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Radical-assisted polymerisation in interstellar ice analogues : formyl radical and polyoxymethylene

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ABSTRACT

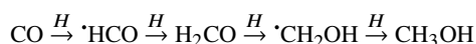
We present new laboratory experiments on the low-temperature formation of COMs such as polyoxymethylene (POM), glycolaldehyde (GA), ethylene glycol (EG) and possibly glyceraldehyde (GCA) and glycerol (GCO) through radical-induced reactivity from VUV photolysis of formaldehyde in Ar and Xe matrices. The radical reactivity and the endogenous formation of COMs were monitored *in-situ* via infrared spectroscopy in the solid state and post photolysis with temperature programmed desorption (TPD) using a quadrupole mass spectrometer. Based on experimental finding and quantum calculations, we elaborate a formation pathway for formaldehyde polymerisation induced by radicals ($\cdot\text{HCO}$ and $\cdot\text{CH}_2\text{OH}$) that support the POM formation in cometary environments. In addition, fragmentation patterns obtained from the sublimation of short chain-length POM are consistent with data collected by the Ptolemy instrument on-board the *Rosetta* mission and strengthen the POM identification made by this instrument.

Key words: Astrochemistry – Molecular processes– Comets:general– Methods: laboratory: molecular

1 INTRODUCTION

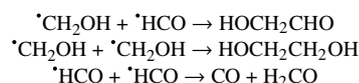
About 180 organic molecules were observed in the interstellar medium. They are generally simple molecules but almost 50 of them have 6 or more atoms and are called Complex Organic Molecules, hereafter COMs (Woods et al. 2013). Despite the large molecular diversity and complexity observed in space in most cases we are unaware of their formation mechanism. However, it is generally accepted that COMs form on the surfaces of icy dust grains. This idea is supported by laboratory results and astrochemical simulations, indicating that surface and bulk reactions induced by atom addition/abstraction, cosmic rays, VUV-photons, and by thermal processing are efficient processing for COMs formation in space.

In cold and dark environments (molecular clouds or dense cores), non-energetic processing mainly consists of atomic and hydrogenation reactions. One of the most well known reaction is the CO hydrogenation (Watanabe et al. 2003, 2004; Fuchs et al. 2009; Minissale et al. 2016):

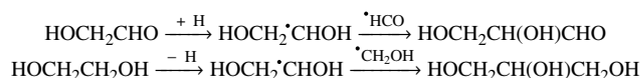


Along H-atom addition reactions on CO, two important radicals are formed $\cdot\text{HCO}$ and $\cdot\text{CH}_2\text{OH}$. It has been recently proven that direct recombination $\cdot\text{HCO}$ and $\cdot\text{CH}_2\text{OH}$ free radicals form glycolalde-

hyde (GA, HOCH_2CHO), ethylene glycol (EG, $\text{HOCH}_2\text{CH}_2\text{OH}$) (Butscher et al. 2015; Fedoseev et al. 2015; Chuang et al. 2015, 2017; Fedoseev et al. 2017). However, Butscher et al. (2017) recently showed that $\cdot\text{HCO}$ in astrophysical conditions is not able to dimerise into glyoxal (CHOCHO) but instead forms CO and H_2CO :



Similar mechanisms based non-energetic processing and radical recombination in interstellar ice analogues can also result in the formation of biologically relevant molecules such as glyceraldehyde (GCA, $\text{HOCH}_2(\text{OH})\text{CHO}$) and glycerol (GCO, $\text{HOCH}_2\text{CH}(\text{OH})\text{CH}_2\text{OH}$) (Fedoseev et al. 2017). In this reaction scheme, carbon backbone increase is realized through recombination of CO-bearing radicals such as $\cdot\text{HCO}$ and $\cdot\text{CH}_2\text{OH}$ on $\text{HOCH}_2\cdot\text{CHOH}$ intermediate radical formed from H-atom addition/abstraction on GA and EG:



Energetic processing (photons, electrons, X-rays) that are prominent during later stages of the star and planet formation process can also trigger reactive intermediates and radicals formation. It has been extensively shown that laboratory experiments on CH_3OH -rich ice analogues exposed to such energetic processing

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can efficiently produce a large diversity of COMs including amino acids and sugars. (Caro et al. 2002; Öberg et al. 2009; Abou Mrad et al. 2014, 2016; Kaiser et al. 2015; Henderson & Gudipati 2015; Chuang et al. 2017).

COMs are also detected in comets that harbor the most pristine material in our solar system that may have preserved their interstellar heritage (Goesmann et al. 2015; Wright et al. 2015; Biver et al. 2014; Crovisier et al. 2004; Huebner 1987) as recently revealed by the *Rosetta* mission. Twenty five minutes after lander Philae's initial comet touchdown onto comet 67P/Churyumov-Gerasimenko, COSAC instruments performed analyses that displayed 16 organic compounds -including CHO-bearing molecules such as glycolaldehyde (GA) and ethylene glycol (EG)- while Ptolemy instrument detected regular mass distributions suggesting the presence of a radiation-induced POM-like polymer (Wright et al. 2015). This backs up analysis made with the PICCA instruments on comet Halley during the *Giotto* mission where a similar regular pattern of peaks has been observed, which was also tentatively assigned at that time to POM-like structures (Huebner 1987). Even if the POM detection by both *Rosetta* and *Giotto* missions is controversial (Mitchell et al. 1989; Altwegg et al. 2017), the presence of POM in comets (Schutte et al. 1993) cannot be excluded. First, it is so far the best candidate to explain the distributed source of CO and formaldehyde observed in coma (Cottin & Fray 2008; Fray et al. 2006). It is also supported by laboratory experiments. POM and POM-like polymers can be easily formed from the heating of interstellar/cometary ice analogs containing H₂CO, H₂O, and a nucleophile (NH₃, CN⁻) (Schutte et al. 1993; Danger et al. 2014; Noble et al. 2012; Vinogradoff et al. 2011; Duvernay et al. 2014) or from UV irradiation of interstellar/cometary ice analogues containing formaldehyde (Butscher et al. 2016).

Thus, by exposing energetic and non-energetic processing on ice analogues, COMs formation is indeed observed but almost no reaction intermediates are identified this way (Schutte et al. 1993; Butscher et al. 2016; Bennett & Kaiser 2007). To solve this problem, we resort to the cryogenic matrix technique. Already used before for the characterisation of radicals of astrophysical interest (Milligan & Jacox 1969; Hiraoka et al. 1995; Pirim & Krim 2011; Lee et al. 2015), it consists in trapping the starting material in a rare gas matrix and submitting it to energetic processes. This way, all species are isolated from one another in a cold environment. This technique has two advantages. First, the way the molecules are isolated prevents them from rotating freely so that most of rotational contributions are removed and no intermolecular interactions are possible if molecules are present in monomeric form in their cage. The resulting spectrum becomes close to that of gas phase without rotational structure. Also, since trapped molecules are alone in their cage, reactive species are not able to find a partner to react with, so when radical species are produced, their lifetime and stability in the rare gas matrix are greatly increased, making their characterisation easier with infrared spectroscopy. Then, by quickly removing the rare gas cages, radicals and reactive species are free to react and recombine (Butscher et al. 2015, 2017).

The objective of the present work is to go further in the understanding of the reactivity of small organic radicals. Using cryogenic matrix technique, we specifically study the formaldehyde polymerisation initiated by radicals that might explain some observations made on comet 67P/Churyumov-Gerasimenko by the Ptolemy instrument on-board *Rosetta*'s lander Philae (Wright et al. 2015) or on Halley comet by the *Giotto* mission. The experimental setup is described in section 2 and the results of each step of our experiments as well as theoretical calculations are detailed in section 3. Discus-

sion includes a confrontation with astrophysical observations and implications.

2 EXPERIMENTAL AND COMPUTATIONAL METHODS

2.1 Experimental details

Argon and Xenon (99.999 % of purity) used as matrix rare gas (Rg) were purchased from Air Liquide. Formaldehyde was purchased as a polymer from Sigma Aldrich (99.95 % of purity) and was heated to about 90°C to produce the gas-phase monomer. For references, glycolaldehyde was purchased as a dimer from Sigma Aldrich (99.95 % of purity) and heated under vacuum to about 80°C to produce gas-phase monomer. Ethylene glycol was purchased from Sigma Aldrich. The experimental AHIA system has already been described precisely in a previous article (Butscher et al. 2015).

All experiments described are performed in a high vacuum chamber with a background pressure of 10⁻⁸ mbar at 295 K and 10⁻⁹ mbar at 13 K. The temperature is controlled using a model 21 CTI cold head, a resistive heater and a Lakeshore 331 temperature controller. Products were mixed in 2/1000 ratio of H₂CO/Rg in a Pyrex vacuum line using standard manometric techniques. Gaseous mixtures were deposited at a rate of 5 10⁻² mbar s⁻¹ on a gold-plated surface kept at 30 K for xenon experiments and 13 K for argon experiments. Glycolaldehyde and ethylene glycol solid films can be obtained by directly dosing them onto the sample holder in order to get spectroscopic references. POM used for spectroscopic reference is produced directly onto sample holder by a thermally-induced formaldehyde polymerisation catalysed by ammonia (Duvernay et al. 2014; Schutte et al. 1993).

Radicals are produced from the VUV photolysis of formaldehyde H₂CO in a rare gas matrix. The VUV photons are generated from a microwave induced low pressure H₂ plasma using a microwave magnetic applicator (Boreal Plasma) fed by a microwave generator (Optos instruments). In a such low pressure H₂ plasma (20 10⁻³ mbar), the VUV emission is dominated by Lyman alpha photons at 121.6 nm but molecular H₂ emission at 160 nm cannot be excluded. The VUV flux is transmitted from the plasma chamber to the vacuum chamber through an MgF₂ window. The photon flux has been measured using the O₂ → O₃ actinometry method (Cottin et al. 2003) to *c.a.* 2.5 10¹³ photons cm⁻² s⁻¹, which is 10¹⁰ times larger than the UV secondary flux in dense molecular clouds.

The sample are monitored using a Bruker Tensor 27 FTIR spectrometer with MCT detector between 4000 and 600 cm⁻¹ in reflection mode (reflection angle <5°) which can be considered as double transmission technique since the infrared beam angle of incidence is closed to the normal. Each spectrum was averaged over 20 scans with a 0.5 cm⁻¹ resolution, except for the background averaged over one hundred scans with the same resolution. The quantification of products observed in the solid film after rare gas desorption is based on infrared analysis. The column density \mathcal{N} (molecule cm⁻²) can be determined by integrating either the absorbance $A(\nu)$ or the optical depth $\tau(\nu)$, according to the equations: $\mathcal{N} = \frac{\int \tau(\nu)}{2\cos(5^\circ)\mathcal{F}} = \frac{\ln(10) \int A(\nu)}{2\cos(5^\circ)\mathcal{F}}$, where \mathcal{F} is the band strength (cm molecule⁻¹) and ln(10) is needed to convert the integrated absorbance to optical depth. The column density of GA is obtained from the IR band at 1746 cm⁻¹ ($\mathcal{F} = 2.6 \times 10^{-17}$ cm molecule⁻¹), the amount of EG is obtained from the IR band at 1089 cm⁻¹ with a strength $\mathcal{F} = 3.9 \times 10^{-18}$ cm molecule⁻¹ (Hudson et al. 2005), and the amount of POM is obtained from the IR band at 902

cm^{-1} with a strength $\mathcal{F} = 5 \times 10^{-19}$ cm molecule $^{-1}$ (Schutte et al. 1993). It has to be noted that the values of the band strengths depend on the nature, composition, and temperature of the ice in which they are found, and this dependence is a major source of uncertainties when evaluating the column densities of frozen molecules.

All the experiments described below were done in three steps : (i) Radical formation from VUV photolysis of radical precursor formaldehyde; (ii) Rare gas matrices annealing; (iii) Rare gas desorption at 35 K for argon and 85 K for xenon. After rare gas desorption, a ramp of 4 K min $^{-1}$ is applied while the ion current is recorded from RGA quadrupole mass spectrometer (MKS Microvision-IP plus) to obtain a full temperature programmed desorption (TPD) profile, relative to various products. The ionisation source was a 70 eV impact electronic source and the mass spectra were recorded between 1 and 100 amu in a full scan.

2.2 Computational details

Density Functional Theory (DFT) and coupled-cluster calculations were performed using the Turbomole V7.2 package. The structures of the reactants, transition states and products were fully optimised using the Becke's three-parameter hybrid functional (B3LYP) (Becke 1993) in an unrestricted formalism, along with the dev2-TZVP basis set (Weigend & Ahlrichs 2005). These structures were characterised by vibrational harmonic frequencies and zero-point energies (ZPE) calculated at the same level of theory using the aoforce module (Deglmann et al. 2004).

For all stationary points, single-point calculations were carried out at the explicitly correlated Unrestricted Coupled Cluster Single and Double with inclusion of a perturbative estimation for triple excitations (UCCSD(T)-F12) (Tew & Klopper 2010) level along with the cc-pVTZ basis set (Dunning 1989). Explicitly correlated (F12) methods allow rapid orbital convergence and can lead to near basis set limit accuracy using small basis sets, with a small extra computational cost compared to conventional methods (Hättig et al. 2012). They are very useful for benchmarking, especially in the field of thermochemistry, and have proven to be highly accurate for astrochemically relevant systems (Stein et al. 2015).

3 RESULTS AND DISCUSSION

3.1 Experimental results

3.1.1 Formaldehyde photolysis at $\lambda > 120$ nm : radicals formation in argon and xenon matrices

A gaseous mixture of formaldehyde and argon in a 2/1000 ratio is deposited on a cold surface (13 K in our case) at low pressure (10^{-8} mbar) with a controlled flow rate. In Ar matrix, the main bands of H₂CO are observed at 2864, 2798, 1742 and 1499 cm $^{-1}$ (negative bands in Fig. 1, Table 1). These absorptions correspond to asymmetric and symmetric CH stretching mode, C=O stretching mode and CH bending mode respectively and are down-shifted with ^{12}C to ^{13}C isotopic change (Milligan & Jacox 1969). The corresponding values in Xe matrix are listed in Table 1. We assume that formaldehyde is trapped as monomer inside the matrix since the infrared signatures of formaldehyde dimer are not detected after the deposition (Nelander 1980). In this work, we tried to figure out the reactivity of formyl radical $\cdot\text{HCO}$ when the main reactivity partner is formaldehyde H₂CO. To make it possible, we performed an VUV irradiation ($\lambda > 120$ nm) at low fluence ($2.7 \cdot 10^{18}$ photons cm $^{-2}$ corresponding to 30 min of irradiation) of H₂CO trapped in

Table 1. Infrared absorption bands and assignments of H₂CO in Ar and Xe matrices at 13 K.

Ar		Xe		Assignments
^{12}C	^{13}C	^{12}C	^{13}C	
3464	3389	3449	3375	$2\nu(\text{CO})$
2997	2954	2980	2944	$2\delta(\text{CH})$
2864	2846	2837	2822	$\nu(\text{CH}) - a$
2798	2793	2771	2767	$\nu(\text{CH}) - s$
2719	2708	2701	2691	
1742	1704	1735	1697	$\nu(\text{CO})$
1499	1499	1491	1491	$\delta(\text{CH})$
1245	1235	1238	1232	$\rho(\text{CH})$
1168	1156	1163	1151	$\omega(\text{CH})$

ν : stretching; δ : bending; ρ : rocking; ω : wagging.

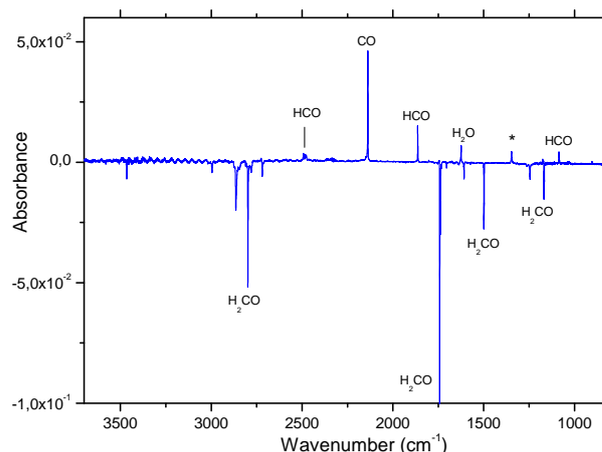


Figure 1. Difference infrared spectrum after VUV photolysis (30 min, $\lambda > 120$ nm) at 12 K in Ar matrix. Negative bands stand for H₂CO, while positive bands stand for photo-products. Bands labeled with stars refer to unknown species.

rare gas matrices. This way, only 7% of formaldehyde is consumed and photo-products are then produced.

Figure 1 displays the difference spectrum showing the effects of VUV irradiation in Ar matrix. The negative bands in Fig. 1 are due to the consumed formaldehyde and the positive bands to the newly formed products. Table 2 displays IR assignments of products observed after ^{12}C and ^{13}C formaldehyde photolysis in both Ar and Xe matrices. $\cdot\text{HCO}$ radical is clearly identified in ^{12}C experiments by the bands in Ar matrix at 2482 cm $^{-1}$, 1863 cm $^{-1}$ and 1085 cm $^{-1}$, corresponding to its CH stretching, C=O stretching and CHO bending vibrational modes respectively (Milligan & Jacox 1969) (Table 2, Fig. 1). The corresponding value in Xe matrix and for ^{13}C experiments are listed in Table 2. CO and CO₂ are also visible and characterised by their bands in Ar matrix at 2138 cm $^{-1}$ and 2345 cm $^{-1}$ respectively, corresponding to the CO stretching mode (Abe et al. 1999; Schriver et al. 2000). Finally, a band at

Table 2. Infrared absorption bands and assignments of produced species after ^{12}C and ^{13}C formaldehyde photolysis ($\lambda > 120$ nm, 30 min) at 12 K in Ar and Xe matrices.

Wavenumbers (cm^{-1})				Modes	Species
Ar	Xe				
^{12}C	^{13}C	^{12}C	^{13}C		
2482	2476	2442	2436	$\nu(\text{CH})$	$\cdot\text{HCO}$
2345	2279	2334	2269	$\nu(\text{CO})$	CO_2
2138	2091	2133	2086	$\nu(\text{CO})$	CO
1863	1823	1857	1817	$\nu(\text{CO})$	$\cdot\text{HCO}$
1624	1624			$\delta(\text{OH})$	H_2O
1346	1346			?	?
1085	1078	1082	1076	$\delta(\text{CHO})$	$\cdot\text{HCO}$
		953	953	$\nu(\text{XeH})$ as	Xe_2H^+
903	903			$\nu(\text{ArH})$ as	Ar_2H^+
		843	843	$\nu(\text{XeH})$ as	Xe_2H^+
		731	731	$\nu(\text{XeH})$ as	Xe_2H^+

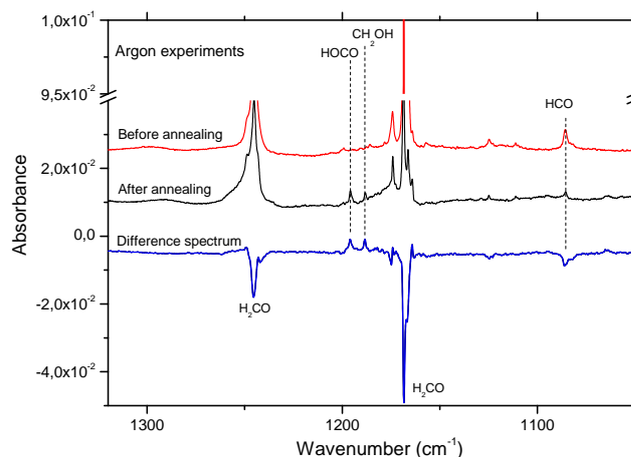
ν : stretching; δ : bending; ρ : rocking; ω : wagging; τ : twisting.

903 cm^{-1} is detected after photolysis in Ar matrix. Its intensity increases with the irradiation duration, proving that it is a product from VUV photolysis. According to Kunttu et al. (1992), this band can be assigned to Ar_2H^+ since small hydrides are known to form ionic complexes in rare gas matrices (Pettersson et al. 2000). When photolysis is performed in Xe matrix three bands are observed at 952, 842, and 730 cm^{-1} assigned to Xe_2H^+ (Pettersson et al. 2000). Thus, the main radical species formed in both Ar and Xe matrices at low fluence in a H_2CO -dominated environment is the formyl radical ($\cdot\text{HCO}$).

3.1.2 Matrix annealing : *in situ* hydrogenation reactions

The second step of our experiment is an annealing step, in which we heat at 30 K (50 K in Xe matrix) during 10 min without desorbing the matrix and then cool down the system back to 13 K. This way, all H-atoms - formed during the formaldehyde photolysis - will diffuse through the matrix and will react whenever a partner is met. New species coming from hydrogenation reactions within the rare gas matrices are then detected on the infrared spectrum. This technique has been already successfully used in our previous work to form the hydroxymethyl radical $\cdot\text{CH}_2\text{OH}$ from *in situ* hydrogenation reactions of formaldehyde (Butscher et al. 2015, 2016). During the annealing small molecules or small radicals may also be able to exit from their matrix cage but because of the high dilution used in our experiments they will enter in a matrix cage that is most likely empty. In these particular conditions radical-radical recombination are unlikely and only products coming from reaction with H-atoms, the most diffusive species, are expected.

Annealing the irradiated sample results in noticeable changes in the difference IR spectrum (Fig. 2). The study is focused on the $1400\text{-}1000\text{ cm}^{-1}$ zone of the difference spectrum, known as the fingerprint zone, to make the assignment easier. New products formed from *in situ* hydrogenation reactions are shown as positive bands

**Figure 2.** Infrared difference spectrum showing the effect of annealing at 30 K after the VUV photolysis ($\lambda > 120$ nm, 30 min) of formaldehyde in Ar matrix. Positive bands relates to formed species and negatives bands to consumed species

on the difference spectrum in Figure 2, whereas negative bands relate to consumed species. Infrared frequencies and assignments of present species after annealing step are listed on Table 3 along with their evolutions during the annealing. This annealing step makes the Ar_2H^+ band at 903 cm^{-1} disappears (the same is true with Xe_2H^+ in Xe matrix experiments), proving that all H-atoms reacted (Butscher et al. 2015; Pettersson et al. 2000). Due to the low fluence used in our experiments, only a low amount of free H-atoms were produced. However, CO is still partially consumed to form $\cdot\text{HCO}$ by hydrogenation reactions since CO band at 2138 cm^{-1} decreases during the annealing (Table 3). This radical will also be partially hydrogenated to form formaldehyde H_2CO . However, as seen in Fig. 2, the $\cdot\text{HCO}$ radical is still present after the annealing in the matrix.

A new radical species appears by the hydrogenation of formaldehyde H_2CO . Hydroxymethyl radical $\cdot\text{CH}_2\text{OH}$ is detected in Ar matrix from its bands at 1358 , 1183 , and 1048 cm^{-1} corresponding to its HOC bending mode, CH bending mode, and HCOH bending mode respectively. These assignments are in good agreement with previous works and are confirmed in ^{13}C and xenon experiments (Butscher et al. 2015; Jacox & Milligan 1973) (Table. 3). A band at 1188 cm^{-1} is also detected. It is assigned to the CH bending mode of $\cdot\text{CH}_2\text{OH}:\text{CO}$ molecular complex (Butscher et al. 2015). As only a few free H-atoms were formed, only a small amount of $\cdot\text{CH}_2\text{OH}$ is produced whereas methanol CH_3OH -the fully saturated molecule in the CO hydrogenation chain- is not detected. Another radical is also detected $\cdot\text{HOCO}$ from its bands located at 1843 cm^{-1} in Ar matrix (Jacox 1988; Ryazantsev & Feldman 2014). A small band at 1195 cm^{-1} (Fig. 2) is tentatively assigned to the HOC bending mode of $\cdot\text{HOCO}:\text{CO}$ molecular complex (Jacox 1988; Ryazantsev & Feldman 2014). At this stage, the present molecules are H_2CO , $\cdot\text{HCO}$, CO , CO_2 , $\cdot\text{CH}_2\text{OH}$, and $\cdot\text{HOCO}$.

3.1.3 Radical induced polymerisation : reactivity with an unsaturated molecule

The last step of our experiments is the rare gas desorption. The rare gas matrices are quickly heated at 35 K for Ar and 85 K for Xe, making the matrix cages disappear forming an amorphous solid film

Table 3. Infrared absorption bands and assignments of present species after annealing step (heating to 30 K in Ar matrix and 55 K in Xe matrix and cooling down back to 12 K).

Wavenumbers (cm ⁻¹)				Modes	Species	Bands evolution*
Ar		Xe				
¹² C	¹³ C	¹² C	¹³ C			
2482	2476	2442	2431	$\nu(\text{CH})$	[•] HCO	↘
2345	2279	2334	2269	$\nu(\text{CO})$	CO ₂	↗
2138	2091	2133	2086	$\nu(\text{CO})$	CO	↘
1863	1823	1857	1817	$\nu(\text{CO})$	[•] HCO	↘
1843	-	-	-	$\nu(\text{CO})$	HOCO	↗
1358	1355	1355	1350	$\delta(\text{HOC})$	[•] CH ₂ OH	↗
1195	-	-	-	$\delta(\text{HOC})$	HOCO:CO	↗
1188	-	-	-	$\delta(\text{CH})$	[•] CH ₂ OH:CO	↗
1183	1176	1176	1162	$\delta(\text{CH})$	[•] CH ₂ OH	↗
1085	1078	1082	1076	$\delta(\text{CHO})$	[•] HCO	↘
1048	-	1044	1040	$\delta(\text{CHOH})$	[•] CH ₂ OH	↗

ν : stretching; δ : bending; *: Infrared bands evolution during the annealing at 30 K.

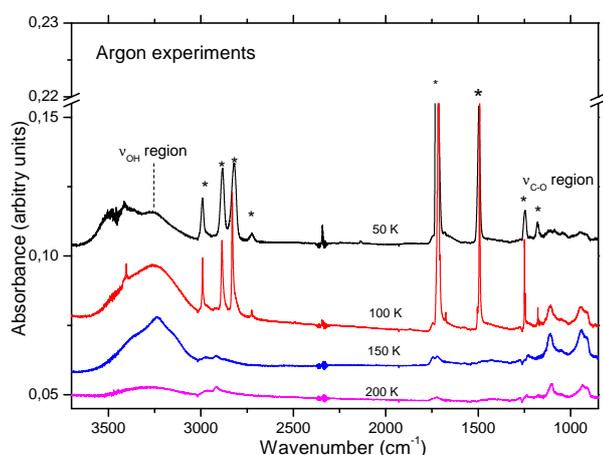


Figure 3. Infrared spectrum of the solid film obtained after argon desorption (35 K) from the sample recorded during the warming from 50 to 200 K (4 K min⁻¹). Bands labeled with stars are due to formaldehyde.

in which freed radicals will then react with other partners from radical/radical and radical/non-radical reactions, forming new species. Figure 3 (top spectrum) displays the solid film infrared spectrum recorded after Ar desorption at 50 K. As expected, it is dominated by H₂CO signatures but new infrared contributions are also visible at 3238 cm⁻¹ in the OH stretching region and at 1116, 941, and 905 cm⁻¹ in the C-O stretching region. (Fig. 3, top spectrum). The solid obtained after Ar desorption at 50 K is warmed at a rate of 4 K min⁻¹. The infrared spectra recorded at specific temperatures (100, 150, and 200 K) are displayed in Figure 3. Between 50 and 100 K, the most striking differences are the increase of bands located at 1116, 941, and 905 cm⁻¹ in the CO stretching region and the refinement of infrared absorption bands of formaldehyde mainly due to

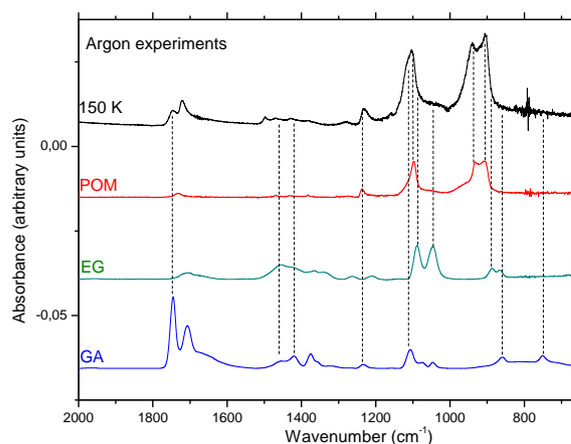


Figure 4. Infrared spectrum of the solid film recorded at 150 K after Ar desorption from the sample at 35 K. It is compared to reference spectra of polyoxymethylene, ethylene glycol, and glycolaldehyde

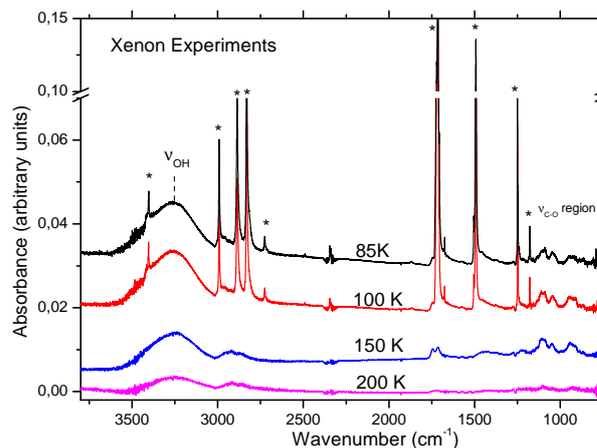


Figure 5. Infrared spectrum of the solid film obtained after xenon desorption (85 K) from the sample recorded during the warming from 85 to 200 K (4 K min⁻¹). Bands labeled with stars are due to formaldehyde.

crystallisation of the solid film. At 150 K, the bands of formaldehyde are not present due to its desorption from the sample holder (Fig. 3). In order to determine the new-formed species, attributions are made on the 2000-700 cm⁻¹ area that is a more characteristic region for identification. In this figure, the IR spectrum recorded at 150 K after formaldehyde desorption is compared with reference spectra of polyoxymethylene (POM), ethylene glycol (EG, HOCH₂CH₂OH), and glycolaldehyde (GA, C(O)HCH₂OH) (Fig. 4). As seen on Figure 4 (top spectrum), the bands at 1116, 941, and 905 cm⁻¹ can easily be identified as C-O stretching modes of formaldehyde polymer polyoxymethylene (POM) (Schutte et al. 1993; Duvernay et al. 2014). Apparently, the low amount of radicals ([•]HCO, [•]CH₂OH, etc) is sufficient to initiate chain-propagating H₂CO polymerization reactions right after the rare gas sublimation but also during the warming. Finally the bands of POM start to decrease around 180 K and totally disappear after 250 K (Fig. 3). This is due to POM sublimation and thermal decomposition into formaldehyde (Duvernay et al. 2014).

Glycolaldehyde (GA, C(O)HCH₂OH) and ethylene glycol (EG, HOCH₂CH₂OH) are also detected in this experiment. Im-

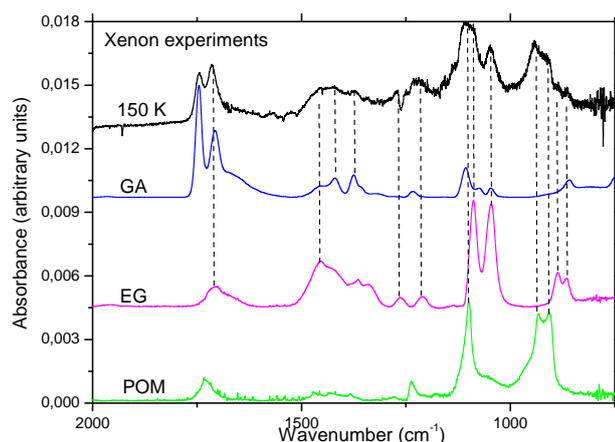


Figure 6. Infrared spectrum of the solid film recorded at 150 K after Xenon desorption from the sample at 85 K. It is compared to reference spectra of polyoxymethylene, ethylene glycol, and glycolaldehyde. See experimental part for details.

Table 4. Infrared absorption bands and assignments of present species in the solid film at 50 K.

Wavenumbers (cm ⁻¹)		Assignments	Molecules
¹² C	¹³ C		
3238	3235	$\nu(\text{OH})$	EG + GA + POM
2992	2958	$2\delta(\text{CH})$	H ₂ CO
2863	2958	$\nu(\text{CH})$	H ₂ CO
2821	2816	$\nu(\text{CH})$	H ₂ CO
2343	2275	$\nu(\text{C}=\text{O})$	CO ₂
2136	2090	$\nu(\text{CO})$	CO
1746	1716	$\nu(\text{C}=\text{O})$	GA
1725	1690	$\nu(\text{C}=\text{O})$	H ₂ CO
1699	1661	?	GA
1495	1495	$\delta(\text{CH})$	H ₂ CO
1247	1236	$\rho(\text{CH})$	H ₂ CO
1178	1165	$\omega(\text{CH})$	H ₂ CO
1116	1096	C-O	POM
1089	1060	$\nu(\text{C}-\text{O})$	EG
1047	/	$\nu(\text{C}-\text{O})$	EG
941	923	$\nu(\text{C}-\text{O})$	POM
905	885	$\nu(\text{C}-\text{O})$	POM

ν : stretching; δ : bending; ρ : rocking; ω : wagging.

portantly, these species were not detected in rare gas matrix before rare gas sublimation indicating that COMs observed in the solid films come from radical reactivity during the rare gas sublimation. The GA C=O stretching mode at 1746 cm⁻¹ is clearly visible. Two bands at 1089 and 1047 cm⁻¹ hint that EG is also present in the short-irradiation experiment. Nevertheless, it is also interesting to note that glyceraldehyde (GCA) and glycerol (GCO) have similar frequencies and they could also contribute to these bands. All

Table 5. GA, EG, and POM column densities (mol cm⁻²) observed in solid films after VUV photolysis and rare gas desorption.

	Ar experiments		
	GA	EG	POM
50 K	$2 \pm 0.2 \times 10^{15}$	$4 \pm 0.3 \times 10^{15}$	$4 \pm 0.2 \times 10^{15}$
150 K	$2 \pm 0.2 \times 10^{15}$	$4 \pm 0.3 \times 10^{15}$	$13 \pm 0.2 \times 10^{15}$
	Xe experiments		
	GA	EG	POM
85 K	$1 \pm 0.3 \times 10^{15}$	$20 \pm 0.1 \times 10^{15}$	$3 \pm 0.4 \times 10^{15}$
150 K	$1 \pm 0.3 \times 10^{15}$	$20 \pm 0.1 \times 10^{15}$	$4 \pm 0.2 \times 10^{15}$

frequencies and attributions are listed on Table 4. As previously suspected, infrared absorptions of methanol are not visible on the spectra of this experiment. It is a direct consequence of the limited production of H-atoms from the photolysis of formaldehyde H₂CO.

We calculated the column densities of GA, EG, and POM (Table 5). At 50 K (ie after the Ar sublimation) EG and POM are the dominant products with a column densities of $4 \pm 0.3 \times 10^{15}$ mol cm⁻² each whereas the column densities of GA is $2 \pm 0.2 \times 10^{15}$ mol cm⁻². At 150 K as seen in Fig. 4, POM is efficiently formed and becomes the main product whereas there is no evolution on GA and EG column densities during the warming. This indicates that GA and EG are mainly formed during Ar sublimation when cage opening released $\cdot\text{HCO}$ and $\cdot\text{CH}_2\text{OH}$ radicals that quickly react forming GA and EG (Butscher et al. 2015; Fedoseev et al. 2015; Chuang et al. 2015, 2017; Fedoseev et al. 2017). After the rare gas sublimation is complete, remaining radicals in low amount (below the detection limit) are isolated in a formaldehyde-dominated environment inhibiting radical-radical recombination but allowing chain-propagating H₂CO polymerisation reactions as the temperature is increased.

This is confirmed with xenon experiments that allow matrix cage sublimation at higher temperature than in argon. Figure 5 displays the solid film infrared spectrum recorded after Xe desorption at 85 K. The same products as previously are detected but now EG is the most abundant product (see Table 5). During the warming from 85 K to 150 K no evolution is observed on GA and EG column densities and only a small variation of POM (Fig. 6 and Table 5). Since the xenon desorption occurs at higher temperature than in Ar, radical-radical recombination is more efficient due to higher radical mobilities. The dramatic consumption of radical initiators -especially $\cdot\text{CH}_2\text{OH}$ - inhibits formaldehyde polymerisation in the solid film explaining the small POM formation during the warming. This may also indicate that $\cdot\text{CH}_2\text{OH}$ radical is a more important radical in chain-propagating H₂CO polymerization reactions than $\cdot\text{HCO}$ radical. This point will be clarified in the theoretical part.

Attributions were also confirmed using Temperature Programmed Desorption (TPD) experiment with a quadrupole mass spectrometer. The objective of such an experiment is to identify the desorbing molecules by mass spectrometry. Indeed, desorption temperature and fragmentation pattern are inherent to a molecular structure. Figure 7 shows the TPD graph from 125 K to 300 K, after the formaldehyde desorption. At least seven desorption peaks are observed during the TPD experiments indicating that more than the three previously assigned products have been formed in the experiments (Fig. 7).

The first desorption peaks at 160 K can be assigned to GA from signal at m/z 60 and m/z 43. (Fedoseev et al. 2017). EG is also

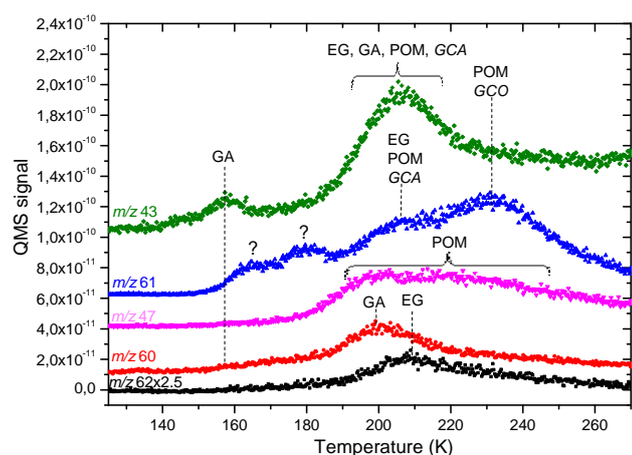


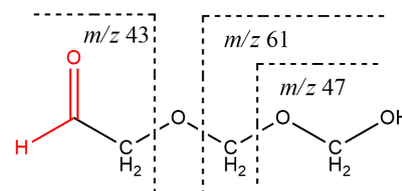
Figure 7. Temperature Programmed Desorption (TPD) spectra of the solid film formed after rare gas desorption step of the photolysis at 12 K ($\lambda > 120$ nm, 30 min) of $\text{H}_2\text{CO}/\text{Ar} = 2/1000$ matrix. m/z 43: CHOCH_2^+ ; m/z 47: HOCH_2O^+ ; m/z 60: $\text{HOCH}_2\text{CHO}^+$; m/z 61: $\text{HOCH}_2\text{CH}_2\text{O}^+/\text{HOCH}_2\text{OCH}_2^+$; m/z 62: $\text{HOCH}_2\text{CH}_2\text{OH}^+$. GA: m/z 61, 43; EG: m/z 62, 61, 43; POM: m/z 61, 47, 43; GCA: m/z 61, 43; GCO: m/z 61 43. GCA and GCO detections are only tentative.

easily assigned from its characteristic desorption around 210 K for m/z 62. These two products were previously detected using infrared spectroscopy. The third product detected by infrared spectroscopy POM is also confirmed by the QMS analysis from its characteristic fragments at m/z 47, 43, and 61 (Duvernay et al. 2014). It has to be noted that m/z 47 is only due to POM fragmentation whereas m/z 43 and 61 are also common fragments to other molecules such as GA, EG. This explain why the evolution of m/z 47 is different from the one observed for m/z 43 and 61. In ^{13}C experiment, m/z 43 becomes m/z 45 and m/z 47 becomes m/z 49. These matches different fragments of polyoxymethylene (POM). This is shown in Figure 8 that displays different fragmentation patterns for polyoxymethylene (POM) according to terminal groups. For example the fragment at m/z 43 indicates that the polymerisation process has been initiated/terminated by $\cdot\text{HCO}$ radical while the fragment at m/z 47 indicates $\cdot\text{CH}_2\text{OH}$ radical was the initiator/terminator. Also, an interesting point is their desorption temperature ranging from 180 K to 250 K (m/z 47). It depends on the length of the polymer and increases when the polymer grows (Duvernay et al. 2014). Two small peaks at 165 K and 180 K on the signal m/z 61 are coming from the desorption on unknown products.

The desorption peak starting at around 185 K for m/z 43 cannot be explained by the desorption of POM only. We also observe in this temperature range co-desorption with GA at 200 K (m/z 60) and EG at 210 K (m/z 62). These two species contribute to the m/z 43 signal observed between 180 K and 220 K since POM, GA, and EG share this mass fragment. Co-desorption of glyceraldehyde ($\text{HOCH}_2\text{CH}(\text{OH})\text{CHO}$) as reported by Fedoseev et al. (2017) cannot also be excluded since the desorption temperature for this compound is observed at 215 K and the m/z 43 and 61 are the most intensive peaks (Fedoseev et al. 2017). Thus, the co-desorption observed between 180 K and 220 K can be explained by the fact that GA, EG, and other COMs have been trapped in POM and are driven away from the sample holder during the POM desorption.

Finally, the last desorption peak is observed around 235 K for m/z 61, m/z 47, and m/z 43. The presence of m/z 47 indicates POM desorption but again other contribution cannot be excluded. Com-

Fragmentation pattern for POM with HCO end group



Fragmentation pattern for POM with CH₂OH end group

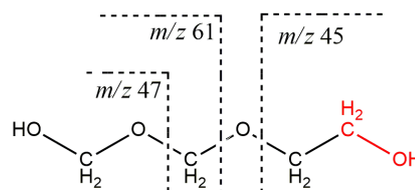


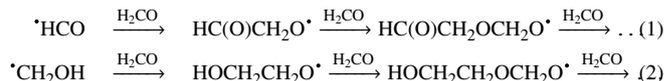
Figure 8. Polyoxymethylene fragmentation pattern according to end-groups.

plex molecules such as glycerol (GCO, $\text{HOCH}_2\text{CH}(\text{OH})\text{CH}_2\text{OH}$) may also contribute to the desorption peak observed at 235 K (Fedoseev et al. 2017) for m/z 61. The difficulty of mass assignments comes from the fact that EG, POM but also GCA, and GCO all share the same characteristic fragments m/z 43 and 61. The suspected formation of both GCA and GCO can be explained by the mechanism proposed by Fedoseev et al. (2017). Both GA and EG -two products observed in our experiments- can participate in H-induced reactions, producing two carbon bearing radical intermediates HOCH_2CHOH . This later, in turn, can react in a barrierless radical-radical recombination with $\cdot\text{HCO}$ or $\cdot\text{CH}_2\text{OH}$ radicals resulting in the formation of GCA and GCO respectively (Fedoseev et al. 2017). In a such mechanism, sugar formation (e.g. GCA and GCO) are competitive reactions with formaldehyde polymerisation (e.g. POM formation). However, more experiments are needed to confirm the GCA and GCO formation in our experiments.

Thus, after radical recombination, detected products are polyoxymethylene (POM), glycolaldehyde (GA), ethylene glycol (EG) and possibly glyceraldehyde (GCA) and glycerol (GCO).

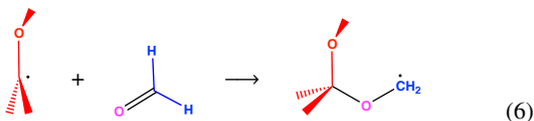
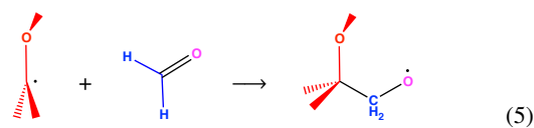
3.2 Theoretical results

In this section, we aim at providing a possible formation mechanism of POM, using state-of-the-art quantum calculations. We considered the two following polymerisation reactions :



For both reactions (1) and (2), the initiation step can *a priori* proceed via the two addition reactions : either a C-C bond (steps (3) and (5)) or a C-O bond (steps (4) and (6)) is formed.





3.2.1 Initiation steps for the $\text{·HCO/·CH}_2\text{OH} + \text{H}_2\text{CO}$ polymerisation

The results for the initiation steps (3), (4), (5) and (6) at the UCCSD(T)-F12 level are shown on Figure 9, where the reaction path for reaction (1) appears on the right-hand side of the figure and the one for reaction (2) appears on the left-hand side. For both reactions, the C–O bond formation appears to be much less favoured than the C–C bond formation, with barriers as high as $\approx 60 \text{ kJ}\cdot\text{mol}^{-1}$ for the former and up to $\approx 30 \text{ kJ}\cdot\text{mol}^{-1}$ for the latter. Interestingly, the formyloxymethyl radical $\text{HC(O)OCH}_2\cdot$ is thermodynamically more stable than the formylmethoxy radical $\text{HC(O)CH}_2\text{O}\cdot$ (the same is true although less pronounced for the $\text{·CH}_2\text{OH}$ counterpart). Under the current experimental conditions (@ 35 K) however, the "C-O" products are unlikely to be formed. Thus, only the initiation step (3) and (5) for reactions (1) and (2) respectively will be considered in the following. For this step, reaction (2) is more likely to happen than reaction (1) at low temperatures, with a barrier of $\approx 12 \text{ kJ}\cdot\text{mol}^{-1}$ for the former and $\approx 30 \text{ kJ}\cdot\text{mol}^{-1}$ for the latter. However, it is difficult to estimate if a barrier of $\approx 12 \text{ kJ}\cdot\text{mol}^{-1}$ is low enough under the experimental conditions (@ 35 K) so that the polymerisation reaction can be initiated. A comparison at the same level of calculation with the hydrogenation reaction of CO, known to occur at temperatures as low as 3 K (Pirim & Krim 2011) will be informative with regard to this question and will be discussed in section 3.2.3. Even if we were to assume that the initiation step was possible, one has to check if the next formaldehyde additions proceed through lower energy barriers than the first one, that is if the initiation step is the rate-determining step for the formaldehyde polymerisation reaction. A careful inspection of the propagation steps for both reactions (1) and (2) will help providing an answer.

3.2.2 Propagation steps for the $\text{·HCO/·CH}_2\text{OH} + \text{H}_2\text{CO}$ polymerisation

We calculated the reaction paths for the next two propagation steps of reactions (1) and (2). The reaction paths at both B3LYP and UCCSD(T)-F12 levels of theory are shown for each reaction in Figure 10 and Figure 11, respectively, and the corresponding energy values are reported in table 6. For each path, the reference energy corresponds to the sum of energies of the ·HCO (resp. $\text{·CH}_2\text{OH}$) radical and of $n \text{ H}_2\text{CO}$ moieties ($n \in [1 - 3]$) calculated separately, corrected by the ZPE at the B3LYP level. Each step of the reaction paths corresponds to the addition of a formaldehyde molecule. Among the possible conformers of the products, only the most stable one was kept for the reaction path.

As can be seen on Figure 10 and Figure 11, both DFT and coupled cluster calculations show the two following important results : (a) the polymerisation of formaldehyde initiated by both ·HCO and $\text{·CH}_2\text{OH}$ radicals is thermodynamically favourable and (b) the energy barriers of the propagation steps all lie lower (or nearly) than

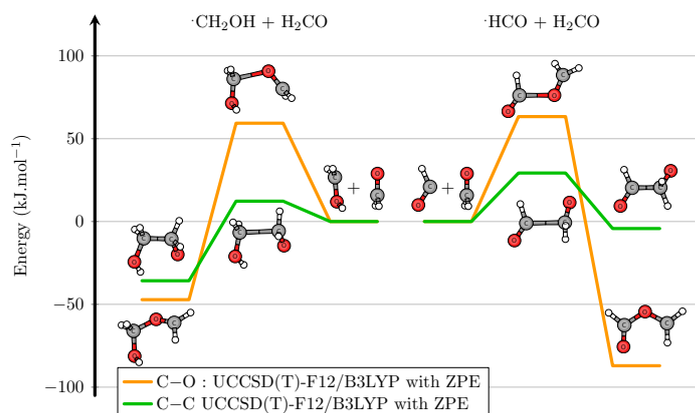


Figure 9. Left: $\text{·CH}_2\text{OH} + \text{H}_2\text{CO}$ reaction paths for the C–C bond formation (green) and C–O bond formation (yellow) in the initiation steps (5) and (6). Right: $\text{·HCO} + \text{H}_2\text{CO}$ reaction paths for the C–C bond formation (green) and C–O bond formation (yellow) in the initiation steps (3) and (4) respectively (See text).

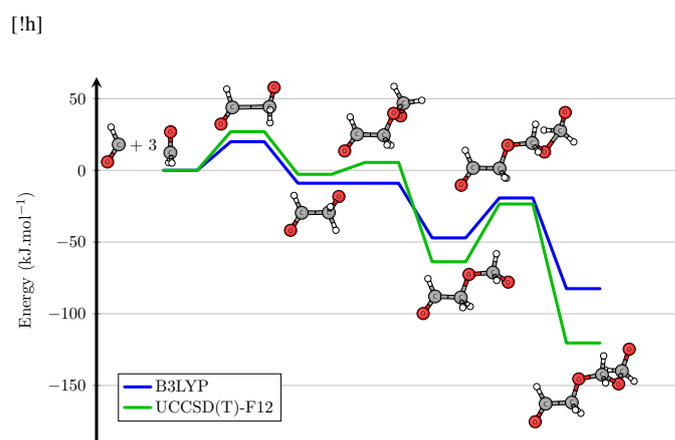


Figure 10. Reaction path of the initiation step (C–C bond formation) and the first two propagation steps for reaction (1) (See text).

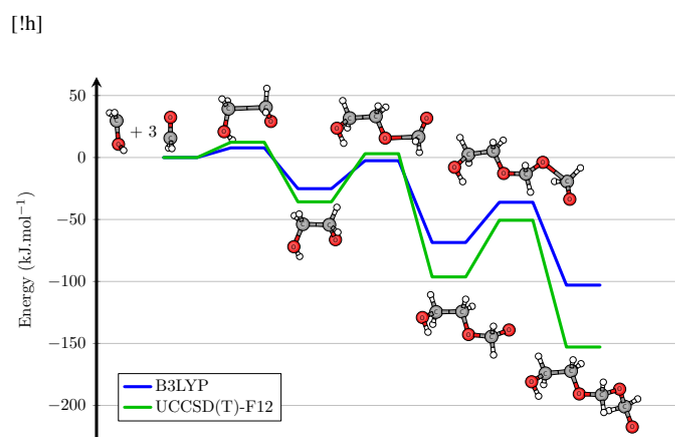


Figure 11. Reaction path of the initiation step (C–C bond formation) and the first two propagation steps for reaction (2) (See text).

[!h]

Table 6. B3LYP and UCCSD(T)-F12 energies (ZPE corrected, in $\text{kJ}\cdot\text{mol}^{-1}$) of reactants, transition states and products of the first three steps of polymerisation reactions of formaldehyde from $\cdot\text{HCO}$ and $\cdot\text{CH}_2\text{OH}$ radicals.

Species of reaction (1)	B3LYP ($\text{kJ}\cdot\text{mol}^{-1}$)	UCCSD(T)-F12 ($\text{kJ}\cdot\text{mol}^{-1}$)
$\cdot\text{HCO}+\text{H}_2\text{CO}$	0.00	0.00
TS1	+20.0	+27.0
$\text{HC}(\text{O})\text{CH}_2\text{O}\cdot$	-8.98	-2.77
TS2	-8.99	+5.52
$\text{HC}(\text{O})\text{CH}_2\text{OCH}_2\text{O}\cdot$	-47.1	-63.7
TS3	-19.3	-23.5
$\text{HC}(\text{O})(\text{CH}_2\text{O})_2\text{CH}_2\text{O}\cdot$	-82.5	-120.3
Species of reaction (2)		
$\cdot\text{CH}_2\text{OH}+\text{H}_2\text{CO}$	0.00	0.00
TS1	+7.77	+12.3
$\text{HOCH}_2\text{CH}_2\text{O}\cdot$	-25.2	-35.8
TS2	-2.56	+3.06
$\text{HOCH}_2\text{CH}_2\text{OCH}_2\text{O}\cdot$	-68.6	-96.3
TS3	-36.1	-50.6
$\text{HOCH}_2(\text{CH}_2\text{O})_2\text{CH}_2\text{O}\cdot$	-102.9	-152.9

the initial energy of the reactants, showing that the rate-determining step for the polymerisation is namely the initiation step. If the first barrier can be overcome, the polymerisation reaction would proceed and would be driven by the formation of even more thermodynamically stable species, which is consistent with the detection of POM in our experiments. Before focusing on the rate-determining step, it is interesting to analyse the differences between the DFT and coupled cluster results on one hand and on the differences between both reaction paths on the other hand, from the values of Table 6.

For both reactions, the DFT method underestimates energy barriers as well as exothermicities compared to the coupled cluster ones. The only exception regarding exothermicities appears for the formylmethoxy radical $\text{HC}(\text{O})\text{CH}_2\text{O}\cdot$, which is found to be about $6 \text{ kJ}\cdot\text{mol}^{-1}$ less stable and almost isoenergetic with the reactants at the coupled cluster level. Apart from this case, the differences between DFT and coupled cluster values are particularly well pronounced for the second propagation step with a difference in the energy barrier up to $14 \text{ kJ}\cdot\text{mol}^{-1}$ and in the exothermicity up to $22 \text{ kJ}\cdot\text{mol}^{-1}$ for reaction (2). At the UCCSD(T)-F12 level, the energy release of each propagation step remains more or less constant ($\approx 60 \text{ kJ}\cdot\text{mol}^{-1}$), and this for each reaction, whereas they range from 35 to $43 \text{ kJ}\cdot\text{mol}^{-1}$ at the DFT level.

Regarding the differences between the two reaction paths, both methods predict more stable products in reaction (2) than in reaction (1): a plausible explanation is the possibility of an hydrogen bond between the O-H group brought by the hydroxymethyl radical and the oxygen of formaldehyde, which stabilises the newly formed radical and is absent in the case of the formyl radical. This is particularly obvious for the hydroxymethylmethoxy radical $\text{HOCH}_2\text{CH}_2\text{O}\cdot$ which is stabilised by $\approx 36 \text{ kJ}\cdot\text{mol}^{-1}$ with respect to its reactant, whereas the formylmethoxy radical $\text{HC}(\text{O})\text{CH}_2\text{O}\cdot$ is only stabilised by $\approx 3 \text{ kJ}\cdot\text{mol}^{-1}$. Another feature of both reaction paths is that the first energy barrier is lower for reaction (2), indicating that this reaction could possibly be the dominant mechanism for the formation of POM.

[!h]

Table 7. B3LYP and UCCSD(T)-F12 energies (ZPE corrected, in $\text{kJ}\cdot\text{mol}^{-1}$) of reactants, transition states and products of the hydrogenation reaction of CO

	B3LYP ($\text{kJ}\cdot\text{mol}^{-1}$)	UCCSD(T)-F12 ($\text{kJ}\cdot\text{mol}^{-1}$)
H+CO	0.00	0.00
TS	1.88	11.2
$\cdot\text{HCO}$	-85.3	-56.5

3.2.3 Interpretation of the experimental results

The calculated reaction paths for both reactions (1) and (2) have shown that these reactions are good candidates to explain the polymerisation of formaldehyde into POM, provided the first energy barrier can be overcome. Since the DFT method is known to underestimate energy barriers, we will refer in the following discussion to the more reliable values obtained by the UCCSD(T)-F12 method. The lowest barrier was found to be $12.3 \text{ kJ}\cdot\text{mol}^{-1}$ for the addition of formaldehyde on the hydroxymethyl radical (See Table 6). In order to estimate if this barrier can be overcome at 35 K, one can compare it to the barrier of the elementary reaction of hydrogenation of CO. Pirim and Krim (Pirim & Krim 2011) have recently shown that the reaction of non-energetic H-atoms with CO can occur to form formyl radicals at temperatures as low as 3 K. We calculated the energy barrier for this reaction at DFT and UCCSD(T)-F12 levels and the results are reported in Table 7. As can be seen, if one would trust the DFT values, one would conclude that the small energy barrier of $1.88 \text{ kJ}\cdot\text{mol}^{-1}$ and the high exothermicity are compatible with the experimental observations. The UCCSD(T)-F12 energy barrier is however much higher ($11.2 \text{ kJ}\cdot\text{mol}^{-1}$) and very close to the energy barrier of reaction 2. If one trusts the coupled cluster value, this gives an indication that such a $\approx 10 \text{ kJ}\cdot\text{mol}^{-1}$ barrier is obviously not contradictory with the fact that this reaction efficiently occurs at very low temperature. Hence, one can put forward the following hypotheses to explain the formation of POM:

- There is a possible "cage effect" of the desorbing Ar atoms that could artificially stabilise the first transition state and thus lower the reaction barrier. It seems however difficult to support experimentally this idea by probing the Ar matrix at its desorbing temperature in order to have an idea on the local environment of the reactants.
- Tunneling effects are known to be important at very low temperature, mainly for light species. In the case of the hydrogenation of CO, the tunneling of H-atom certainly enhances the reactivity. In our case, tunneling effect would correspond to a C-C bond formation. The imaginary frequencies of the transition states, $263i \text{ cm}^{-1}$ and $286i \text{ cm}^{-1}$ for the initiation steps of reactions (1) and (2), respectively, indicate that, in the framework of the simple standard Eckart model, the barriers are not thin enough for the tunneling to be important. However, important tunneling effects for heavy atoms are known to occur at very low temperature (Borden 2016; Meisner & Kästner 2016; Goumans & Andersson 2010) and tunneling has recently been recognised as the third reactivity paradigm (Schreiner 2017). It would certainly be interesting to investigate further the role of tunneling in the POM formation in the light of these recent findings.
- More likely, the energy release from the first step of reaction (2)

or from a fast radical-radical recombination reaction to form EG or GA could possibly be transferred to the reactants of reaction (1) and enable the barrier to be overcome.

Finally, the termination step can occur either with a recombination with a $\cdot\text{HCO}$ or $\cdot\text{CH}_2\text{OH}$ radical. This recombination step between two radicals is expected to proceed without any barrier and should thus be fast. The length of the POM chain would then depend on the amount of the remaining radicals as the Ar desorption is fully over. This is consistent with the mass spectra recorded between 150 and 230 K (Fig. 7) that clearly shows chain-length distribution as well as both $\cdot\text{HCO}$ and $\cdot\text{CH}_2\text{OH}$ end groups.

4 ASTROCHEMICAL IMPLICATIONS

The presence of polyoxymethylene in comets is an exciting story that started in 1986. During the Halley comet flyby by the *Giotto* mission, the PICCA instrument (Positive Ion Cluster Composition Analyser) recorded mass spectra displaying a periodic pattern with repeating units of m/z 16:14 consistent with an $-\text{O}-$ and $-\text{CH}_2-$ addition/loss. At that time, this pattern was identified as being compatible with formaldehyde polymers, also referred to as polyoxymethylene (POM) (Huebner 1987). Later, Mitchell et al. (1989) showed that this regular pattern is generally characteristic of any kind of CHO-bearing molecules that include POM-like structures. Recently, a similar pattern was found by the Ptolemy instrument of the *Rosetta* mission (Wright et al. 2015) strengthening the POM detection in cometary nucleus. However, this is to be tempered since recent re-investigation of Ptolemy data and comparison with data from two other mass spectrometers on-board the *Rosetta* mission, COSAC and ROSINA suggest that the detection of polyoxymethylene by Ptolemy instrument cannot be confirmed (Altwegg et al. 2017). They calibrated the ROSINA flight spare instrument with commercially available polymer. POM was thermally desorbed in a vacuum chamber and then connected to ROSINA flight spare instrument. From the fragmentation pattern they observed, they concluded that thermal desorption of POM mostly produce monomer (m/z 28 30), and only very little dimer or longer chains and it does not produce the observed periodic pattern with repeating units of m/z 16:14 as observed on Ptolemy instrument. In addition, the ROSINA results obtained in laboratory are not compatible with the suggestion made by Wright et al. (2015) that POM is responsible for high peak on m/z 91 and peaks between m/z 55 and 62. We would like to make comments on these results. It is well known that commercially available POM thermally decompose at temperature higher than 40°C into formaldehyde. Commercially available POMs mainly consist of long chain-length polymers that cannot sublime easily. Instead, at temperature higher than 40°C, they start to depolymerise into H_2CO monomer (Duvernay et al. 2014; Le Roy et al. 2012). However, the mass spectra of formaldehyde oligomers (*i.e.* short chain-length POMs) are really different from the ones of commercially available POMs and present high similarities with data recorded by Ptolemy instrument supporting POM identification in cometary nuclei made by both PICCA and Ptolemy instruments (Duvernay et al. 2014). Results from this study also bring some new elements that strengthen the likely formation of POM on cometary nuclei based on formaldehyde polymerisation initiated by radicals. The figure 12 displays the mass spectra comparison between the POM desorbing from our sample holder at 260 K with some peaks extracted from the Ptolemy spectra (Wright et al. 2015). Those peaks at m/z 31, 43, 45, 47, 57, 59, 61, 73 have been proposed to be polyoxymethylene (Wright et al. 2015) originating from the surface of the comet 67P.

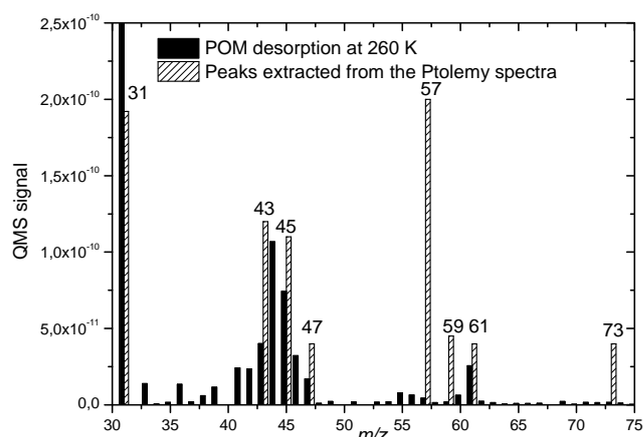


Figure 12. Mass spectrum of POM recorded at 260 K compared with extracted peaks from the Ptolemy spectra. m/z 44 is related to CO_2 contamination.

These data fit well the mass spectra of volatile POM recorded at 260 K except three masses at m/z 57, 59, and 73. Mass spectra recorded by Ptolemy instrument do not reflect a single idealized compounds with its ends terminated by H-atoms. Rather they indicate a number of different terminations with chain running as $-\text{CH}_2\text{O}-$. For instance, peaks at m/z 57, 59 and 73 on Ptolemy spectrum result POM fragmentation with $-(\text{CO})\text{CH}_3$ termination that is unlikely in our conditions since the only possible terminations are $-\text{CHO}$ or $-\text{CH}_2\text{OH}$. This may explain the absence of peaks at m/z 57, 59, and 73 on our sample. The question therefore remains whether POMs are an important component of comets and more laboratory/theoretical studies and space missions will be needed to close this passionate debate.

5 CONCLUSION

We report new laboratory experiments on the low-temperature solid state formation of formaldehyde polymer polyoxymethylene through radical-induced reactivity from VUV photolysis of formaldehyde in Ar matrix. We show, based on experimental finding and quantum calculations, the ability of free radicals to react at low temperature with others on the double bonds of unsaturated molecules initiating COMs formation such as GA, EG, and formaldehyde polymerisation. This formation mechanism based on radical-induced reactivity is likely in interstellar ices or cometary grains and supports the presence of COMs and POM in cometary nuclei. In addition the fragmentation pattern we observe with mass spectroscopy for short chain-length POM polymers is consistent with the data collected by the Ptolemy instrument on-board the *Rosetta* mission.

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REFERENCES

- Abe H., Takeo H., Yamada K. M., 1999, *Chemical physics letters*, 311, 153
- Abou Mrad N., Duvernay F., Theulé P., Chiavassa T., Danger G., 2014, *Analytical chemistry*, 86, 8391
- Abou Mrad N., Duvernay F., Chiavassa T., Danger G., 2016, *Monthly Notices of the Royal Astronomical Society*, 458, 1234
- Altwegg K., et al., 2017, *Monthly Notices of the Royal Astronomical Society*, 469, S130
- Becke A., 1993, *J. Chem. Phys.*, 98, 5648
- Bennett C. J., Kaiser R. I., 2007, *The Astrophysical Journal*, 661, 899
- Biver N., et al., 2014, *Astronomy & Astrophysics*, 566, L5
- Borden W. T., 2016, *Wiley Interdisciplinary Reviews: Computational Molecular Science*, 6, 20
- Butscher T., Duvernay F., Theule P., Danger G., Carissan Y., Hagebaum-Reignier D., Chiavassa T., 2015, *Monthly Notices of the Royal Astronomical Society*, 453, 1587
- Butscher T., Duvernay F., Danger G., Chiavassa T., 2016, *Astronomy & Astrophysics*, 593, A60
- Butscher T., Duvernay F., Rimola A., Segado-Centellas M., Chiavassa T., 2017, *Physical Chemistry Chemical Physics*, 19, 2857
- Caro G. M., et al., 2002, *Nature*, 416, 403
- Chuang K.-J., Fedoseev G., Ioppolo S., van Dishoeck E., Linnartz H., 2015, *Monthly Notices of the Royal Astronomical Society*, 455, 1702
- Chuang K.-J., Fedoseev G., Qasim D., Ioppolo S., van Dishoeck E., Linnartz H., 2017, *Monthly Notices of the Royal Astronomical Society*, 467, 2552
- Cottin H., Fray N., 2008, *Space Science Reviews*, 138, 179
- Cottin H., Moore M. H., Bénilan Y., 2003, *The Astrophysical Journal*, 590, 874
- Crovisier J., Bockelée-Morvan D., Biver N., Colom P., Despois D., Lis D. C., 2004, *Astronomy & Astrophysics*, 418, L35
- Danger G., Rimola A., Mrad N. A., Duvernay F., Roussin G., Theule P., Chiavassa T., 2014, *Physical Chemistry Chemical Physics*, 16, 3360
- Deglmann P., May K., Furche F., Ahlrichs R., 2004, *Chemical physics letters*, 384, 103
- Dunning T. H., 1989, *The Journal of Chemical Physics*, 90, 1007
- Duvernay F., Danger G., Theulé P., Chiavassa T., Rimola A., 2014, *The Astrophysical Journal*, 791, 75
- Fedoseev G., Cuppen H., Ioppolo S., Lamberts T., Linnartz H., 2015, *Monthly Notices of the Royal Astronomical Society*, 448, 1288
- Fedoseev G., Chuang K.-J., Ioppolo S., Qasim D., van Dishoeck E. F., Linnartz H., 2017, *The Astrophysical Journal*, 842, 52
- Fray N., Bénilan Y., Biver N., Bockelée-Morvan D., Cottin H., Crovisier J., Gazeau M.-C., 2006, *Icarus*, 184, 239
- Fuchs G., Cuppen H., Ioppolo S., Romanzin C., Bisschop S., Andersson S., van Dishoeck E., Linnartz H., 2009, *Astronomy & Astrophysics*, 505, 629
- Goesmann F., et al., 2015, *Science*, 349, aab0689
- Goumans T. P. M., Andersson S., 2010, *Monthly Notices of the Royal Astronomical Society*, 406, 2213
- Hättig C., Klopper W., Köhn A., Tew D. P., 2012, *Chemical Reviews*, 112, 4
- Henderson B. L., Gudipati M. S., 2015, *The Astrophysical Journal*, 800, 66
- Hiraoka K., Yamashita A., Yachi Y., Aruga K., Sato T., Muto H., 1995, *The Astrophysical Journal*, 443, 363
- Hudson R. L., Moore M. H., Cook A. M., 2005, *Advances in Space Research*, 36, 184
- Huebner W., 1987, *Science*, 237, 628
- Jacox M. E., 1988, *The Journal of chemical physics*, 88, 4598
- Jacox M. E., Milligan D. E., 1973, *Journal of Molecular Spectroscopy*, 47, 148
- Kaiser R. I., Maity S., Jones B. M., 2015, *Angewandte Chemie International Edition*, 54, 195
- Kunttu H., Seetula J., Räsänen M., Apkarian V., 1992, *J. Chem. Phys.*, 96, 5630
- Le Roy L., Briani G., Briois C., Cottin H., Fray N., Thirkell L., Poulet G., Hilchenbach M., 2012, *Planetary and space science*, 65, 83
- Lee Y.-F., Chou W.-T., Johnson B. A., Tabor D. P., Sibert III E. L., Lee Y.-P., 2015, *Journal of Molecular Spectroscopy*, 310, 57
- Meisner J., Kästner J., 2016, *Angewandte Chemie International Edition*, 55, 5400
- Milligan D. E., Jacox M. E., 1969, *J. Chem. Phys.*, 51, 277
- Minissale M., Moudens A., Baouche S., Chaabouni H., Dulieu F., 2016, *Monthly Notices of the Royal Astronomical Society*, 458, 2953
- Mitchell D., et al., 1989, *Advances in Space Research*, 9, 35
- Nelander B., 1980, *J. Chem. Phys.*, 73, 1026
- Noble J., Theule P., Mispelaer F., Duvernay F., Danger G., Congiu E., Dulieu F., Chiavassa T., 2012, *Astronomy & Astrophysics*, 543, A5
- Öberg K. I., Garrod R. T., Van Dishoeck E. F., Linnartz H., 2009, *Astronomy & Astrophysics*, 504, 891
- Pettersson M., Khriachtchev L., Roozeman R.-J., Räsänen M., 2000, *Chemical Physics Letters*, 323, 506
- Pirim C., Krim L., 2011, *Phys. Chem. Chem. Phys.*, 13, 19454
- Ryazantsev S. V., Feldman V. I., 2014, *The Journal of Physical Chemistry A*, 119, 2578
- Schreiner P. R., 2017, *Journal of the American Chemical Society*, 139, 15276
- Schrivier A., Schriver-Mazzuoli L., Viginas A. A., 2000, *Vibrational Spectroscopy*, 23, 83
- Schutte W., Allamandola L., Sandford S., 1993, *Icarus*, 104, 118
- Stein C., Weser O., Schröder B., Botschwina P., 2015, *Molecular Physics*, 113, 2169
- Tew D. P., Klopper W., 2010, *Molecular Physics*, 108, 315
- Vinogradoff V., Duvernay F., Danger G., Theulé P., Chiavassa T., 2011, *Astronomy & Astrophysics*, 530, A128
- Watanabe N., Shiraki T., Kouchi A., 2003, *The Astrophysical Journal Letters*, 588, L121
- Watanabe N., Nagaoka A., Shiraki T., Kouchi A., 2004, *The Astrophysical Journal*, 616, 638
- Weigend F., Ahlrichs R., 2005, *Physical Chemistry Chemical Physics*, 7, 3297
- Woods P. M., Slater B., Raza Z., Viti S., Brown W. A., Burke D. J., 2013, *The Astrophysical Journal*, 777, 90
- Wright I., Sheridan S., Barber S., Morgan G., Andrews D., Morse A., 2015, *Science*, 349, aab0673