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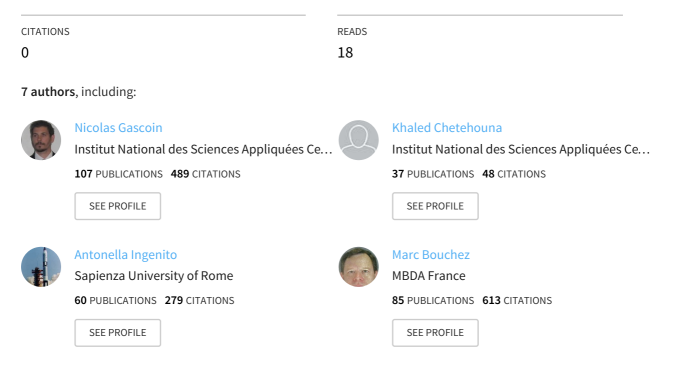
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# Experimental Study of Pyrolysis-Combustion Coupling in a Regeneratively Cooled Combustor: System Dynamics Analysis

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## Abstract

Scramjets are suitable for hypersonic flight, but their use requires the ability to ensure their thermal protection. In this context, a remotely controlled fuel-cooled combustor, suitable for the experimental analysis of the pyrolysis-combustion coupling characterizing a regeneratively cooled combustion chamber when a hydrocarbon propellant is used, has been designed. Similitude rules were used. Ethylene is used as fuel, air as oxidizer, with an equivalence ratio between 1.0 and 1.5 and a fuel injection pressure between 1 and 10 bar. Experiments are realized by varying operating conditions, to determine their impact on combustor heat transfer dynamics. Previous numerical results have been confirmed. A hysteresis effect has been demonstrated. It has been observed that system response time to fuel mass flow rate increases is lower (of about 40 to 50%) than system response time to fuel mass flow rate decreases and that a rise in equivalence ratio from 1.0 to 1.25 produces an increase in system response intensity that is, following the operating conditions, in the range from 90% to 170%. It has also been seen that an increase in equivalence ratio from 1.0 to 1.5 produces a raise of the temperature of the fuel-coolant (of about 40 to 50%), due to the increase in the emissivity of the flame.

# **Keywords**

- ➤ Scramjet
- Hydrocarbon Fuel Pyrolysis

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- Combustion-Pyrolysis Coupling
- Regenerative Cooling
- Parametrical Analysis

Nome	Nomenclature			
$C_p$	=	heat capacity, J.kg <sup>-1</sup> .K <sup>-1</sup>		
D	=	diameter, m		
Н	=	height, m		
k	=	thermal conductivity, W.m <sup>-1</sup> .K <sup>-1</sup>		
L	=	length, m		
'n	=	mass flow rate, kg.s <sup>-1</sup>		
Р	=	pressure, Bar		
S	=	cross section, m <sup>2</sup>		
t	=	time, s		
Т	=	temperature, K		
v	=	velocity, m.s <sup>-1</sup>		
Greel	Greek Symbols			
З	=	emissivity, dimensionless		
φ	=	equivalence ratio of fuel to oxidizer, dimensionless		
μ	=	dynamic viscosity, Pa.s		
ρ	=	density, kg.m <sup>-3</sup>		
σ	=	Boltzmann's constant, W.m <sup>-2</sup> .K <sup>-4</sup>		
Subsc	Subscripts			
сс	=	cooling channel		
f	=	fuel fed to the cooling channel		
g	=	combustion gases		
g,th	=	combustion gases in the vicinity of the thermocouple		
hf	=	fuel at cooling channel outlet, before injection in the burner		

# **1. Introduction**

th

For decades, hypersonic flight has challenged researchers. Indeed, both reusable aircrafts and space vehicles able to reach velocities over Mach 5 may provide significant commercial benefits. Hypersonic flight is also suitable for military applications, as the development of hypersonic missiles [1]. Because of the operational limitation of gas turbine engines to speeds generally below Mach 4, the propulsion systems for high-speed flight must integrate engines capable of propelling the vehicle when its velocity becomes hypersonic. To this end, Scramjets, which are air-breathing jet engines suitable for atmospheric flight, represent a very effective solution [2-4].

Yet, when flight velocities increase beyond supersonic speeds, the difficulties accompanying aircraft design grow, particularly with respect to vehicle thermal protection, which becomes a critical problem notably with regard to the engine combustion chamber, exposed to very high temperatures, higher of all known structural materials capability [5-8]. The combustor wall must also face a very high heat load (at Mach 8 the average heat flux passing from the combustion gases to the combustor wall has been estimated to be about 1.7 MW.m<sup>-2</sup>) [9]. Furthermore, the heat flux distribution characterizing Scramjet engines can result in hot spots where heat flux is expected to exceed 5.7 MW.m<sup>-2</sup> [7].

Several cooling techniques can be used to protect the combustor internal surfaces, in order to maintain its structures at a limited and safe temperature. In general, they are divided into passive ones, where no coolant is used, and active ones, where a coolant is used [10,11]. Active cooling techniques consist in the injection of a coolant through cooling channels located in the combustion chamber wall. As at hypersonic speeds air stagnation temperature is too high to use incoming air as an effective coolant, the fuel becomes the primary cooling source, providing both coolant capability for heat removal and heat sink capacity for storing the absorbed heat [7,12]. In this sense, endothermic hydrocarbon jet fuels represent very attractive coolants. Their heat sink potential is both a sensible one, through fuel heating (in the order of 1-1.5 kJ.kg<sup>-1</sup>), and a chemical one, through decomposition heat-absorbing reactions that can occur when the fuel is heated at temperatures over 750 K (in the order of 1.5-3.5 kJ.kg<sup>-1</sup>) [7,13].

One of the most common active cooling technique using the fuel as coolant is regenerative cooling. The fuel flows through cooling channels located between the inner and the outer walls of the engine, before being injected in the combustor; a counter-flow heat exchange between the fuel-coolant and the burned gases is thus established [5,14]. Regenerative cooling is particularly effective when an endothermic propellant is used as fuel-coolant as fuel decomposition in the cooling channels also generates light species, mainly hydrogen, ethane, ethylene, acetylene, whose ignition delay times are very low when compared to the initial heavy propellant. Thus, fuel combustion kinetics becomes faster and the performance of the engine improves [13-15].

The design of a regeneratively cooled engine is very challenging, as the definition of an engine regulation strategy allowing to master the various conditions encountered during the flight requires the ability to manage several issues [13-17]. First, the design of the combustion chamber and of the cooling channels. Second, the evaluation of the heat fluxes passing from the combustion gases to the combustor internal walls and of the highest heat flux regions. Third, the choice of the materials that are more suitable for combustor structures. Forth, the understanding of the complex pyrolysis reactions characterizing fuel decomposition and of the coupling between fuel pyrolysis and fuel combustion (which affect each other). Finally, when the engine is fueled with hydrocarbons, the cooling system of the coking activity typical of hydrocarbons thermal decomposition.

These issues have been studied in the framework of various programs [4-6,7-14,17,22-42]. Many numerical studies, directed at estimating the heat flux density passing from the combustion gases to the combustor wall, the highest heat flux combustor regions and the heat flux absorbed by the pyrolyzing fuel are available [4-6,9,10,12-14]. The design of the regeneratively cooled combustion chambers, the definition of the shape of the cooling channels and the choice of materials appropriate for endothermic fuel cooled structures have been investigated, mainly by means of numerical calculations, since the 1990s [5,22-26]. Numerical investigations have also been performed to determine the effect of the shape and of the number of the cooling channels on combustion chamber walls, coolant temperature and fuel-coolant pressure drop [5,24]. The physical properties of composite materials suitable for the manufacturing of the combustor and of the cooling channels walls have been thoroughly studied [17,27-30]. The possibility of employing carbon composite materials instead of steel and copper alloys has been investigated [28-30].

Studies have also been performed on hydrocarbons thermal pyrolysis [31-35]. Within the framework of the PROMETHEE project, experiments on hydrocarbon jet-fuels pyrolysis have been run, to identify the main products of hydrocarbons decomposition under supercritical conditions (fuel pressure up to 80 bar and fuel temperature up to 1200 K). The main cracking products have been identified for various operating conditions of pressure, temperature and residence time [31]. In this context, it is worth mentioning the studies of Daniau and Sicard [32] and Vincent-Randonnier et al. [33], who showed that fuel pyrolysis in a regeneratively cooled engine is normally a kinetically controlled process. The COMPARER project, developed by the University of Orleans and by MBDA-France, aimed at identifying methods suitable to measure some of the most important parameters characterizing hydrocarbon-fueled regeneratively cooled engines, such as fuel mass flow rate and fuel composition at combustor inlet. Within its framework, it is important to evoke the studies of Abraham et al. [34] and Gascoin et al. [35].

Hydrocarbon fuels coking activity has also being deeply investigated [36-41]. Song et al. [36] and Chin and Lefebvre [37] studied the effects on jet fuels cooking activity of fuel and wall temperatures, fuel mass flow rate, velocity and pressure. They observed that fuel temperature and velocity are the most important factor affecting deposits formation. Sobel et al. [38] and Spadaccini et al. [39] investigated coke deposition in heated aircraft fuels by considering a wide temperature range, going from 420 to 820 K, in order to analyze the contribution to carbon deposition of both thermal-oxidative and pyrolytic deposits. They found out that for a liquid fuel the variation of fuel pressure has a minor effect on coke formation. Maurice et al. [7] and Gascoin et al. [40] indicated that there is a residence time influence on pyrolytic deposition levels beyond the influence associated with conversion level. Finally, the analysis of the effect of the operating conditions of the cooling system, realized by considering the coupling among flow, heat transfer and chemical reactions in the cooling channels, has been carried out by Bao et al. [10].

However, to the author's knowledge, none of published studies focused on the most important aspect characterizing regeneratively cooled combustors, i.e. the strong coupling between fuel decomposition and fuel combustion, which makes the definition of an engine regulation strategy very hard. This is a very important point, as even the effect of the most important operating parameter, i.e. fuel mass flow rate, on engine thrust is not easy to determine. In fact, engine thrust depends both upon fuel mass flow rate and fuel composition, which are linked to each other, as fuel composition vary with fuel residence time in the cooling channels that, in turn, vary with fuel mass flow rate. A mass flow rate increase would probably result in a residence time decrease and, consequently, fuel decomposition could be less enhanced. Thus, the propellant injected in the combustor would contain less low ignition delay time species. The global effect could even be a drop in the engine thrust or even flame extinction [13,14,43-45].

To the author's knowledge, the only work proceeding along this path is the one of Gascoin et al. [13], in which the impact of fuel mass flow rate on important parameters such as burned gases temperature and fuel-coolant temperature and residence time in the cooling channels has partially been defined by means of numerical calculations. This work has highlighted the complexity of the matter, showing the existence of hysteresis phenomena that make system control even more challenging. Now, the following tasks must be accomplished: 1) the already acquired numerical knowledge must be validated by means of experiments; 2) the characteristic times of all the involved physical and chemical phenomena, which are still unknown, must be determined and compared to those specific of engine control, to understand with which degree of effectiveness operating parameters variations can influence engine thrust; 3) the regeneratively cooled engine potential in terms of heat recovery and of utilization on long-live aircraft systems (more than 2000 hours) when a hydrocarbon fuel is used must be fully understood.

The objective of this study is to validate the numerically acquired knowledge and to understand the dynamics of a regeneratively cooled Scramjet fueled with an endothermic hydrocarbon, aiming at developing an engine control strategy suitable for a vahicle flying at hypersonic velocities. Thanks to an innovative experimental set-up, consisting of a lab-scale fuel-cooled combustor suitable for the experimental study of fuel combustion – fuel pyrolysis coupling, some of the most relevant issues characterizing this cooling technology have been experimentally investigated [14]. Experiments are run using ethylene as fuel-coolant and air as oxidizer. The effect of two operating parameters, i.e. fuel mass flow rate and fuel to oxidizer equivalence ratio, is analyzed.

The evaluation and the comprehension of the effects of fuel pyrolysis-fuel combustion and the identification of the most important system characteristic times are the main goals of this work. The effects of fuel mass flow rate and fuel to oxidizer equivalence ratio on the heat flux density passing from the combustion gases to the combustor wall, the sensible heat flux absorbed by the fuel-coolant, the heat loss due to the exhaust gases and the heat loss due to heat transfer from the combustor external surface to the surrounding environment and on the heat transfer efficiency of the combustor will be presented in a future work.

In the authors' opinion, this study will improve the knowledge of the scientific community on the thermal management and dynamic control of a Scramjet combustion chamber equipped with a regenerative cooling system.

### 2. Materials and Methods

#### 2.1. Description of the experimental set-up

The bench used to run all the experiments consists in an automated combustion chamber which enables the study of the dynamics of a regenerative cooled engine. It allows the experimental investigation of the coupling between fuel pyrolysis and fuel combustion and of the coking activity of the fuel. The combustor, whose height is of 0.5 m, is composed of two stainless steel co-axial tubes having respectively a diameter of 0.1 m (the internal one) and of 0.3 m (the external one) and a thickness of 1 mm. An insulation blanket, made of a ceramic refractory, whose density is 128 kg.m<sup>-3</sup> and whose thermal conductivity is 0.33 W.m<sup>-1</sup>.K<sup>-1</sup>, is located between them. The cooling channels are reproduced by a single rolled-up stainless steel tube having a length of 40 m, an internal diameter of 1 mm and a thickness of 1.1 mm, which passes in the combustion chamber and is placed on the internal surface of its internal wall.

A scheme of the experimental set-up is given in fig. 1. The path followed by the fuel fed to the combustor, which is ethylene, is represented in blue. As shown in fig. 1, this hydrocarbon enters the rolled-up stainless steel cooling channel at position 1, at the top of the combustor. Then, it flows into this channel, absorbing part of the heat flux passing from the combustion gases to the internal surface of the combustion chamber wall. Thanks to this, its temperature raises reaching values that are sufficiently high to permit its thermal decomposition. As a result of this decomposition, fuel composition at position 2 (where the rolled-up cooling channel leaves the combustion chamber) is unknown. The mixture resulting from the pyrolysis of the ethylene originally fed to the combustor is finally injected in the burner.

The burner used to generate the flame is a Five North American SPB5 pilot burner, provided with a venturi jet mixer body which enables to mix the fuel and the oxidizer (which is air at ambient pressure) in order to produce a pre-mixed combustion, having a nominal

capacity of 6 kW. The choice of a premixed flame was made to facilitate the numerical study of the system, as a diffusion flame is much more complex to reproduce with CFD calculations. Before being injected in the burner, the fuel passes through a steel permeation cell (indicated as *PC* in fig. 1), containing a stainless steel porous disk which can collect the particles of coke generated by the decomposition of the fuel and separate them as a function of their granulometry. Each disk has a diameter of 3 cm and a thickness of 3 mm, presenting an overall porosity of 30%; the average diameter of the pores is 4.1  $\mu$ m [46]. The properties of the coke that accumulates in permeation cell could be later analyzed by GC-MS analysis.

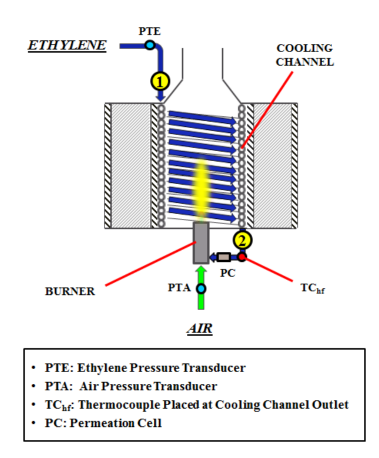


Figure 1. Scheme of the regeneratively cooled combustion chamber

The ethylene and the air that are used to carry out the tests are stocked at high pressure (above 100 bar) and are fed to the combustor by using two high-precision mass flow controllers, respectively a Bronkhorst F-201CM-10K-RDA-88-K for ethylene (with a range of 0-0.2 g.s<sup>-1</sup>) and a Bronkhorst F-202AV-M20-RDA-55-V for air (with a range of 0-5 g.s<sup>-1</sup>). A pressure transducer with a range between 0 and 100 bar (Baumer CTX155B115, indicated as PTE in fig. 1) and a pressure transducer with a range between 0 and 10 bar (Baumer CTX255B255, indicated as PTA in fig. 1) are used to measure respectively the pressure of the

ethylene entering the cooling channel and of the air entering the burner. Temperatures in the combustion chamber are measured by chromel–alumel type K thermocouples, which can be shifted from the wall to the axis of the combustor. Their positions are shown in fig. 2, where the base of the combustion chamber is used as reference. A chromel–alumel type K thermocouple (indicated as  $TC_{hf}$  in fig. 1) is located at the outlet of the cooling channel from the combustor, allowing to measure the temperature of the heated fuel. The accuracy of the thermocouples, provided by TC Thermocouples, is 0.75% of the full scale. As, following the presence of radiation errors, the temperature of a flame measured by means of thermocouples may significantly differ from the real one, all the measured temperatures have been corrected in order to obtain the real ones. The approach used to perform these corrections is detailed in paragraph 2.3.

The pressure transducers and the thermocouples are connected to a data acquisition system (National Instrument PXIe-1073) by means of two acquisition cards, respectively a National Instrument PXI-6229 for the pressure transducers (16-Bit, 32 analog inputs, 48 digital inputs/outputs) and a National Instrument PXIe-4353 for the thermocouples (32-Bit, 32 thermocouple input channels).

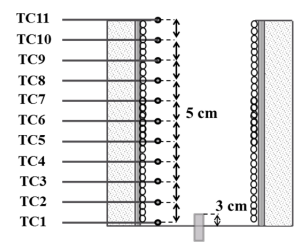


Figure 2. Scheme representing the position of the thermocouples measuring the combustion gases temperature

The experimental bench is completely automated; it can be controlled from a personal computer with an application created by using NI LabVIEW, which allows to open and to close each valve, to ignite and to turn off the burner, to start and to stop data acquisition. A video camera ( $1280 \times 720$  pixels and 15 images/s) is used to record a video of the flame during each test. Further details on the test bench have been given in a previous work [13].

#### 2.2. Test Methodology

Tests are performed under both stationary and transient conditions. The pyrolysis of ethylene and consequent effects on heat transfer from the combustion gases to the combustor wall and to the cooling channel are investigated by varying two operating parameters that are fuel mass flow rate and fuel to oxidizer equivalence ratio. This equivalence ratio is modified by varying the ratio between ethylene mass flow rate and air mass flow rate. A brief review of the tests which are presented in this work is given in table 1.

The sequence of each test is programmed before its start using the LabVIEW application. Tests are carried out by varying a single input parameter, in order to determine its impact on the dynamics of the test bench. During each experiment, this parameter is increased twice and then decreased twice, while keeping the others constant. This enables catching the hysteresis phenomena whose presence has previously been detected by means of numerical calculations that have been detailed in a previous work [13]. Operating conditions are varied only when steady state has been achieved. For example, let us consider the case number 1. The test is started with a fuel mass flow rate of 0.020 g.s<sup>-1</sup>, a fuel inlet pressure of 1 bar, and an equivalence ratio of 1.0; when steady state is achieved, the fuel mass flow rate is increased from 0.020 to 0.030  $g.s^{-1}$  to let the system reach a new stationary state. In the same way, the fuel mass flow rate will be later increased to 0.040  $g \cdot s^{-1}$ , then reduced twice respectively to 0.030 and 0.020 g.s<sup>-1</sup>, to come back to the initial operating conditions. Meanwhile, fuel pressure and equivalence ratio are not modified. The combustor is assumed to be at steady state when the variation rate of the temperature of the fuel at the outlet of the cooling channel becomes lower than 0.1 K.min<sup>-1</sup>. Thus, during each experiment, the test bench achieves steady state five times. For simplicity purposes, they will be numbered consecutively, beginning from steady state 1 and finishing with steady state 5.

Test case	Fuel mass flow rate (g.s <sup>-1</sup> )	Equivalence ratio	Fuel Inlet Pressure (bar)
1	0.020-0.030-0.040	1.50	1.0
2	0.020-0.024-0.028	1.00	1.0
3	0.020-0.024-0.028	1.25	1.0
4	0.020-0.024-0.028	1.50	1.0
5	0.024	1.00-1.25-1.50	5.0
6	0.020-0.024-0.028	1.00	4.0
7	0.020-0.025-0.029	1.25	5.0

8	0.020-0.024-0.029	1.50	4.5
9	0.016	1.25	5.00-8.0-11.0

Table 1. Test cases for different operating conditions

At the start of each test, all the thermocouples are placed at 1 cm from the wall of the combustor, at position P1, with the exception, as already explained, of the two thermocouples indicated respectively as TC2 and TC7, which are placed at 5 cm from the wall, on the axis of the combustion chamber, at position P3 (fig. 3-a). When a steady state is achieved, the thermocouples which are at position P1 are first shifted at position P2 (at 3 cm from the wall, fig. 3-b), then at position P3 after the stabilization of the measured temperatures (fig.3-c). Before varying the input parameter whose effect is studied, the thermocouples which had been displaced are moved back to their original positions (fig. 3-d).

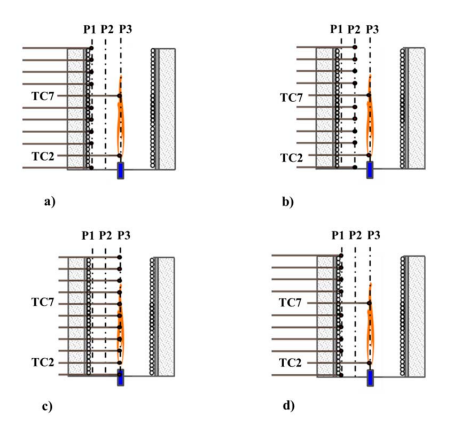


Figure 3. Sequence representing how thermocouples are moved during each test

#### 2.3. Data Post-Processing

After completing the tests, some post-processing has been done to ensure the reliability of experimental data. For example, because of the presence of radiation errors, the temperature of a flame measured by thermocouples may significantly differ from the real combustion gases temperature [47]. Radiation corrections of measured temperatures are not easy to perform; indeed, the quantification of radiation errors depends on various parameters such as the emissivity of the thermocouple, the temperature and the emissivity of the gases, the velocities of the gases in the vicinity of the thermocouple. To obtain the real burned gases temperatures, a steady-state heat transfer model, which can be applied to a bare-bead thermocouple, has been consider [49]. The following equation has been used:

$$h_{th} \cdot \left(T_g - T_{th}\right) = \varepsilon_{th} \cdot \sigma \cdot \left(T_{th}^4 - T_{w,int}^4\right) \tag{1}$$

where h<sub>th</sub> is the convective heat transfer coefficient in the vicinity of the thermocouple,  $T_g$  is the real gas temperature,  $T_{th}$  is the measured temperature,  $\varepsilon_{th}$  is the thermocouple emissivity (which is considered equal to 0.8 [47]),  $\sigma$  is the Stefan–Boltzmann constant,  $T_{w,int}$  is the temperature of the surrounding combustor wall. The above equation was chosen because it is widely used to correct radiation errors affecting combustion gases temperature measures [47, 49-52]. To calculate h<sub>th</sub>, the thermodynamic properties, the transport properties, and the velocities of the combustion gases must be known. The physical properties of the combustion gases have been determined by calculation on the basis of the composition and have been defined as a polynomial function of temperature. Burned gases composition is obtained by numerical calculations with the IdealGasReactor module of the CANTERA package [53] using as chemical reaction mechanism the one developed by Dagaut et al. [54]. The thermodynamic and transport properties of each constituent of the burned gases are taken from NIST website and from literature [55,56]. The velocities of the combustion gases in the combustor are determined with CFD software Fluent, considering a 2-D axisymmetric domain. Turbulence is modeled by using the standard k- $\varepsilon$  model, with standard wall functions near wall treatment [57,58]. For engineering purposes, this approach saves computational resources, as the viscosity affected near wall region does not need to be resolved. Grid independence has been verified.

The convective heat transfer coefficient  $h_{th}$  has been calculated using the equation proposed by Whitaker et al. [59]:

$$h_{th} = \frac{k_{g,th}}{D_{th}} \cdot \left[ 2 + \left( 0.4 \cdot 0.06 \cdot Re_{th}^{0.5} \right) Re_{th}^{0.5} + Re_{th}^{0.67} \cdot Pr^{0.4} \right]$$
(2)

where Reth and Prth are defined as follows:

$$Re_{th} = \frac{\rho_{g,th} \cdot D_{th} \cdot v_{g,th}}{\mu_{g,th}} \tag{3}$$

$$Pr_{th} = \frac{c_{p,gth} \cdot \mu_{g,th}}{k_{g,th}} \tag{4}$$

In the above equations,  $k_{g,th}$  is the gas thermal conductivity,  $D_{th}$  is the thermocouple diameter,  $\rho_{g,th}$  is the gas density,  $\mu_{g,th}$  is the gas viscosity,  $C_{p,gth}$  is the gas specific heat,  $v_{g,th}$  is gas velocity. All the properties are calculated in the proximity of the thermocouple.

## 3. Results and discussion

# **3.1. Influence on Combustor Dynamics of Fuel Mass Flow Rate: Study of a First Full Length Test**

The analysis of the influence of the operating parameters, fuel mass flow rate (indicated as  $\dot{m}_f$ ) and equivalence ratio (indicated as  $\varphi$ ), on the dynamics of the combustor represents a key challenge.

The schemes of fig. 4-a and 4-b, showing the effects of  $\dot{m}_f$  and  $\varphi$  on combustor dynamics, illustrate the complexity of this issue, due to the strong interaction between the physical and chemical phenomena that are involved. Fig. 4-a shows that  $\dot{m}_f$  increases result in a decrease of fuel residence time in the cooling channel (indicated as  $t_f$ ), leading to a loop whose effect on combustor dynamics are very difficult to predict. Similar considerations can be done on the scheme of fig. 4-b, showing that  $\varphi$  increases result in flame emissivity increases that, in turn, result in a raise of the heat flux passing from the combustion gases to the combustor wall. Hence, the temperature of the fuel flowing in the cooling channel would increase, leading the system to the same loop of fig. 4-a.

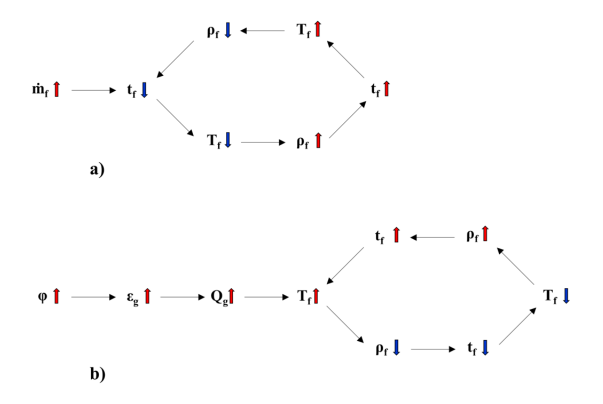


Figure 4. Schemes showing the influence of  $m_f$  and  $\varphi$  on combustion chamber dynamics

Fig. 4-a and fig. 4-b properly illustrate the difficulty of defining an engine regulation strategy permitting to assure the required thrust over the entire flight while securing a proper thermal management.

To facilitate the access of the reader to the wide data set that was collected during the experiments, a detailed analysis of one of the test cases is proposed (test case number 6 reported in table 1, having a duration of 449.2 min). A summary of the conditions under which this test has been conducted is given in table 2. The experiment is run by varying a single input parameter, i.e. fuel mass flow rate, following the experimental procedure given in section 2. As it can be seen from table 2,  $m_f$  is varied between 0.020 and 0.028 g.s<sup>-1</sup>. In order to assure a constant equivalence ratio ( $\varphi$ =1), each time ethylene mass flow rate is varied, air mass flow rate is varied as well. The rate of variation of ethylene mass flow rate is set to 4.0 kg.s<sup>-2</sup> whereas the rate of variation of air mass flow rate is set to 12.5 kg.s<sup>-2</sup>. The test is started at  $t_0$ =4.2 min and is stopped at  $t_f$ =453.4 min.

Time (min)	Change in thermocouples position	Change in fuel mass flow rate (g.s <sup>-1</sup> )	Change in air mass flow rate (g.s <sup>-1</sup> )
4.2	Start of the test sequence		
143.8	P1 to P2	-	-

147.2	P2 to P3	-	-
154.2	P3 to P1	0.020 to 0.024	0.30 to 0.36
223.2	P1 to P2	-	-
227.0	P2 to P3	-	-
235.1	P3 to P1	0.024 to 0.028	0.36 to 0.42
299.1	P1 to P2	-	-
302.0	P2 to P3	-	-
307.6	P3 to P1	0.028 to 0.024	0.42 to 0.36
374.0	P1 to P2	-	-
376.9	P2 to P3	-	-
384.5	P3 to P1	0.024 to 0.020	0.36 to 0.30
442.4	P1 to P2	-	-
445.3	P2 to P3	-	-
453.4		End of the test Sequence	

Table 2. Time events during the operation of test case number 6

The temperatures measured during test number 6 are reported, as a function of time, in fig. 5 and in fig. 6, representing respectively the burned gases temperature and the fuel-coolant temperature measured at the outlet of the cooling channel (measured by thermocouple TC<sub>hf</sub> and indicated as  $T_{hf}$ ). To avoid crowding, fig. 5 only provides the temperatures measured by thermocouples TC2, TC4, TC6, TC8, and TC10, respectively at 3, 13, 23, 33, and 43 cm from the extremity of the burner, which is placed at 3 cm from the base of the combustor.

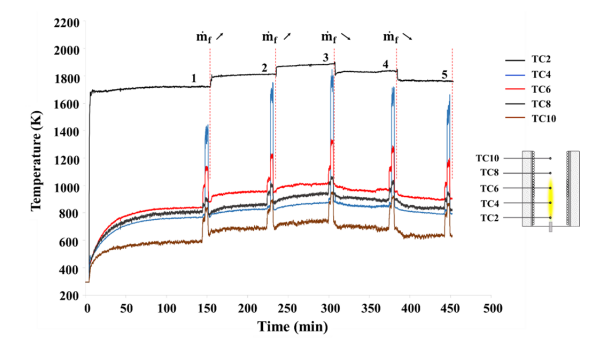


Figure 5. Variation of combustion gases temperatures with time for test 6

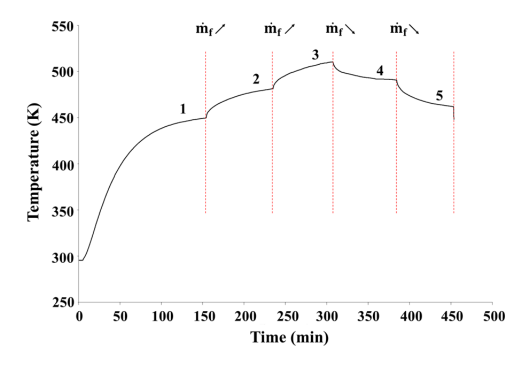


Figure 6. Variation of  $T_{hf}$  with time for test 6

Both fig. 5 and 6 indicate that a raise of  $\dot{m}_f$  leads to a raise of both burned gases and fuel-coolant temperaturs, while its drop leads to their decrease. This is consistent with literature works [13,14]. Figures 5 and 6 also show that the time needed to reach the first steady state, which is of 139.6 min, is much higher than the times required to reach the next ones, which are between 58.4 (time to reach steady state 5) and 68.8 min (time to reach steady state 2). This is a consequence of the very high volumetric heat capacity of the combustion chamber. In fact, since at the start of each test the combustor is at ambient temperature, its heating needs a very long time. Then, when operating conditions are modified, the time required for the achievement of the following steady state is much lower because the system has already accumulated a significant amount of thermal energy.

It is important to observe that the maximum temperatures reached by the fuel-coolant in the cooling channel are probably higher than those shown in fig. 6, which are measured at the outlet of the rolled-up tube from the combustor (position 2 in fig. 1). Indeed, as shown below in fig. 8 in this paragraph, due to the peculiar temperature profile characterizing the combustion chamber, in the vicinity of its inner wall the temperatures achieved by the combustion gases tend to be higher at half of the combustor height, between 20 and 25 cm from its base. Consequently, it is approximately at half of the height of the combustor that the temperature reached by the fuel-coolant in the rolled-up cooling channel achieves its maximum value. Before reaching its outlet, where its temperature is measured by means of

thermocouple  $TC_{hf}$ , the fuel passes through a colder region of the combustion chamber where it cools down because of the heat exchange between the fluid and the relatively colder combustor zones through which the rolled-up tube is passing [60].

The authors have tried to model the evolution of the fuel flowing in the cooling channel by means of numerical calculation carried out with the Plug Flow Reactor module of the CHEMKIN package. These calculations, which will be presented in a future paper, have shown that the maximum temperatures reached by the fuel-coolant are between 350 K and 450 K higher that the ones measured at its outlet. This explains why fuel decomposition has been observed despite the temperature measured by thermocouple  $TC_{hf}$  is always lower than 750 K (which is too low to get ethylene pyrolysis [61-63]).

 $T_{hf}$  values at steady state are shown in fig. 7. A hysteresis effect can be observed both by comparing steady state 1 and steady state 5 and by comparing steady state 2 and steady state 4. Indeed, fig. 7 shows that even in the presence of the same fuel mass flow rate (0.020 g.s<sup>-1</sup>), the outlet fuel-coolant temperature at steady state 5 ( $T_{hf,5}$  = 463.1 K) is higher than at steady state 1 ( $T_{hf,1}$  = 448.7 K). The same applies with respect to steady state 2 and steady state 4, being fuel mass flow rate the same (0.024 g.s<sup>-1</sup>) and the outlet fuel-coolant temperatures different (respectively  $T_{hf,2}$  = 480.0 K and  $T_{hf,4}$  = 491.4 K). This phenomenon, which results from the gap existing between the different time constants characterizing heat transfer by conduction, by convection and by radiation (which are, in the context of a fuel-cooled combustor, all involved), had already been numerically predicted [13]. Obviously, it further complicates the definition of a satisfactory engine control strategy as it makes it very difficult to develop analytical relationships between the physical and chemical parameters that must be taken into account in developing such a strategy.

 $T_{hf}$  increases and decreases that occur as a result of  $\dot{m}_f$  variations will be the object of a future work, where the most important heat fluxes characterizing the combustor (the heat flux density passing from the combustion gases to the combustor wall, the heat flux density absorbed by the fuel-coolant in the cooling channels, etc.), calculated by experimental data post-processing, will be presented.

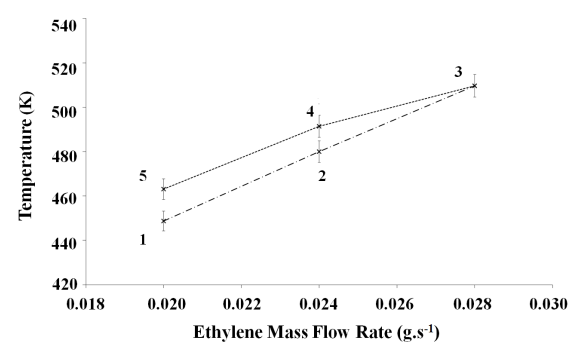


Figure 7. Variation of  $T_{hf}$  with time for test 6

In fig. 8, the combustion gases temperatures along the axis are given for steady state conditions from 1 to 5. For all the five plateaus, the higher temperature zone along the axis (position P3) is situated near the burner region (between 5 and 20 cm away from the base). As already mentioned above in this paragraph, in the zone between the axis and the wall (position P1 and P2), the temperatures tend to be higher at half of the combustor height (around 25 cm away from its base).

These findings are consistent with the results of previous experimental studies realized on a similar configuration [64]. In fact, as the burner is not equipped with any swirl, combustion gases recirculation in the downstream region of the combustion chamber is very weak; consequently, in this zone heat transfer from the flame to the gases beside it is essentially due to radiation. As these gases weekly absorb propagating radiation, their temperatures are, when compared to the combustion gases, quite low.

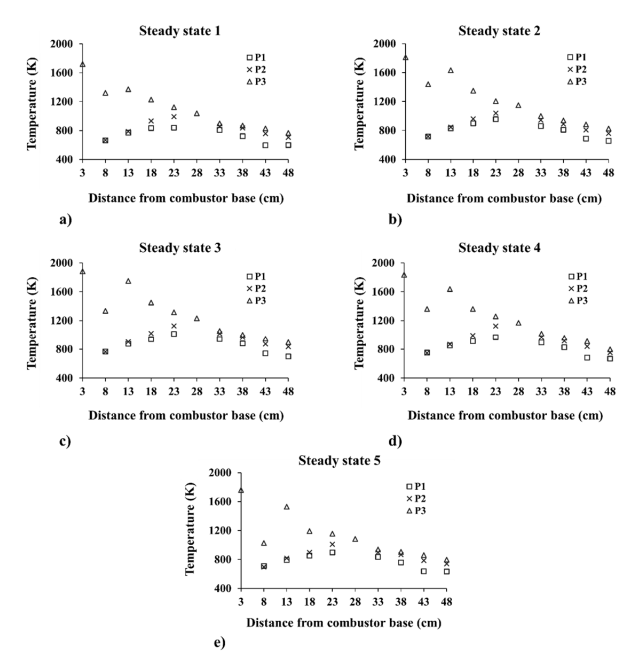


Figure 8. Flame temperature profile along the axis at steady state for test 6

Data of fig. 8 also indicates that the temperature radial variations are greater near the region of the burner. Indeed, when moving from position P3 to position P1, the average radial variation for the five plateaus is 668.9 K between 10 and 15 cm from the base whereas it is 190.2 K between 40 and 45 cm. This means that the fuel-coolant flowing in the cooling channel is mostly heated in the last part of the cooling channel, i.e. in the burner region.

One of the main parameters in the study of a fuel-cooled combustor is fuel-coolant residence time in the cooling channels. In the framework of this study,  $t_f$  has been calculated with the following equation:

$$t_f = \frac{L_{cc}}{v_f} = \frac{L_{cc} \cdot \rho_f \cdot S_{cc}}{\dot{m}_f} \tag{5}$$

In equation (5),  $L_{cc}$  and  $S_{cc}$  are respectively the length and the cross section of the cooling channel and  $\rho_f$  is the fuel-coolant density.

Fuel residence time  $t_f$  cannot be easily determined as it depends on fuel-coolant density that, in turn, depends on fuel composition and temperature, which are only known respectively at the inlet and at the outlet of the rolled-up tube. To determine fuel properties evolution along the tube, numerical calculations with ConstPressureReactor module of the CANTERA package have been performed. To this end, cooling channel pressure has been supposed to be equal to the average one between its inlet and its outlet. This allows to evaluate the variation of the fuel-coolant density and then to calculate its residence time by equation 5).

Results are given in fig. 9 as a function of fuel mass flow rate, for steady state plateaus from 1 to 5. The estimated values are between 6.75 and 7.07 s. Despite a relative uncertainty in the calculated results of about  $\pm 12\%$ , due to the assumptions made on pressure and to the difficulty of modeling the pyrolysis of ethylene, these outcomes permit to observe that  $\dot{m}_f$  variations only weekly affect  $t_f$ . This result can be explained by the fact that, as this experiment has shown,  $\dot{m}_f$  increases always result in  $\rho_f$  increases whereas  $\dot{m}_f$  decreases always result in  $\rho_f$  decreases. Consequently, as it can be understood from equation 5),  $\dot{m}_f$  variations do not influence  $t_f$ .

This outcome is very relevant. It is anyway important to note that probably it would not be valid if the fuel were a liquid as, if so, fuel mass flow rate variations would not result in fuel density changes.

In fig. 10, the rate of variation of the fuel-coolant temperature at cooling channel outlet (represented by the derivative of fuel-coolant temperature, indicated as  $dT_{hf}/dt$ ) is given as a function of time.

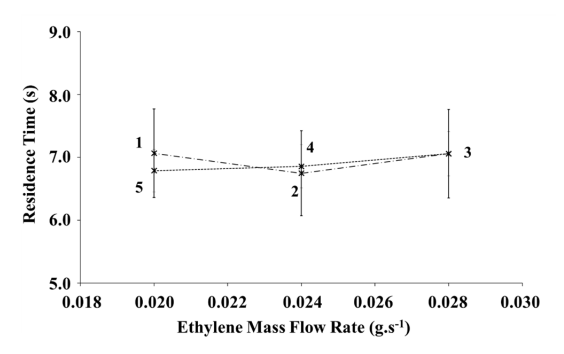


Figure 9. Fuel residence time  $t_f$  as a function of mf at steady state for test 6

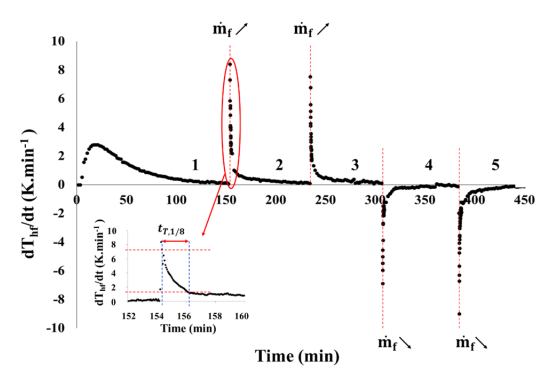


Figure 10. Variation of  $dT_{hf}/dt$  with time for test 6

During the first 10 minutes this parameter increases, reaching a maximum of 2.95 K.min<sup>-1</sup>, following the global heating of the combustion chamber. Then, it remains stable for approximately 5 minutes before starting to decrease, until the achievement of a steady state. Fig. 10 indicates that  $dT_{hf}/dt$  is strongly affected by  $\dot{m}_f$  variations. For example, when  $\dot{m}_f$  is

increased from 0.020 to 0.024 g.s<sup>-1</sup>,  $dT_{hf}/dt$  raises from 0.07 to 8.39±1.26 K.min<sup>-1</sup> whereas when  $\dot{m}_f$  is increased from 0.024 to 0.028 g.s<sup>-1</sup>, it raises from 0.08 to 7.51±1.13 K.min<sup>-1</sup>. In any case, if the absolute values are considered, it can be remarked that during the transient evolution of the system they always gradually decay, as the system tends to a steady state. It is important to stress that it is quite difficult to establish the reliability of above given  $dT_{hf}/dt$  variations as their determination relies on the acquisition of a single data point. Thus, with a higher sampling frequency, the results could be different.

The analysis of the derivative of fuel-coolant temperature  $dT_{hf}/dt$  also allows to estimate system response time to fuel mass flow rate variations (indicated as  $t_{resp}$ ). In this sense, the data of table 3 show that  $t_{resp}$  is lower than  $t_f$  (of about 40 to 50%) when fuel flow rate is increased whereas it is very close to  $t_f$  when fuel flow rate is decreased.

In fig. 11, the time derivative of fuel pressure at cooling channel inlet (indicated as  $dP_f/dt$ ), is presented as a function of time.

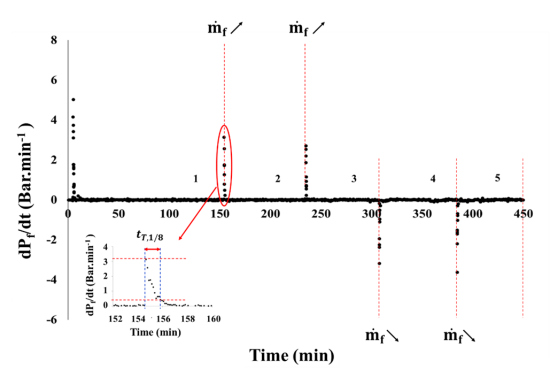


Figure 11. Variation of  $dP_f/dt$  with time for test 6

Complementarily, table 4 provides the empirical values corresponding to the times needed for  $dT_{hf}/dt$  and for  $dP_{f}/dt$  to reach one in eight of the maximum values achieved after mass flow variations (they are indicated respectively as  $t_{T,1/8}$  and  $t_{P,1/8}$ ). It is clear from table 4 that the dynamics of the hydraulic flow in the cooling channel are much faster than the thermal ones (of a factor of about 10).

Time (min)	Fuel mass flow rate (g.s <sup>-1</sup> )	$t_{resp}\left(\mathbf{s}\right)$	$t_f(\mathbf{s})$
154.2	0.020 to 0.024	3.54±1.06	6.78±0.61
235.1	0.024 to 0.028	3.48±1.04	6.85±0.62
307.6	0.028 to 0.024	6.98±1.95	7.06±0.65
384.5	0.024 to 0.020	7.02±2.11	7.07±0.61

Table 3. Comparison between system response time and fuel residence time in the cooling channel for test 6

Time (min)	Change in ethylene mass flow rate (g.s <sup>-1</sup> )	<i>t</i> <sub>7,1/8</sub> (s)	<i>t<sub>P,1/8</sub></i> (s)
154.2	0.020 to 0.024	133.8	30.0
235.1	0.024 to 0.028	259.8	28.8
307.6	0.028 to 0.024	239.4	33.0
384.5	0.024 to 0.020	231.6	30.0

Table 4. Times  $dT_{hf}/dt$  and  $dP_{f}/dt$  need to reach 1/8 of the maximum values after  $\dot{m}_{f}$  variations

#### 3.2. Influence of Equivalence Ratio

Due to the strong effect on fuel-coolant heating of the combustion gases temperature and emissivity, the influence of equivalence ratio  $\varphi$ , which can affect both, needs also to be investigated. To this end, the results of tests number 2, number 3, and number 4 are compared. Again, these tests are carried out by varying a single input parameter, i.e.  $\dot{m}_f$ , between 0.020 and 0.028 g.s<sup>-1</sup>.

To study the influence of  $\varphi$ , three different equivalence ratio values are chosen, respectively 1.0 for test number 2, 1.25 for test number 3, and 1.5 for test number 4. The rates of variation of ethylene and air mass flow rate are the same as in experiment 6. With regard to case 4 ( $\varphi$ =1.5), data are only available for steady state plateaus from 1 to 3. Indeed, during this experiment, the decomposition of the fuel has originated coke deposits that have caused the occlusion of the cooling channel and the interruption of the test before its end. Hence, plateaus 4 and 5 are not available.

The values reached by  $T_{hf}$  at steady state by the fuel-coolant at the outlet of the cooling channel for these three experiments are given in fig. 12.

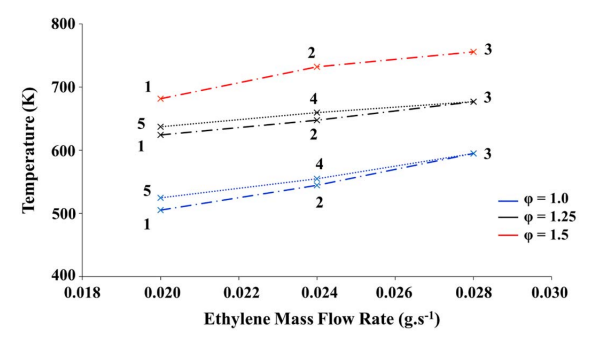


Figure 12. T<sub>hf</sub> at steady state for tests 2, 3, and 4

It can be seen that they linearly increase with  $\varphi$ . This cannot be justified on the basis of fuel residence time variations, as it can be seen from table 5, which gives for the three test cases the average values of this parameter along the test. The data of table 5 indicate that  $t_f$  higher values are reached in the case of a stoichiometric combustion, when  $T_{hf}$  values are the lower ones. Indeed  $T_{hf}$  increases with  $\varphi$  can rather be explained by considering that even a small increase in this parameter results in a substantial increase in the heat flux passing from the burned gases to the combustor wall, due to the formation of soot which determines the growth of flame emissivity and of the radiative heat flux [65-67]. These outcomes will be widely discussed in a future paper.

Test case	φ	$t_f(\mathbf{s})$
2	1	$3.78\pm0.41$
3	1.25	$3.12\pm0.38$
4	1.5	$3.16\pm0.37$

Table 5. Average residence times at steady state for tests 2, 3, and 4

Fig. 13, which illustrates, respectively for case 2, 3, and 4, some pictures of the flame, confirms that an increase in the fuel to oxidizer equivalence ratio results in an increase in flame luminosity. It shows that the flames of case 2 and case 3, due to incandescence of

the soot generated under fuel-rich conditions, are yellow, while the flame of case 4 is blue as no radiating soot is produced [67].

In table 6,  $dT_{hf}/dt$  variations occurring after  $\dot{m}_f$  variations (which are referred to respectively as  $dT_{hf}/dt_1$ ,  $dT_{hf}/dt_2$ ,  $dT_{hf}/dt_3$  and  $dT_{hf}/dt_4$ ) are given. The data of this table indicate that all these derivatives strongly increase with the increasing of  $\varphi$ .

These outcome is of major importance as they show that test bench dynamics are strongly influenced by the ratio between fuel and oxidizer mass flow rates. Under fuel-rich conditions, when, as it is discussed in the second part of the paper, heat transfer by radiation becomes predominant compared to convection heat transfer, system response in terms of fuelcoolant temperature variation is more accentuated.

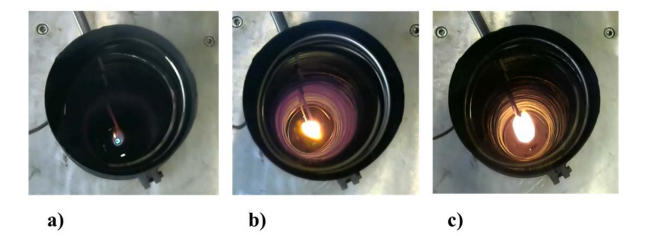


Figure 13. Pictures of the flame for  $\varphi=1.0$  (a),  $\varphi=1.25$  (b), and  $\varphi=1.5$  (c)

Differently from the fuel heating rate, no evidence of an influence of  $\varphi$  on the rate of variation of fuel pressure  $dP_f/dt$  is observed. This shows that the hydraulic part is mostly influenced by the coolant flow rate, at least more than by the equivalence ratio.

Test case	φ	$\frac{dT/dt _{max}}{(\text{K.s}^{-1})}$	$\frac{dT/dt _1}{(\mathbf{K}.\mathbf{s}^{-1})}$	$dT/dt _2$ (K.s <sup>-1</sup> )	$\frac{dT/dt}{3}$ (K.s <sup>-1</sup> )	$\frac{dT/dt}{(\mathbf{K.s}^{-1})}$
2	1.0	2.78	$13.69\pm2.05$	$10.65\pm1.60$	$-10.68 \pm 1.60$	$-12.12 \pm 1.82$
3	1.25	6.86	$30.85\pm4.63$	$27.17\pm4.08$	$-22.12 \pm 3.32$	$-24.60 \pm 3.69$
4	1.5	9.60	$33.12\pm4.97$	$36.66 \pm 5.50$	-	-

Table 6. Fuel heating rate variations following fuel flow rate variations for tests 2, 3, and 4

# 4. Conclusions

Regenerative cooling by the on-board hydrocarbon fuel is a widely applied cooling technique suitable to secure the thermal protection of supersonic combustion air-breathing vehicles flying above Mach 5. Due to the complexity of the phenomena which are involved (heat transfers, phase transitions, chemical reactions), this technology needs to be investigated by an experimental approach permitting to validate the numerically acquired knowledge and to understand its dynamics. Hence, a regeneratively cooled combustor has been designed. Experiments are performed under both stationary and transient conditions using ethylene as fuel and air as oxidizer. The effect of two major parameters, i.e. the fuel mass flow rate and the equivalence ratio, on flame temperature field, coolant pressure and coolant temperature has been determined. It has been shown that a higher fuel mass flow rate results in an increase in flame temperatures, coolant temperature and coolant pressure. This clearly demonstrates that the management of engine cooling by the fuel is a real challenge because it is absolutely counter-intuitive in classical heat exchangers to observe that coolant temperature increases with coolant flow rate. It has also been demonstrated that equivalence ratio increases lead to a raise of coolant temperature, due to a raise of the radiative heat flux passing from the burned gases to the combustor wall. This point is of particular interest for engineering design because it illustrates the need to control the burning process and its homogeneity in order to avoid local fuel-rich conditions. A hysteresis effect due to heat transfer dynamics has been found; this is in good agreement with previous numerical predictions. The characteristic times of the commands (few seconds to change fuel mass flow rate and equivalence ratio) and of system response are far smaller than the characteristic time of thermal stabilization in this channel (in the range from about 140 to 240 s corresponding to a renewal of the channel of approximately 50-70 times). It has also been observed that the raise of fuel flow rate has a greater quantitative impact on the coolant temperature (which rises) in case of fuel-rich conditions compared to stoichiometric ones. In a future work, the effects of fuel mass flow rate and fuel to oxidizer equivalence ratio on the heat flux density passing from the combustion gases to the combustor wall, the sensible heat flux absorbed by the fuel-coolant, the heat loss due to the exhaust gases and the heat loss due to heat transfer from the combustor external surface to the surrounding environment will be analyzed. The heat transfer efficiency of the combustor will be defined and its dependence upon fuel mass flow rate and fuel to oxidizer equivalence ratio will be presented.

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