

20 **ABSTRACT**

21

22 We have used the Titan Haze Simulation (THS) experimental set-up at NASA Ames' COSmIC
23 facility to produce four laboratory analogs of Titan's aerosols, or tholins. These tholin samples
24 were produced from four gas mixtures, representative of Titan's atmosphere, of initial
25 compositions $\text{N}_2:\text{CH}_4$ (90:10 and 95:5) and $\text{N}_2:\text{CH}_4:\text{C}_2\text{H}_2$ (89.5:10:0.5 and 94.5:5:0.5) at 150 K,
26 using a plasma discharge in the stream of a jet-cooled gas expansion. Here we present an ex-situ
27 X-ray absorption near-edge structure (XANES) spectroscopy analysis of these four tholin samples.
28 The C- and N-XANES spectra show the presence of various functional groups (aromatic carbon,
29 imines, nitriles, etc.) whose abundances are correlated with (i) the relative proportions between N_2
30 and CH_4 , and (ii) the presence or absence of C_2H_2 in the initial mixtures. In particular, mixtures
31 containing C_2H_2 result in the formation of tholins consisting of more aromatic structures and
32 displaying larger relative amounts of imines and nitriles, but lower overall nitrogen content.
33 XANES spectroscopy also allowed for the determination of the elemental C/N ratios for the four
34 tholins, which were all found to be low (< 2.5) and consistent with those measured for tholins
35 produced in other, independent experimental set-ups. The C/N ratios of tholins produced from gas
36 mixtures that contained C_2H_2 (2.2–2.4) were found to be about twice as large as those of tholins
37 produced from gas mixtures that did not contain C_2H_2 (0.9–1.3). These new experimental results
38 from the COSmIC/THS experiment demonstrate the impact initial precursors have on the nitrogen
39 chemistry during the formation of tholins, which may help in the interpretation of observations
40 and the development of models of Titan's atmosphere.

41

42 *Keywords:*

43 Titan, atmosphere

44 Tholins

45 Organic materials

46 XANES

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48

49 1. Introduction

50

51 Titan, the largest moon of Saturn, is the only other object in the Solar System besides Earth that
52 possesses a dense atmosphere (1.5 bar at the surface) whose composition is dominated by
53 molecular nitrogen (N_2 ; 95–98%), with methane (CH_4 ; 2–5%) as the second most abundant gas
54 (Hanel et al., 1981; Strobel and Shemansky, 1982; Lindal et al., 1983; Fulchignoni et al., 2005;
55 Niemann et al., 2010). A complex organic chemistry between N_2 and CH_4 , induced by solar
56 ultraviolet (UV) photons and energetic particles, results in the formation of a large variety of
57 organic compounds that include hydrocarbons such as acetylene (C_2H_2), ethylene (C_2H_4), ethane
58 (C_2H_6), propyne (C_3H_4 ; also called methylacetylene), benzene (C_6H_6), nitriles such as hydrogen
59 cyanide (HCN), hydrogen isocyanide (HNC), cyanoacetylene (HC_3N), acetonitrile (CH_3CN), ethyl
60 cyanide (C_2H_5CN), and cyanogen (C_2N_2), as well as larger, more complex molecules that are
61 believed to include aromatic and heteroaromatic compounds such as propane (C_3H_8), butane
62 (C_4H_{10}), and polycyclic aromatic hydrocarbons (see Hörst, 2017; Barnes et al., 2021; and
63 references therein for a complete review).

64 This complex organic chemistry taking place in the atmosphere also leads to the production
65 of solid particles that form haze layers and play an important role in Titan's atmospheric dynamics,
66 climate, as well as surface composition and processes. Because of its N_2 -based dense atmosphere
67 and its CH_4 cycle that resembles the hydrological cycle on Earth, Titan is often compared to the
68 early Earth (Trainer et al., 2004; Lunine and Atreya, 2008; Roe, 2012; He and Smith, 2014). In
69 recent years, Titan's atmosphere and surface were monitored in detail by the NASA *Cassini*
70 mission, which orbited the Saturnian system from 2004 to 2017 (Matson et al., 2002, 2003;

71 Mitchell, 2006), as well as the ESA *Huygens* probe, which plunged into Titan's atmosphere and
72 landed on its surface in 2005 (Lebretron et al., 2005; Owen, 2005; Zarnecki et al., 2005).

73 Many laboratory studies have simulated Titan's organic atmospheric chemistry in the past
74 four decades, using various energy sources to induce the dissociation and ionization of N_2 and CH_4 .
75 These experiments result in the formation of refractory organic materials that are analogs of Titan's
76 aerosols, often referred to as 'tholins' (Sagan and Khare, 1979; Cable et al., 2012; Coll et al., 2013;
77 He et al., 2017; Hörst, 2017; Sciamma-O'Brien et al., 2017; Sebree et al., 2018). Tholins have
78 been studied using a wide variety of techniques to determine their chemical compositions as well
79 as their physical, chemical, and optical properties (e.g., Cable et al., 2012; Raulin et al., 2012;
80 Brassé et al., 2015; Maillard et al., 2021; Schultz et al., 2021; and references therein). Among other
81 findings, these studies have shown that tholins are macromolecular materials consisting mainly of
82 carbon, nitrogen, and hydrogen, distributed among a mixture of aliphatic and aromatic structures.

83 In the present study, we produced tholins using the Titan Haze Simulation (THS)
84 experimental set-up at NASA Ames' COsmic SIMulation Chamber (COSmIC) facility from four
85 initial gas mixtures that contained mainly N_2 , together with 5% or 10% of CH_4 , with or without
86 the addition of 0.5% of C_2H_2 (see Section 2.1). We then analyzed these tholins ex situ using X-ray
87 absorption near-edge structure (XANES) spectroscopy to obtain information on their chemical
88 compositions and elemental C/N ratios. After describing the experimental protocol and analytical
89 technique to produce and analyze our tholin samples, we compare our findings with those derived
90 for tholins produced in other, independent experimental studies. Finally, we also describe how
91 these new experimental results may have implications for the interpretation of observational data
92 of Titan's atmosphere, especially during seasonal changes that result in the modification of the
93 relative abundances of specific species.

94

95 **2. Experimental Methods**

96

97 *2.1. Production of tholins with COSmIC/THS*

98

99 Tholin samples were produced using the COSmIC/THS experimental set-up, whose mode of
100 operation has been described in detail in previous publications (Sciamma-O'Brien et al., 2014,
101 2017; Salama et al., 2018). Briefly, a supersonic, adiabatic expansion is used to (i) cool down a
102 gas mixture, flowing at 2,000 standard cubic centimeters per minute (sccm), to Titan-like
103 temperatures (150 K), and (ii) decrease the gas pressure to 30 mbar (Broks et al., 2005; Biennier
104 et al., 2006). A plasma discharge (typically, -700 to -1000 V) generated in the stream of that
105 expansion is then used as an energy source to induce chemistry between the different molecular
106 species present in the jet-cooled gas mixture (Sciamma-O'Brien et al., 2014), hence simulating the
107 electron bombardment of Titan's atmosphere when passing through Saturn's magnetosphere.
108 Although Titan's atmospheric chemistry is induced by both photolysis and radiolysis, it is
109 experimentally advantageous to use a plasma discharge as the energy source instead of a UV lamp,
110 as it allows to dissociate N_2 , hence simulating the nitrogen chemistry taking place in Titan's
111 atmosphere.

112 Resulting from this chemistry, larger species are formed in the gas phase, eventually
113 leading to the formation of solid particles that can be jet-deposited onto substrates placed in the
114 stream of the expansion. These solid particles, or tholins, are considered analogs of Titan's aerosols,
115 which result from similar processes occurring in Titan's upper atmosphere. Figure 1 shows (a) a
116 schematic of the pulsed discharge nozzle, which is at the core of the COSmIC/THS chamber and

117 is used to generate the pulsed planar plasma expansion (adapted from Broks et al., 2005), and (b)
118 a photograph of the COSmIC/THS pink N₂-based plasma expansion during solid sample
119 production (note: this picture shows an experiment in which a KBr window and two silicon wafers
120 placed 5 cm from the slit were used as substrates for the deposition). Previous studies have
121 demonstrated the effect of different gaseous precursors present in the initial gas mixture on the
122 formation of both gas-phase (Sciamma-O'Brien et al., 2014; Raymond et al., 2018) and solid-
123 phase (Sciamma-O'Brien et al., 2017) products in the COSmIC/THS chamber, particularly in the
124 cases where acetylene (C₂H₂) is added to N₂:CH₄ gas mixtures, which are traditionally used to
125 simulate Titan's atmospheric chemistry.

126 In the present study, tholins were produced from four different gas mixtures, namely,
127 N₂:CH₄ (90:10), N₂:CH₄:C₂H₂ (89.5:10:0.5), N₂:CH₄ (95:5), and N₂:CH₄:C₂H₂ (94.5:5:0.5). The
128 abundances of CH₄ relative to N₂ in the gas mixtures chosen for our experiments are representative
129 of the upper limit or higher than the relative CH₄ abundance observed with *Cassini* and *Huygens*
130 instruments as well as ground-based and space observatories (Hörst, 2017, and references therein).
131 Nevertheless, studying these gas mixtures present a couple of advantages: (i) these gas mixtures
132 have been used extensively in previous laboratory studies to simulate Titan's atmosphere, and the
133 results obtained here can therefore be compared with a large number of other experiments,
134 including those performed in our laboratory (Sciamma-O'Brien et al., 2014, 2017), and (ii) due to
135 the truncated chemistry occurring in our experimental set-up, starting with a higher relative
136 abundance of CH₄ allows for higher tholin production yield and therefore thicker samples, which
137 results in higher-quality analytical measurements. Finally, the purpose of adding 0.5% of C₂H₂ in
138 half of the gas mixtures is to study chemical processes which occur at later stages of the chemistry
139 that leads to the tholin formation in COSmIC, as the residence time of the gases in the chamber in

140 the plasma discharge is short and truncates the chemistry (see below). The addition of C₂H₂ also
141 allowed us to compare our results with previous experiments (Sciamma-O'Brien et al., 2014, 2017).

142 The gas mixtures were introduced into the COSmIC chamber, exposed to a pulsed plasma
143 discharge generated with a -800-V high voltage used as the energy source to induce the chemistry,
144 and deposited for 1–2 hours (to ensure very little material was deposited; see Section 3.5) onto
145 silicon nitride (Si₃N₄) substrates placed 5 cm away from the slit, i.e., at the same distance as the
146 substrates shown in Fig. 1(b). The voltage chosen for these experiments, which is higher than the
147 voltage used in previous experiments (Sciamma-O'Brien et al., 2014, 2017), allowed for a higher
148 production of grains in a shorter amount of time. **We expect the chemistry to be slightly different**
149 **with -800 V than with -700 V, however, this does not impact the results of this study whose**
150 **main goal is to compare 4 tholin samples produced under the same experimental conditions**
151 **at -800 V.** The solid deposits were then collected in a glove box connected to the vacuum chamber
152 and stored under an argon atmosphere in sealed containers with desiccant until further analysis.
153 The four tholin samples were therefore exposed to air only at the time of their characterization by
154 XANES spectroscopy, as described below.

155

156 2.2. XANES analysis of tholins

157

158 X-ray absorption near-edge structure (XANES) spectroscopy is a synchrotron-based soft X-ray
159 micro-analytical technique that can be used to assess both the functional groups present in a sample
160 and the relative elemental abundances between the elements studied. XANES spectroscopy at the
161 carbon (C), nitrogen (N), and oxygen (O) 1s edges has been extensively used to study organic
162 compounds in extraterrestrial samples such as meteorites, interplanetary dust particles (IDPs), and

163 cometary grains returned by the NASA Stardust mission (Bajt et al., 1996; Keller et al., 2004;
164 Sandford et al., 2006; Cody et al., 2008; Wirick et al., 2009; Matrajt et al., 2010; De Gregorio et
165 al., 2013). XANES spectroscopy has also been used to study laboratory refractory residues
166 produced from the simultaneous deposition and UV irradiation of astrophysical ice mixture
167 analogs (H_2O , CH_3OH , CO , NH_3 , etc.) at low temperature (< 20 K) (Nuevo et al., 2011), as well
168 as laboratory refractory organic materials produced from the UV irradiation or 1.2-keV electron
169 bombardment of $\text{N}_2:\text{CH}_4:\text{CO}$ (100:1:1) ice mixtures at 15 K, analog to the surface materials on
170 Pluto and other trans-Neptunian objects (TNOs) and hereafter referred to as Pluto/TNO tholins
171 (Materese et al., 2014, 2015). However, prior to this study, XANES spectroscopy analysis of Titan
172 tholins has been reported only once for a sample produced from a $\text{N}_2:\text{CH}_4$ (95:5) gas mixture in
173 another experimental set-up that uses a radiofrequency (RF) plasma to induce chemistry (Kuga et
174 al., 2014). XANES spectroscopy is thus well suited for the study of the Titan tholins produced
175 with COSMIC/THS, as the tholin production rate is extremely low in our experiments, which
176 makes it difficult to use more commonly used combustion methods to obtain information on their
177 elemental composition.

178 The tholin samples produced in this study were deposited on Si_3N_4 windows, which are
179 suitable for XANES measurements. XANES measurements were conducted using the scanning
180 transmission X-ray microscope (STXM) mounted on beamline 5.3.2.2 of the Advanced Light
181 Source of the Lawrence Berkeley Laboratory. The STXM provides photons in the 250–700 eV
182 range with a 10^7 photons s^{-1} flux (Kilcoyne et al., 2003). XANES spectra of the tholin samples
183 were acquired in the 275–580 eV energy range in order to span the carbon (275–340 eV), nitrogen
184 (385–440 eV), and oxygen (520–580 eV) 1s X-ray absorption edges, with an energy step size of
185 0.1 eV in the fine structure regions, and 1–2 eV in the pre- and post-edge regions (Cody et al.,

186 2008; Nuevo et al., 2011). All STXM data were reduced and analyzed using the aXis 2000
187 software.¹

188 The C-, N-, and O-XANES spectra provided information on the functional groups present
189 in the tholin samples. In addition, the XANES spectra measured over the full 275–580 eV energy
190 range can be used to derive the relative elemental abundances between C, N, and O for each tholin
191 sample. This is done by computationally fitting the full-range XANES spectra of all tholins
192 spanning the C, N, and O 1s edges with the atomic mass absorption coefficients (Henke et al.,
193 1993) to estimate the overall increase in absorption above the ionization C, N, and O edges relative
194 to the absorption below these ionization edges. This is possible for low-Z elements, for which
195 absorption in the soft X-ray range is dominated by photo-ionization cross sections, and for which
196 atomic scattering is very weak (Stöhr, 1992). These abundances were then used to determine the
197 C/N and C/O ratios for each tholin sample, using the approach described in Cody et al. (2008) and
198 Nuevo et al. (2011).

199

200 **3. Results and Discussion**

201

202 *3.1. Carbon XANES spectra*

203

204 The normalized C-XANES spectra of all four tholins (Fig. 2) show the presence of a variety of
205 functional groups. **Table 1 lists the main bands present in all tholin samples. They are**
206 **associated with** aromatic carbon (band C1 at ~285.2 eV), imines (C=N; band C2 at ~285.9 eV),
207 and nitriles (C≡N; band C3 at ~286.8 eV) (see Fig. 2 and Table 1). The relative proportions

¹ aXis 2000 is written in Interactive Data Language (IDL) and available for free for non-commercial use from
<http://unicorn.mcmaster.ca/aXis2000.html>.

208 between these bands and the rest of the spectra vary significantly from one sample to another,
209 although they appear to be more intense (relative to other bands) for tholins produced from the two
210 gas mixtures containing acetylene (C_2H_2) (Fig. 2). Higher formation yields for aromatic
211 compounds when acetylene is present are consistent with (i) previous mass spectrometry
212 experiments conducted on COSmIC which showed that the addition of C_2H_2 in gas mixtures in
213 which Ar or N_2 is the main component leads to the production of benzene in the gas phase
214 (Contreras and Salama, 2013; Sciamma-O'Brien et al., 2014), and (ii) the fact that C_2H_2 is a known
215 precursor of benzene in astrophysical environments (Frenklach and Feigelson, 1989; Cherchneff
216 et al., 1992; Jäger et al., 2009; Zhao et al., 2016). Two other bands, assigned to amide groups
217 and/or benzene (band C4 at 287.8–288.3 eV) and amorphous carbon, alcohols, and/or urea groups
218 (band C5 at 288.9–289.7 eV), are also present in all tholins with varying intensities. Note that if a
219 band is assigned to more than one functional group or structure, it may indicate the presence of
220 either only one of these groups or several of them. In the case of the band C4, the presence of
221 amide groups is supported by the band O3 (see Fig. S1 and Table S3 in the Supplementary
222 Material). However, the contribution of benzene to this band, and thus its presence in the tholins,
223 should not be ruled out, in particular in the tholins produced from the gas mixtures that contained
224 acetylene, because acetylene is a well-known precursor to benzene (Frenklach and Feigelson, 1989;
225 Cherchneff et al., 1992; Jäger et al., 2009; Zhao et al., 2016). Similarly, although the presence of
226 alcohols and/or urea groups (band C5) is supported by the band O4 (alcohols) and a band at 532.1–
227 532.2 eV (carboxyls) in Fig. S1 and Table S3, the presence of amorphous carbon should not be
228 ruled out. The presence of O-bearing groups is believed to be mainly due to oxidation of the tholin
229 samples when exposed to air during the transfer to the STXM vacuum chamber to conduct XANES

230 measurements, rather than from plasma-induced chemistry in the COSmIC/THS experiments, as
231 discussed in more detail in Section 3.5.

232 The C-XANES spectra of all four tholin samples also show the presence of multiple minor
233 bands assigned to a wide variety of chemical functions (Table S1, Supplementary Material). Most
234 of these assignments are tentative, as the intensities of most of these bands are small. Among these,
235 bands at ~ 283 , 284.6 , ~ 286 , and ~ 290.5 eV are associated with the presence of amorphous carbon
236 (Table S1), which is mixed with aromatic carbon structures (band C1, 285.0 – 285.3 eV; Table 1)
237 and aliphatic carbon chains (bands at ~ 287 – 288.5 eV; Table S1). The bands assigned to aliphatic
238 carbon chains appear to be the strongest in the spectrum of the tholin sample produced from the
239 $N_2:CH_4$ (90:10) mixture. A complete list of the bands observed in all four tholins is shown in Table
240 S1. Other minor bands consistent with the presence of different types of aromatic materials are
241 also observed in all tholins, including nitrogen-bearing heterocyclic compounds, but are stronger
242 for tholins produced from mixtures containing C_2H_2 ; these bands are consistent with the presence
243 of C- and N-bearing compounds similar to graphite (~ 285.2 and ~ 291.5 eV), pyridine (C_5H_5N ;
244 ~ 286.9 and ~ 290.6 eV), and fullerene (C_{60} ; ~ 290.6 eV) (Table S1). However, it is difficult to
245 estimate the relative intensities between aromatic hydrocarbons and heterocyclic aromatic
246 hydrocarbons because most of these bands are weak and overlap with each other as well as with
247 other bands.

248 The C-XANES spectra of both our $N_2:CH_4$ tholins (90:10 and 95:5) are consistent with the
249 only other XANES study of a Titan tholin produced from a $N_2:CH_4$ (95:5) mixture in an RF plasma
250 experimental set-up reported by Kuga et al. (2014). The two most intense bands observed in Kuga
251 et al. (2014)'s C-XANES spectrum at 286.8 and 288.1 eV, assigned to nitriles/ketones and
252 aliphatic carbon/amides, respectively, correspond to the bands C3 and C4 observed in our C-

253 XANES spectra (Fig. 2, Table 1). The C-XANES spectra of our four tholin samples also share
254 strong similarities with those obtained in a XANES study of two laboratory-generated Pluto/TNO
255 tholins produced from the UV photo-irradiation and 1.2-keV electron bombardment of N₂:CH₄:CO
256 (100:1:1) ice mixtures (Materese et al., 2014, 2015). In particular, the C-XANES spectra of our
257 four tholins and these two Pluto/TNO tholins show the presence of aromatic carbon (285.0–285.3
258 eV) and nitriles (399.3–400.0 eV).

259

260 *3.2. Nitrogen XANES spectra*

261

262 The normalized N-XANES spectra for all four of our tholin samples (Fig. 3) are consistent with
263 their C-XANES spectra (Fig. 2, Section 3.1). **Table 1 lists the main bands observed in all tholin**
264 **samples. They** are associated with imines (band N1 at 398.9–399.2 eV in Fig. 3) and nitriles (band
265 N2 at 399.8–400.0 eV), in particular for the tholins produced from mixtures containing **C₂H₂. In**
266 the spectra of tholins produced from mixtures that did not contain acetylene, the band N2 is slightly
267 redshifted (band N2* at 399.3–399.8 eV), indicating that the nitrile groups are mostly linked to
268 aliphatic carbon chains (Pels et al., 1995; Shard et al., 2004; Ray et al., 2005; Leinweber et al.,
269 2007; Cody et al., 2008), as opposed to nitrile groups linked to aromatic hydrocarbons (band N2)
270 (Zhu et al., 1997). Another band (N3 at ~401.3 eV) that is observed in all spectra can be assigned
271 to amino groups (Cody et al., 2011), although the position of this band is also consistent with the
272 presence of amide groups (O=C–NH_x) and mirrors the presence of bands C4 in the C-XANES
273 spectra (Fig. 2 and Table 1) and O3 in the O-XANES spectra (Fig. S1 and Table S3; Supplementary
274 Material).

275 N-XANES spectra also contain several minor bands that are tentatively associated with a
276 wide variety of functional groups or compounds (Table S2, Supplementary Material), including
277 pyridine, pyrrole, different types of nitrogen heterocycles, nitroso groups, nitro groups, and
278 oxidized variants of pyridine (pyridone and pyridine *N*-oxide). However, the assignments of these
279 O-bearing species could not be confirmed by the presence of mirror bands in the O-XANES spectra
280 (Table S3; Supplementary Material). Finally, multiple weak bands at ~400.5, ~401.3, and ~407.0
281 eV are consistent with the presence of nitrogen atoms inserted in graphene structures, supporting
282 the bands associated with graphene-like structures in the C-XANES spectra (Section 3.1, Table
283 S1).

284 As was the case for the C-XANES spectra, the N-XANES spectra of the four tholin samples
285 studied here share similarities with the only other XANES spectrum of another Titan tholin sample
286 produced from a N₂:CH₄ (95:5) gas mixture in a different experimental set-up (Kuga et al., 2014).
287 In particular, the spectrum of our tholin sample produced from the N₂:CH₄ (95:5) gas mixture is
288 very similar to that of Kuga et al. (2014), in which the two main bands observed at 398.9 and 399.7
289 eV, assigned to imines and nitriles, respectively, are comparable to the bands N1 and N2/N2* in
290 our N-XANES spectra. The Pluto/TNO tholins reported by Materese et al. (2014, 2015) also share
291 similarities with our tholin samples, with bands at 399.9 and 401.0 eV associated with nitriles and
292 amides, respectively, comparable to the bands N2/N2* and N3 observed in our N-XANES spectra.
293 In addition, although not explicitly reported in Materese et al. (2014, 2015), a shoulder visible in
294 the N-XANES spectra of the Pluto/TNO tholins around 398–399 eV is consistent with the presence
295 of imines, i.e., comparable to band N1 in our N-XANES spectra.

296

297 *3.3. C/N ratios and nitrogen content*

298

299 The elemental C/N abundance ratios derived from fitting the full-range XANES spectra (Fig. 4)
300 with atomic mass absorption coefficients (Henke et al., 1993; Section 2.2) for the four tholins
301 studied here are summarized in Table 2. The error bars on these ratios are due to the method used
302 to fit the XANES spectra to the atomic mass absorption coefficients (Cody et al., 2008; Nuevo et
303 al., 2011). C/N ratios were found to be low, in the 0.9–2.4 **range**. **These values** are consistent with
304 the C/N ratios **obtained using combustion and pyrolysis techniques** for other tholins produced
305 in experimental set-ups which use energy sources that can dissociate N₂ (i.e., cold plasma or
306 extreme UV radiation **sources**) (Cable et al., 2012). Note that an increase (respectively, decrease)
307 in the C/N ratio can be due to an increase (respectively, decrease) in the relative abundance of C
308 and/or a decrease (respectively, increase) in the relative abundance of N.

309 Our tholins produced **from gas** mixtures without acetylene display lower C/N ratios (0.9–
310 1.3) than those produced from mixtures with acetylene (2.2–2.4). This increase by a factor of 2 in
311 the C/N ratios of the tholin samples produced from the gas mixtures **with C₂H₂** is correlated with
312 an increase in the abundance of aromatic carbon, imines, and nitriles observed in these samples
313 (Fig. 2, Section 3.1). These observations indicate that the addition of acetylene by as small a mixing
314 ratio as 0.5% in the initial gas mixture results in a significant increase in the formation of chemical
315 groups that consist of double and triple covalent bonds (C=C, C=N, and C≡N). In addition, the
316 nitrile band (band N2 in Fig. 3 and Table 1) of the tholin samples produced from mixtures that
317 contained acetylene tends to be shifted to higher energies, implying a larger proportion of nitrile
318 groups attached to aromatic structures, as opposed to aliphatic structures (band N2*) (Section **3.2**).

319 A plausible explanation for these observations resides in the design of the COSmIC/THS
320 experimental set-up, which only allows for the first steps of the chemistry to occur, due to the short

321 residence time ($\sim 3.5 \mu\text{s}$) of the gas in the plasma discharge. This truncated chemistry was
322 demonstrated by a mass spectrometry study of the plasma-generated gas-phase products
323 (Sciamma-O'Brien et al., 2014), and subsequently confirmed by a modeling study of the gas phase
324 (Raymond et al., 2018). These studies indicated that, in the COSmIC/THS experiment, plasmas
325 generated in gas mixtures containing heavier molecular precursors result in the production of
326 heavier molecular products. The presence of heavier precursors in the initial gas mixtures allows
327 to go further into the chemical pathways associated with that precursor, leading to the formation
328 of larger molecules within the short residence time of the gas in the plasma cavity. However, the
329 use of heavier precursors can also lead to a chemistry favoring specific chemical pathways that
330 involve these initial precursors and/or dominant products formed from these precursors.

331 In addition, in an IR spectroscopy study of the solid-phase products formed in the
332 COSmIC/THS set-up (Sciamma-O'Brien et al., 2017), it was observed that both the nitrogen
333 chemistry and the incorporation of N atoms into the solid phase vary depending on the composition
334 of the initial gas mixture. In particular, when acetylene was present in the mixture, the chemistry
335 appeared to favor the formation of aromatic hydrocarbons, in which fewer N atoms were
336 incorporated into the tholins. The IR bands associated with N-bearing functional groups showed
337 that N atoms were added to the periphery of the molecules rather than deeper inside the molecular
338 structure, which is consistent with the findings of the present study. A possible explanation for the
339 higher C/N ratios reported here for tholins produced from mixtures containing acetylene as well
340 as from the IR spectroscopy results reported in Sciamma-O'Brien et al. (2017) is that the high
341 reactivity of C_2H radicals produced from the dissociation of C_2H_2 (Herbst and Woon, 1997;
342 Agúndez and Wakelam, 2013) triggers a chemistry in which the formation of hydrocarbons is
343 favored over that of N-bearing molecules **during** the short $3.5\text{-}\mu\text{s}$ residence time of the gas in the

344 plasma cavity, which results in more aromatic hydrocarbons being formed and less nitrogen being
345 incorporated in a bottom-up-type chemistry starting with smaller molecules.

346 **When comparing the C/N ratios of the tholin samples produced from N₂:CH₄ and**
347 **N₂:CH₄:C₂H₂ gas mixtures with different concentrations of CH₄, we also observe that an**
348 **increase in the concentration of CH₄ (from 5% to 10%) results in a slight decrease in the C/N**
349 **ratio for both the samples produced from N₂:CH₄ mixtures (from 1.3 to 0.9) and those**
350 **produced from N₂:CH₄:C₂H₂ mixtures (from 2.4 to 2.2). Although this observation suggests**
351 **that more N atoms may be incorporated into the tholins when the initial gas mixture contains**
352 **more CH₄, in particular because this change in C/N ratio is observed for both N₂:CH₄ and**
353 **N₂:CH₄:C₂H₂ mixtures, note that the uncertainties associated with the C/N ratio values**
354 **(Table 2) are large enough for the C/N ratios to be considered as constant for a given gas**
355 **composition (N₂:CH₄ or N₂:CH₄:C₂H₂) when the CH₄ concentration increases.**

356 **Our XANES results are also consistent** with a previous IR study of our COSmIC/THS
357 tholins (Sciamma-O'Brien et al., 2017), **in which** we observed lower C/N ratios and smaller
358 amounts of nitriles in tholins produced from the mixtures that did not contain C₂H₂. **The lower**
359 **level of incorporation of nitrogen in the tholins formed from N₂:CH₄:C₂H₂ mixtures is also**
360 **consistent with the findings of a recent study of the surface energy of tholins produced in the**
361 **COSmIC set-up from N₂:CH₄ (95:5) and N₂:CH₄:C₂H₂ (94.5:5:0.5) gas mixtures (Li et al.,**
362 **2021). The effects of varying the concentration of CH₄ and/or adding acetylene as a precursor in**
363 **the initial gas mixtures therefore reveal significant differences in how nitrogen is incorporated into**
364 **the solid phase of our tholin samples.**

365

366 *3.4. Comparison with C/N ratios from the literature*

367
368 From a general standpoint, it is difficult to conduct a rigorous comparison of the C/N ratios
369 determined for the tholins of our study with other Titan tholins produced in previous,
370 independent studies reported in the literature (e.g., McDonald et al., 1994; Coll et al., 1995,
371 1997, 1999, 2001; McKay, 1996; Sarker et al., 2003; Tran et al., 2003, 2008; Imanaka et al.,
372 2004; Curtis et al., 2005; Ferris et al., 2005; Somogyi et al., 2005; Bernard et al., 2006; Szopa
373 et al., 2006; Neish et al., 2008; Quirico et al., 2008; Sekine et al., 2008; Carrasco et al., 2009,
374 2016; Imanaka and Smith, 2010; Sciamma-O'Brien et al., 2010; Cable et al., 2012), as these
375 tholins were produced using a wide range of experimental set-ups with many varying
376 parameters (gas mixture composition, energy source, pressure, temperature, etc.) that could
377 all have an impact on the final measured C/N ratios.

378 To the best of our knowledge, before the study presented here with the COSmIC
379 experimental set-up, only two other experimental set-ups were used to conduct systematic
380 C/N ratio studies of tholins produced with different concentrations of methane. McDonald
381 et al. (1994) used a continuous flow plasma experiment at room temperature with a pressure
382 of 1.5 mbar to produce tholins from N₂:CH₄ gas mixtures with 0.1% and 10% of CH₄ and
383 did observe a decrease in the C/N ratio when increasing the CH₄ mixing ratio (C/N ratios of
384 1.49 and 0.75 for CH₄ concentrations of 0.1% and 10%, respectively), which is in agreement
385 with the trend we observe in the COSmIC study presented here (C/N ratios of 0.9 and 1.3 for
386 CH₄ concentrations of 5% and 10%, respectively). The opposite trend was reported for
387 tholins produced in another experimental set-up (PAMPRE) at room temperature and 1–2
388 mbar pressure in a continuous flow plasma experiment (Quirico et al., 2008; Sciamma-
389 O'Brien et al., 2010; Carrasco et al., 2016), where a higher C/N ratio was observed for tholins

390 produced with higher concentration of CH₄, with C/N ratios of 1.67 and 2.5 for CH₄
391 concentrations of 5% and 10%, respectively (Carrasco et al. 2016). These opposite trends in
392 published data show the complexity of comparing the results obtained from different
393 experiments.

394 For instance, if we look at the only other experimental set-up (PAMPRE) that was
395 used to conduct systematic studies in which tholins were produced from concentrations of
396 CH₄ of 5% and 10% and their C/N ratios were determined (Sciamma-O'Brien et al., 2010;
397 Carrasco et al., 2016), an increase by a factor of 1.5 in the C/N ratio of the PAMPRE tholins
398 was observed when increasing the concentration of CH₄ in the initial N₂:CH₄ gas mixture
399 from 5% to 10%. This trend is the opposite of what we observe in the COSmIC tholins,
400 where the C/N ratio decreases by a factor of 1.4 when the CH₄ concentration increases from
401 5% to 10% (Table 2, Section 3.3). In the PAMPRE experiment, however, a continuous
402 plasma is used, and the set-up is designed so that the aerosols remain in levitation and grow
403 in the plasma discharge for a long time (~100 s; Alcouffe et al., 2010). In contrast, in the
404 COSmIC experiment, a pulsed plasma is used in the stream of an accelerated gas expansion,
405 and the residence time of the gas in the plasma cavity is ~3.5 μs, which means that the
406 residence time of the aerosols, which are carried by the accelerated pulsed gas expansion, is
407 shorter than 3.5 μs (Sciamma-O'Brien et al., 2014). The very short residence time in
408 COSmIC results in truncated chemistry and aerosol growth (Sciamma-O'Brien et al., 2014,
409 2017; Raymond et al. 2018), while the longer residence time in PAMPRE allows the
410 chemistry to progress much farther. This difference could change the way nitrogen is
411 incorporated into the solid phase and explain the opposite trends observed in the PAMPRE
412 and COSmIC experiments.

413 Another difference between the PAMPRE and COSmIC experimental set-ups that
414 could explain the opposite trends observed for the C/N ratios is the operating gas
415 temperature in the reaction chamber (room temperature for PAMPRE vs. 150 K for
416 COSmIC), which could result in different chemical pathways, changing the nitrogen
417 incorporation into the solid phase. The significant effect of the operating gas temperature on
418 the tholin C/N ratios has been demonstrated by Coll et al. (1999). In their experiments, Coll
419 et al. produced tholins via plasma chemistry at room temperature and 100–150 K from
420 N₂:CH₄ (98:2) gas mixtures at low pressure (1.5 mbar). The C/N ratios determined for the
421 resulting tholins were 1.73 at room temperature and 2.82–2.83 at 100–150 K. This study thus
422 highlights an important effect of the gas temperature on the chemistry and in particular the
423 nitrogen incorporation, and may explain why the evolution of the C/N ratio as a function of
424 the concentration of CH₄ in the initial mixture in our experiments is different from what is
425 observed in the PAMPRE experiments.

426 Other factors like the gas pressure and the energy flux inducing the chemistry most
427 likely also contribute to the differences observed in the chemistry and the nitrogen
428 incorporation into the solid phase between the PAMPRE and COSmIC experimental set-ups.
429 This demonstrate how difficult comparative analyses can be between different experimental
430 set-ups. Only systematic and purposeful comparative studies where specific parameters are
431 varied to observe their effects on the properties of the tholins (optical constants, physical and
432 bulk properties, composition) will shed some light on the differences observed between
433 different experimental set-ups. All three experimental set-ups that have conducted C/N ratio
434 studies of tholins produced in different gas mixtures have however demonstrated that the

435 **initial composition has an effect on the way nitrogen is incorporated into the solid phase,**
436 **which is a piece of the puzzle to unveil the complexity of Titan's chemistry.**

437

438 *3.5. Oxygen-bearing compounds and C/O ratios*

439

440 Although oxygen was not present in any of the initial starting gas mixtures, its contribution to the
441 full-range XANES spectra is non negligible. Indeed, Fig. 4 shows that the intensity of the O edge
442 is 5–8 and 2–4 times smaller than the intensities for the C and N edges, respectively. This means
443 that a measurable amount of oxygen was incorporated into the tholins during the few minutes they
444 were transferred onto the sample holders to be placed into the STXM device for XANES analysis.
445 Oxidation of the tholin samples is supported by the presence of multiple oxygen-bearing functional
446 groups in the C-XANES and N-XANES spectra of all tholins, in particular amides and alcohols
447 (Table 1), as well as other types of carbonyls (Tables S1 and S2). The corresponding normalized
448 O-XANES spectra of the four tholins produced in this study are shown in Fig. S1, while the
449 complete list of the band assignments is given in Table S3, both in the Supplementary Material.

450 The presence of O-bearing groups in all our tholin samples may be explained by the
451 structure of the tholin samples produced in these experiments. Because materials analyzed with
452 the STXM instrument need to be thin enough for the X-rays to penetrate through the full thickness
453 of the samples, the tholins studied here were produced from experiments that lasted 1–2 hours,
454 resulting in substrates covered with scattered individual grains, 100–500 nm in diameter
455 (depending on the initial mixture), separated from each other on the substrate. In contrast, during
456 longer experiments, in which the tholins are typically recovered after 10–40 hours of sample
457 production, the grains jet-deposited on the substrate stack on top of each other, forming a more

458 homogenous layer of material (Sciamma-O'Brien et al., 2017). Consequently, oxidation will affect
459 a larger fraction of the samples produced from shorter experiments, as the ratio between the surface
460 exposed to oxidation and the volume of material is much higher for samples consisting of
461 individual grains (i.e., shorter experiments) than for samples consisting of a homogenous deposit
462 (i.e., longer experiments). **In Carrasco et al. (2016), the layer of oxidation of a Titan tholin
463 sample produced from an oxygen-free N₂:CH₄ (90:10) initial mixture was measured to be
464 about 20 nm thick. Considering spherical grains with a diameter of 500 nm for our tholin
465 samples produced from gas mixtures containing C₂H₂ (Sciamma-O'Brien et al., 2017), a 20-
466 nm-thick oxidation layer would correspond to an oxidation level of about 22% of the total
467 volume. However, the tholin sample produced by Carrasco et al. (2016) was not kept under
468 inert atmosphere and was therefore exposed to air for a long period of time before
469 characterization. Since our tholin samples were kept under an argon atmosphere and
470 exposed to air for only a few minutes before being analyzed, the thickness of the oxidation
471 layer is expected to be smaller than 20 nm. Nonetheless, because the tholin samples produced
472 in our XANES study have a large surface-to-volume ratio, their XANES spectra are more
473 susceptible to higher relative levels of oxidation, as confirmed by the high abundances of O-
474 bearing groups (Fig. S1, Table S3, Supplementary Material).**

475 Kuga et al. (2014) also reported the presence of O-bearing functional groups in the C- and
476 N-XANES spectra of their Titan tholin sample produced from an oxygen-free N₂:CH₄ (95:5) initial
477 mixture. They observed bands associated with amides, alcohols, and ethers, which are also present
478 in all our tholin samples (Tables 1, S1, and S2). Previous tholin studies showed that unsaturated
479 chemical groups in the molecular structure of tholins produced by cold plasmas are dominated by

480 nitrogen-bearing moieties (i.e., imines, nitriles), which may explain why tholins are easily prone
481 to hydrolysis and therefore oxidation (Cable et al., 2012).

482 Finally, the elemental C/O abundance ratios (and their associated error bars) derived for
483 the four tholins studied here (see Sections 2.2 and 3.3.) are summarized in Table S4
484 (Supplementary Material). These ratios were determined to range from 1.9 to 3.3, with higher C/O
485 ratios for tholins produced from initial mixtures containing acetylene. A previous scanning
486 electron microscopy (SEM) study of COSmIC/THS tholin samples (Sciamma-O'Brien et al., 2017)
487 demonstrated that tholins produced from $N_2:CH_4:C_2H_2$ gas mixtures consist of grains with much
488 larger diameter than tholins produced from simpler $N_2:CH_4$ mixtures (500 nm vs. 100 nm). This
489 implies that the surface-to-volume ratio of the tholin samples produced from C_2H_2 -containing
490 mixtures is much smaller than for tholins produced in mixtures without C_2H_2 , which results in less
491 oxidation of the samples when exposed to air. The observation of lower C/O ratios for the tholins
492 produced in C_2H_2 -containing mixtures are therefore consistent with the hypothesis that oxidation
493 most probably occurred when the tholins were exposed to air during their transfer to the sample
494 holder to be analyzed with the STXM instrument rather than during the experiments inside the
495 COSmIC/THS chamber.

496

497 *3.6. Applications to Titan's atmosphere*

498

499 The XANES study at the C and N edges of the tholin samples produced in our COSmIC/THS
500 experimental set-up from four different initial gas mixtures, namely, $N_2:CH_4$ (90:10 and 95:5) and
501 $N_2:CH_4:C_2H_2$ (89.5:10:0.5 and 94.5:5:0.5), showed that the relative abundances between the C-
502 and N-bearing products that are formed vary with both the presence or absence of C_2H_2 and

503 **possibly** the relative proportion of CH₄ in the initial gas mixtures. XANES spectra of these tholins
504 also showed that mixtures containing C₂H₂ favored the formation of aromatic compounds and
505 larger relative amounts of imines and nitriles compared with when C₂H₂ is not present.

506 These important results can be used to infer some connections with what can be expected
507 for the production mechanisms of the aerosols that are responsible for the haze layers in Titan's
508 atmosphere. Indeed, the variations observed for the composition and C/N ratios of our four tholin
509 samples can be interpreted as potential indicators of variations in the composition of Titan's haze,
510 as predicted by photochemical models (Wilson and Atreya, 2003; Lavvas et al., 2008a, 2008b).
511 Our experimental results **confirm that** the nitrogen content of Titan's aerosols **is expected to vary**
512 over seasonal and astronomical timescales (Lunine and Atreya, 2008; Roe, 2012).

513 The nitrogen content of Titan's aerosols is also expected to vary as a function of the altitude
514 and latitude at which the aerosols are formed, particularly during seasonal changes that affect the
515 atmospheric circulation and result in a molecular enrichment at the poles. **For example, molecules**
516 **such as C₂H₂, C₂H₄, C₂H₆, C₆H₆, HC₃N, and HCN were observed in higher concentrations at**
517 **the south pole after the northern spring equinox, where they are expected to have an impact**
518 **on the composition of the aerosols formed at these latitudes** (e.g., Teanby et al., 2012; Vinatier
519 et al., 2015; Coustenis et al., 2016; Mathé et al., 2020). Our experimental results demonstrate that
520 a change in the atmospheric composition due to these seasonal and local enhancements would
521 influence the chemistry and the resulting aerosol production mechanisms. The low C/N ratios
522 measured for all our tholin samples are also consistent with the hypothesis that Titan's aerosols
523 act as a sink for N atoms in the atmosphere, as supported by the data collected by *Huygens'* Aerosol
524 Collector Pyrolyser (Israël et al., 2005).

525

526 4. Conclusions

527

528 We produced Titan tholins in the COSmIC/THS experimental set-up using a plasma discharge in
529 the stream of a jet-cooled expansion for four different initial gas mixtures: N₂:CH₄ (90:10), N₂:CH₄
530 (95:5), N₂:CH₄:C₂H₂ (89.5:10:0.5), and N₂:CH₄:C₂H₂ (94.5:5:0.5). The tholin samples were
531 collected under inert atmosphere after 1–2 hours of deposition and were analyzed with XANES
532 spectroscopy at the C, N, and O edges. The absorption bands identified in the XANES spectra of
533 these tholin samples indicate the presence of various chemical structures and groups, including
534 aromatic carbon, imines, and nitriles. The relative abundances between these bands vary with (i)
535 the presence or absence of C₂H₂, and (ii) **possibly** the concentration of CH₄ in the initial mixtures.
536 COSmIC/THS experiments conducted with initial mixtures in which C₂H₂ was added resulted in
537 the formation of tholins that contain more aromatic compounds and larger relative amounts of
538 imines and nitriles compared with tholins produced from mixtures that did not contain C₂H₂.
539 XANES spectra of all tholins also show the presence of O-bearing species that resulted from the
540 oxidation of the tholins when exposed to **air**. **Furthermore**, XANES analysis allowed for the
541 determination of the elemental C, N, and O composition of our four tholin samples, and the
542 subsequent estimate of their C/N and C/O ratios. C/N ratios were all found to be low (< 2.5), in
543 agreement with previous, independent studies of tholins produced with energy sources that can
544 dissociate N₂. **C/N ratios were found to be slightly lower when the concentration of CH₄ in**
545 **the gas mixture was higher. Finally**, the C/N ratios of tholins produced from gas mixtures that
546 contained **C₂H₂** were found to be twice as large as of those produced from mixtures that did not
547 contain **C₂H₂**, **demonstrating the impact of the composition of the initial gas on the**
548 **composition of the resulting solid-phase particles, in particular the nitrogen incorporation.**

549 The differences in chemical (aromatic vs. aliphatic compounds, nitriles, etc.) and elemental
550 (C/N ratios) compositions, observed for our COSmIC/THS tholins produced from gas mixtures
551 with different concentrations of CH₄ and in the absence/presence of C₂H₂, highlight differences in
552 how nitrogen is incorporated into the solid phase: when C₂H₂ is absent from the initial mixture,
553 nitrogen is incorporated deeper in the molecular structure and nitrile groups are preferably attached
554 to aliphatic structures, while when C₂H₂ is present, nitrogen is added to the periphery of aromatic
555 molecules, for example in the form of nitrile groups.

556 Our experimental results can also guide the analysis of observations of Titan's atmosphere
557 and the development of models. Indeed, the differences in chemical and elemental compositions
558 of our tholins illustrate how a change in composition at the location where aerosols are formed in
559 Titan's atmosphere may lead to variations in the nitrogen content and composition of Titan's
560 aerosols. This variability depends on several parameters which need to be taken into account in
561 models, such as the location (altitude, latitude), the season, and the astronomical timescales, as
562 they can affect the atmospheric circulation, the concentration of specific molecules in different
563 parts of the atmosphere, and therefore the chemistry and product formation.

564

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566

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911

912 **Table 1**

913 Assignments of the main bands observed at the carbon (C1–C5, Fig. 2) and nitrogen (N1–N3, Fig.
 914 3) 1s edge XANES spectra of all four tholins. A more complete list of possible assignments of all
 915 the bands (including minor bands) in these C- and N-XANES spectra are listed in Tables S1 and
 916 S2 (Supplementary Material), respectively. The band N2* corresponds to the band N2 that is
 917 redshifted in tholins produced from gas mixtures that did not contain acetylene (Fig. 3). C and N
 918 atoms involved in their respective XANES bands are marked in bold.

	Band	Energy (eV)	Assignment	Transition	References*
<i>Carbon edge</i>	C1	285.0–285.3	Aromatic C=C	1s → π*	1,2,3
	C2	285.7–286.0	Imine C=N	1s → π*	4,5
	C3	286.8–287.0	Nitrile C≡N	1s → π*	2,4,6,7,8
	C4	287.8–288.3	Amide O=C–NH_x	1s → π*	2,9
	C4	287.9–288.2	Benzene [Ar] C	1s → π*	10
	C5	288.9–289.2	Amorphous C (<i>sp</i> ³)	1s → 3p/σ*	10
	C5	288.9–289.4	Alcohol CH_x–OR	1s → 3p/σ*	11,12,13
	C5	288.9–289.7	Urea group (NH _x) ₂ C=O	1s → π*	2
<i>Nitrogen edge</i>	N1	398.2–399.1	Imine C=N	1s → π*	2,7,14
	N2*	399.3–399.8	Nitrile (aliphatic) C≡N	1s → π*	2,5,14,15,16
	N2	399.8–400.0	Nitrile (aromatic) C≡N	1s → π*	17
	N3	401.0–401.3	Amide O=C–NH_x	1s → π*	2,3,14,18
	N3	401.0–401.4	Heterocyclic nitrogen	1s → π*	15
	N3	401.1–401.4	Amine C–NH_x	1s → 3p/σ*	3

919

920 *References: **(1)** Stöhr (1992); **(2)** Cody et al. (2008); **(3)** Cody et al. (2011); **(4)** Dhez et al. (2003); **(5)**
921 Shard et al. (2004); **(6)** Kikuma et al. (1998); **(7)** Apen et al. (1993); **(8)** Zhou et al. (2009); **(9)** Urquhart
922 and Ade (2002); **(10)** Díaz et al. (2001); **(11)** Ishii and Hitchcock (1988); **(12)** Sham et al. (1989); **(13)**
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924 et al. (1997); **(18)** Gordon et al. (2003).

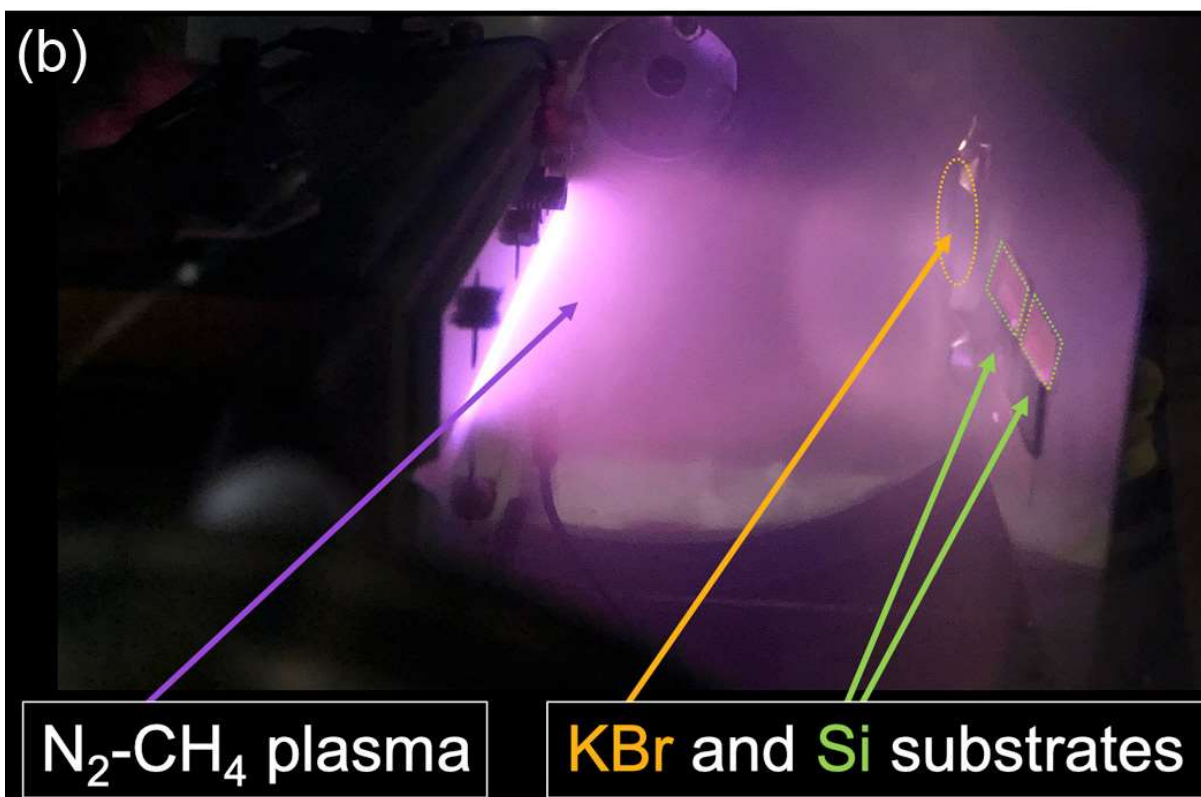
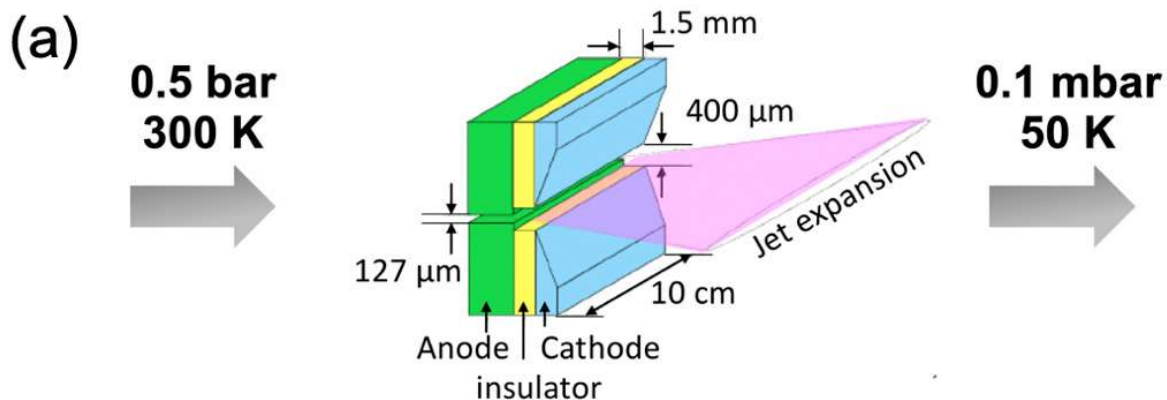
925

926 **Table 2**

927 Elemental C/N ratios determined from fitting the full-range XANES spectra of all tholins spanning
 928 the C, N, and O 1s edges (Fig. 4) with the atomic mass absorption coefficients (Henke et al., 1993,
 929 see Sections 2.2 and 3.3 for details). Error bars are due to the fitting method used (Cody et al.,
 930 2008; Nuevo et al., 2011).

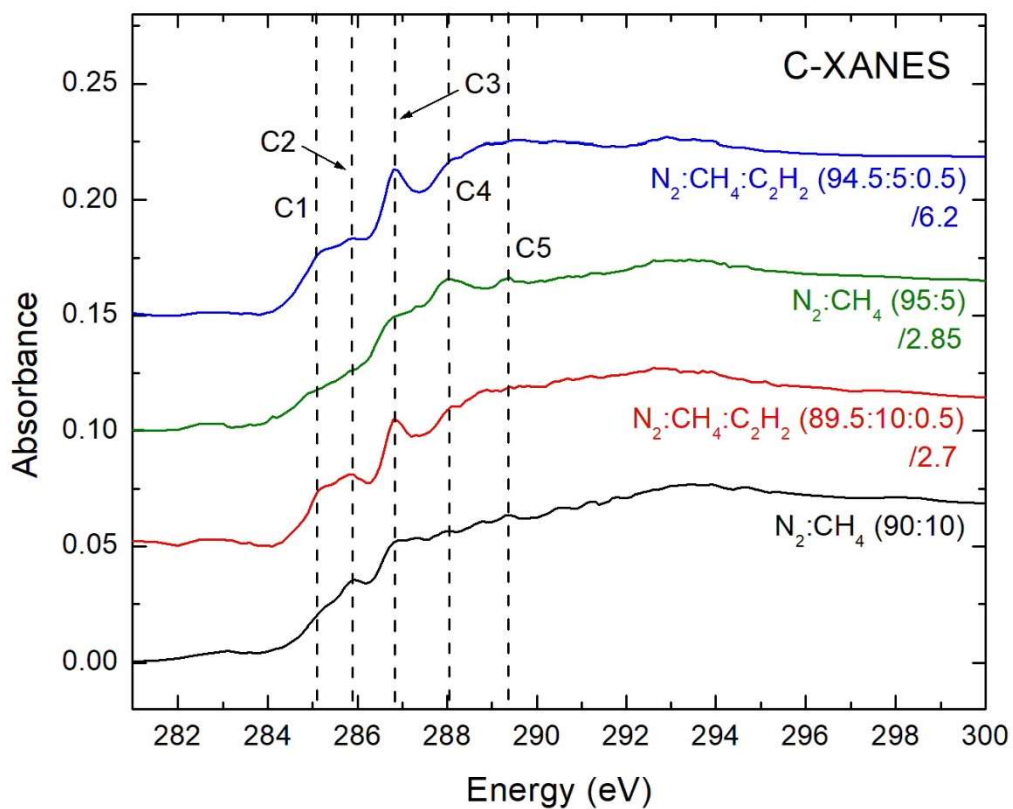
Initial gas mixture	C/N Ratio
N ₂ :CH ₄ (90:10)	0.9 ± 0.3
N ₂ :CH ₄ :C ₂ H ₂ (89.5:10:0.5)	2.2 ± 0.4
N ₂ :CH ₄ (95:5)	1.3 ± 0.2
N ₂ :CH ₄ :C ₂ H ₂ (94.5:5:0.5)	2.4 ± 0.4

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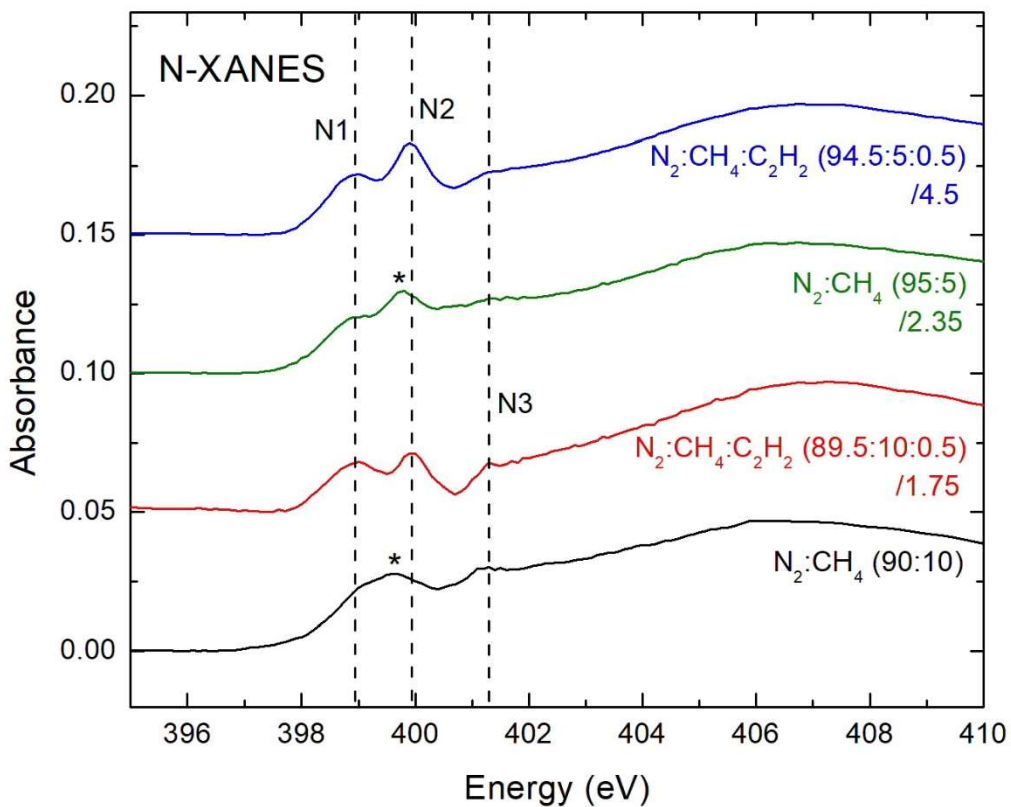
932

933 **Fig. 1.** (a) Schematic of the COSmIC/THS adiabatic expansion of the gas allowing to achieve low
 934 temperatures and pressures that are representative of cold atmospheres such as that of Titan (from
 935 Broks et al., 2005). The plasma discharge is the energy source used to induce the chemistry in the
 936 expansion, by simulating the chemistry induced by charged particles in planetary atmospheres. (b)
 937 Actual photograph of the expansion and plasma in the COSmIC/THS chamber, which also shows
 938 how the so-formed solid particles ('tholins') are deposited on different substrates. Si₃N₄ substrates
 939 were used for the experiments reported here.

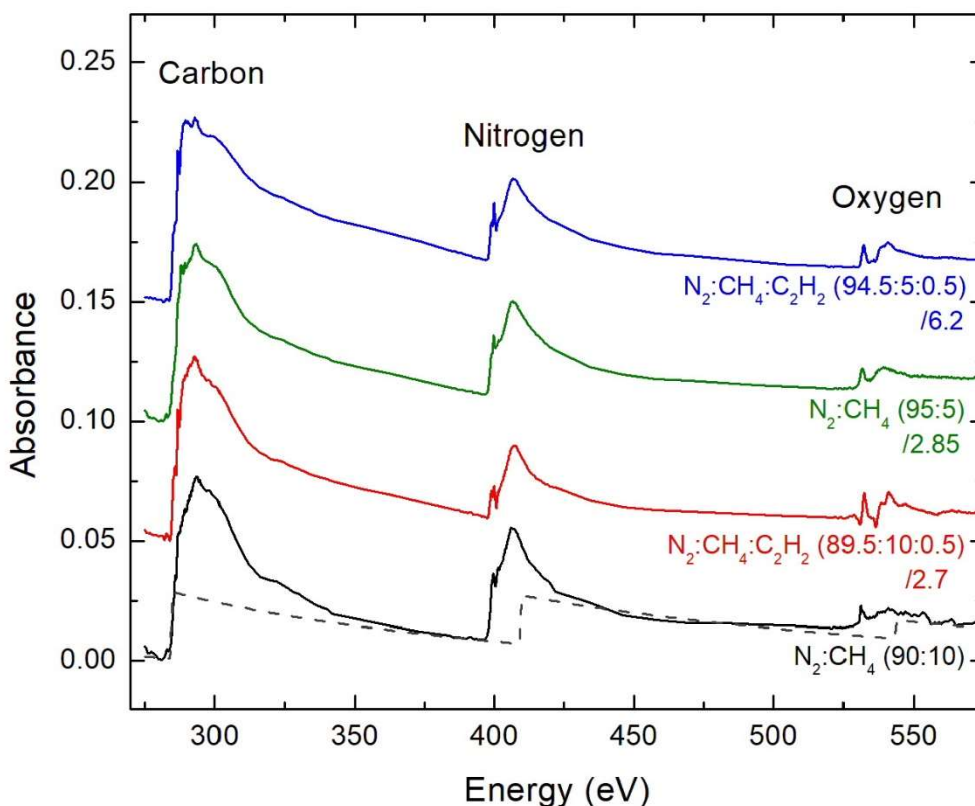


940
 941 **Fig. 2.** XANES spectra at the carbon 1s edge of the four tholin samples produced in the present
 942 study. **The intensities of the spectra are normalized to the C-edge peak intensity of the N₂:CH₄**
 943 **(90:10) spectrum. Normalization factors are provided below each gas mixture composition**
 944 **on the plot.** The main bands (C1–C5) are highlighted with dashed lines and their assignments are
 945 given in Table 1. Minor bands are listed in Table S1 (Supplementary Material). Absorbances of
 946 the spectra are normalized and offset for clarity.

947



948
 949 **Fig. 3.** XANES spectra at the nitrogen 1s edge of the four tholin samples produced in the present
 950 study. **The intensities of the spectra are normalized to the N-edge peak intensity of the $N_2:CH_4$**
 951 **(90:10) spectrum. Normalization factors are provided below each gas mixture composition**
 952 **on the plot.** The main bands (N1–N3) are highlighted with dashed lines and their assignments are
 953 given in Table 1. Minor bands are listed in Table S2 (Supplementary Material). The bands marked
 954 with an asterisk correspond to band N2 that is redshifted in energy and referred to as band N2*
 955 (Table 1, Section 3.2). Absorbances of the spectra are normalized and offset for clarity.
 956



957
 958 **Fig. 4.** Full-range XANES spectra spanning the C, N, and O 1s edges (275–575 eV) of the four
 959 tholin samples produced in the present study. **The intensities of the spectra are normalized to**
 960 **the C-edge peak intensity of the N₂:CH₄ (90:10) spectrum. Normalization factors are**
 961 **provided below each gas mixture composition on the plot.** These spectra were used to determine
 962 the C/N and C/O ratios of each sample, after fitting the spectra with atomic mass absorption
 963 coefficients (see Sections 3.3 and 3.5, respectively). An example of such computational fit is
 964 shown for the spectrum of the tholin produced from the N₂:CH₄ (90:10) gas mixture (dashed grey
 965 curve). Detailed XANES spectra at the C and N 1s edges are shown in Figs. 2 and 3, respectively,
 966 and the assignments of the bands at the C and N edges are given in Table 1. Detailed XANES
 967 oxygen 1s edge spectra are shown in Fig. S1 and the assignments of the bands are given in Tables

968 S1 and S2 (Supplementary Material). Absorbances of the spectra are normalized and offset for
969 clarity.

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977 **The Titan Haze Simulation (THS) experiment on COSmIC.**

978 **Part III. XANES study of laboratory analogs of Titan tholins**

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983 Michel Nuevo, Ella Sciamma-O'Brien, Scott A. Sandford, Farid Salama,

984 Christopher K. Materese, and A.L. David Kilcoyne

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Supplementary Material

989

990 **Table S1**

991 Complete list of possible assignments of all the main (C1–C5) and minor bands observed at the XANES
 992 carbon 1s edge spectra of all tholins (Fig. 2). Bands C1–C5 are the same as those labeled in Fig. 2 and Table
 993 1. Because of the small intensities of the minor bands, their assignments are tentative. Note that assignments
 994 do not necessarily indicate the presence of specific compounds; rather, they are indicative of chemical
 995 bonds that are consistent with such compounds. C atoms involved in the XANES bands are marked in bold.
 996 [Ar], aromatic ring; R, carbon chain.

Band	Energy (eV)	Assignment	Transition	References*
	282.8–283.2	Amorphous C		1,2
	283.5–283.9	<i>p</i> -Benzoquinone C=C–C=O	1s → π*	3
	284.1–284.3	Quinone C=C–C=O	1s → π*	4
		Protonated aromatic [Ar]C–H	1s → π*	4
	284.6–284.7	Amorphous C (<i>sp</i> ²)	1s → π*	5,6
	284.9–285.4	Alkylated aromatic [Ar]C–R	1s → π*	7,8
		Protonated aromatic [Ar]C–H	1s → π*	7,8
C1	285.0–285.3	Aromatic C=C	1s → π*	9,10,11
	285.2–285.4	Graphite (C <i>sp</i> ²)	1s → π*	12,13,14,15
C2	285.7–286.0	Imine C=N	1s → π*	16,17
	285.7–286.2	Aromatic carbonyl [Ar]–C=O	1s → π*	18
	285.8–286.2	Amorphous C	p5	12
		<i>p</i> -Benzoquinone C=C–C=O	1s → π*	3
	286.0–286.2	Conjugated ketone C=C–C=O	1s → π*	10,11
C3	286.8–287.0	Nitrile C≡N	1s → π*	10,15,16,19,20
		Pyridine C=N	1s → π*	21
		Ketone C=O	1s → π*	4,22,23
	286.8–287.1	Aliphatic C–C–H	1s → 3p/σ*	15
	287.0–287.5	Aromatic alcohol [Ar]C–OH	1s → π*	3,4
		Aliphatic ketone C–C=O	1s → π*	24,25
	287.1–288.0	Enol C=C–OH	1s → π*	10
	287.2–288.3	Aliphatic C–C	1s → 3p/σ*	10,22,24,26,27
	287.3–287.7	Aliphatic C–H	1s → 3p/σ*	11
	287.7–288.3	Aromatic carbonyl [Ar]–C=O	1s → π*	18
C4	287.8–288.3	Amide O=C–NH _x	1s → π*	10,23
C4	287.9–288.2	Benzene [Ar]C	1s → π*	6

	288.0–288.8	Carboxylic acid O=C–OH	1s → π*	6,23,24,28
	288.6–289.0	Diamond (C <i>sp</i> ³)	1s → 3p/σ*	12
C5	288.9–289.2	Amorphous C (<i>sp</i> ³)	1s → 3p/σ*	6
C5	288.9–289.4	Alcohol CH _x –OR	1s → 3p/σ*	4,24,25
C5	288.9–289.7	Urea group (NH _x) ₂ C=O	1s → π*	10
	289.4–289.7	Secondary alcohol R ₂ CH–OH	1s → 3p/σ*	11
	289.5–290.1	Amorphous C (<i>sp</i> ³)	1s → 3p/σ*	6
	290.1	Alkene C=C	1s → 3p/σ*	29
		Carbamoyl NH _x (C=O)O–R	1s → π*	10
	290.3–294.6	Carbonate R–O(C=O)O–R	1s → π*	10,30
	290.4–290.8	Pyridine C–N (<i>sp</i> ²)		15
	290.6–290.8	Amorphous C	1s → 3p/σ*	6
		<i>p</i> -Benzoquinone C=C–C=O	1s → π*	3
		Fullerene C ₆₀	1s → 3p/σ*	31
	291.2–291.9	Graphite C–C	1s → 3p/σ*	6
	293.4–293.8	<i>p</i> -Benzoquinone C=C–C=O	1s → σ*	3
		<i>p</i> -Benzoquinone C=C–C=O	1s → π*	3

997

998 *References: (1) Gago et al. (2005); (2) Ray et al. (2006); (3) Francis and Hitchcock (1992); (4) Lehmann
999 et al. (2005); (5) Gutiérrez et al. (1995); (6) Díaz et al. (2001); (7) Hitchcock et al. (1987); (8) Robin et al.
1000 (1988); (9) Stöhr (1992); (10) Cody et al. (2008); (11) Cody et al. (2011); (12) Jimenez et al. (2003); (13)
1001 Ray et al. (2005); (14) Abrasonis et al. (2008); (15) Zhou et al. (2009); (16) Dhez et al. (2003); (17) Shard
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1003 Bhattacharyya et al. (2000); (22) Hitchcock and Brion (1980); (23) Urquhart and Ade (2002); (24) Ishii and
1004 Hitchcock (1988); (25) Sham et al. (1989); (26) Hitchcock et al. (1986); (27) Hitchcock and Ishii (1987);
1005 (28) Ishii and Hitchcock (1987); (29) Ziethen et al. (2002); (30) Flynn et al. (1997); (31) Terminello et al.
1006 (1991).
1007

1008 **Table S2**

1009 Complete list of possible assignments of all the main and minor bands observed at the XANES nitrogen 1s
 1010 edge spectra of all tholins (Fig. 3). Bands N1–N3 are the same as those labeled in Fig. 3 and Table 1.
 1011 Because of the small intensities of the minor bands, their assignments are tentative. Note that assignments
 1012 do not necessarily indicate the presence of specific compounds; rather, they are indicative of chemical
 1013 bonds that are consistent with such compounds. N atoms involved in the XANES bands are marked in bold.
 1014 [Py], pyridinic ring; R, carbon chain; [Ar], aromatic ring.

Band	Energy (eV)	Assignment	Transition	References*
	398.2–399.0	Pyridine C=N	1s → π*	1,2
N1	398.2–399.1	Imine C=N	1s → π*	3,4,5
	399.0–399.3	Heterocyclic nitrogen (‘center’)	1s → π*	1
	399.2–399.6	Amine C–NH ₂	1s → 3p/σ*	1
N2*	399.3–399.8	Nitrile (aliphatic) C≡N	1s → π*	1,2,4,5,6
	399.6–399.9	Pyridine amine [Py]–NH ₂		7,8,9
N2	399.8–400.0	Nitrile (aromatic) C≡N	1s → π*	8
	399.9–400.2	Nitroso group C–N=O		1
	400.2–400.5	Pyridine		1
	400.2–400.6	Pyrrole		1
	400.5–401.2	Nitrogen in graphene		10
	400.6–400.8	Protonated imine C=NH ⁺	1s → π*	11
	400.6–401.1	Pyridone [Py]C=O		1,9
	400.9–401.2	Nitrile (aromatic) C≡N	1s → π*	8
	400.9–401.7	Quaternary nitrogen NR ₄ ⁺		1,8
N3	401.0–401.3	Amide O=C–NH _x	1s → π*	4,5,12,13
N3	401.0–401.4	Heterocyclic nitrogen (‘valley’)	1s → π*	1
N3	401.1–401.4	Amine C–NH _x	1s → 3p/σ*	12
	401.3–401.6	Nitrile (aliphatic) C≡N	1s → π*	8
	401.3–401.7	Nitrogen in graphene		14
	401.4–401.9	Amide (peptide) O=C–NHR	1s → π*	4,5,12,13
	401.4–402.1	Pyridone [Py]C=O		1,9
	401.6–401.9	Urea group (NH _x) ₂ C=O	1s → π*	15
	402.1–402.6	Amine C–NH _x	1s → 3p/σ*	5
	402.3–404.0	Pyrrole		7,8,9,12
	402.5–402.8	Oxidized pyrrole		7

402.8–403.2	Urea group (NH _x) ₂ C=O	1s → π*	5
	Carbazole		7
402.9–403.4	Pyridine <i>N</i> -oxide [Py]–N ⁺ –O ⁻		1,8
403.2–403.5	Porphyrin		7
403.3–403.5	Nitrile C≡N	1s → π*	9
	Diacylimide [R–(C=O)] ₂ NH		16
403.4–403.5	Pyridone [Py]C=O		9
403.4–403.8	Nitro group C–NO ₂	1s → π*	12
403.5–404.0	Aromatic nitro group [Ar]–NO ₂	1s → π*	4,12,17
403.8–403.9	Pyridine		8
403.8–404.0	Carbazole		8
404.8–405.0	Aromatic amine [Ar]–NH ₂	1s → 3p/σ*	7,8
	Pyridine <i>N</i> -oxide [Py]–N ⁺ –O ⁻		8
405.3–405.5	Nitrate NO ₃ ⁻	1s → π*	4
405.9–406.2	Nitro group C–NO ₂	1s → π*	1
406.0–406.5	Saturated amine NR ₃	1s → 3p/σ*	7,8
407.0–407.2	Pyridine	1s → σ*	10
	Nitrogen in graphene	1s → σ*	10
407.4–407.8	Saturated amine NR ₃	1s → 3p/σ*	7,8

1015

1016 *References: (1) Pels et al. (1995); (2) Ray et al. (2005); (3) Apen et al. (1993); (4) Leinweber et al. (2007);
 1017 (5) Cody et al. (2008); (6) Shard et al. (2004); (7) Mitra-Kirtley et al. (1993); (8) Zhu et al. (1997); (9) Xiao
 1018 et al. (2005); (10) Hellgren et al. (2001); (11) Hennig et al. (1998); (12) Cody et al. (2011); (13) Gordon et
 1019 al. (2003); (14) Grant et al. (1994); (15) Urquhart et al. (1995); (16) Lessard et al. (2007); (17) Turci et al.
 1020 (1996).

1021

1022 **Table S3**

1023 Complete list of possible assignments of all main (O1–O4) and minor bands observed at the XANES
 1024 oxygen 1s edge spectra of all tholins (Fig. S1). Because of the small intensities of the minor bands, their
 1025 assignments are tentative. Note that assignments do not necessarily indicate the presence of specific
 1026 compounds; rather, they are indicative of chemical bonds that are consistent with such compounds. O atoms
 1027 involved in the XANES bands are marked in bold. [Ar], aromatic ring; PVMK, poly(vinyl methyl ketone);
 1028 PEAMA, poly(ethylene-*alt*-maleic acid); PAA, poly(acrylic acid); Asp, aspartic acid; Glu, glutamic acid;
 1029 PVP, poly(4-vinylphenol); PVA, poly(vinyl alcohol); Ser, serine; Thr, threonine; Tyr, tyrosine; Hyp,
 1030 hydroxyproline.

Band	Energy (eV)	Assignment	Transition	References*
	530.0	<i>p</i> -Benzoquinone [Ar]C=O	1s → π*	1
		9-Anthraldehyde C=O	1s → π*	2
	530.0–530.2	9-Acetylanthracene O=C–CH ₃	1s → π*	2
		PVMK O=C–CH ₃	1s → 3p/σ*	2
O1	531.0	Ketone C=O	1s → π*	3,4
		Formaldehyde C=O	1s → π*	5,6
		Benzaldehyde [Ar]C=O	1s → π*	7
		1-Naphthoic acid C=O	1s → π*	2
		PEAMA C=O	1s → π*	2
		PAA C=O	1s → π*	2
O2	531.6	Ethyl benzoate O=C–O–C ₂ H ₅	1s → π*	7
O3	532.1–532.2	Amide O=C–NH _x	1s → π*	8
	532.1–532.3	Carboxyl O–C=O	1s → π*	3,9
		Methyl formate O=C–O–CH ₃	1s → π*	7
	532.2–532.3	Amino acids O–C=O	1s → π*	10
	532.9–533.1	Phthalic anhydride O–C=O		2
		1-Naphthoic acid O=C–O–H		2
	533.1	<i>p</i> -Benzoquinone [Ar]C=O	1s → π*	1
	533.5	Benzaldehyde [Ar]C=O	1s → 3s/σ*	7
		Terephthalaldehyde C=O	1s → 3s/σ*	7
		PAA O=C–O–H		2
O4	534.1–534.2	Alcohol H _x C–O–H	1s → 3p/σ*	3,11
O4	534.1–534.2	Carbonate CO ₃	1s → π*	12
		Asp, Glu O=C–O–H	1s → π*	10

534.2	Ethyl benzoate $\mathbf{O}=\mathbf{C}-\mathbf{O}-\mathbf{C}_2\mathbf{H}_5$	$1s \rightarrow 3s/\sigma^*$	7
	Ethyl benzoate $\mathbf{O}=\mathbf{C}-\mathbf{O}-\mathbf{C}_2\mathbf{H}_5$	$1s \rightarrow \pi^*$	7
534.7	PVP $[\text{Ar}]\mathbf{C}=\mathbf{O}$		2
534.7–535.0	Enol $\mathbf{C}=\mathbf{C}-\mathbf{O}-\mathbf{H}$	$1s \rightarrow 3p/\sigma^*$	3
534.9–535.3	Methyl formate $\mathbf{O}=\mathbf{C}-\mathbf{O}-\mathbf{CH}_3$	$1s \rightarrow 3s/\sigma^*$	7
	Methyl formate $\mathbf{O}=\mathbf{C}-\mathbf{O}-\mathbf{CH}_3$	$1s \rightarrow \pi^*$	7
535.4–535.6	Formaldehyde $\mathbf{C}=\mathbf{O}$	$1s \rightarrow 3s/\sigma^*$	5,6
536.2	Formaldehyde $\mathbf{C}=\mathbf{O}$	$1s \rightarrow 3p/\sigma^*$	5,6
	Terephthalaldehyde $\mathbf{C}=\mathbf{O}$	$1s \rightarrow 4p/\sigma^*$	7
536.9–537.0	Methyl cellulose		2
	Methyl formate $\mathbf{O}=\mathbf{C}-\mathbf{O}-\mathbf{CH}_3$	$1s \rightarrow 3p/\sigma^*$	7
	PVA $\mathbf{C}-\mathbf{O}-\mathbf{H}$		2
537.3–537.6	Benzaldehyde $[\text{Ar}]\mathbf{C}=\mathbf{O}$	$1s \rightarrow 5p/\sigma^*$	7
	Ethyl benzoate $\mathbf{O}=\mathbf{C}-\mathbf{O}-\mathbf{C}_2\mathbf{H}_5$	$1s \rightarrow 3p/\sigma^*$	7
537.6	Formaldehyde $\mathbf{C}=\mathbf{O}$	$1s \rightarrow 4p/\sigma^*$	5,6
538.2–538.3	<i>p</i> -Benzoquinone $[\text{Ar}]\mathbf{C}=\mathbf{O}$	$1s \rightarrow \pi^*$	1
538.2–539.3	Ser, Thr, Tyr, Hyp $\mathbf{C}-\mathbf{O}-\mathbf{H}$	$1s \rightarrow \sigma^*$	10
540.7–540.9	Ethyl benzoate $\mathbf{O}=\mathbf{C}-\mathbf{O}-\mathbf{C}_2\mathbf{H}_5$	$1s \rightarrow \sigma^*$	7
540.9–541.0	Methyl formate $\mathbf{O}=\mathbf{C}-\mathbf{O}-\mathbf{CH}_3$	$1s \rightarrow \sigma^*$	7
541.9–548.6	Amino acids $\mathbf{O}-\mathbf{C}=\mathbf{O}$	$1s \rightarrow \sigma^*$	10
542.5–542.7	<i>p</i> -Benzoquinone $[\text{Ar}]\mathbf{C}=\mathbf{O}$	$1s \rightarrow \sigma^*$	1
544.1	Formaldehyde $\mathbf{C}=\mathbf{O}$	$1s \rightarrow \sigma^*$	5,6
	Benzaldehyde $[\text{Ar}]\mathbf{C}=\mathbf{O}$	$1s \rightarrow \sigma^*$	7
	Terephthalaldehyde $\mathbf{C}=\mathbf{O}$	$1s \rightarrow \sigma^*$	7
546.1–546.2	Ethyl benzoate $\mathbf{O}=\mathbf{C}-\mathbf{O}-\mathbf{C}_2\mathbf{H}_5$	$1s \rightarrow \sigma^*$	7
546.9–547.2	Methyl formate $\mathbf{O}=\mathbf{C}-\mathbf{O}-\mathbf{CH}_3$	$1s \rightarrow \sigma^*$	7
553.9–554.1	<i>p</i> -Benzoquinone $[\text{Ar}]\mathbf{C}=\mathbf{O}$	$1s \rightarrow \sigma^*$	1

1031

1032 *References: (1) Francis and Hitchcock (1992); (2) Kim et al. (2011); (3) Cody et al. (2008); (4) Yabuta et
1033 al. (2014); (5) Hitchcock and Brion (1980); (6) Robin et al. (1988); (7) Hitchcock et al. (1992); (8) De
1034 Gregorio et al. (2011); (9) Gordon et al. (2003); (10) Frati et al. (2020); (11) Ishii and Hitchcock (1988);
1035 (12) Madix et al. (1988).

1036

1037 **Table S4**

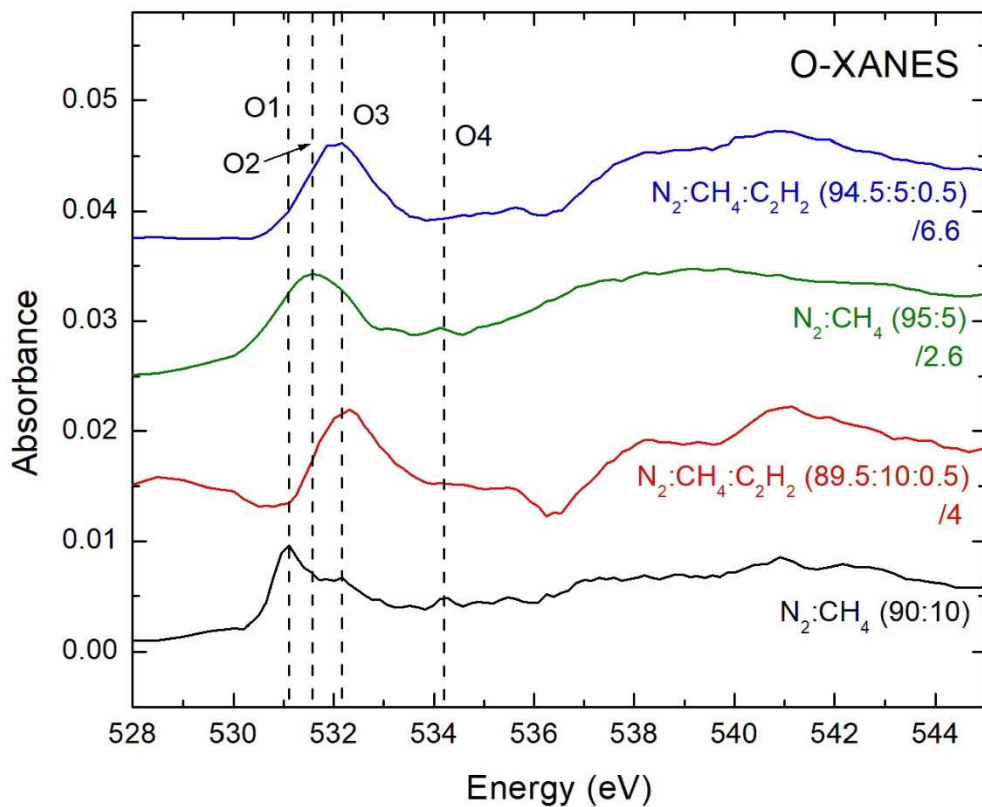
1038 Elemental C/O ratios determined from fitting the full-range XANES spectra of all tholins spanning the C,
1039 N, and O 1s edges (Fig. 4) with the atomic mass absorption coefficients (Henke et al., 1993; see Sections
1040 2.2 and 3.3 for details). Error bars are due to the fitting method used (Cody et al., 2008; Nuevo et al., 2011).

Initial gas mixture	C/O Ratio
N ₂ :CH ₄ (90:10)	2.2 ± 0.9
N ₂ :CH ₄ :C ₂ H ₂ (89.5:10:0.5)	2.7 ± 0.6
N ₂ :CH ₄ (95:5)	1.9 ± 0.4
N ₂ :CH ₄ :C ₂ H ₂ (94.5:5:0.5)	3.3 ± 0.7

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1044
 1045 **Fig. S1.** XANES spectra at the oxygen 1s edge of the four tholin samples produced in the present study.
 1046 **The intensities of the spectra are normalized to the O-edge peak intensity of the $N_2:CH_4$ (90:10)**
 1047 **spectrum. Normalization factors are provided below each gas mixture composition on the plot. The**
 1048 **main bands (O1–O4) are highlighted with dashed lines and their assignments are given in Table S3, together**
 1049 **with the assignments of minor bands. Absorbances of the spectra are normalized and offset for clarity.**
 1050

1051 **Supplementary Material References**

1052

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