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Hydrocarbon pyrolysis with a methane focus: a review on the catalytic effect and the coke production

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ABSTRACT

Hydrocarbon pyrolysis has been widely studied since the 1900’s for applications in aerospace as a fuel and/or coolant or for use with fuel cells and hydrogen production with a catalyst. In this context, the role of heterogeneous reactions with homogeneous phase chemistry is unclear despite the fact that it is obviously at the heart of coupled physico-chemical phenomena. In addition, the thermal formation of solid carbon particles -coke, which can be deposited on the structure, impacts the heterogeneous reactions. The aim of this work is to review the available literature on hydrocarbon pyrolysis involving reactions with solid surfaces and coke particles. The influential parameters such as the nature of the fluid, the temperature (up to 2000 K), the pressure (up to 100 bars), the residence time (µs order to min order), the reactor type (plug flow, batch, perfectly stirred reactor) and the type of catalyst (inert, metallic or more complex such as zeolites) are discussed. Then, a link between catalicity and coke production is addressed. This literature survey focuses in particular on methane because of the growing interest regarding the potential for hypersonic applications.
1. Introduction

Within the scope of hydrocarbon pyrolysis, various parameters and conditions have been studied, among which: the nature of the reactive fluid (methane [1], propane [2] and other even more complex fluids, such as JP-7 [3]), temperature (from 400 K [4] to more than 2500 K [5]), pressure (from below ambient pressure [6] up to 20 MPa and even more [7]), type of reactor (shock tube [8], flow reactor [9],...
solar tube [10], diluted [11] or pure [12] conditions. These experiments and the subsequent results have enabled simple to complex kinetic mechanisms to be constructed in order to describe the chemistry in detail and to perform numerical simulations in a multi-physics configuration. The numerous related works have progressively constituted an accumulation of results which are now diluted in the open literature. Only a few reviews are available and those for methane can be cited [13–15].

The surface catalytic effect, which is inherent to the reactor, is an additional complex phenomenon to be considered during hydrocarbon pyrolysis. A number of parameters (the nature of the catalyst, of the pyrolyzed fluid, several products, the ambient conditions, the ratio surface/volume -S/V, etc.) play a role and have been initially explored experimentally. From a numerical point of view, proposing kinetic schemes to describe heterogeneous reactions causes a rapid increase in the complexity of the models. Because heterogeneous reactions may be of importance, this justifies details and complex mechanisms being taken into account. Nevertheless, studies on the kinetics of reactions inherent in a large surface (i.e. the inner wall of the reactor) are quite rare. Rather than coating the inner surface of the reactor to study such catalytic effects, an alternative is to place a solid material, a catalyst, inside an inert reactor. Although the chemistry may be quite different, notably regarding the S/V, as suggested by Gordon [16], such data represent a precious source of information regarding heterogeneous chemistry.

In parallel, another critical phenomenon is the generation of solid particles. Pyrolysis leads to the production of heavy aromatic compounds which can agglomerate to form a particular type of solid carbon which can both dissolve in the fluid and condense on the reactor wall. Such particles can be classified according to several criteria. As an example, some studies distinguish coke according to its origin: pyrolytic [17], catalytic [18] or also asphaltic [19]. The production of such solid carbon depends on numerous parameters (temperature, pressure, residence, time, surface effect, oxygen content) and its formation has been widely studied in the literature, e.g. [20–22]. The particularity of solid carbon is that it can be deposited on the surface as a thin layer, and, in so doing, can, among other things, deactivate the catalytic sites. Consequently, several studies have been conducted in order to avoid such deposits thereby durably sustaining the activity of the catalysts (e.g. [23–25]).
These two phenomena (catalicity and coke production) have generated a large number of data disseminated over the years in the literature. The present paper aims at reviewing the articles related to the catalytic effect and coke production during the thermal decomposition of hydrocarbons (mostly small ones and notably methane). These two issues are linked in several ways: firstly, the catalytic activity can, among other things, modify the production of coke particles (in both ways: enhancement and limitation). Secondly, coke can also have an impact on the catalicity of a material. Indeed, it can stick to the surface of a material and in so doing decrease or even inhibit its effect. The context of this survey is that of regenerative cooling during hypersonic flights, i.e. the endothermic degradation of the fuel which acts as a heat sink. Catalicity and coke production can be critical with respect to the cooling method. Indeed, heterogeneous reactions could occur between the solid material and the fuel. The interactions with the solid materials may either increase or decrease the fuel degradation and this has both direct and indirect consequences on the internal convective cooling of the permeable solid material by the fuel. The direct relationship is due to the pyrolysis rate. When the rate increases, the endothermic effect is higher and the cooling efficiency is thus greater. The indirect relationship is that of the intrinsic formation of coke (coke particles stick inside the pores of the solid material) which is higher for higher pyrolysis rates. This coke layer decreases the heat transfer between the solid and the fluid. As a consequence, cooling efficiency is lower in the case of coke formation. One can also notice that the permeation process (fluid flow) is modified by the coke particles in case of clogging [26,27].

A wide review has been proposed of former works specifically concerning methane pyrolysis [15]. This is expected to cover the field of catalicity and coking phenomenon to provide a complete situation of hydrocarbon pyrolysis, particularly as applied to methane which is, as suggested by Davis et al., a possible efficient fuel (advantageous standard enthalpy of combustion, low price, relatively low coking rate) [28]. This study is mainly devoted to methane. However, it has been widened to include several other hydrocarbons in order to provide complementary information when necessary.
2. Catalytic effect involved in hydrocarbon pyrolysis

Even in the earliest methane pyrolysis investigations, it was observed that several parameters can modify both the kinetic and product ratio. Hurd and Pilgrim [29] pointed out that the species produced during the first instants of decomposition, the contact time, its relationship with the temperature and then, the surface reactor are all parameters to be taken into consideration. Slater [30] worked on the last point and clearly showed that each material has a specific effect on the decomposition rate. Therefore, in most cases, the surface effect was neither considered nor investigated but merely neglected. The catalyst effect may be considered in different ways, namely: the reactor material (metallic, composite), the treatment of the inner wall (e.g. coating) or by a small solid sample insertion. The following subsections investigate these techniques both in order to differentiate the results and to determine the consequences.

2.1 Catalytic effect considering a massive reactor

2.1.1 Massive reactor: the intrinsic effect of the materials

Studies on the catalytic effect of the inner wall of a reactor are relatively uncommon and were essentially conducted in the 1970's. Before this, few researchers worked on this phenomenon. Thus, Hurd and Pilgrim [29] concluded that the nature of the reactor alters hydrocarbon pyrolysis. They investigated the thermal degradation of butane in a reactor made of iron, nickel or monel (Ni-Cu-Fe alloy) and they noted significant differences with high catalicity for a monel reactor. In comparison, Hurd and Eilers also explored olefin pyrolysis in different metallic reactors and confirmed the high activity of the monel reactor [31]. Regarding iron or nickel, less catalicity was found (decomposition at a lower temperature for the first and modification of the product distribution for the second).

Several researchers explored other materials considering different small hydrocarbons. In [32], in propane, ethylene and propylene pyrolysis Crynes and Albright observed little activity for nickel and stainless steel reactors while a non-negligible catalicity for low carbon steel reactors was demonstrated. They also performed tests inside mixed reactors (respectively 30 % and 70 % length of low-carbon steel...
and stainless steel) and noticed that a higher activity of low-carbon steel could be observed only if this material was spatially placed after the stainless steel part. Given such results, the authors suggested the role of the surface in the initiation and termination for free radical chain reactions. They also suggested that tubes of small size (i.e. with a high S/V ratio) present enhanced catalytic activity and inherent surface reactions compared with bigger tubes but this is counter-balanced by the nature of the fuel which is considered during the experiment (ethane or propane). Ghaly and Crynes further explored propylene pyrolysis considering more materials [33]. More details are given and notably the nature of the catalicity of each material. There are certain contradictions compared with the previous results. Thus, for the stainless steel reactor, the wall activity is expressed via a higher production of carbon. The low-carbon steel and nickel reactors had a fairly similar profile and demonstrated an increase of activity during the first stage of pyrolysis (coke and hydrogen yield increased) and then it fell and reached a steady-state after one hour. Such a trend suggested a progressive deactivation of the surface by the of solid carbon. Inconel and incoloy reactors showed less activity. Ghaly and Crynes concluded that the wall activity was essentially due to the presence of iron which acts as a catalyst for carbon and hydrogen production. However, the presence of chromium in some alloys inhibits its activity. Therefore, a surface containing an active carbon layer which carries iron atoms (e.g. low-carbon steel) may show higher activity. Concerning nickel, it is assumed that the active sites have a lower affinity with hydrocarbons than iron. Finally, alloys containing both iron and nickel may consequently be less effective than pure metal. Dunkleman and Albright worked on paraffin (ethane and propane) pyrolysis and compared metallic and Vycor glass reactors [2,34]. Significant differences were found between metallic and Vycor reactors but it clearly appeared that the surface reactions had less importance for propane pyrolysis in comparison with ethane experiments. Globally, they noticed that the hydrogen amount and coke formation were lower for Vycor reactors than for metallic ones. In the 2000’s, the works of the PRISME laboratory on the catalytic activity of massive reactors constituted recent and quite unique data [35–37]. For example, in [37], dodecane pyrolysis was investigated in two different tubular reactors (stainless steel and titanium). Greater pyrolysis activity was noted for the titanium reactor at the same temperature.
than for a stainless steel one. Indeed the yields of the major products (ethane, ethylene and propane) were multiplied by a factor of 2 while the conversion rate increased from 40 % to 60 %. Nevertheless, it was strange to note that a higher coke deposit was found for the stainless steel tube.

These first results demonstrated the mix effect of the catalicity of a reactor. For the same material with the same fluid, substantial differences could be found. Additionally, the catalicity does not only consist in reducing the temperature of hydrocarbon pyrolysis but can also clearly influence certain reaction pathways. Parameters such as time, type of reactor or S/V ratio are of the utmost importance with regard to the catalytic phenomenon.

2.1.2 Treatment of the inner surface

The treatment of the reactor surface also constituted part of the research investigations on catalicity. In [32], in several cases the reactor was pre-treated (oxygen, hydrogen sulfide, steam, sulfur, anhydrous hydrogen bromide and nitric oxide). Oxygen treatment appeared to drastically affect the product distribution and to be an efficient catalyst for ethylene, propylene and carbonaceous solids (Figure 1).

![Figure 1. Effect of oxygen pre-treatment. From Crynes and Albright [32]](image)

With regard to hydrogen treatment, it limited the activity while steam had a moderate influence (the conversion increased from 32 % to 35 %). Treatment with hydrogen sulfide inhibited the catalytic
effect and was similar to a passivation (subsequent oxygen treatment is ineffective). Other treatments did not significantly affect the hydrocarbon conversion. Ghaly and Crynes explored propylene degradation and confirmed the higher activity for an oxygen pretreated stainless steel reactor but they also noticed that such pretreatment moderated the activity of a low-carbon steel wall (oxygen acts a protective oxide film) [33]. For nickel, this treatment presented no effect for the first 10 minutes before undergoing a great increase due to the partial reduction of the oxide surface layer (by the action of the carbon and hydrogen formed) which led to the formation of a more active nickel. For inconel and incoloy, the results were close to those of oxygen-treated stainless steel. Finally, they found quite different results regarding hydrogen sulfide pretreatment. Thus, a mixed effect was noted for stainless steel (inhibition and then promotion) while a similar effect compared with oxygen was observed for low-carbon steel (passivation). Dunkleman et al. confirmed the passivation effect of hydrogen treatment on the degradation of small paraffin [2,34]. They explained such an effect by the reduction of the surface reactions.

The pretreatment of a reactor has a clear impact on catalicity but such effects can clearly differ depending on the materials treated and the treatment itself.

2.1.3 Coated reactor, the example of zeolites

Catalicity should be also studied by considering coated reactors. In this domain, several investigations have been conducted on particular materials such as zeolites or certain of its derivatives [38–41] but generally on heavy hydrocarbons. Zeolites have the great advantage of allowing fuel thermal degradation at lower temperatures whilst reducing the production of coke. The works of Li. et al. consisted in coating the inner wall of a stainless steel tube by a mixture of different zeolites (HZM-5) and a ceramic-like binder [38]. The pyrolysis of two jet fuels including dodecane was performed under supercritical conditions and within a continuous flow. They demonstrated that the heat sink capacity and conversion were enhanced by the zeolite coating with an increase of about 25 % for the heat sink capacity while dodecane conversion rose from 66 % to 80 %. Zhao et al. [39] conducted a similar study
with Palladium inside the HZM-5 coating. Pd/HZSM-5 coating presented a better performance than HZM-5 with a higher rate of conversion (increase of about 10%) and a promotion of hydrogen yield (about twice as much). The heat sink was also enhanced, even more with the rise of the temperature. Meng et al. contributed to the zeolite coating studies by performing supercritical pyrolysis of n-dodecane inside an HZSM-5 coated stainless steel reactor [40]. The authors observed that the conversion of dodecane was higher when the reactor was coated but only up to a certain temperature. Thus, for temperatures of over 848 K, conversion becomes equivalent and even lower than for a nude reactor. This fact is explained by the progressive deactivation of catalytic sites by the filamentous coke. Liu and coworkers further investigated HZMS-5 zeolites under supercritical conditions [42]. They directed their study on the characterization of zeolites and highlighted the relative importance of the particle size. They found that coating a tube with nanoscale zeolites presented certain advantages and notably that of the enhancement of the catalytic activity (more than 1 time). Nevertheless, adhesion on the wall is poorer, but mixing nano- and microscale particles permits this problem to be overcome.

The catalicity of zeolite coated reactors is clearly demonstrated and shows interesting abilities namely regarding the reduction of the production of coke. Nevertheless, it must be mentioned that such catalysts are employed only at mid-range temperatures. Consequently, their application in the hottest parts of the cooling channel for jet fuel applications is not currently feasible since zeolites are not active in these conditions.

### 2.2 Effect of small solid catalysts in a pyrolysis reactor

Differentiation of the massive reactor experiments with the catalyst sample is namely motivated by the high importance of S/V ratio or also the contact time with the catalyst. Hence, Gordon [16] was one of the first to experiment its impact on methane pyrolysis. He used a porcelain reactor whose catalytic activity is assumed to be very limited. However, it was demonstrated that this parameter clearly enhances methane decomposition. The pyrolysis rate was even multiplied by a factor of two to three (for an S/V ratio multiplied by 3.5). Later, other research confirmed that this ratio was more or less
critical for conversion and product distribution but that its importance was directly linked to the reactor material [2,32–34].

2.2.1 Unsupported metallic materials

Chronologically, metallic catalysts were investigated first. These catalysts were inserted in reactor considered to be inert (e.g. quartz). The term “unsupported” refers to this inactivity. Fang and Yeh [43] considered 24 metal oxide catalysts deposited on a silica gel. Methane pyrolysis was performed at 1400 K and at 2.10⁻³ MPa in a quartz reactor. The effectiveness of the metals was quantified with regard to C₂ selectivity and decomposition activity. They found that catalysts generally increase methane decomposition but some of them can also decrease it with respect to quartz (e.g. Bi₂O₃/SiO₂). Based on their observations, they suggested a primary mechanism including the catalyst interaction. Van Der Zwet et al. conducted a similar study in the presence of different aluminum oxides [44]. The effect of the S/V was confirmed and demonstrated a clear impact on product distribution. The increase in the S/V leads to a progressive decrease in all the species excluding coke and tar (Figure 2).

![Figure 2. Selectivity to C₂H₄, C₂H₂, light aromatics and coke/tar as a function of surface to volume ratio at 25 % conversion and 1398 K (reactor volume = 10⁻⁶ m³) [44].](image)

However, the conversion of methane was poorly affected by increasing the catalytic surface. To explain such a fact, the authors assumed that the radical or dislocation sites of the surface may promote
hydrogen abstraction. Moreover, due to the phenomenon of autocatalysis, it may have also an influence on termination (by capturing from the gas phase the species involved in the autocatalysis). Finally, they suggested that a large catalytic surface (aluminum oxide) is favorable to the production of coke and hydrogen while gaseous and liquid products are preferentially produced in a reactor with a low S/V ratio. Wolf et al. [45] explored the effect of a platinum catalyst and modeled it over the temperature range of 300 K-700 K. Starting with the experimental results of Belgued et al. [46], they created a mechanism composed of 39 elementary surface reactions involving 14 surface species. The kinetic model accounts for the thin monolayer of carbonaceous overlayer which quickly covers the surface of the catalyst. It includes successive steps from the reactant adsorption to the product desorption. Wolf et al. pointed out that the dominance of a reaction on others depends on the contact time. They presented Ethylidyne (CCH₃) as the determinant species of the mechanism, due to its surface decomposition to C and CH₃ and its influence on ethane production.

Zeolites have been also explored as small solid catalysts [e.g. 46–48]. Thus, Xian et al. worked on the decomposition of dodecane over HZSM-5 under subcritical and supercritical conditions [48]. They notably found that the activity of the catalyst decreases with the pressure increase and consequently leads to a lower conversion. Such a trend opposes the other results found in the literature (e.g. Dardas and coworkers [49]) and demonstrates, once again, the importance of the reactant and the nature/properties of the catalyst.

2.2.2 Unsupported carbonaceous materials

With regard to non-metallic catalysts, several studies have been performed using a carbon based catalyst. Hence, Muradov used different types of carbon [50,51]. He determined that graphite and carbon black showed the least activity whereas the initial methane decomposition rate was the highest in the presence of glassy carbon. Lee and coworkers investigated other carbon catalysts (coconut shell and coal) at 850 K [52]. Regarding the nature of the catalyst, they did not observe significant difference in their activity. However, they noted that such a catalyst has good activity at the beginning of the process.
but is rapidly deactivated by the deposition of coke which blocks the pores. Pinilla et al. [53] tested two activated carbons (carbon black) and found a similar trend i.e. high activity but limited as the process progresses but they also noticed that the carbon produced by the degradation of methane can also show a catalytic activity which slightly compensates for the deactivation. Bai et al. also explored the activity of commercial activated carbon and compared it with alumina [54]. They observed less (due to the higher activation energy) and different (autocatalytic reaction of carbon production) catalytic activity for the metallic catalyst. They concluded that the methane conversion could be simplified by a two step scheme with the formation of carbon nuclei followed by growth in carbon crystallites. Moliner and coworkers further investigated activated carbon and brought supplementary information about the main characteristics which influence its activity [55]. They concluded that three elements have to be considered: the surface chemistry, the size and the distribution of the pores, and the presence of oxygenated compounds.

2.2.3 Supported materials

It has been demonstrated that not only does the catalyst have an impact on chemistry but the support (i.e. the reactor) can also have a catalytic activity which leads to heterogeneous catalicity. The term “supported” refers to this phenomenon. Ferreira-Aparicio et al. specifically studied the interactions between CH₄ and the surface of several catalysts [56]. They exposed the effect of different metallic catalysts (cobalt, nickel, ruthenium, rhodium, iridium and platinum) supported on alumina or silica, thereby demonstrating the importance of the support. Indeed, more hydrogen is produced by the catalysts supported on alumina than those on silica. Ruthenium and iridium were the least efficient catalysts for retaining carbon on their surface while the presence of rhodium on alumina support enhances the production of a more reactive coke (amorphous). Takenaka et al. further investigated the support effect on the nickel activity and on its catalytic lifetime [57]. It was observed that SiO₂, TiO₂ and graphite supports enhanced methane conversion. Al₂O₃ and MgO had the shortest lifetime. In parallel, they studied the characteristics of coke according to the support and they pointed out certain
differences namely regarding the size and the thickness of the carbon filaments. However, Zein et al. [58] found quite different results. They investigated the catalytic decomposition of methane and the subsequent production of H\textsubscript{2} and of carbon. Nickel was chosen as the catalyst and TiO\textsubscript{2}, Al\textsubscript{2}O\textsubscript{3}, MgO and SiO\textsubscript{2} were selected as the support. The catalysis was performed at atmospheric pressure and for a temperature range of 823 K-1173 K. Methane was mixed with argon (1:1 mole) and injected in a stainless steel reactor where 1 g of the catalyst was placed in its center. They concluded that titanium oxides were the best support (activity maintained after 2 hours) followed by MgO. The SiO\textsubscript{2} and moreover, the Al\textsubscript{2}O\textsubscript{3} supports had the shortest lifetime. Finally, they proposed a mechanism similar to the preceding one with regard to carbon and H\textsubscript{2} production:

\begin{align}
\text{CH}_4 + S &\leftrightarrow \text{CH}_4 - S \quad (1) \\
\text{CH}_4 - S + S &\leftrightarrow \text{CH}_3 - S + H - S \quad (2) \\
\text{CH}_3 - S + S &\leftrightarrow \text{CH}_2 - S + H - S \quad (3) \\
\text{CH}_2 - S + S &\leftrightarrow \text{CH} - S + H - S \quad (4) \\
\text{CH} - S + S &\leftrightarrow C - S + H - S \quad (5) \\
C - S + S &\leftrightarrow C + S \quad (6) \\
2H - S + S &\leftrightarrow H_2 + 2S \quad (7)
\end{align}

Zadeh and Smith studied the pyrolysis of methane in presence of supported cobalt catalysts (Co-SiO\textsubscript{2}, K-Co-SiO\textsubscript{2}, Co-Al\textsubscript{2}O\textsubscript{3} and K-Co-Al\textsubscript{2}O\textsubscript{3}) [59]. A mixture of CH\textsubscript{4} and Ar (respectively 95 % and 5 %) was injected in quartz fixed-bed microreactor heated at 723 K. They observed the so-called Metal Support Interaction (MSI) which consists in the migration of the SiO\textsubscript{2} support onto the cobalt catalyst. The MSI increases the methane decomposition activity and the facilitation of the migration of carbonaceous species from the catalyst surface to the support. This phenomenon was promoted by the decrease of the initial Co loading. The temperature increase promoted the CH\textsubscript{4} conversion and the metal coverage. As was mentioned in the previous part, the support also has an effect: conversion is greater for an alumina support than for a silica support. Finally, the addition of potassium was noticed to promote the species migration from the metal to the support.
Numerous studies have also been conducted on zeolite supported catalysts. Several combinations with different metals have been tested. Xu and Lin have produced a very complete and interesting review on this subject [60]. Globally, it appears that the molybdenum supported on HZSM-5 zeolite presents the best association. Hence, further investigations have been conducted on this combination and by modifying the catalyst preparation. Hence, for example, the works of Solymosi et al. who explored methane degradation with unsupported and supported (notably zeolites ZSM-5) molybdenum compounds can be cited [61–63]. According to the nature of the catalyst, its production method, and the nature of the support, substantial differences were observed regarding the pyrolyzed products. The authors also confirmed the progressive deactivation of the catalyst due to the carbon deposition.

### 2.2.4 Catalicity used for specific purposes

Numerous works on the decomposition of methane with metallic catalysts have been conducted over the past years, but with specific goals. The production of CO$_2$-free hydrogen is possibly the most important one [64,65]. The works of Shah et al. could be cited as an example [66]. They pyrolyzed methane and other different pure and binary metallic catalysts (Fe, Pd, Mo, Ni) supported on alumina at a temperature ranging from 673 K to 1173 K. They pointed out the importance of the pretreatment of such materials: for example, methane is decomposed at a temperature of above 1123 K for a non-treated iron catalyst while the degradation begins 200 degrees lower after oxidization and even lower (723 K) if the catalyst has been previously reduced in a flow of hydrogen and then carburized. They also demonstrated that a binary catalyst (Fe-Mo, Fe-Pd, or Fe-Ni) always possesses higher activity (i.e. production of hydrogen) than a pure one. The production of carbon (e.g. filament) is also a widely studied topic. By way of example, González et al. explored the formation of carbon nanotubes using unsupported nanoparticles of nickel as a catalyst [67]. In such studies, the temperature is quite low and the aim is to obtain a fairly low conversion of methane coupled with a moderate carbon growth dynamic in order not to deactivate the catalyst. They tested the adjunction of copper in order to enhance the activity and concluded that such a procedure was inefficient contrary to other experiments with
supported nickel catalysts. The third objective is the production of higher (i.e. heavier) hydrocarbons like aromatics.

Thus, Holmen proposes an interesting overview of the research conducted in this domain [65]. He points out that the zeolites associated with metals (preferentially molybdenum) are good catalysts and initially enable acetylene and then aromatic compounds (notably benzene) to be formed. All in all, if the interest of such works is undeniable, their purpose is quite far from the present one. For this reason, we do not further review the numerous publications available in the open literature. A very valuable literature survey of Amin et al. [68] is helpful to get a better view on the works which are devoted to the three specific objectives mentioned above. The works of Ahmed and coworkers or Abbas and Wan Daud also represent an abundant source of information in this domain [69,70].

3. Investigation on the coke deposits

During the final steps of the pyrolysis of hydrocarbons, solid particles with high carbon content are produced. The so-called “coke” can be deposited on the inner wall of the reactor and it can be found under various natures; Albright et al. described seven different types of coke [71]. Thus, the term “coke” reassembles several different particles, which could lead to some confusion. Consequently, in this paper the definition proposed by Fitzer et al. [72] shall be employed. Coke is defined as a solid with a high content of the carbon element which is structurally in the non-graphitic state. Hereafter, the terms “coke” and “solid carbon” should be considered as equivalent.

Considering the hypersonic application considered in this work, knowledge of coke production is required since its deposit on the surface of the reactor wall has a critical impact on the decomposition of the fuel by changing the thermal equilibrium, modifying the surface catalytic effect and limiting the permeation and the associated cooling of the structure. Particularly, high pressure conditions (and even supercritical states) should be considered when dealing with coke formation. Unfortunately, no work dealing with coke production during supercritical methane pyrolysis was found. It can be noticed that
other supercritical fluids have been explored in the open literature and the reader could refer to the following reference for further information [21,73–75].

3.1 Characterization of the coke particles and parameters of influence

Determining the coke characteristics and the parameters which enhance or inhibit its production is important in overcoming the drawback of fuel pyrolysis when used aboard aerospace vehicles. According to the literature, several parameters of importance have a clear influence on coke production. Among them, can be cited: the residence time, the nature of the fluid, the pressure or also the geometry of the reactor [76–83].

Blekkan et al. [76] studied tar and coke formed during methane pyrolysis in a tubular reactor. First, they highlighted the critical role of acetylene, benzene and PAH formation. They distinguished two types of solid carbon particles: tars containing a high proportion of heavy PAH (e.g. pyrene), come from liquids and stick to the reactor wall, and soot, formed in the gas phase from reactive species like acetylene. They pointed out a first parameter of influence on coke production which is the residence time. It was observed that coke selectivity reaches about 80% for a residence time of 1 second while it is only 20% for 0.5 s. Guéret et al. confirmed the role of residence time and added more details. But contrary to Blekkan et al., they distinguish three types of coke and also work on the temperature influence [77]. Globally, they classified the solid carbon deposit as follows: graphitic type which is produced whatever the residence time (it seems to act as an initial layer) and is predominant for temperatures of over 1423 K; amorphous (carbon black) whose production increases with the residence time and is maximal between 873 K and 1423 K; and soot which is produced for short residence times (lower than 0.6 s) and at low temperatures (573 K – 873 K). Albright and Marek confirmed the critical importance of the residence time (considering longer periods than the previous one), but also added two other parameters which are the geometrical features of the surface and the time of operation [78]. They pyrolyzed acetylene and toluene in a Vycor glass tubing reactor where two rectangular specimens were placed. The amount of solid carbon and the nature of the coke were different depending on the residence
time, the length of the pyrolysis run, the position of the specimen and its nature. Regarding the coking rate, two maxima could be reached for different residence times (3 s and then 9 s). They assumed that this signals the existence of two different mechanisms as a function of the production of certain precursors. The morphology of the coke significantly varies with the residence time. Thus, spherical coke particles whose size increases with the residence time and filamentous carbon which is produced preferably for a second order residence time were observed. Geometrical factors, such as the way the specimen is inserted in the reactor or the S/V ratio, play a significant role and this impacts the coke quantity, the particle size and the coking rate. In addition, the morphology changed significantly along with the progress of the coke production (increase of the diameters of both the spherical and filamentous coke). The nature of the specimen (incoloy, stainless steel, quartz, and vycor) also affected the coke production with a higher value for the metallic one. The authors suggested that heterogeneous reactions can enhance or limit the coke deposition and its nature. This will be discussed later. Finally, Albright and Marek [78] also observed that the length of the experiment can also modify the morphology of the carbon particles. This could signal a modification of the coking mechanism during the progress of the production process. Dunkleman and Albright demonstrated and quantified the influence of the surface [34]. They estimated the deposition rate for three reactors and determined the following ascending order: vycor glass, incoloy and then stainless steel. Additionally, it was observed that the S/V ratio has an impact, i.e. the higher S/V, the larger the carbon deposit. A mechanistic model of 18 reactions was developed in complement to their experimental data. Due to the very limited results, the model did not integrate surface reactions but “suppressed” them by correcting the initial reaction rates of the “classical” reaction pathway and simulating a surfaceless experiment. Such a correction permitted good agreement to be obtained between numerical predictions and experimental results. Altin and Elser [79,80,84] also explored the impact of the surface and compared the deposition of carbon, from JP-8 pyrolysis, on several surfaces and notably on two superalloys (Inconel 600 and Inconel X). The carbon deposition was 15 times higher for Inconel 600 than for Inconel X. Filamentous carbon coated by a carbonaceous layer remained on Inconel 600 while both crystalline and amorphous carbons were
deposited on Inconel X. Such differences are explained by the presence of several elements (e.g. Ti or Al) which act as carbon deposit inhibitors. Gascoin et al. conducted similar investigations [20] by performing dodecane pyrolysis experiments in stainless steel, low carbon steel and titanium tubular reactors. They provided the characteristics of the coke produced thanks to SEM observations, EDS, X-ray and FTIR analysis. The coke has different aspect as a function of reactor nature (e.g. for stainless steel tube, millimetric aggregates constituted from micrometric particles were mainly formed inside the fluid flow while for steel reactors, the deposit appeared as very dry sticky shavings). They found spatial heterogeneous deposit along the reactor. The increase of the residence time or of the pressure enhances the formation of coke while the production of solid carbon has been linked to methane formation. In [81], Reyniers et al. confirmed the role of pressure in the increase of the carbon productions. Lucas and Marchand pyrolyzed methane in a tubular reactor while modifying the temperature, the pressure and the flow rate [82]. They observed that the temperature and flow rate did not impact the density of the coke particles, the increase of pressure led to a decrease in the density. Naturally, the studies mentioned above are just a fraction of the numerous works which are available in the literature (e.g. studies on the effect of the surface on coke production, Ref.[85–90]). As explained by Oberlin in her excellent review [83], a considerable quantity of research has been performed under a very wide panel of conditions. This is why the results are so mixed and, sometimes, contradictory. This is why the first part of this paper aims at isolating the works which present the main information related to the influent parameters on coke production. Thus, it has to be noted that the increase of the pressure, the temperature and the residence time favor coke production. Regarding a hypersonic application, the first two parameters are not easy to control efficiently. On the contrary, acting on the residence time in order to shorten it but also using specific materials are parameters which appear, to a certain extent, more controllable.

3.2 Modeling the coke formation and its coupled impact on surface deactivation

Guéret et al. [77] studied the solid carbon production from the methane pyrolysis and they proposed an extensive literature survey. Their conclusion was that coke formation begins with the
formation of radicals from hydrocarbon degradation. Then, condensation and polymerization reactions lead to the production of some “complexes” (intermediates molecules in the formation of coke). At this point, there are two possibilities: on the one hand, adsorption of complexes at the surface and formation of an anisotropic pyrolytic carbon layer or on the other, condensation and polymerization of these in the gas phase. This last pathway is faster than the first one and generates a dispersed carbon with an isotropic structure (the so-called carbon black). Lucas and Marchand supplied a first approach for modeling the coking mechanism in case of methane pyrolysis [82]. They determined that C₂ hydrocarbons (acetylene and ethylene) and benzene were the most represented light species while naphthalene, anthracene, phenanthrene, acenaphtylene, pyrene and fluoranthrene were found to be the most common PAHs. Based on those results, Lucas and Marchand proposed the following simple mechanism divided into 4 steps:

\[
\text{CH}_4 \xrightarrow{\text{Fast}} \text{C}_2 \xrightarrow{\text{STEP 1}} \text{C}_6\text{H}_6 \xrightarrow{\text{STEP 2}} \text{PAH} \xrightarrow{\text{STEP 3}} \text{Carbon} \quad (8)
\]

They noted that the C₂ species had an essential role in the formation of benzene and of different PAHs. Considering the kinetics of the reactions, they assumed that methane dissociation coefficient had practically no influence on the production of C₂ because of the rate of the first step. On the contrary, a low dissociation coefficient of methane would decrease the formation of benzene and PAH while for a higher one, the production of benzene and PAH would initially increase before diminishing. The authors observed that carbon was more actively produced by reactions involving anthracene than the other PAHs. This simple mechanism was used and extended by Benzinger et al. [91]. They divided the coking phenomenon into five steps beginning with the cracking of methane:

STEP 1: From methane to ethane:

\[
\text{CH}_4 \xrightarrow{\text{Fast}} \text{CH}_3 + \text{H} \quad (9)
\]
\[
\text{CH}_3 + \text{CH}_4 \xrightarrow{} \text{C}_2\text{H}_6 + \text{H} \quad (10)
\]
\[
2\text{CH}_4 \xrightarrow{} \text{C}_2\text{H}_6 + \text{H}_2 \quad (11)
\]

STEP 2: From ethane to ethylene:


\[
\text{C}_2\text{H}_6 \rightarrow \text{C}_2\text{H}_4 + \text{H}_2 \quad (12)
\]

\[
\text{CH}_3 + \text{C}_2\text{H}_6 \rightarrow \text{CH}_4 + \text{C}_2\text{H}_5 \quad (13)
\]

\[
\text{C}_2\text{H}_5 + \text{H} \rightarrow \text{C}_2\text{H}_4 + \text{H} \quad (14)
\]

STEP 3: From ethylene to acetylene:

\[
\text{C}_2\text{H}_4 \rightarrow \text{C}_2\text{H}_2 + \text{H}_2 \quad (15)
\]

\[
\text{CH}_3 + \text{C}_2\text{H}_4 \rightarrow \text{CH}_4 + \text{C}_2\text{H}_3 \quad (16)
\]

\[
\text{C}_2\text{H}_3 \rightarrow \text{C}_2\text{H}_2 + \text{H} \quad (17)
\]

STEP 4: From acetylene to benzene:

\[
3\text{C}_2\text{H}_2 \rightarrow \text{C}_8\text{H}_6 \quad (18)
\]

STEP 5: Growth of solid carbon (\(\text{C}_\infty\)):

\[
\text{C}_\infty + \text{C}_6\text{H}_6 \rightarrow \text{C}_\infty + 3\text{H}_2 \quad (19)
\]

So, methane first cracks into \(\text{C}_2\) hydrocarbons. The condensation of acetylene leads to benzene and finally to the growth of carbon particles. Even if other compounds contribute to the coking phenomenon, benzene remains the most important intermediate species. Becker and Hüttinger followed their investigations on coking and performed experiments [92]. The authors observed the following elements. First, they found that the \(\text{C}_1\)-species can form coke directly. Secondly, the deposit rate shows an exponential increase with residence time. Thirdly, the increase of the initial partial pressure leads to the raise of \(\text{C}_1\) to \(\text{C}_6\) species with a plateau for ethane, ethylene and acetylene. Finally, hydrogen decreases coke production by saturating the free radical active sites. This leads to a critical fall of acetylene and of benzene formation. Based on those elements, the authors proposed a basic coke formation scheme where coke can be directly formed from numerous hydrocarbons including methane itself. It was also demonstrated that ethylene is not only formed from ethane, acetylene not only from ethylene and benzene not only from acetylene. These last assertions clearly reveal that a realistic coking mechanism involves a high number of reactions. Albright and Marek proposed a relatively more
complex model [93]. They combined three distinct -but linked- mechanisms. The first one (the so-called mechanism 1) considers the catalytic effect of the reactor and involves metal-catalyzed reactions. The coke produced by such a mechanism is filamentous and often contains metal. Acetylene is a predominant precursor while metal carbides are considered as intermediate compounds and iron and nickel as catalysts. Such coke is produced at temperatures ranging from 673 K to 1323 K. Mechanism number 2 highlights the importance of aromatics species (Ar-H) as intermediate in the formation of coke. A majority of those compounds come from acetylene. The following schematic sequence describes the production of solid carbon:

\[
\text{reaction} \quad n\text{Ar} - H \xrightarrow{-H_2} \text{tar} \xrightarrow{\text{nucleation}} \text{tar droplets} \xrightarrow{-H_2} \text{semitar droplets} \xrightarrow{-H_2} \text{coke} \quad (20)
\]

This coke is metal-free and does not appear below 973 K. The last mechanism describes the growth of the coke layer involving microspecies (generally acetylene) with the free radicals (methyl, ethyl, phenyl or benzyl radicals) on the coke surface forming aromatic rings. Hence, the authors concluded that acetylene is essential in the production of solid carbon as it produces the majority of coke precursors. Reyniers et al. confirmed this assumption of three combined mechanisms thanks to experimental data [81]. They conducted coking experiments considering feedstock from methane to complex fuels like kerosene; in presence of H2O or N2. Their main conclusions were: the light species like methane and ethane produce more coke than heavier molecules; more coke is found during the pyrolysis of isoparaffin than of normal paraffin; the presence of steam relatively inhibits the coke deposit by removing it at temperatures of over 1123 K; the solid carbon yield rises with the increase in pressure. Based on these observations, Reyniers et al. assumed that the carbon layer deposition depends on the contribution of three mechanisms. The first one is similar to the mechanism 1 described by Albright and Marek [93]. It consists in a heterogeneous catalytic mechanism which is predominant in the initial phase with a high rate of carbon deposition. Figure 3 presents a schematic view of this deposition.
This carbon layer is a porous structure where active sites permit coke deposition via a non-catalytic mechanism. Indeed, the covering of the metal surface leads to a progressive decrease in coke formation by this catalytic method. The second mechanism, described by the author as the "Heterogeneous Noncatalytic Mechanism", is the predominant one. It consists of the decomposition of hydrocarbons into aromatic structures and then into graphitic coke filaments. The polyaromatic layer is progressively dehydrogenated by free radicals which are present in the gas phase. The number of possible reactions is very large and depends on the nature of the feedstock. The last mechanism is described as homogeneous and noncatalytic. It involves the formation of PAH in the gas phase. These molecules agglomerate into tar droplets which generally adhere to the surface and are incorporated in the coke layer. This pathway prevails for heavy liquid feedstock and at temperatures of over 1173 K.

Since acetylene and PAH are the keys for the production of solid carbon, it could be of interest to take a quick look at this (e.g. [94–97]). The works of Böhm et al. [94,95] could be cited as an example for the present review. They studied the growth of PAH and the formation of soot experimentally during the pyrolysis of acetylene and of benzene at high temperature and pressure. The PAH formation model that they developed was based on three reaction pathways. First, the H-abstraction is associated with C$_2$H$_2$ addition to aromatics (e.g. acetylene addition to naphthalene forming phenanthrene and pyrene): the so-called HACA route. Then, combination reactions involve phenyl with benzene, the so-called ring-ring...
condensation. Finally, cyclopentadienyl is recombined (i.e. naphthalene may be formed without benzene). As a conclusion, PAH growth is more dependent on the ring-ring condensation than on the HACA route for short residence times. Benzene is consequently the most efficient precursor in comparison with acetylene. On the contrary, for longer residence times, the benzene consumption infers the dominance of acetylene as PAH precursor. The HACA route becomes essential. Equally, the optimal soot production is reached when these two mechanisms work “hand in hand”. Finally, the cyclopentadienyl pathway is important during acetylene pyrolysis.

3.3 Catalicity and coke

As is mentioned in the previous parts, the presence of a catalyst (reactor surface or solid) can modify the course of coke formation (enhancement or limitation) [2,33,34,44,56,78–80,84–90]. In addition, in the presence of a catalyst, another type of coke can be produced i.e. catalytic coke. If this is generally observed as different forms, catalytic coke is mostly formed as a filamentous carbon [98,99]. Schematically and considering a metallic surface, the production of filamentous carbon can be summarized as follows: first, the hydrocarbon is adsorbed on the surface; reactions occur and some coke particles are formed; then, carbon atoms are progressively dissolved and diffuse through the metal particles; an accumulation occurs which leads to a certain tension; the metal particles are removed from the surface and will act as active sites in the production of catalytic coke; a carbon filament is progressively formed with metal particles at its topside. Equally, the coke could affect catalicity generally by covering the reactive sites (i.e. catalytic deactivation). First, it must be mentioned that several reviews have been proposed on catalytic deactivation namely due to coke deposition, particularly during the last 70s and 80s. Precious information can be found and reference may be made to the following works for complements: Rostrup-Nielsen and Trimm [100]; Butt and Billimoria [101]; Trimm [102] or more recently Bartholomew [103]. Secondly, to the authors’ knowledge, no works are available regarding catalytic methane pyrolysis with coke production and catalyst deactivation. Consequently, this review has been extended to other hydrocarbons through the presentation of the
works of Guisnet and Magnoux [104–106]. In Ref.[104,105], they compared the thermal degradation of n-heptane with 4 different zeolites (USHY, HMOR, HZSM5 and HERI). They evaluated the coking impact on deactivation, which varies according to the nature of the zeolite. HMOR, HERI and, to a lesser extent, USHY deactivates quickly contrary to HZSM5 whose deactivation is slow. The deactivation is about 40 times greater for HERI, 15 times for HMOR, 3 times for USHY than for HZSM5. The structure and the size of the zeolite pores on the one hand and the density of acid sites on the other are responsible for this difference between zeolites. The modes of deactivation are the limitation of the access (for low coking rates, particularly predominant with HSZM5) or the total blockage of the access (predominant for HMOR and HERI) of the catalytic sites. In Ref.[106], Guisnet and Magnoux presented some propositions in order to limit this unwanted deactivation. They isolated 3 points: adapting the pore structure by preferentially chose zeolites without trapping cavities; adjusting the acidity of catalytic sites in order to obtain good homogeneity and finally, setting efficient operating conditions in order to avoid the blockage of pores. Paweewan et al.[107] also worked on “zeolite deactivation” during n-hexane degradation. They provided the main characteristics of coke particles and they explained its effect on deactivation which was quite different compared with the previous case. Indeed, its effect was attributed to “selective site poisoning” (i.e. the removing of the predominant active sites) instead of pore blockage.

Froment and coworkers[108–116] studied the deactivation of numerous catalysts (zeolites, metal) with regard to coke deposition. They developed models in order to predict this phenomenon by assessing the concept of site coverage and pore blockage. The particularity of this model is that time is not taken into account. For example, the deactivation function $\varphi_A$ (regarding the site coverage) for a given reaction (e.g. $A \rightarrow B$) is defined as:

$$\varphi_A = \left( \frac{C_t - C_{CI}}{C_t} \right)^{n_A} \quad (21)$$

with $n_A = 1$ (single site main reaction) or 2 (dual site main reaction), $C_t =$ total concentration of active sites and $C_{CI} =$ concentration of sites covered with coke or coke precursor.
As a consequence, the rate $r_A$ of this reaction is modified as follows:

$$r_A = r_A^0 \varphi_A$$  \hspace{1cm} (22)

with $0 \leq \varphi_A \leq 1$ and $r_A^0$ is the initial rate in absence of coke.

It must be noted that the deactivation of the catalyst directly affect the coke formation so that Eq. 22 is also valid for the coke rate. Since these equations are not sufficient for global deactivation to be taken into account, namely regarding the probability of pore blockage, Froment et al. explained that it is necessary to develop a model using probability theory and considering the geometry of the catalysts as a network of pores. For example, in Ref.[110] they suggested a complex model of four different catalyst networks. In Ref.[112], they improved this model by taking into account the diffusion limitation. In the early 2000s, Froment published a complete work on catalyst deactivation [116]. He divided his approach according to three levels: the active sites, the catalyst particle and the reactor (referring to the previous equations used for particle level and adding equations which describe the temperature and the concentration evolutions).

Among other works from the 90s and 2000s (e.g. Ref. [48,117–122]), Beltramini et al.[117] studied coke production and the deactivation effect on a bifunctional catalyst (metallic and acidic) according to different metal contents. They highlighted the different trends between metallic and acid sites, the first one being more affected by coke production (i.e. quick deactivation) and acting as a regulator on the coke production because of its involvement in the precursors of coke. Forzatti and Lietti proposed a review on catalyst deactivation dealing with its main causes [119]: poisoning, coking or sintering phenomena. They exposed an example of catalyst deactivation considering a fluid catalytic cracking model. Finally, Reyniers et al. [120,121] pointed out other important notions regarding the coke. They demonstrated its influence on several chemical reactions during the pyrolysis of several hydrocarbons: n-hexane, 2,2,4 triMe-pentane, propene, and i-butene. They showed the influence of the gas phase composition on the coking rate, notably the positive effect of olefins. The coke molecules can form carbenium ions and they act as hydride donors and, consequently, they interact and they intervene.
in the formation of catalytic products. For example, regarding the hexane degradation, the formation and
the presence of coke favors the selectivities of C₁ and C₂ hydrocarbons. Finally, the reaction rates can be
positively or negatively influenced by the coke formation according to the reaction type, the coke
content and the acid sites properties.

4. Conclusions

Pyrolysis involves a complex kinetic mechanism in the homogeneous phase whatever the initial
fuel composition to be considered and this becomes even more challenging when considering
heterogeneous reactions with a surface reactor and the formation of solid coke particles. Through the
literature survey presented in this work, a wide variety of experimental conditions has been presented.
The understanding of all the coupled phenomena which occur is not simple and numerous parameters
have to be considered. The nature of the inner wall drastically affects the course of thermal degradation.
Regarding this aspect, several elements are of importance. If the nature of the catalyst seems to be an
obvious factor, others should be mentioned and must be considered and in particular: the temperature,
the pressure, the nature of the fuel and the S/V ratio. Equally, interesting parallel parameters have been
assessed such as the pretreatment of the reactor or the support of the catalyst which can significantly
modify the catalytic activity. Several kinetic models have been developed. Studying the interaction
between gas species and active sites has become more complex by considering its potential deactivation
namely by solid carbons produced during pyrolysis. The so-called coke is an unwanted product which
can be of different natures, diluted in the gas phase and deposited at the surface of the reactor wall. In so
doing the catalytic activity is drastically reduced which impacts the kinetics of the reaction. Several
works have demonstrated the possibility of decreasing such an undesired process namely by pre-treating
the surface or setting efficient conditions.

Studying such effects is incontrovertible regarding the context of regenerative cooling thanks to
the endothermic degradation of fuel during hypersonic flights. If several studies are available, it is
undeniable that there remains a clear lack in this domain with regard to methane and namely regarding
the interaction between this fluid and its pyrolyzed product with materials of interest (i.e. composites). The operating conditions should be intensively explored within the scope of limiting coke production.

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6. References


