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# Pyrolysis in Porous Media: Numerical Analysis and Comparison to Experiments.

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Only limited studies are available experimentally to investigate hydrocarbon fuel pyrolysis, which can be of practical interest for the active cooling of head loaded components in aerospace vehicles such as combustors in rocket engines. The numerical simulation is an additional way to investigate the related phenomena (heat and mass transfer with chemistry). After a first code validation with experiments based upon inert gases, the code is further extended towards permeation with reactive dodecane and a reasonable agreement is found. The experimental accuracy of the permeability's determination is confirmed numerically to be within  $\pm 30\%$ . Numerically it is shown that this accuracy is due to strong flow spatial heterogeneities. The border effect of the test cell is found to be related to the permeable medium thickness whereas the temperature field is correlated to the reaction zone. Two different kinetic mechanisms are used to investigate the chemistry effect on the heat and mass transfer. They provide also a better analysis of the fuel pyrolysis and of the products' formation.

## Nomenclature

$\bar{c}_p$	=	heat capacity at constant pressure ( $\text{J}\cdot\text{mol}^{-1}\cdot\text{K}^{-1}$ )
$C_2$	=	inertial resistance factor ( $\text{m}^{-1}$ )
$E_A$	=	activation energy ( $\text{J}\cdot\text{mol}^{-1}$ )
$\Delta H$	=	enthalpy of reaction ( $\text{J}\cdot\text{mol}^{-1}$ )
$k_{eff}$	=	effective thermal conductivity of the medium ( $\text{W}\cdot\text{m}^{-1}\cdot\text{K}^{-1}$ )
$K_{D/F}$	=	Darcian ( $\text{m}^2$ )/ Forchheimer's coefficients (m)
$\dot{m}$	=	mass flow rate ( $\text{kg}\cdot\text{s}^{-1}$ )

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$P$	=	pressure (Pa)
$Q_R$	=	heat of reaction ( $\text{J}\cdot\text{mol}^{-1}$ )
$S_i$	=	sink term (Pa)
$\vec{v}$	=	velocity vector ( $\text{m}\cdot\text{s}^{-1}$ )
$Y_j$	=	species mass fraction ()
$\alpha$	=	permeability factor ( $\text{m}^2$ )
$\mu$	=	dynamic viscosity (Pa.s)
$\rho_{f/s}$	=	density of the fluid / solid ( $\text{kg}\cdot\text{m}^{-3}$ )

## I. Introduction

Studying porous flow is of interest in numerous fields of research and more specifically in the present framework of active cooling intended to be used on highly thermally loaded structures [1, 2]. Transpiration cooling through a porous structure is highly efficient as shown by Langener *et al.*[3,4]. This technology is found in rocket engines and chambers for example [5,6]. This reduces the temperature of the solid permeable materials used to sustain the large heat load encountered by the combustor [7,8]. Carbon- or oxide-based composite materials are preferred for thermal, mechanical and chemical reasons [9]. In addition, when using the fuel as a coolant, the so-called regenerative technique, a pyrolysis is generally encountered for fluid temperature over 800 K [10]. This generates endothermic reactions which contribute to the heat flux absorption. The formation of a thin film inside the combustion chamber, the so-called film cooling technique, is another parameter participating in the structure cooling [11]. The main drawbacks of these coupled solutions are the multiphysics' interactions which modify transiently the permeabilities of the structure and the cooling efficiency [7]. The fuel cracking produces light and heavy species which are coke precursors [12]. The formation of solid particles modifies the fluid filtration and consequently the heat and mass transfers [13].

Experimental chemical studies generally focus on the fuel pyrolysis as a function of the fuel nature, of the operating conditions or of the reactor's nature [14-18]. Some numerical works are also available [19,20]. Other studies are related to the heat and mass transfers involved in the active cooling by means of computing [21] and of testing [1, 2]. Nevertheless, there is only rare studies aiming at grouping both experimental and numerical approach to provide a deep analysis of the multiphysics' phenomena (thermal, fluid, mechanical and chemical points) [7]. The complementarity of these two kinds of work is due to the fact that numerical studies still face strong difficulties in solving massively detailed kinetic mechanism and in handling porous flow with microscopic description over full length scales and over long timescales. On the opposite, it is difficult experimentally to determine microscopic or local parameters such as the temperature or the species concentration and to measure the heat flux or the residence time. For this reason, a European collaboration has been settled to establish a numerical modelling of reactive porous flow with semi-detailed chemistry and taking the other physical parameters into account (heat and mass transfers, multi-diffusion, porous flow modelling, compressibility effects, real fluid properties). The authors here aim at comparing the numerical results to the experimental ones on dodecane pyrolysis cases in order to determine the model sensitivity and accuracy. Numerical data are of major importance for the analysis of the experimental results. Dodecane is used to compare with experiments but methane should be used in latter steps for rocket combustors application.

## II. Material and Methods

### A. Numerical simulation

A 2-D geometry with 14,242 structured cells (Figure 1) models the test cell to be detailed in next section. The solver handles compressible steady-state configurations. The Navier-Stokes equations (Eq. 1 to Eq. 3) are modified taking into account the porosity effect of the sample, the consequent pressure drop through the medium and the effective thermal conductivity of the porous zone (fluid plus porous material). The Forchheimer's equation is used to simulate the pressure drop via a sink term in the momentum equation (Eq.4), where  $\alpha$  and  $C_2$  are respectively  $2.061 \cdot 10^{-13} \text{ m}^2$  and  $4924000 \text{ m}^{-1}$  (obtained experimentally [22]).

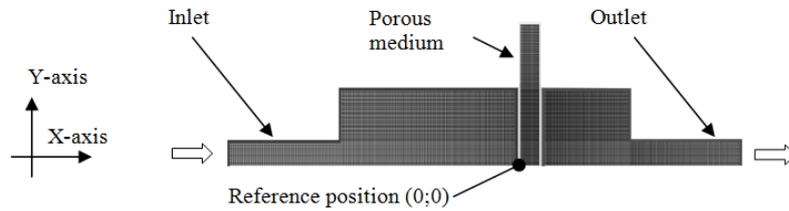
$$\frac{\partial(\gamma\rho_f\vec{v})}{\partial t} + \nabla \cdot (\gamma\rho_f\vec{v}\vec{v}) = -\gamma\nabla P + \nabla \cdot (\gamma\vec{\tau}) + \gamma\rho_f\vec{g} - S_i \quad (1)$$

$$\frac{\partial(\gamma\rho_f)}{\partial t} + \nabla \cdot (\gamma\rho_f\vec{v}) = 0 \quad (2)$$

$$\frac{\partial}{\partial t}(\gamma\rho_f E_f + (1-\gamma)\rho_s E_s) + \nabla \cdot (\vec{v}(\rho_f E_f + P)) = \nabla \cdot [k_{eff}\nabla T - (\sum_i h_i \vec{J}_i) + (\bar{\tau} \cdot \vec{v})] + S_f^h \quad (3)$$

$$S_i = -\frac{\mu}{\alpha}v_i + C_2 \frac{1}{2}\rho|v|v_i \quad (4)$$

The pressure-based solver guarantees a faster and stable convergence. The boundary conditions correspond to constant: inlet mass flow rate, pressure outlet and wall temperature. The conditions measured experimentally are used directly in the software to simulate identical test cases. The simulation settings change depending on the case under study: nature of the fluid (nitrogen or kerosene surrogate), operating temperatures (from 300 K to 1200 K) and mass flow rate (60 mg.s<sup>-1</sup> to 3.98 g.s<sup>-1</sup>). Additional details will be provided when necessary for each test case (section III).



**Figure 1 Computational domain layout.**

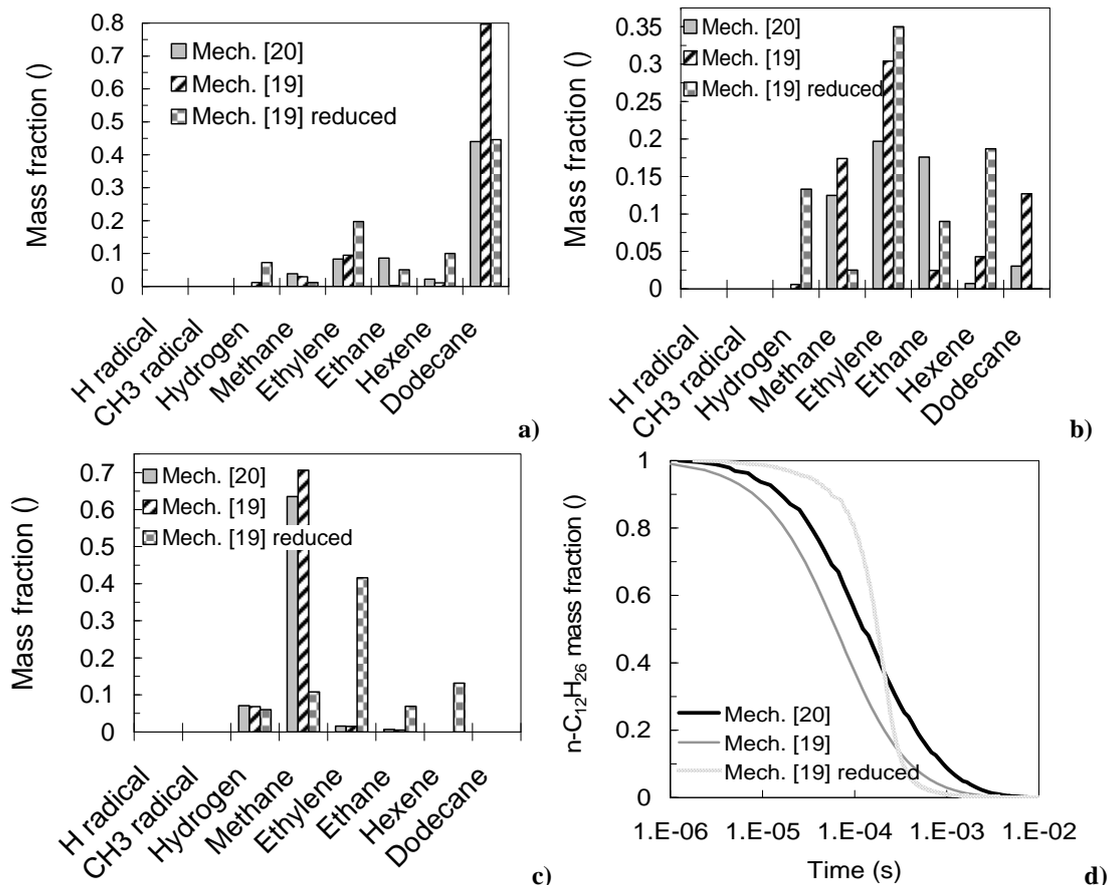
A detailed kinetic mechanism (56 species and 289 reactions) simulates the dodecane pyrolysis [22]. The present authors have reduced this mechanism (15 species, 20 reactions) for computation cost reason with limited loss of accuracy. Two primary reactions (“1” and “2” in Table 1) allow the cracking of dodecane producing ethylene (C<sub>2</sub>H<sub>4</sub>) and two radicals but-1-yl (pC<sub>4</sub>H<sub>9</sub>) and n-propyl (nC<sub>3</sub>H<sub>7</sub>). These two reactions controlling the pyrolysis process correspond to a high activation energy ( $\approx 3.68 \cdot 10^5$  J.mol<sup>-1</sup>). The sensitivity analysis [22] for n-dodecane pyrolysis shows that the fuel conversion rate is controlled by C-C fission of n-dodecane and H-abstractions by H and CH<sub>3</sub>. So the present authors primary considered adding at least the reactions allowing the formation of these two radicals (monatomic hydrogen via backward reaction “4” and “5” and methyl radical via “12”). The presence of H and CH<sub>3</sub> accelerates the pyrolysis rate via propagation reactions for reactions “6” → “10” and “13” → “17”. The ramification is indirect via dissociation of light species as hydrogen H<sub>2</sub> (reaction “11”) and ethane C<sub>2</sub>H<sub>6</sub> (reaction “20”). The reactions involving directly the interaction between the n-dodecane and the radicals H and CH<sub>3</sub> include both H-abstraction and  $\beta$ -scission of n-C12-alkyl-radical (C<sub>12</sub>H<sub>25</sub>).

By comparing this reduced mechanism under perfectly stirred reactor configurations at 3.5 MPa with the original one [22] and with a very detailed one (271 species and 1449 reactions) [23], the present reduced scheme is judged adequate for the specific conditions used in this paper despite important discrepancies (Figure 2). The overall performance is generally poor but they are acceptable in the specific thermal range 900 K-1200 K at 35 bar and for residence time comparable to those in the system. At 900 K and after 0.7 s of simulation (which is roughly the residence time found later in the cell, see section C), the conversion of dodecane is adequately simulated by the reduced scheme (Figure 2a) while the initial detailed one of You *et al.* [22] presents a factor 2 compared to the accurate reference mechanism of Herbinet *et al.* [23]. A factor 2 is found on ethylene between the reference and the reduced scheme, which shows that the products distribution is predicting with a limited accuracy. After 10 s at 900 K (constant pressure of 3.5 MPa), ethylene and ethane are reproduced fairly contrary to some species like hexene (Figure 2b). At 1200 K and 35 bar, after 10 s, the discrepancies are still important between the reference and the reduced mechanisms, mainly concerning the methane formation while the original mechanism of You *et al.* [22] gives satisfactory results. The major compound found in the reduced mechanism is ethylene (Figure 2c). Looking at the dynamics of the pyrolysis (Figure 2d), the original mechanism presents a shift on the log- time scale compared to the reference mechanism. The reduced mechanism presents a mean behaviour around the reference one. As a consequence, in order to perform first kinetic computations within 2-D geometry, the reduced mechanism is used but one should pay attention to the limited accuracy in terms of products formation. This mechanism has been optimised for representing the dodecane consumption and it cannot accurately represent the formation of radicals, of light species like hydrogen and of middle-weight species like hexene. This reduced mechanism is a compromise

which should be later improved by increasing the number of species and of reactions and, as a consequence, the computation cost.

**Table 1 Reduced kinetic mechanism used in this paper.**

[1]	$\text{NC}_{12}\text{H}_{26} \Rightarrow 3\text{C}_2\text{H}_4 + 2\text{nC}_3\text{H}_7$	[12]	$2\text{CH}_3 \rightleftharpoons \text{H} + \text{C}_2\text{H}_5$
[2]	$\text{NC}_{12}\text{H}_{26} \Rightarrow 2\text{C}_2\text{H}_4 + 2\text{pC}_4\text{H}_9$	[13]	$\text{NC}_{12}\text{H}_{26} + \text{CH}_3 \Rightarrow 4\text{C}_2\text{H}_4 + \text{pC}_4\text{H}_9 + \text{CH}_4$
[3]	$\text{C}_2\text{H}_4 + \text{C}_2\text{H}_5 \rightleftharpoons \text{pC}_4\text{H}_9$	[14]	$\text{NC}_{12}\text{H}_{26} + \text{CH}_3 \Rightarrow \text{C}_4\text{H}_8 + 2\text{C}_2\text{H}_4 + \text{pC}_4\text{H}_9 + \text{CH}_4$
[4]	$\text{C}_6\text{H}_{12} + \text{H} = \text{C}_2\text{H}_4 + \text{pC}_4\text{H}_9$	[15]	$\text{NC}_{12}\text{H}_{26} + \text{CH}_3 \Rightarrow \text{C}_3\text{H}_6 + \text{C}_6\text{H}_{12} + \text{nC}_3\text{H}_7 + \text{CH}_4$
[5]	$\text{C}_5\text{H}_{10} + \text{H} = \text{C}_2\text{H}_4 + \text{nC}_3\text{H}_7$	[16]	$\text{NC}_{12}\text{H}_{26} + \text{CH}_3 \Rightarrow \text{C}_5\text{H}_{10} + 2\text{C}_2\text{H}_4 + \text{nC}_3\text{H}_7 + \text{CH}_4$
[6]	$\text{NC}_{12}\text{H}_{26} + \text{H} \Rightarrow 4\text{C}_2\text{H}_4 + \text{pC}_4\text{H}_9 + \text{H}_2$	[17]	$\text{NC}_{12}\text{H}_{26} + \text{CH}_3 \Rightarrow \text{C}_6\text{H}_{12} + \text{C}_2\text{H}_4 + \text{pC}_4\text{H}_9 + \text{CH}_4$
[7]	$\text{NC}_{12}\text{H}_{26} + \text{H} \Rightarrow \text{C}_4\text{H}_8 + 2\text{C}_2\text{H}_4 + \text{pC}_4\text{H}_9 + \text{H}_2$	[18]	$\text{C}_2\text{H}_6 + \text{CH}_3 \rightleftharpoons \text{C}_2\text{H}_5 + \text{CH}_4$
[8]	$\text{NC}_{12}\text{H}_{26} + \text{H} \Rightarrow \text{C}_3\text{H}_6 + \text{C}_6\text{H}_{12} + \text{nC}_3\text{H}_7 + \text{H}_2$	[19]	$\text{CH}_3 + \text{CH}_2 \rightleftharpoons \text{C}_2\text{H}_4 + \text{H}$
[9]	$\text{NC}_{12}\text{H}_{26} + \text{H} \Rightarrow \text{C}_5\text{H}_{10} + 2\text{C}_2\text{H}_4 + \text{nC}_3\text{H}_7 + \text{H}_2$	[20]	$2\text{CH}_3(+\text{M}) \rightleftharpoons \text{C}_2\text{H}_6(+\text{M})$
[10]	$\text{NC}_{12}\text{H}_{26} + \text{H} \Rightarrow \text{C}_6\text{H}_{12} + \text{C}_2\text{H}_4 + \text{pC}_4\text{H}_9 + \text{H}_2$		LOW / 1.7700E+50 -9.670 6220.00/
[11]	$2\text{H} + \text{M} \rightleftharpoons \text{H}_2 + \text{M}$		TROE/ 0.5325 151.0 1038.0 4970.0 /
	$\text{H}_2 / 0.00/$		$\text{H}_2 / 2.00/ \text{CH}_4 / 2.00/ \text{C}_2\text{H}_6 / 3.00/$
<b>Species:</b> n-dodecane, hex-1-ene, pent-1-ene, but-1-ene, propylene, ethane, ethylene, methane, tripropylmethylene			
<b>Radicals:</b> but-1-yl, n-propyl, ethyl, methyl, monoatomic hydrogen			



**Figure 2.** 0-D numerical computations with three kinetic mechanisms at 900 K and 3.5 MPa after 0.7 s (a) and after 10 s (b) and at 1200 K and 3.5 MPa after 10 s (c) and dodecane pyrolysis as a function of time at 1200 K and 3.5 MPa (d).

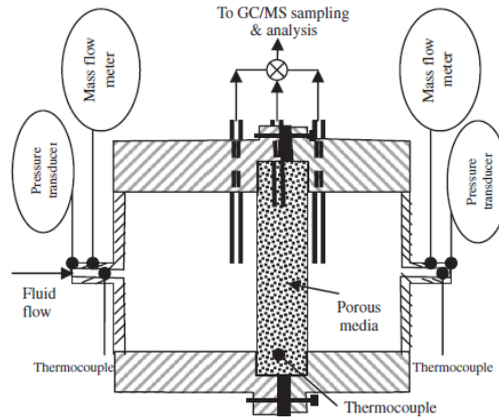
## B. Experimental setup

The COMPARER pyrolysis test bench is used to pressurize and to heat the fuel under flow conditions [24]. Its main characteristics are the following:

- Maximum operating conditions: 1800 K, 80 bar,  $0.6 \text{ g.s}^{-1}$  for liquid fluid and  $6 \text{ g.s}^{-1}$  for gas
- Chemical analysis: Gas Chromatograph/Mass Spectrometer-Flame Ionisation Detector-Thermal Conductivity Detector with 10 sampling loops for *a posteriori* quantitative analysis, Fourier Transform Infra Red spectrometer (inline direct quantitative analysis)
- Sensors: 5 mass flow rates, 5 pressure transducers, over 10 K-type and R-type thermocouples with data acquisition system (16 bits, 48 channels, 0.1 Hz)

A permeation test cell contains the porous sample (**Figure 3**). This cell is inserted inside the furnace of the COMPARER bench and it is connected to the fluid supply system and to the appropriate sensors. The permeable medium bounds the cell in two high and low pressure chambers (upstream and downstream to the porous medium respectively). These chambers are noted HPC and LPC in this paper. An inlet pipe provides the fuel into the system. This cell is connected to a dynamic sampling system to get hot pressurized samples at three location points in the cell. These samples are later cooled and chemically quantified by the GC/MS-FID-TCD. This allows having the longitudinal pyrolysis profile of the fuel under given operating conditions. The FTIR analysis provides real-time quantification at the process outlet on the gas products only after having cooled and expanded the pyrolysis mixture at ambient conditions.

The permeable medium can be changed. In the present work, an isotropic stainless steel material is preferred to composite one to avoid considering complex microstructure (fibres, layers). It is characterized by a porosity around 30 %, a grain diameter of  $14.1 \mu\text{m}$  and a pore diameter of  $4.1 \mu\text{m}$  [25]. Further geometrical information can be found in [25].



**Figure 3. Schematic of the permeation test cell with porous medium and associated measures [25].**

## III. Fuel pyrolysis analysis

### A. Preliminary validation on inert test cases and first experiments analysis

Nitrogen is injected with constant mass flow rate and the inlet pressure is 2.5 MPa. The validation is made on the pressure drop through the porous material. Several temperatures have been tested from 300 K to 600 K. The thermal conductivity of the stainless steel - nitrogen system ( $k_{eff}$ ) is evaluated as a function of temperature using an averaged method between Parallel model and Maxwell Upper Limit [26].

The simulations performed with the initial  $C_2$  value presented some discrepancies compared to the experiments. Thus, the authors have corrected the  $C_2$  value by a multiplication factor of 1.25 (new  $C_2 = 6155000 \text{ m}^{-1}$ ) because the Forchheimer accuracy is  $\pm 30 \%$  [25]. As a result, the computations are in good agreement with the experimental results (Figure 4). Note that, since the Forchheimer's coefficient is linked to the square of the velocity, the changes in  $C_2$  have a negligible effect on the pressure drop for low mass flow rates.

The numerical simulation interest is clear because it can help validating the permeabilities values and understanding the reasons of any possible error. For example here, the complex velocity and pressure field upstream to the porous medium (**Figure 5**) generates a recirculation zone, which makes the pressure measurements sensitive to the spatial position. This impacts the experimental estimation of  $K_D$  and  $K_F$ . As a consequence, it is wanted to verify how the pressure measures are appropriate for the kerosene experiments. A similar cold simulation has thus

been done using kerosene surrogate (n-dodecane), which is liquid at ambient temperature ( $T_{\text{inlet}} = 300 \text{ K}$ ). The mass flow rate is  $65 \text{ mg}\cdot\text{s}^{-1}$ . The density is fixed at  $746 \text{ kg}\cdot\text{m}^{-3}$  and the inlet pressure is 3.5 MPa. The high viscosity of the kerosene ( $\mu = 1.37\cdot 10^{-3} \text{ Pa}\cdot\text{s}$ ) and its low injection velocity bounds the recirculation zones at the cell inlet between  $x \approx -0.025 \text{ m}$  and  $x \approx -0.018 \text{ m}$  (Figure 6). This means that the flow invests the porous medium almost uniformly and, in this case, it is not so important where the pressure sensors are located.

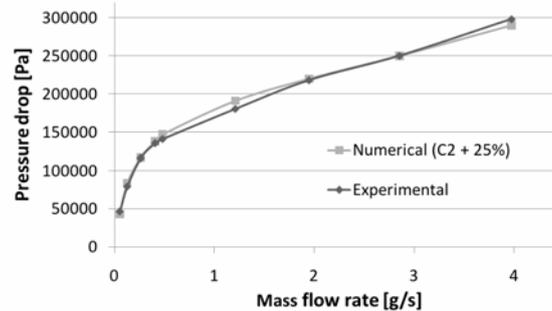


Figure 4 Comparison of the numerical / experimental pressure drop as a function of the mass flow rate at 300 K and 2.5 MPa.

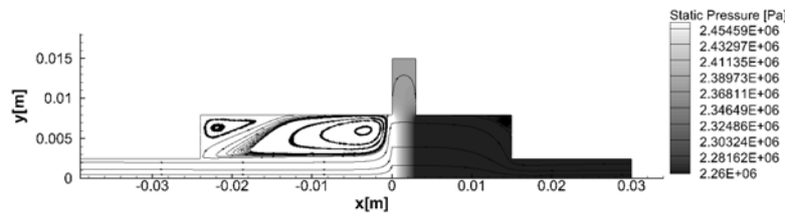


Figure 5 Static pressure distribution with streamlines at 300 K and 2.5 MPa with nitrogen.

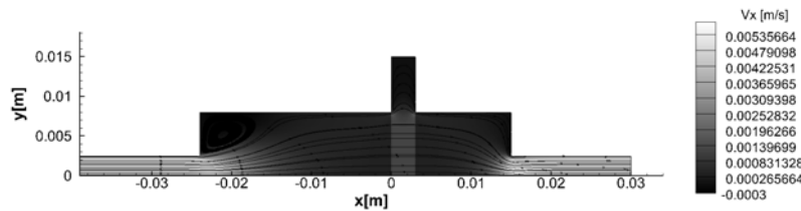


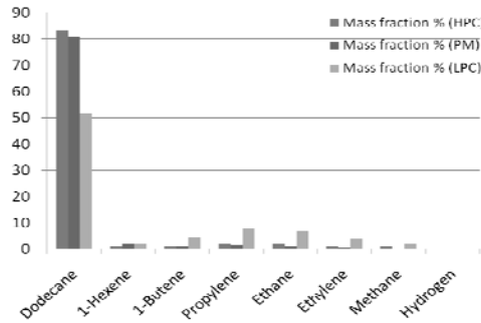
Figure 6 Contour of axial velocity and streamlines for dodecane at 300 K and 3.5 MPa.

## B. Validation with reactive dodecane experiment

The experiments were achieved by Fau *et al.* [7] at 3.5 MPa with a varying mass flow rate between  $60 \text{ mg}\cdot\text{s}^{-1}$  and  $65 \text{ mg}\cdot\text{s}^{-1}$  due to stability problems. The setup furnace temperature was 900 K and the temperature experimentally measured at the outside wall of the cell was 890 K. The fluid temperature measured inside the test cell (in the high pressure chamber) was 820 K. Longitudinal thermal gradients measured experimentally were lower than 10 K roughly between two positions: 3 mm before the porous medium and 3 mm after, following the main fluid flow direction. The experiments (Figure 7) show a high dodecane content upstream to the porous medium (in the **H**igh **P**ressure **C**hamber) and inside the porous medium (83.3 wt.% and 81 wt.% respectively) while a mass fraction around 52 wt.% has been found downstream, in the **L**ow **P**ressure **C**hamber. The experimental quantification of the main species considered by the reduced mechanism (Table 1) are summarized in **Figure 7** for comparison purpose.

Numerically, it is difficult to represent all the experimental fluctuations. As a consequence, several test cases have been computed (Table 2). For all of them, the pressure is 3.5 MPa, the fluid is the dodecane and the reduced kinetic mechanism presented above is used (15 species, 20 reactions). The real gas model enables computing the fluid properties, which requires knowing the species critical properties such as the critical temperature, pressure and volume and the acentric factor. For the first case (Table 2), minimum (Figure 8a) and maximum (Figure 8b) mass fractions at several longitudinal x-axis positions are given for a selection of species. The upstream zone presents a low dodecane content (Figure 8a) due to the recirculation in which quasi static fluid is found. Hexene, pentene and ethylene are found in high proportion (Figure 8b). Comparing these results with the second case (Figure 8c,d) clearly shows that the higher temperature increases the dodecane pyrolysis because the minimum species quantities

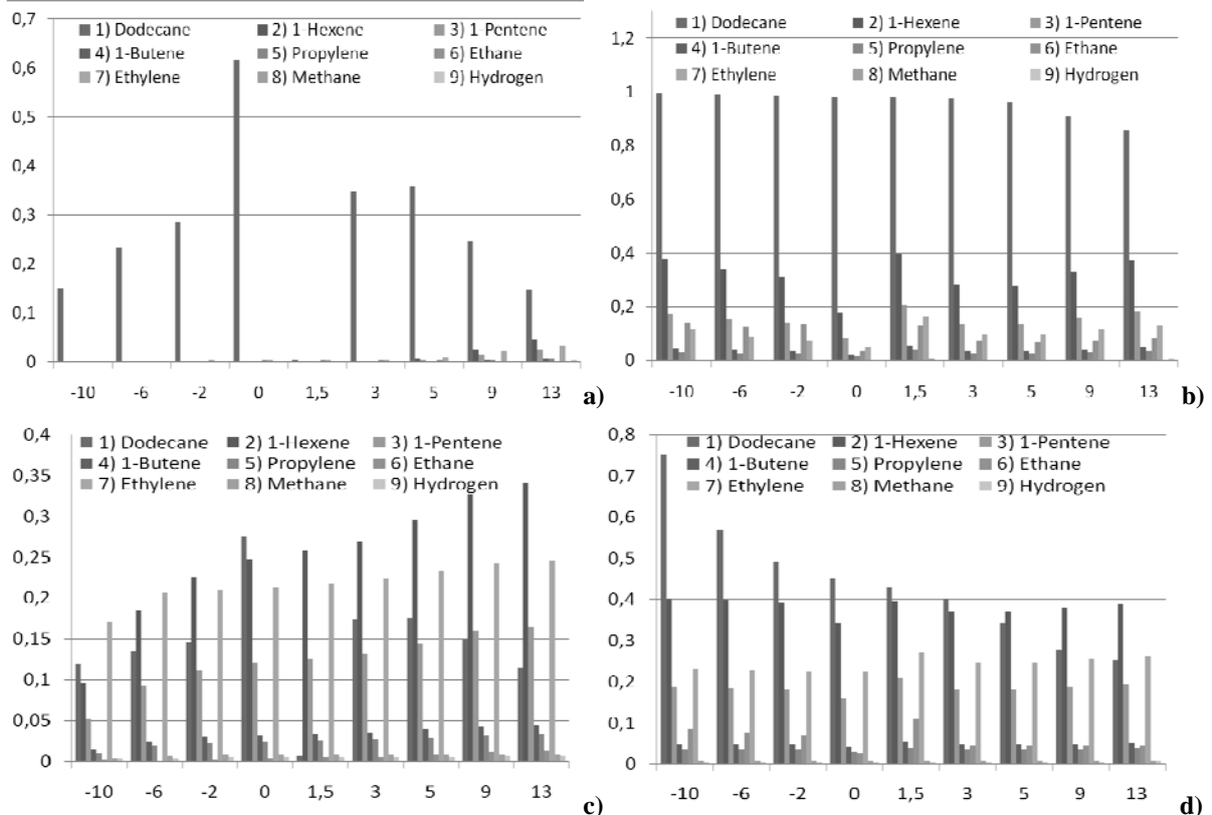
are much higher for this second case (Figure 8c) compared to the first one (Figure 8a). As a consequence, the maximum dodecane quantities are lower for this second case (Figure 8d). The same major products are found. The discrepancies between the two numerical cases are attributed in priority to the heat transfers but the mass flow rate also plays a role which will be investigated in section C.



**Figure 7 Species mass fraction (wt. %) experimentally quantified upstream (HPC), in the porous medium (IPM) and downstream (LPC) during dodecane pyrolysis at  $65 \text{ mg}\cdot\text{s}^{-1}$ , 3.5 MPa and 820 K.**

**Table 2 Numerical test conditions for comparison with fluctuating experiments.**

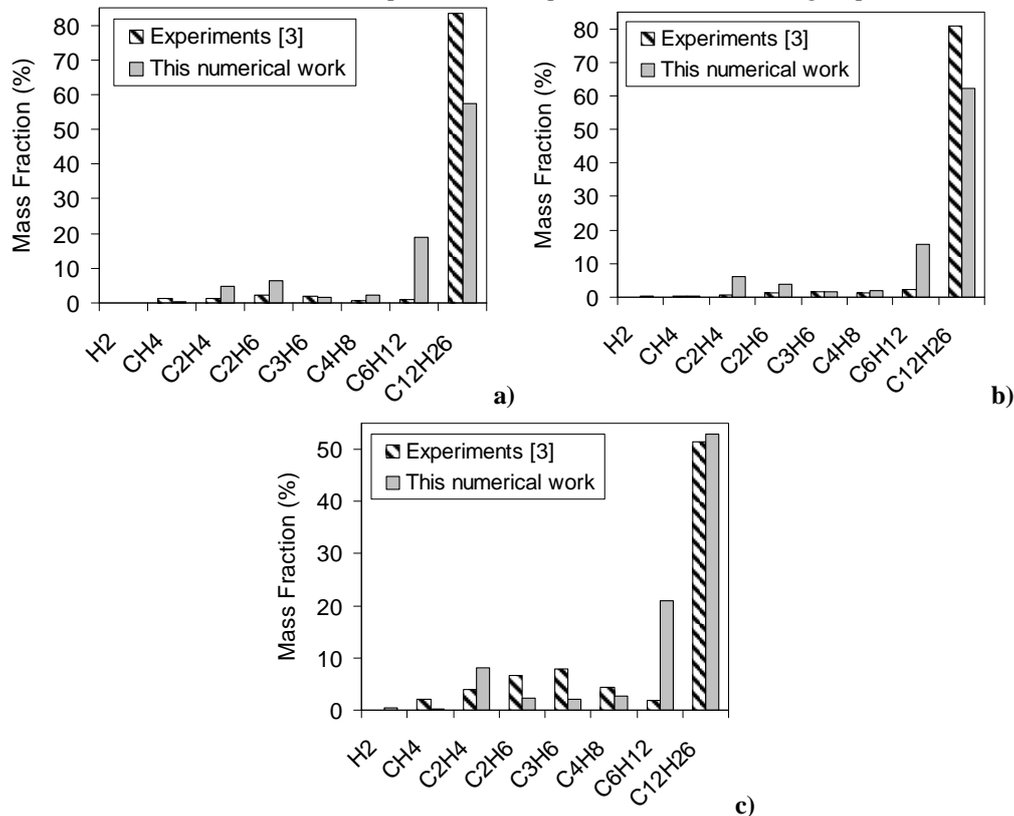
	Inlet fluid temperature (K)	Inlet pipe temperature (K)	Walls of the test cell (except inlet pipe) (K)	Fuel Mass flow rate ( $\text{mg}\cdot\text{s}^{-1}$ )
Case 1	820	820	890	65
Case 2	890	890	890	60
Case 3	1200	1200	1200	60
Case 4	1200	1200	1200	65



**Figure 8 Minimum (a) and maximum values (b) of the species mass fraction computed numerically as a function of the x-coordinate (in mm) for the first case and equivalent data for the second case (c and d).**

The dodecane mass fraction at the cell outlet of case 2 ( $x = 13$  mm just before the outlet pipe) is found between 10 wt.% and 25 wt.%, which is lower than the experimental value. For the case 1, the extreme values are 15 wt.% and 85 wt.%. This includes the experimental value of 52 wt.%. Consequently, the test conditions are very important and slight changes (820 K to 890 K, in conjunction with mass flow rate variations) generate different results which are difficult to validate with the experiments. In addition, the kinetic mechanism has a role in the disagreement found between the numerical and the experimental data. The scheme has been tested (section II.A) and its accuracy is average. The pyrolysis process and the fluid dynamics are closely related because the density changes with the chemical composition and this affects the dynamics into the cell (residence time' and heat transfers' consequence). Hence, the kinetic mechanism becomes a critical parameter and adopting a detailed scheme is preferable.

Finally, because the case 1 is judged to be closer to the experimental conditions, the corresponding results are compared to the experimental ones (Figure 9). Three zones are defined: before the porous medium (for a position  $x = -3$  mm), inside the porous zone ( $x = 1.5$  mm) and at the process outlet ( $x = 30$  mm). The mean conversion rate is over-estimated in the HPC (Figure 9a) and in the porous zone (Figure 9b) by the computations because the spatial average over the  $y$ -axis for each  $x$ -axis position takes the stagnation zones near the wall into account while the experimental sampling locations are closer to the central axis of the cell. The agreement is much better at the bench outlet (Figure 9c). Nevertheless, due to the uncertainties presented above, there are too much parameters acting on the system to be certain that the numerical simulation is validated. This is particularly due to the kinetic scheme. The hexene content is overpredicted by the computations because the secondary mechanism of this species is not considered and it cannot be consumed. This explains the low quantities found for the light species like  $\text{CH}_4$ ,  $\text{C}_2\text{H}_4$ .

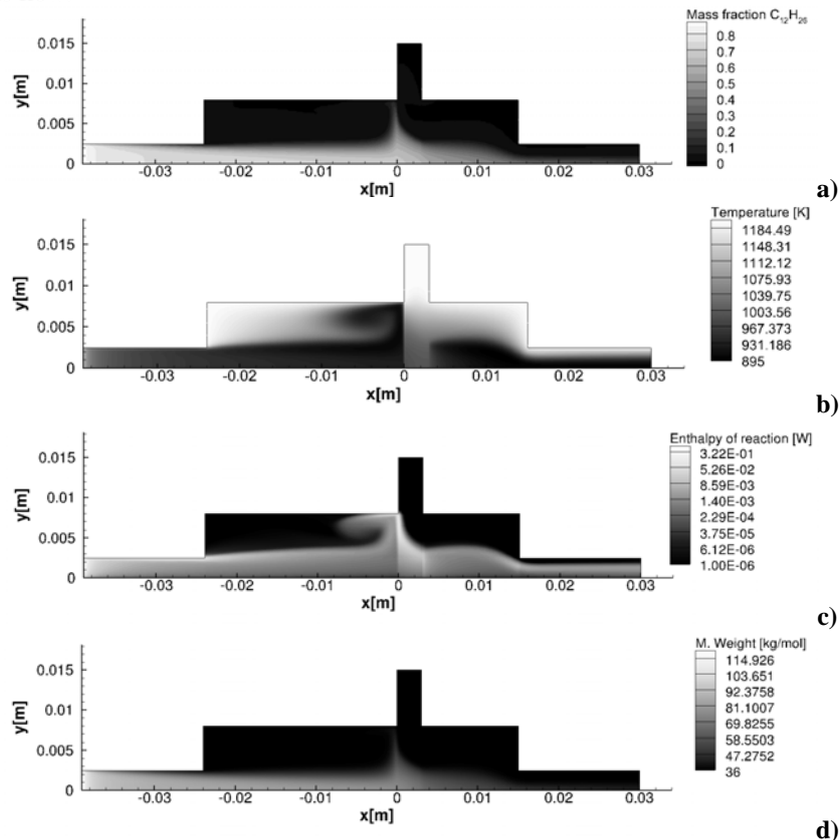


**Figure 9. Comparison of the pyrolysis products composition computed for case 1 and measured experimentally in the HPC (a), inside the porous zone (b) and at the process outlet (c).**

To verify the effect of the chemistry, a two-steps kinetic scheme is used to redo the same test case 2. A reduction of only 17 K of the fluid temperature between the inlet and the outlet is observed and the net heat flux at the wall is 3.2 W. This explains the very low pyrolysis rate at the outlet (remaining dodecane mass fraction of 98 wt. %). Consequently, the experimental values (pyrolysis rate of 48 wt. %) are bounded by the numerical ones obtained with two different kinetic schemes (pyrolysis rate of 80 wt. % for the detailed scheme and of 2 wt. % for the 2-steps scheme). The lower reactivity of the simplified mechanism is due to the radicals, such as methyl and monatomic hydrogen, which are not in the two-steps mechanism while they increase the reaction rate.

### C. Use of the numerical simulation to investigate the reactive porous flow

Considering the third case (Table 2), the conversion rate of dodecane reaches 94 wt. % at the cell outlet (**Figure 10a**). A maximum thermal decrease of about 300 K is found (**Figure 10b**). This temperature fall is directly related to the enthalpy of reaction (**Figure 10c**), thus to the reaction zone. This point is particularly important because following the temperature would give precious indication on the chemistry. This is confirmed by looking at the molecular weight profile (**Figure 10d**). Such a result demonstrates the benefit of the computations compared to the experiments. This also shows that the recirculation zones, where the temperature is maximum (at  $y = 0.008$  m and for  $x = -0.024$  m and  $x = 0.013$  mm), do not really participate to the pyrolysis. This is satisfactory when conducting pyrolysis in porous flow.

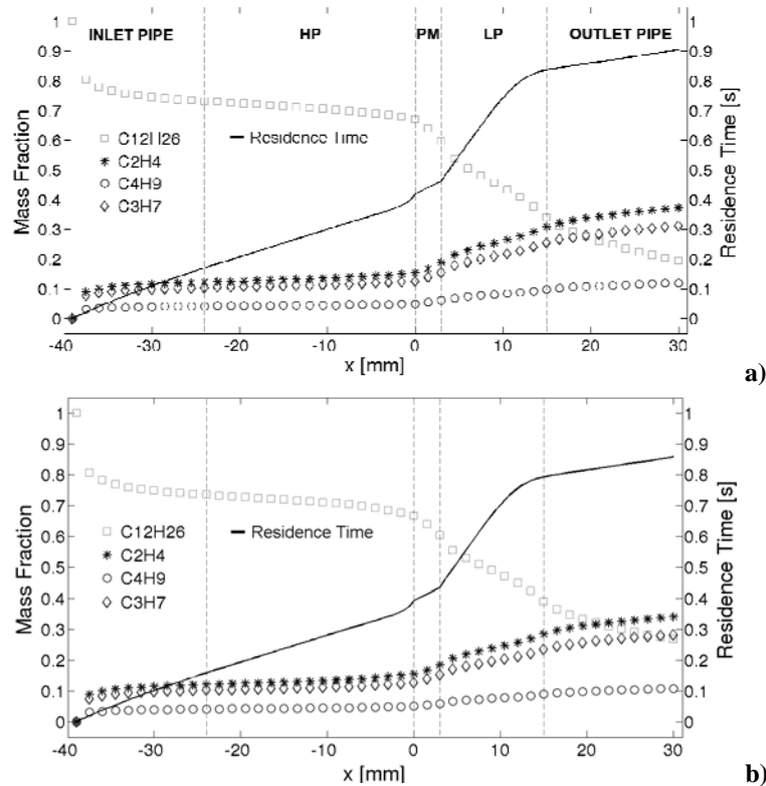


**Figure 10**  $C_{12}H_{26}$  mass fraction (a), temperature distribution (b), enthalpy of reaction (c) and molecular weight (d) for the two-steps mechanism at 1200 K, 3.5 MPa and  $60 \text{ mg}\cdot\text{s}^{-1}$

This information concerning in which part of the system does the pyrolysis take place can be completed. Looking at the fuel consumption along the cell following the streamline on the axis of symmetry, a fast dodecane cracking occurs immediately at the inlet of the cell due the high inlet temperature (Figure 11a). The dodecane mass fraction entering the HPC is about 72 wt. % and it is around 66 wt. % at the porous medium inlet. It is 55 wt. % at its outlet, 28 wt. % at the LPC outlet and 18 wt. % at the pipe outlet. This means that the porous medium only represent 11 wt. % of the converted dodecane compared to the 82 wt. % which are converted in the entire system. Thus, the porous media represents about 14 % of the pyrolysis activity and its thickness (3 mm) corresponds to 4 % of the total length of the system (70 mm). In terms of residence time, the porous medium represents less than 10 %. The slope of the curve giving the residence time shows that the fuel is slowed down in the LPC because the slope increases. The fuel velocity increases in the outlet pipe due to the cross-section reduction.

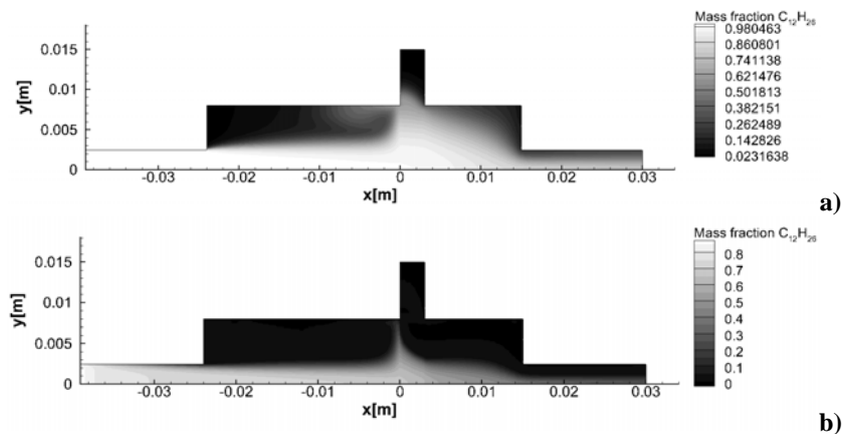
Since the experimental mass flow rate slightly fluctuates between  $60 \text{ mg}\cdot\text{s}^{-1}$  and  $65 \text{ mg}\cdot\text{s}^{-1}$ , its effect has been investigated (case 3 and 4). The total residence time and its distribution are not so different (Figure 11b). Increasing the fuel mass flow rate decreases the pyrolysis rate at the outlet which means that the density and the residence time effect (acting similarly) have a lower impact than the convective heat transfers which increase (80 wt.% and 72 wt.% of pyrolysis rate for cases 3 and 4 respectively). This discrepancy of about 10 % is directly linked to the mass flow rate fluctuations of about 10 %. This demonstrates the direct effect of the fluid flow on the chemical

parameters. Thus, simulating the experiments in which the flow rate varies is difficult numerically where the flow rate is fixed. This contributes to explain the disagreement found in section B.



**Figure 11. Species mass fraction and residence time along the axis of symmetry for the third case at  $60 \text{ mg}\cdot\text{s}^{-1}$  (a) and the fourth case at  $65 \text{ mg}\cdot\text{s}^{-1}$  (b).**

The effect of the temperature (comparison of cases 1 and 4) on the chemistry is represented by Figure 12. The strong chemical differences regarding the 2-D dodecane content (Figure 12a, b) are directly related to the thermal conditions. The residence time distribution is similar for case 1 (Figure 12c) than for cases 3 and 4 (Figure 11). Nevertheless, the residence time is about three times higher in case 1 than in other cases. This demonstrates that the temperature impacts the chemistry and the density, so the residence time, but its importance is much higher. A temperature decrease increases the density and decreases the velocity through the mass conservation. Thus, the residence time is higher but this is not balanced by an increased convective heat flux. This spate of events is important to analyse the results and this cannot be done experimentally, which shows the good complementarity of both tools.



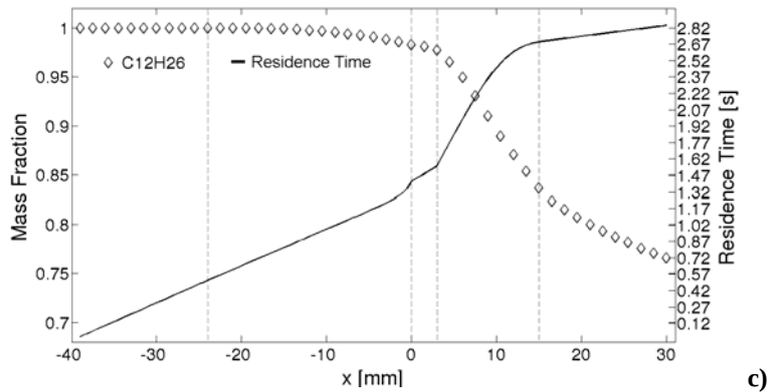


Figure 12. Dodecane mass fraction inside the system for case 1 (a) and case 4 (b) with residence time and mass fraction distribution along the symmetric x-axis (c).

#### D. Specific analysis of coupled phenomena in the porous medium

Comparing the dodecane conversion at the inlet and at the outlet of the porous medium for the fourth case, a coupling between the fluid flow and the chemistry is found (Figure 13a). Close to the symmetric axis ( $y = 0$  mm), the flow speed is the highest and the conversion is limited. The chemical gradient is consequently low (68 wt. % to 60 wt. %). However, for  $y = 4$  mm and higher, the fluid velocity decreases, the residence time and the heat flux increase (mass fraction difference at  $y = 4$  mm of 48 wt. %). Thus, the pyrolysis is impacted by the border effect (wall of the cell). It is possible to plot it as a function of the x-axis, i.e. the porous medium thickness (Figure 13b). The Figure 13b shows that after 1 mm of porous flow, the wall impacts the flow on a distance of 1.5 mm. As a consequence, the radius of the permeable sample should not be lower than 1.5 mm. It should even be much more in order to limit its effect. The thicker the porous material in the present permeation test cell, the bigger should be the diameter in order to limit this border effect. This result is of major importance to design the experiments and to analyse the data. This is related to both dynamic and thermal boundary layers.

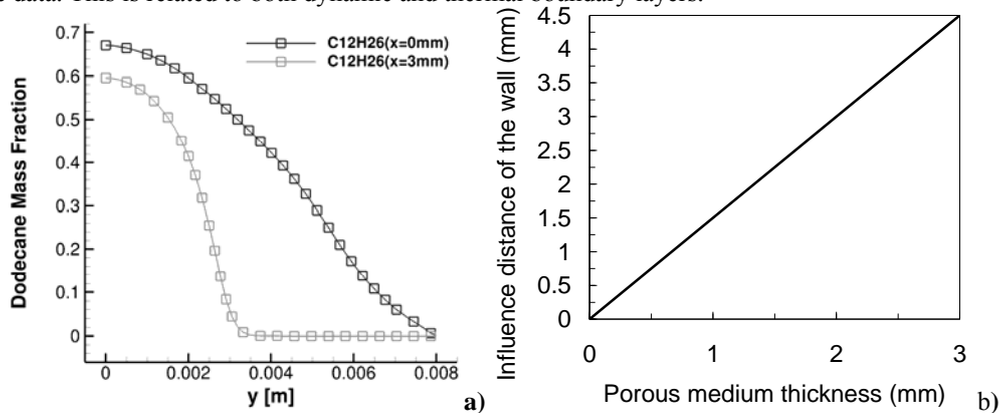
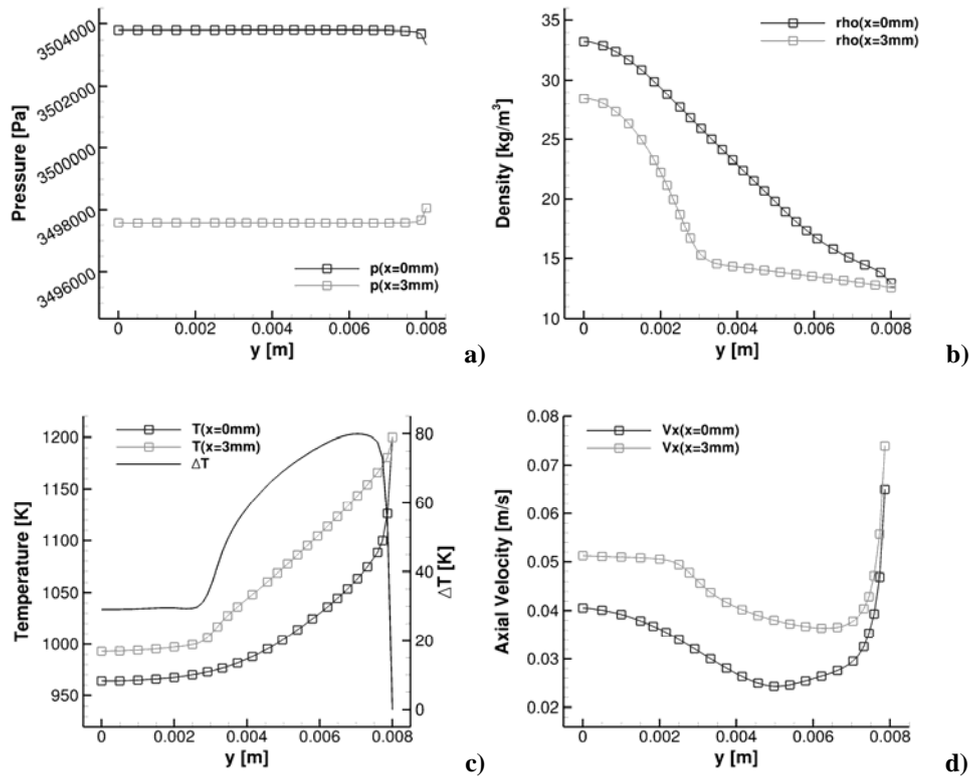


Figure 13. a:  $C_{12}H_{26}$  mass fraction in the crosswise direction at the inlet and at the outlet of the porous medium at 1200 K, 3.5 MPa and  $65 \text{ mg}\cdot\text{s}^{-1}$  and b: effect distance from the wall as a function of medium thickness.

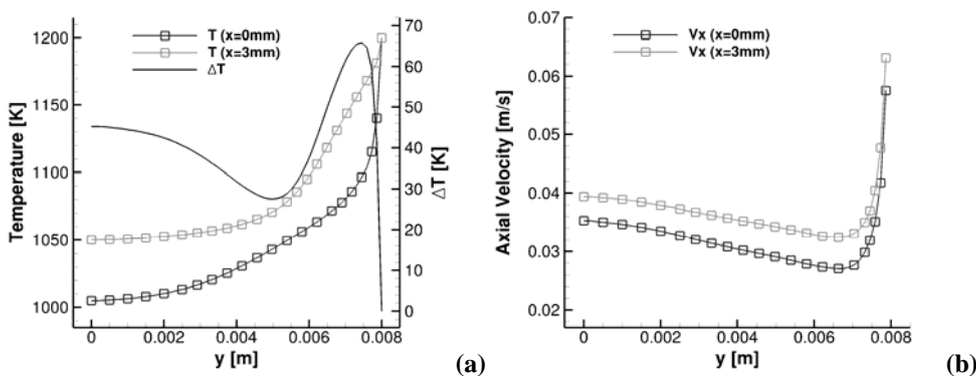
Focusing on the porous zone, Figure 14 shows the profiles of pressure, of density, of temperature and of the axial velocity along the y-coordinate at the inlet and at the outlet surface of the porous medium. The pressure drop, due to the permeable sample, is 6800 Pa and it shows a uniform distribution on both sides of the porous zone. Although the pressure does not change in the crosswise direction (Figure 14a), the density decreases moving from the axis of symmetry toward the walls and from the upstream side to the downstream one (Figure 14b). This is directly due to the heat transfers (Figure 14c) and it acts on the fluid velocity (Figure 14d).

The spatial thermal gradient at the porous medium inlet presents a parabolic shape (Figure 14c) while it tends to be linear at the outlet. This could be due to the software assumptions considering equilibrium between solid and fluid. This may also be realistic in case of low speed flow (the conduction gets more important than the convection). In this latest case, it could be interesting to determine the length required by the fluid to present a complete linear

profile. This would correspond to the thermal establishment length. Another interesting point is the location of the minimum velocity (**Figure 14d**). It is around  $y = 0.005$  m at the porous medium inlet and  $y = 0.007$  m at the outlet. This minimum is spatially shifted due to the thermal or chemical phenomena. Non-reactive hot flow could help understanding the reason of such a speed shift. Thus, a similar test case without reactions has been achieved for comparison purpose (Figure 15). Different temperature and velocity profiles have been obtained in comparison with Figure 14c, d. The difference is attributed to the chemistry, which impacts the molecular weight profile. The velocity profiles at the 0 mm and 3 mm positions are perfectly similar (the velocity amplitude only is shifted but not spatially). As a consequence, the modification of the minimum speed observed on **Figure 14d** is due to the chemical reactions which modify the dynamic boundary layer.



**Figure 14** Pressure (a), density (b), temperature (c) and axial velocity (d) along y-direction at  $x = 0$  mm and  $x=3$  mm at 1200 K, 3.5 MPa and  $65 \text{ mg}\cdot\text{s}^{-1}$ .



**Figure 15** Temperature (a) and axial velocity (b) along y-direction at  $x = 0$  mm and  $x=3$  mm without pyrolysis reactions in the porous medium (1200 K, 3.5 MPa,  $65 \text{ mg}\cdot\text{s}^{-1}$ ).

#### IV. Conclusion

The numerical simulation and the experimental tests are two complementary approaches, which are both suited for the study of fuel pyrolysis in porous media. Experimentally, all possible phenomena are taken into account with their intrinsic complexity and coupling, possibly under extreme conditions (1200 K and 6 MPa). Numerically, the heterogeneity profiles and the effect of the test bench itself (size and design of the permeation cell) have been observed finely despite the validity of the results is subject to the numerical, physical and chemical assumptions. A European Collaboration has been set up to provide a wide expertise in this field and to propose a validated numerical tool using detailed chemistry. A specific validation under no flow conditions has been first achieved for the chemistry since this is a major point in the numerical simulation of the reactive flow. After validation of the CFD simulation under inert conditions, its application to reactive test case presents average agreement with the experimental results. The computation method is judged promising. A modification by 25 % of the Forchheimer's permeability tends to confirm the accuracy which was suggested experimentally ( $\pm 30\%$ ). The pressure sensor position is suggested to be of importance considering the pressure drop measures. At least qualitatively, the relationships between the heat and mass transfers and the chemistry have been considered. The effects of the choice of the kinetic schemes on the multiphysics' simulation have been presented. Detailed mechanisms are clearly necessary. The comparison between numerical and experimental data appears difficult due to the complexity of accurately representing the experimental conditions. The numerical simulation showed strong benefits in understanding a possible border effect related to the permeable sample thickness. The relationship between the temperature field and the reaction zone tends to demonstrate the major impact of chemistry in the heat flux repartition. As a future work, other detailed kinetic mechanisms should be tested to reinforce the chemical modelling and to determine which level of accuracy is necessary for the chemistry to have a good representation of the phenomena. The final aim is to consider methane for rocket engines application.

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