothermal chemical reactions have been selected in order to demonstrate the MHE potentiality to control an exothermal chemical reaction: (i) an acid–base reaction between sulphuric acid (H$_2$SO$_4$) and sodium hydroxide (NaOH) (instantaneous), (ii) an oxidation–reduction reaction between sodium thiosulfate (Na$_2$S$_2$O$_3$) and hydrogen peroxide (H$_2$O$_2$) (fast).

This paper is organized as follows. In Section 2 the multifunctional heat exchanger is described and semi-empirical correlations used for calculation of the Nusselt number and

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**Table 1**

Heat exchange capability for different reactors

| Process                                      | Compact multifunctional heat exchanger (MHE) (a) | Tubular exchanger reactor (b) | Batch reactor with outer heat exchanger (b) | Batch reactor with double envelope (b) |
|----------------------------------------------|-------------------------------------------------|-------------------------------|---------------------------------------------|---------------------------------------|
| Scheme                                       | ![Diagram](image1)                               | ![Diagram](image2)           | ![Diagram](image3)                          | ![Diagram](image4)                    |
| Specific area (m$^2$/m$^3$)                  | 800                                             | 400                           | 10                                          | 2.5                                   |
| Heat transfer capacity (W/m$^2$ K)           | 5000                                            | 500                           | 1000                                         | 400                                   |
| Duty/volume (kW/m$^3$ K)                     | 4000                                            | 200                           | 10                                          | 1                                     |

(a) Results based on offset strip fins at $Re = 2000$.
(b) Trambouze and Euzen [7].
pressure drop are presented. To facilitate the comparison of the performance of this device with similar devices now in operation, we have intentionally used the semi-empirical correlations commonly used in the industry. In Section 2 the two chemical reactions employed in the experiments are also described. Experimental results are then presented and discussed in Section 3. Section 4 summarizes the concluding remarks.

2. Multifunctional heat exchanger and experimental flow loop

Based on compact heat exchanger geometry, a simple and modular MHE was designed and constructed, as shown in Fig. 1. The reaction is carried out in the central rectangular channel, called the reactive channel. The reactive channel is surrounded by two parallel rectangular channels where coolant fluids circulate. The geometrical characteristics of the MHE are summarized as follows: height $H = 5$ mm, width $l = 20$ mm, length $L = 1.47$ m, hydraulic diameter $D_h = 8.00 \times 10^{-3}$ m, flow cross section $S = l \times H = 1$ cm$^2$, primary heat transfer area $S_{\text{primary}} = 2 \times l \times L = 5.88 \times 10^{-2}$ m$^2$, total volume $V = L \times l \times H = 1.47 \times 10^{-4}$ m$^3$.

For this study two geometrical configurations have been considered in the reactive central channel: an empty large aspect ratio duct channel (i.e. without inserts) and offset strip fins (OSF) usually used in compact heat exchangers.

No insert has been placed in the “double jacket” channel. This design is not thermally optimized but this could easily be done by fixing inserts in the coolant flows and brazing the fins on the support.

Fourteen “K” type thermocouples of 0.5 mm bead size have been inserted every 11 cm in the core of the reactive channel, and four other thermocouples of the same type have been placed at the inlet and outlet of the two coolant channels to evaluate the heat balance. Temperatures are quasi uniform in the duct cross-section, as shown by DTS analysis. Therefore, for the global analysis (which is the aim of this work) one thermocouple can give a signal sufficiently representative of the bulk temperature. These temperatures are recorded on a computer through a data acquisition system. In Fig. 2 a schematic diagram of the experimental flow loop in which the MHEs have been tested. More details on the design of the loop can be found in [8]. The two reactants are stored in 50 l tanks at controlled temperature of 25°C, and then pumped to the MHE where they are mixed.

Design rules for the two selected geometrical configurations, e.g. offset strip fins and empty duct channel, are presented in the following sections.

2.1. Finless duct channel

Many thermal-hydraulic correlations on empty (finless) duct channel are reported in the available literature [12,13]. A short review of pressure drop and heat transfer coefficient correlations is given here.

For a straight rectangular cross-section channel, the following correlations allow to estimate the Darcy friction coefficient. For $Re \leq 2300$ Shah and Bhatti [14] proposed for $0 < x^* < 1$, ($x^* = 2b/2a$)

$$A = \frac{96(1 - 1.3553x^* + 1.9467x^2- 1.7012x^3 + 0.9564x^4 - 0.2537x^5)}{Re}$$

where $2b$ is the channel height (m) and $2a$ the channel width (m).

![Fig. 2. Schematic diagram of the flow loop.](image-url)
For $5000 \leq Re \leq 10^7$ the same authors [14] proposed the following correlation for $0 < z' < 1$:

$$A = 4(1.0875 - 0.1125z')f_c$$  \hspace{1cm} (2)

where

$$f_c = \left( \frac{Re}{1.964 \ln Re - 3.8215} \right)^{-2}$$  \hspace{1cm} (3)

The Nusselt number in the laminar flow regime ($0 \leq Re \leq 2300$) is a function of the channel aspect ratio and the thermal boundary conditions (constant wall temperature or constant wall heat flux) [14]. The relevant condition here is the constant wall heat flux. In turbulent flow regime ($5000 \leq Re \leq 10^6$), the well-known Colburn correlation is used

$$Nu = 0.023Re^{0.8}Pr^{1/3}$$ \hspace{1cm} (4)

2.2. Offset strip fins

2.2.1. Physical concepts for OSF design

In this section, the current understanding of the physics of flow and heat transfer in plate-fins heat exchangers will be described. Depending on the fins type the flow passage can be uninterrupted or interrupted, the latter being known for their high thermal efficiency. In this type, the most common geometry is the offset strip fins shown in Fig. 3. Here the fin surface is broken into a number of small sections. For each section, a new leading edge is encountered, and thus a new boundary layer starts to develop, and is then abruptly disrupted at the end of the fin offset length. The objective in such flow passages is to prevent the thickening of the boundary layers and hence provide high heat transfer coefficients. The interruptions create a wake region, and the thermal boundary conditions dominate the heat transfer and friction. For intermediate Reynolds numbers (roughly $400 < Re < 1000$), the flow remains laminar, but unsteadiness and vortex shedding become important [13,14]. For example, at $Re = 850$, boundary layer reconstruction causes roughly a 40% increase in heat transfer over the plain channel. Unfortunately, there is a commensurate increase in the pressure drop due to boundary layer reconstruction and vortex shedding. For Reynolds numbers greater than 1000, the flow becomes turbulent in the array, and chaotic advection may be more efficient in the low Reynolds number turbulent regime. A factor of 2 or 3 increase in heat transfer and pressure drop over plain fins can be obtained as a result of the turbulent mixing. The important variables affecting the wake region identified are the strip length, the fin spacing and the fin thickness. The fin spacing and the strip length are responsible for the boundary layer interactions and wake dissipation; and the fin thickness introduces form drag and also affects the heat transfer performance. Higher aspect ratio, shorter strip lengths and thinner fins are found to provide higher heat transfer coefficients and friction factors.

2.2.2. Semi empirical correlations for OSF thermal-hydraulic performance

To define the appropriate thermal-hydraulic correlations, one needs first to define the flow regime, classically
based on the Reynolds number \( Re = (\rho U D_h)/\mu \) where \( U \) is the bulk flow velocity (m s\(^{-1}\)), \( \rho \) the fluid density (kg m\(^{-3}\)), \( \mu \) the fluid viscosity (Pa s) and \( D_h \) the hydraulic diameter (m). For the OSF geometry, a photography of the OSF inserts used in the MHE and the geometrical parameters are given in Fig. 3.

The friction factor \( f \) (fanning factor) is linked to the pressure drops \( \Delta P \) per unit length \( L \) (length of the heat exchanger) by

\[
f = (\Delta P D_h) \left( \frac{4L}{\frac{1}{2} \rho U_0^2} \right)
\]

(5)

The heat transfer factor \( j \) (Colburn factor) is related to the temperature difference over the heat transfer surface by

\[
j = \frac{NuD_h}{(ReD_h Pr^{1/3})}
\]

where the Nusselt number is defined by \( Nu_{Dh} = \frac{hD_h}{\lambda} \) and the Prandtl number \( Pr = \frac{CP}{\lambda} \), with \( CP \) the specific heat capacity (J kg\(^{-1}\) °C\(^{-1}\)) and \( \lambda \) the thermal conductivity of fluid (W m\(^{-1}\) °C\(^{-1}\)).

Correlations proposed by Manglik and Bergles [10] have been developed from the data of Kays and London, [13], in the following range of parameters: 0.646 ≤ \( D_h \) ≤ 3.414 (mm), 0.135 ≤ \( w/h_s \) ≤ 1.034 (mm), 0.012 ≤ \( e/l_s \) ≤ 0.060 (mm), 0.038 ≤ \( e/w \) ≤ 0.132 (mm).

The hydraulic diameter is defined as:

\[
D_h = \frac{4wh_h l_s}{2(wl_s + wh_l_s + eh_l_s + ew)}
\]

(6)

According to authors, the range of validity of these correlations is 300 < \( Re_{Dh} \) < 10,000.

\[
f = 9.6243Re_{Dh}^{-0.7422} \left( \frac{w}{h_s} \right)^{-0.1856} \left( \frac{e}{l_s} \right)^{0.3053} \left( \frac{e}{w} \right)^{-0.2659}
\]

\[
\times \left[ 1 + 7.669 \times 10^{-8} Re_{Dh}^{4.249} \left( \frac{w}{h_s} \right)^{0.920} \left( \frac{e}{l_s} \right)^{3.767} \left( \frac{e}{w} \right)^{0.236} \right]^{0.1}
\]

(7)

\[
j = 0.6522Re_{Dh}^{-0.5403} \left( \frac{w}{h_s} \right)^{-0.1541} \left( \frac{e}{l_s} \right)^{0.1499} \left( \frac{e}{w} \right)^{-0.0678}
\]

\[
\times \left[ 1 + 5.269 \times 10^{-5} Re_{Dh}^{1.340} \left( \frac{w}{h_s} \right)^{0.504} \left( \frac{e}{l_s} \right)^{0.456} \left( \frac{e}{w} \right)^{-1.055} \right]^{0.1}
\]

(8)

Using these correlations, all the experimental results are predicted with an error less than 20%. The dimensions of the offset strip fins used in this study are summarized as follows: \( e = 0.3 \) mm. \( w = 3.0 \) mm, \( l_s = 1.6 \) mm, \( h_s = 5.0 \) mm and \( D_h = 2.0 \) mm.

2.3. Description of the investigated chemical reactions

2.3.1. Acid–base reaction

The acid–base reaction, between sulphuric acid and sodium hydroxide, is a strong acid/strong base reaction and is often considered as a total and irreversible reaction. It is instantaneous and not very temperature sensitive. All components are miscible and flow is a single liquid phase flow. The reaction scheme is as

\[
\text{H}_2\text{SO}_4 + 2\text{NaOH} \rightarrow 2\text{H}_2\text{O} + \text{Na}_2\text{SO}_4
\]

(9)

The reaction rate law, which is a second-order law (first-order law for each reactant), can be written as

\[
r = k_{\text{CH}_3\text{SO}_3}c_{\text{NaOH}}
\]

(10)

with: \( c_{\text{CH}_3\text{SO}_3} \) the sulphuric acid concentration (mol l\(^{-1}\)), \( c_{\text{NaOH}} \) the sodium hydroxide concentration (mol l\(^{-1}\)) and

\[
k = 1.4 \times 10^{11} \text{mol}^{-1} \text{ s}^{-1} \text{ at } 25 ^\circ\text{C}
\]

The reaction enthalpy is \( \Delta H_r = -92.4 \text{ kJ per mol of sodium hydroxide}. \) This reaction is not very exothermic. Nevertheless, its instantaneous kinetic leads to a very local exothermicity in the process. This reaction is therefore, very useful to investigate MHE abilities in extracting a very local exothermicity.

2.3.2. Oxidation–reduction reaction

The oxidation–reduction reaction between the sodium thiosulfate and the hydrogen peroxide is interesting for its demonstrative characteristic [15–17]. This reaction is irreversible, strongly exothermal and meets a fast chemical kinetics.

The reaction scheme is as follows

\[
2\text{Na}_2\text{S}_2\text{O}_3 + 4\text{H}_2\text{O}_2 \rightarrow 2\text{Na}_2\text{S}_2\text{O}_6 + \text{Na}_2\text{SO}_4 + 4\text{H}_2\text{O}
\]

(11)

The reaction rate law, which is a second-order law (first-order law for each reactant), can be written as

\[
r = k c_{\text{S}_2\text{O}_3^-} c_{\text{H}_2\text{O}_2}
\]

(12)

with: \( c_{\text{S}_2\text{O}_3^-} \) the sodium thiosulfate concentration (mol l\(^{-1}\)), \( c_{\text{H}_2\text{O}_2} \) the hydrogen peroxide concentration (mol l\(^{-1}\)),

\[
k = k_0 \text{exp}(E/RT)
\]

the reaction rate constant (1 mol\(^{-1}\) s\(^{-1}\)), \( k_0 \) the frequency factor (1 mol\(^{-1}\) s\(^{-1}\)), \( E \) the activation energy (J mol\(^{-1}\)), \( R = 8.314 \text{ J K}^{-1} \text{ mol}^{-1} \) the perfect gas constant, \( T \) the temperature (K).

Kinetics data available in the literature concerning this reaction are summarized as follows:

\[
k = 2.13 \times 10^{10} \text{exp}(-8196/T)
\]

[15],

\[
k = 1.63 \times 10^{10} \text{exp}(-8140/T)
\]

[16] and

\[
k = 8.13 \times 10^{11} \text{exp}(-9156/T)
\]

[17]. The kinetics constant of the reaction follows an Arrhenius law and so is very sensitive to temperature. However, the hydrogen peroxide degrades at temperatures higher than 100 °C (\( T_{\text{Boiling}} = 107 ^\circ\text{C} \)). This reaction is more exothermal than the acid–base reaction. The available data on the enthalpy of this reaction are summarized as follows: \( \Delta H_r = -568 \text{ kJ per mol of Na}_2\text{S}_2\text{O}_3 \) [15] and \( \Delta H_r = -612 \text{ kJ per mol of Na}_2\text{S}_2\text{O}_3 \) [16].

3. Results and analysis

The chemical reaction evolution depends on four parameters: (i) chemical reactant concentration, (ii) micro-mixing level [18–20], (iii) reactive temperature, e.g.
global heat transfer coefficient \([15–17]\), (iv) residence time in the process.

In order to simplify the interpretation of the results, the input conditions for chemical reactant concentrations are constant for all the runs. The three other parameters are conditioned by the process conditions, i.e. the reactive volume flow rate \(Q_{V,R}\), and the coolant volume flow rate \(Q_{V,C}\).

Thus, several tests have been carried out to evaluate the respective influence of the reactive and the coolant volume flow rates on the micro-mixing level, the global heat transfer coefficient and the residence time.

Test runs have been carried out with and without offset strip fins in the MHE, the empty channel being considered as the base-line configuration.

### 3.1. Acid–base reaction

For all test runs, the chemical reaction concentrations are:
\[c_A = c_{\text{H}_2\text{SO}_4} = 0.5 \text{ mol l}^{-1}\] and \(c_B = c_{\text{NaOH}} = 1.0 \text{ mol l}^{-1}\).

The sum of the volume flow rates of reactants A and B represents the total reactive volume flow rate \(Q_{V,R}\) and \(Q_A = Q_B\). In total five series of experiments have been carried out, all tests characteristics are summarized in Table 2.

Comparison between the test cases 1 and 2 shown in Fig. 4 allows to discern the effects of coolant volume flow rate for a MHE configuration without fins. The coolant volume flow rates are 400 l h\(^{-1}\) and 200 l h\(^{-1}\) respectively for tests 1 and 2, leading to global heat transfer coefficients of respectively 410 W m\(^{-2}\) K\(^{-1}\) and 370 W m\(^{-2}\) K\(^{-1}\) respectively for test 1 and test 2. As shown in Fig. 10, the reactive temperature reaches a sharp peak due to the local exothermicity and then decreases slowly along the MHE. The global heat transfer coefficient seems to have a negligible influence on the exothermal peak magnitude, but has a significant influence on the reactive temperature decrease after the peak.

Fig. 5 compares the reactive temperature rise for tests 1 and 3, showing the effect of reactive volume flow rate (Table 2) also carried out without fins. It shows that an increase in reactive volume flow rate at constant coolant volume flow rate leads also to an increase in the global heat transfer coefficient, but at the same time, it leads to an increase of mixing via the turbulence and a decrease in the residence time. The combined consequence of these effects is that for a given residence time, one observes a faster decrease of the temperature when the reactive volume flow rate increases from a lagrangian point of view. However, the outlet temperature of the MHE is higher for test run 3 than 1, meaning that the negative effect of the residence time reduction dominates the positive effect of the

### Table 2

| Test run | 1   | 2   | 3   | 4   | 5   |
|----------|-----|-----|-----|-----|-----|
| Geometric configuration | Empty duct channel | Empty duct channel | Empty duct channel | OSF | OSF |
| \(Q_{V,R}\) (l h\(^{-1}\)) | 50  | 50  | 100 | 50  | 100 |
| \(Q_{V,C}\) (l h\(^{-1}\)) | 400 | 200 | 400 | 400 | 400 |
| \(h_R\) (W m\(^{-2}\) K\(^{-1}\)) | 470 | 470 | 1840| 2900| 5200|
| \(h_C\) (W m\(^{-2}\) K\(^{-1}\)) | 3680| 1840| 3680| 3680| 3680|
| \(H_g\) (W m\(^{-2}\) K\(^{-1}\)) | 410 | 370 | 1160| 1500| 1950|
| Reactive Reynolds number \(Re^a\) | 1300| 1300| 2700| 1160| 2320|

\(^a\) \(Re\) is calculated with fluid properties estimated at the reactive average temperature.

![Fig. 4. Effect of coolant volume flow rate on reactive temperature rise in MHE.](image-url)
flow rate. Finally, these first results show, for a reactive empty duct channel configuration, that the MHE is able to extract the heat generated by the chemical reaction.

The geometry influence on heat extraction from chemical reactions has been investigated by inserting, in the reactive channel, un-brazed offset strip fins. As shown in Table 2, for the same volume flow rate conditions, the OSF insertion leads to an increase of global heat transfer coefficient compared to empty duct channel. This heat transfer performance improvement with OSF configuration is due to interrupted flow passages which do not allow the boundary layers to thicken and thus results in high heat transfer coefficients associated with thin boundary layers. Therefore, it is logical to expect that for the same reactive flow rate and the same residence time, the OSF configuration allows to decrease significantly the reactive temperature rise compared to empty duct channel, as is shown in Fig. 6, where the temperature rise for test runs 1, 3, 4 and 5 are compared. Comparison of test runs 1 and 5 shows that the outlet reactive temperatures are roughly equal while residence time is lower with the OSF configuration. In other words, for a given outlet reactive temperature, the process compactness can be improved by OSF insertion. Experiments 3 and 5 have the same reactive and the same coolant flow rates. Fig. 6 shows that the reaction temperature peak in the test run 5 (with OSFs) is lower than that of the test run 3 (empty duct channel), but has the same slope. The same observation is also valid for the results of experiments 1 and 4. It implies that the extra micro-mixing due to the OSF inserts has not increased the reaction kinetics but has provided a faster heat extraction from the reaction.

### 3.2. Oxidation–reduction reaction

Contrary to the acid–base reaction which reacts instantaneously near the reactant mixing area, this chemical reac-

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**Fig. 5.** Effect of coolant volume flow rate on reactive temperature rise in MHE.

**Fig. 6.** Effect of the geometrical configuration on cooling of chemical reaction.
tion reacts gradually along the MHE. Moreover, the chemical reaction kinetics depends on the temperature. Thus, the chemical reaction evolution is closely linked to the micro-mixing level and the heat transfer coefficient. In order to discriminate both effects, experiments have been carried out in a first step without heat extraction, in quasi adiabatic conditions since the heat transfer takes place in a diffusive mode out of the reactor, so that micro-mixing effects are seen on chemical reaction evolution. In a second step, the global heat transfer effect on chemical reaction is investigated by the use of the active coolant flow.

For all test runs, chemical reaction concentrations are as follows: \( c_A = c_{\text{Na}_2\text{SO}_4} = 0.4 \text{ mol l}^{-1} \), \( c_B = c_{\text{H}_2\text{O}_2} = 0.8 \text{ mol l}^{-1} \).

The sum of the volume flow rates of reactants A and B represents the total reactive volume flow rate \( Q_{V,R} \) where \( Q_A = Q_B \). All test characteristics of this series of experiments are summarized in Table 3.

Without heat removal, the reactive temperature increases and this leads to an acceleration of the chemical reaction kinetics, and thus a conversion rate increase in the process outlet. Moreover, adiabatic conditions allow to compute the conversion rate at a given position in the MHE.

\[
X_A(x) = \frac{T(x) - T(0)}{T_{100\%}}
\]

with \( T(x) \) is the temperature at the position \( x \) (°C) and \( T_{100\%} = (\Delta H_{r} C_{A,0}/(\rho C_p)) \) is the theoretical temperature for a 100% conversion rate (°C).

In these expressions, fluid properties have been estimated at the average temperature \((T_{100\%} + T(0))/2\). The \( T_{100\%} \) calculation is then realized by an iterative method.

Tests 5, 6 and 7 have been carried out in the “finless” channel configuration with different reactive volume flow rates (12, 25 and 50 l h\(^{-1}\)), which correspond respectively to Reynolds numbers of 400, 670 and 1200. As shown in Fig. 7, the comparison of these three tests results show that the chemical reaction progress is roughly the same for a given residence time. Indeed, since for each test the chemical reaction kinetics is fixed by equal initial chemical reactant concentrations and since reactant mixing level is also fixed by laminar flow regime, one could assume that the global chemical reaction kinetics is equal for all tests as is observed in Fig. 7. However, it is difficult to deduce, from these results, whether the limiting factor is the chemical kinetics or the reactant mixing.

Then the influence of the reactor geometry has been tested by inserting the fins, which corresponds to tests 10 and 11 and which results are plotted in Fig. 8. The main difference between these two tests is the reactive volume flow rate which is 12 l h\(^{-1}\) for test 10 and 25 l h\(^{-1}\) for test

### Table 3
Test characteristics for oxidation–reduction reaction

| Test run | Geometric configuration | \( Q_{V,R} \) (l h\(^{-1}\)) | \( Q_{V,C} \) (l h\(^{-1}\)) | \( h_R \) (W m\(^{-2}\) K\(^{-1}\)) | \( h_C \) (W m\(^{-2}\) K\(^{-1}\)) | \( H_g \) (W m\(^{-2}\) K\(^{-1}\)) | Reactive Reynolds number \( Re^a \) |
|----------|-------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| 5        | Empty duct channel      | 12              | 0               | 470             | 0               | 0               | 400             |
| 6        | Empty duct channel      | 25              | 0               | 470             | 0               | 0               | 670             |
| 7        | Empty duct channel      | 50              | 0               | 470             | 0               | 0               | 1200            |
| 8        | Empty duct channel      | 12              | 0               | 470             | 0               | 0               | 1200            |
| 9        | Empty duct channel      | 12              | 0               | 470             | 0               | 0               | 1200            |
| 10       | Empty duct channel      | 12              | 0               | 1840            | 0               | 0               | 1200            |
| 11       | Empty duct channel      | 25              | 0               | 5560            | 0               | 0               | 1200            |
| 12       | Empty duct channel      | 25              | 0               | 700             | 0               | 0               | 1200            |
| 13       | Empty duct channel      | 25              | 0               | 940             | 0               | 0               | 1200            |

\( a \) \( Re \) is calculated with fluid properties estimated at the reactive average temperature.

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![Fig. 7. Reactive conversion rate against residence time in adiabatic conditions with empty duct channel.](image-url)
This figure shows that an increase of reactive volume flow rate leads to a conversion rate increase for a given residence time. It seems thus reasonable to deduce that the global chemical reaction kinetics is controlled by reactant mixing and not by chemical reaction kinetics. This geometrical configuration allows to decrease the Reynolds number for laminar-turbulent transition by generating unsteadiness and vortex shedding downstream of the fin at $Re = 400$ compared to the empty duct channel at $Re = 2300$.

It should be underlined that the chemical reaction kinetics is not constant along the process. Indeed, two different mechanisms are competing: the temperature increase due to the exothermal reaction, and the decrease of the reaction rate due to the accelerated consumption of the reactants. These two effects are difficult to dissociate from these results (Fig. 8).

Chemical reaction progress also depends on the residence time. The reactive flow rate increase in the test run 11 leads to a decrease of the residence time. Thus, one could observe a lower outlet conversion rate for test run 11 than test 10. It means, as for acid–base reaction, that the negative effect of the residence time reduction dominates the positive effect of the flow rate and thus micromixing increase.

Finally, the two different geometrical configurations have been compared in terms of conversion rate. Fig. 9 represents the conversion rate versus residence time. Comparison of test run 6 (without fins) and test run 11 (OSF configuration) reveals that for a given residence time, conversion rate is enhanced thanks to OSF insertion which increases reactant mixing. Indeed, test run 6 is carried out at $Re = 670$, e.g. at laminar flow regime in empty duct channel, while test run 11 is carried out at $Re = 1000$ where the flow becomes turbulent in the array for an OSF configuration. These results confirm that the chemical reaction progress is limited by the reactant mixing.

In order to study the global heat transfer coefficient effect on chemical reaction evolution, the reactant micromixing and the residence time have been fixed by equal reactant volume flow rates for all test runs. Thus, two addi-
tional tests (8 and 9) have been realized with reactive volume flow rate of 12 l h\(^{-1}\) and two coolant volume flow rates (respectively 100 and 8001 h\(^{-1}\)) in order to modify the global heat transfer coefficient in empty duct channel configuration. Fig. 10 shows for these tests (5, 8 and 9) reactive temperature rise versus the position in the MHE. It is discernible that the reactive temperature decreases when the global heat transfer coefficient increases. The reactive temperature decrease implies a slowing down of chemical reaction progress since the chemical reaction speed depends on, among others, temperature. Consequently, these tests also show that the MHE is able to control the chemical reaction evolution thanks to a secondary fluid flow and high level of heat transfer. Moreover, with a high global heat transfer coefficient of 430 W m\(^{-2}\) K\(^{-1}\) in test 9, the reactive temperature is maintained at its initial temperature. In this case, the reaction has been frozen.

As for empty duct channel configuration, in order to study the global heat transfer coefficient effect of OSFs on chemical reaction evolution, the reactant micro-mixing and the residence time have been fixed, thanks to the equal reactant volume flow rates for all tests. In addition to the adiabatic test run 10, two tests (12 and 13) have been realised with reactive volume flow rate of 12 l h\(^{-1}\) and different coolant volume flow rates (respectively 100 and 8001 h\(^{-1}\)) in order to modify global heat transfer coefficient via the utility side. As is shown in Table 3, for the same volume flow rate conditions, the OSF insertion increases the global heat transfer coefficient of the reactive side compared to the empty duct channel. Fig. 11 shows for the tests 10, 12 and 13 the reactive temperature versus the position in the MHE. As for empty duct channel configuration, the reactive temperature decreases when the global heat transfer coefficient increases. Even though chemical reaction progress is enhanced by reactant micro-mixing improvement due to the OSF insertion as shown previously, which leads to a higher exothermicity, the MHE is able to control the chemical reaction evolution by a secondary fluid flow.

![Fig. 10. Effect of heat transfer coefficient on reactive temperature rise with empty duct channel.](image)

![Fig. 11. Effect of heat transfer coefficient on reactive temperature in MHE with OSF inserts.](image)
Moreover, this geometrical configuration allows to ensure at the process outlet a reactive temperature decrease of 45 °C between tests 10 and 13 compared to 35 °C between tests 5 and 9 for the simple channel configuration. With a high global convective heat transfer coefficient of 941 W m⁻² K⁻¹ in the test 13, the reactive temperature is maintained at its initial inlet temperature.

4. Conclusions

In this study, the potentiality of multifunctional heat exchangers to monitor exothermal chemical reactions through an open loop control of the reactive temperature is experimentally stated.

Actually many traditional designs such as tubular heat exchanger-reactors or classical batch reactors are often limited by their heat transfer performance for strong exothermal chemical reactions. This poor heat transfer ability forces to work with low reactant concentrations in order to avoid thermal runaway in the process.

Even without thermal optimisation (a two-fold heat transfer increase would easily be attainable with brazing), high global heat transfer coefficients can be obtained simply by adequate coolant or reactive volume flow rate. Moreover, experimental results show that OSF insertion can provide a significant mass and heat transfer intensification which lead to a considerable compactness of MHE.

Using this process allows to increase considerably inlet reactant concentrations without thermal runaway risk and consequently to enhance both chemical reaction conversion rate and yield. Nevertheless, the MHE limitations for exothermal and instantaneous chemical reactions are undeniable, since it is not able to remove instantaneously the heat generated by a very local exothermicity.

As a conclusion, this work shows that heat transfer intensification is an efficient concept to improve multifunctional heat exchangers potentiality in continuous chemical engineering processes.

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