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Modeling of synchrotron-based laboratory simulations of Titan's ionospheric photochemistry

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Abstract

The APSIS reactor has been designed to simulate in the laboratory with a VUV synchrotron irradiation the photochemistry occurring in planetary upper atmospheres. A N2—CH4 Titan-like gas mixture has been studied, whose photochemistry in Titan’s ionospheric irradiation conditions leads to a coupled chemical network involving both radicals and ions. In the present work, an ion–neutral coupled model is developed to interpret the experimental data, taking into account the uncertainties on the kinetic parameters by Monte Carlo sampling. The model predicts species concentrations in agreement with mass spectrometry measurements of the methane consumption and product blocks intensities. Ion chemistry and in particular dissociative recombination are found to be very important through sensitivity analysis. The model is also applied to complementary environmental conditions, corresponding to Titan’s ionospheric average conditions and to another existing synchrotron setup. An innovative study of the correlations between species concentrations identifies two main competitive families, leading respectively to saturated and unsaturated species. We find that the unsaturated growth family, driven by C2H2, is dominant in Titan’s upper atmosphere, as observed by the Cassini INMS. But the saturated species are substantially more intense in the measurements of the two synchrotron experimental setups, and likely originate from catalysis by metallic walls of the reactors.

1. Introduction

Titan’s ionosphere extends above 800 km of altitude, with local pressures no higher than 10^{-5}–10^{-6} mbar. In such diluted conditions, the efficient organic chemistry detected by mass spectrometers onboard the Cassini orbiter was unexpected, and revived our understanding of the organic chemistry occurring in Titan’s atmosphere [1–18]. Large condensable molecules responsible for the nucleation leading to solid atmospheric aerosols are now known to be produced in the ionosphere [9,19–22]. The reason for the chemical growth in spite of the low molecular density has been found in the chemical coupling between radicals and reactive charged species present in the ionosphere, i.e. positive ions, negative ions and electrons [22].

The complexity of the chemistry is however far from being charted, as illustrated by the absence of data on the dissociative recombination products for species larger than four carbon and/or nitrogen atoms [23–25], or even for modest size ions of importance in Titan’s ionosphere, such as CH3NH_2 [17]. The initial steps of molecular growth following EUV photolysis of N2/CH4/C2H2 mixtures were studied, at the second time-scale, in an ion trap by Thissen et al. [26]. No substantial growth was observed in these conditions, unless C2H2 was present in the initial mixture. On the other hand, at the much longer time-scale of Titan’s ionosphere, a correlation analysis of the positive ions densities recorded by Cassini’s Ion and Neutral Mass Spectrometer (INMS) enabled Westlake et al. [18] to infer small growth units having one or two carbon atoms.

To complete and more easily approach in the laboratory this partially ionized chemistry starting with N2/CH4, a new generation of photochemical reactors has been developed [27,28]. In this case, VUV synchrotron sources are used to simulate the VUV part of the solar spectrum, the major source of ionization in Titan’s ionosphere. A first series of results was obtained with the “Atmospheric Photochemistry Simulated by Synchrotron” (APSIS) experiment using a mass spectrometer for neutrals detection, and compared with Cassini’s data [28]. The trends on the neutral species are in...
agreement with Titan’s observations: an efficient organic growth and the formation of nitrogen-bearing products.

However no analytical device enabled the direct detection of the ions in the reactive medium. Knowing their central role in the organic growth in Titan’s atmosphere, our aim is to design a fully coupled model to theoretically probe the ion content in the APSIS reactor and interpret the results observed on the neutral species. Building an exhaustive chemical model involving all the known and partially known processes, as well as their uncertainties to perform uncertainty propagation and sensitivity analysis, prevents from a complex description of the dynamics within the APSIS reactor.

As a first approach, the study is performed with a 1-cell model, run in reference conditions corresponding to the experiments reported in Peng et al. [28]. These results are used to validate the model and perform uncertainty and sensitivity analysis to highlight prominent processes explaining the observed chemical growth. Complementary model runs have been performed for different conditions of the physical parameters (irradiation spectrum, pressure, temperature) and compared to the reference one to infer the impact of these parameters in laboratory simulations and different environments. Finally, the model has been run in conditions similar to the experiments of Imanaka & Smith [27] for comparison with APSIS, and in Titan-like conditions.

The model is established in Section 2, describing the main characteristics of the APSIS reactor, the chemical scheme, the chemistry-transport 1-cell model, and the generation of electron-impact mass spectra. The probabilistic description of the uncertain chemical parameters, and notably new wavelength-dependent photolysis branching ratios, and the method to handle it are also developed in this section. Application of the model to the APSIS reactor is presented in Section 3.1. The stationary state mole fractions of the products are analyzed and compared to experimental data. Exploration of the Monte Carlo samples generated for uncertainty analysis is performed with several objectives: (i) identification of the main uncertainty sources, in terms of reaction type and specific reactions (key reactions); (ii) detection of species communities in the chemical network to identify dominant growth pathways. This section is completed with the analysis of simulations in different physical conditions (irradiation spectrum, pressure and temperature). The following sections are devoted to the analysis of simulations of Imanaka & Smith’s reactor (Section 3.2), and to a proxy of Titan’s ionospheric conditions (Section 3.3). The conclusion outlines the main results of this research in relation with the detailed chemical modeling of N2/CH4 plasmas and the representativity of laboratory simulations of Titan’s ionospheric chemistry.

2. The N2/CH4 photochemical plasma model

2.1. The APSIS reactor

The APSIS reactor is described in detail in Peng et al. [28]. It is a stainless-steel parallelepiped with internal dimensions (length × width × height) of 500 mm × 114 mm × 92 mm. The experimental conditions chosen for the modeling are reported in Table 1. A reactive gas mixture N2/CH4 = 90/10 (purity > 99.999%, Air Liquide) is flowed into the reactor at a gas flow of 7 standard cubic centimeter per minute (sccm), controlled by a MKS mass flow controller. A primary pumping unit ensures a stationary flow of reactive gas whose partial pressure reaches 4.5 mbar and a residence time of the gas mixture of about 180 s.

The APSIS chamber is coupled windowless to the DISCO beamline at SOLEIL synchrotron facility [29], using a differential pumping system [30]. A non-reactive carrier gas (He) is continuously injected between the beamline and the APSIS chamber in order to prevent the reactive mixture to be diffusing out of the reactor into the differential pumping system and leading to a total pressure of 7.0 ± 0.1 mbar in the reactor. The experiments are conducted at room temperature.

In situ measurement of the gas phase composition is achieved using a Pfeiffer QME 200 quadrupole mass spectrometer (MS). The MS detector has a resolution of 1 u and covers the 1–100 u range. Electron energy is 70 eV. Gas is sampled by a capillary tube mobile along the reactor length and probing at about 1 cm from the irradiated column. This tube is long enough to reduce the conductance between the reactor chamber and the MS, with the consequence that only stable molecules can be detected with this setup.

The photon flux of the DISCO beamline (a few 10^11 ph s^{-1} for 0.1 nm bandwidth) leads to a N2 dissociation ratio of less than 10^{-4}. The signal at m/z 28 of N2 is therefore constant enough throughout the experiments to be considered as a fixed reference, and it is used to normalize the mass spectra (Section 2.4).

2.2. The reaction scheme

The present chemical scheme incorporates the complete neutral chemistry, i.e. photodissociation, bimolecular thermal reactions and trimolecular thermal recombinations, from Hébrard et al. [31,32], with new developments for the photodissociation branching ratios of CH4, N2, H2, C2H2, C2H4, C2H6 and HCN, following Gans et al. [33], as described in Section 2.2.1. In addition, the model implements the ion chemistry database established by Carrasco et al. [34] and Plessis et al. [23], including photo-ionizations, ion–molecule reactions and dissociative recombinations involving C, H and N. Oxygen chemistry is excluded, since O2 is just a trace gas in the reactor and, according to GC–MS analysis, does not play a significant role in APSIS photochemistry [28]. The single electron source in the APSIS reactor being photoelectrons, with insufficient energy to ionize the medium, electron impact reactions are not included.

The present chemical scheme considers only positive, singly charged ions. The DISCO spectrum intensity is vanishing below 50 nm, which prevents the double ionization of N2 [35] and CH4 [36], and therefore a sizeable production of dications. Negative ions have been identified as interesting vectors for the formation of heavy particles in Titan’s ionosphere [14] and chemical plasmas [37], but their formation by attachment of thermal electrons involves essentially minor radical species and proceeds at very small rates [14].

In total, the model includes 244 species (125 neutrals and 119 ions) reacting with each other through 1708 reactions (33 photolysis, 415 bimolecular reactions, 82 termolecular reactions, 574 dissociative recombinations, 604 ion–molecule reactions). All the parameters describing the rate constants of these processes are uncertain (some of them unknown). They are represented by random variables with prescribed distributions, following Hébrard et al. [31,32], Carrasco et al. [34] and Plessis et al. [23].

The present model is original for two reasons: (1) it implements a new consistent treatment of the separation between photolysis cross-section and branching ratios, which are typically measured in different experiments (see next section); and (2) it is the single ion–neutral coupled model implementing the state-of-the-art dissociative recombination scheme developed by Plessis et al. [23], which for the first time includes all available data for this process, notably the partial measurements of branching ratios.

2.2.1. Representation of photolysis cross-sections and branching ratios

In previous works, photolysis rate constants were treated as lognormally distributed uncertain parameters [38,32,31,39]. In Peng et al. [40] and Gans et al. [33], the separation of photolysis cross-sections and branching ratios was proposed for several
reasons, the main one being that these data are typically measured separately, and that each has a specific type of measurement uncertainty, with an essential sum-to-one constraint for the branching ratios [41]. This separation has already been systematically implemented by our team for the ion–neutral reactions [34,41,42], the dissociative recombination reactions [23], and recently for the neutral pathways of CH₄ photolysis [33]. This line was pursued here for the main photolytic processes, and their neutral and ionic products. Moreover, the elaboration of informed wavelength-dependent representations for branching ratios is all the more important, considering the flat spectrum of the DISCO beamline, in contrast with the Ly-α-dominated solar spectrum in the wavelength range of interest [33].

2.2.1.2. Branching ratios. The rather uniform spectrum of the DISCO beamline [28], calls for a continuous representation of photolysis branching ratios. However, photolysis branching ratios are often known at sparse discrete wavelengths, and large uncertainty results from the inter-/extrapolation out of these reference points. The case of CH₄ has been treated by Gans et al. [33], who developed a novel wavelength-dependent probabilistic-tree model to account for all existing information about neutral fragmentation channels. For the photochemical model of the APSIS experiments with ionization VUV photons, one has to complete this scheme with the ionic channels. This is true for all photolyzed species, and we extended the method proposed in Gans et al. [33].

The heterogeneity of information sources on branching ratio data is a major problem: the available data are typically composed of: (i) the nominal data and the corresponding uncertainties for the neutral channels of a large set of species in Hébrard et al. [31]; (ii) the SWRI nominal data for both neutral and ionic channels for only a few species, i.e. CH₄, N₂, H₂, C₂H₂, C₂H₄, C₂H₆ and HCN, [44]; and (iii) for CH₄, the probabilistic representation of Gans et al. [33]. There are overlaps between these data, and some contradictions as well. A set of branching ratios for ionic and neutral products, with the corresponding uncertainties, was designed as follows:

- For the species without photoionization channel in the wavelength range of the experiment, nominal data and uncertainties are taken from Hébrard et al. [32,31];
- For N₂, H₂, C₂H₂, C₂H₄, C₂H₆, and HCN, nominal data in the SWRI database are used, and uncertainties for neutral channels are taken from Hébrard et al. [31]. Ionic channels have no recommended uncertainty. Considering that there is generally a much higher precision for ion detection techniques than for neutrals or radicals, we use a nested scheme to preserve as much uncertainty structure as possible:

$$\text{A} + \text{hv} \rightarrow \begin{cases} \text{Neutral Channel 1} \\ \text{Neural Channel 2} \\ \text{Ionic Channel 1} \\ \text{Ionic Channel 2} \end{cases}$$

where $b_{\text{neu}}$ and $b_{\text{ion}}$ are respectively the total neutral and ionic contributions and both have a relative uncertainty of $\Delta b_{\text{neu}}$, $b_{\text{neu1}}, b_{\text{neu2}}$ ... and $b_{\text{ion1}}, b_{\text{ion2}}$ ... are the branching ratios of the neutral and ionic channels, respectively, and $\Delta b_{\text{neu1}}, \Delta b_{\text{neu2}}$ ... and $\Delta b_{\text{ion1}}, \Delta b_{\text{ion2}}$ ... are their corresponding relative uncertainties. For lack of data and according to their typical values in the literature [45], we assign the branching ratios of all ionic channels a 20% relative uncertainty. Their total contribution is typically more accurate and its relative uncertainty was estimated to $\Delta b_{\text{ion1}} = 3\%$.

- The representation of the branching ratios for the photolysis of CH₄ is based on the same scheme. We apply the representation described in Gans et al. [33] to the neutral part, with a lower wavelength limit extended to 80 nm, where the neutral channels become negligible in Huebner and Link [44]. The remaining part of the representation is the same as in the previous case.

To ensure the continuity in the interpollation/extrapolation of the branching ratio data for wavelength intervals (e.g. branching ratios provided in SWRI [44] or recommended in Hébrard et al. [31]), we need to preserve the correlation between wavelengths in a branching ratio sample. To address the nature of these wavelength-interval data, we use the generalized Dirichlet distribution (Dirg), which can be easily parameterized by nominal values and

Table 1
Summary of the conditions for the simulations in this study. *Iₚ*: integrated photon flux (from 50 to 200 nm); *L*: reactors length, *A*: reactor cross-section (width x height); *P*: total pressure; *T*; temperature; *n*: residence time; *N₂*: CH₄: composition of reactant gas; $\rho_{\text{CH}_4}$: methane relative consumption at stationary state.

<table>
<thead>
<tr>
<th>Scenario</th>
<th>Spectrum</th>
<th>$I$ (ph m⁻² s⁻¹)</th>
<th>L (m)</th>
<th>A (cm²)</th>
<th>P (Pa)</th>
<th>T (K)</th>
<th>$\tau$ (s)</th>
<th>N₂:CH₄</th>
<th>$\rho_{\text{CH}_4}$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>APSIS</td>
<td>DISCO</td>
<td>$2 \times 10^{12}$</td>
<td>0.5</td>
<td>105</td>
<td>700</td>
<td>300</td>
<td>284</td>
<td>90:10</td>
<td>0.07</td>
</tr>
<tr>
<td>60 nm</td>
<td>DISCO</td>
<td>$2 \times 10^{12}$</td>
<td>0.5</td>
<td>105</td>
<td>700</td>
<td>300</td>
<td>284</td>
<td>90:10</td>
<td>0.07</td>
</tr>
<tr>
<td>82 nm</td>
<td>DISCO</td>
<td>$2 \times 10^{12}$</td>
<td>0.5</td>
<td>105</td>
<td>700</td>
<td>300</td>
<td>284</td>
<td>90:10</td>
<td>0.07</td>
</tr>
<tr>
<td>82 nm</td>
<td>Solar</td>
<td>$2 \times 10^{12}$</td>
<td>0.5</td>
<td>105</td>
<td>700</td>
<td>300</td>
<td>284</td>
<td>90:10</td>
<td>0.07</td>
</tr>
<tr>
<td>Low $P_{\text{w}}$</td>
<td>DISCO</td>
<td>$2 \times 10^{12}$</td>
<td>0.5</td>
<td>105</td>
<td>700</td>
<td>300</td>
<td>284</td>
<td>90:10</td>
<td>0.07</td>
</tr>
<tr>
<td>Low $T$</td>
<td>DISCO</td>
<td>$2 \times 10^{12}$</td>
<td>0.5</td>
<td>105</td>
<td>700</td>
<td>300</td>
<td>284</td>
<td>90:10</td>
<td>0.07</td>
</tr>
<tr>
<td>“I &amp; S”</td>
<td>DISCO</td>
<td>$2 \times 10^{12}$</td>
<td>0.5</td>
<td>105</td>
<td>700</td>
<td>300</td>
<td>284</td>
<td>90:10</td>
<td>0.07</td>
</tr>
<tr>
<td>“Titan”</td>
<td>Solar</td>
<td>$2 \times 10^{12}$</td>
<td>0.5</td>
<td>105</td>
<td>700</td>
<td>300</td>
<td>284</td>
<td>90:10</td>
<td>0.07</td>
</tr>
</tbody>
</table>

* Ref.: Imanaka and Smith [27].
focus on using the full neutral-ion coupled chemical scheme, and, in order to be able to manage the parametric uncertainties, we have to consider a simplified 1-cell geometry, assuming no transport and a uniform spatial distribution of the species. However, the gas inflow and outflow are taken into account, and radiative transfer is considered through Beer–Lambert-type photoabsorption, but in the assumption of a uniform gas (Fig. 2). This section provides the details of model design and implementation.

2.3. Total balance and transport

It was observed that the pressure in the APSIS reactor varies very weakly during the time course of experiments and presents only a slight decrease in the direction of the gas transport [28], indicating a nearly uniform motion of the gas in the reactor. In consequence, we define a constant and uniform gas advection velocity \( \nu_{ad} \) (m s\(^{-1}\)) in the reactor. Assuming the ideal gas law, \( \nu_{ad} \) is calculated from the reactive gas inflow rate \( q_{in} \) (mol s\(^{-1}\)) and the reactive gas partial pressure \( P_{ad} \) (Pa) in the reactor by

\[
\nu_{ad} = \frac{q_{in} \cdot RT}{P_{ad} \cdot A},
\]

where \( A \) (m\(^2\)) is the reactor cross-section. The carrier gas (He) inflow rate \( q_{car} \) is the counterpart of the reactive gas in their contribution to the total pressure \( P \)

\[
q_{in} = (P - P_{ad}) \frac{A}{RT}.
\]

We assume that at the exit of the reactor, the gas is pumped out also at \( \nu_{ad} \). So as to ensure, in absence of reactions, the mass balance in the reactor. Hence, the residence time of the reactive gas \( \tau_r \) (s) is

\[
\tau_r = L/\nu_{ad}.
\]

where \( L \) (m) is the reactor’s length. In the presence of reactions, the concentration variations due to transport and chemistry should be considered simultaneously. Thus, the differential of concentration for the \( i \)th species, \( c_i \) (mol m\(^{-3}\)), can be written as

\[
\frac{dc_i}{dt} = Q_i + R_i,
\]

where \( t \) is the reaction time, \( Q_i \) and \( R_i \) (mol m\(^{-3}\) s\(^{-1}\)) are the concentration variations due to transport and chemistry, respectively.

Considering the single cell in the model, the term \( Q_i \) can be written as

\[
Q_i = \frac{\Phi_{in,i} - \Phi_{out,i}}{L},
\]

where \( \Phi_{in,i} \) and \( \Phi_{out,i} \) (mol m\(^{-2}\) s\(^{-1}\)) are respectively the influx and the outflux of the \( i \)th species: \( \Phi_{in,i} = \dot{f}_{in,i} \times q_{in}/A \), \( \Phi_{out,i} = \dot{f}_{out,i} \times q_{in}/A \), \( \dot{f}_{in,i} = q_{in}/A \) and \( \dot{f}_{out,i} = 0 \forall i \neq \{N_2, CH_4, He\} \), where \( \dot{f}_{in,i} \) and \( \dot{f}_{out,i} \) are the proportions of the reactants in the incoming gas. For all species, one has \( \Phi_{out,i} = 0 \forall \dot{f}_{out,i} \).

2.3.2. Radiative transfer and photochemistry rate constant

The photochemistry rates depend on the actinic flux \( I(\lambda,i) \) (ph m\(^{-2}\) s\(^{-1}\) nm\(^{-1}\)) at wavelength \( \lambda \) and distance \( x \) from the entrance of the light beam in the reactor (optical path). Considering Beer–Lambert absorption, we have

\[
I(\lambda,i) = I_0(\lambda) \exp\left(-\int_0^x \tau_i(x,\lambda) \, dx \right),
\]

where \( I_0(\lambda) \) describes the photon flux at the entrance, \( \tau_i(x,\lambda) \) is the wavelength-dependent optical depth of the \( i \)th absorber. It depends on the species concentration \( c_i(x) \) along the optical path and the photoabsorption cross-section \( \sigma_i(\lambda) \) (m\(^2\)mol\(^{-1}\)):

\[
\tau_i(x,\lambda) = \sigma_i(\lambda) \int_0^x c_i(x) \, dx.
\]
The photolysis rate constant of the jth channel of the ith absorber at position x, is given by

\[ J_{ij}(x) = \int l(x, \lambda) \sigma_i(\lambda) b_j(\lambda) d\lambda, \]  

(10)

where \( b_j(\lambda) \) is the branching ratio at wavelength \( \lambda \) of this channel.

As there is a single cell in the model design of the present study, an average value of the photolysis rate constant has to be used. \( J_{ij}(x) \) at the cell center, \( J_{ij}(x) \), is commonly chosen in atmospheric modeling studies [38], but this is not appropriate for our model, where the rate constant can strongly vary along the optical path. Instead, we calculate the average of \( J_{ij}(x) \) along the reactor length,

\[ \frac{1}{L} \int_0^L J_{ij}(x) dx. \]  

(11)

Based on the assumption of uniform gas density, one can easily derive

\[ \frac{1}{L} \int_0^L J_{ij}(x) dx = \int l(x, \lambda) \sigma_i(\lambda) b_j(\lambda) \frac{1 - e^{-\sum \sigma_i(\lambda)}}{\sum \sigma_i(\lambda)} d\lambda. \]  

(12)

To evaluate this integral, the wavelength is discretized with a resolution of 1 nm, adapted to the spectral resolution of the cross-sections database of Hébrard et al. [31]. The additional photolysis data used in this study, from the SWRI database [44], are interpolated on this wavelength grid using a Gauss–Legendre quadrature [46].

2.3.3. Integration method

The implicit-explicit Runge–Kutta–Chebyshev (IRKC) method [47], capable of integrating systems of partial differential equations (PDEs), is applied to the present model. It handles the equations by both an explicit part \( F_E(t, y) \) and an implicit part \( F_I(t, y) \), i.e.

\[ y'(t) = F_E(t, y) + F_I(t, y). \]  

(13)

For a transport–reaction model, it is suggested that \( F_E \) and \( F_I \) respectively describe the parts of a cell’s differential which are respectively dependent (e.g. diffusion and photolysis) and independent (e.g. thermal reactions) on other cells. The latter, \( F_I \), is usually regarded as a system of ordinary differential equations (ODEs) and can be very stiff, which is the case for our coupled ion–neutral model. The IRKC method is able to cope with the problem of stiffness in \( F_I \) [48].

The present model has no internal transport. In theory, the whole model could therefore be considered as a set of ODEs and solved in \( F_I \) or by other ODE solvers. However, the photolysis is treated in \( F_E \), as in transport–reaction models, because we treat the radiative transfer explicitly, and complex evaluation of Jacobian elements for the photolysis makes it impractical to regard the photolysis rate equations as ODEs.

Besides, IRKC enables to account for diffusion: for further studies in higher dimension (1D or 2D), IRKC will be well adapted to a transport–reaction model. The part of thermal reactions, independent on other cells, could be treated implicitly and separately in each cell, so that it could be repeatedly implemented by the program. This leads to small storage requirements and good computing performance.

Tests were performed to check the convergence and stability of the stationary state of the present model integrated by the IRKC method. The stationary state is very robust to either perturbations of initial conditions or the choice of propagator tolerance parameters.

2.3.4. Uncertainty management

As the chemical parameters of the model are intrinsically uncertain, one has to evaluate the impact of this uncertainty of the input parameters on the model outputs (mole fractions of products). The formula of the propagation of distributions [49]

\[ p(Y = y) = \int dX_1 dX_2 \ldots dX_n p(X_1, X_2, \ldots, X_n) \times \delta(y - f(X_1, X_2, \ldots, X_n)), \]  

(14)

provides a transformation from the joint probability density function \( p(X_1, X_2, \ldots, X_n) \) of all input parameters, to that of the output(s) \( Y = f(X_1, X_2, \ldots, X_n) \). From \( p(Y) \), one can compute all the necessary statistics.

It is rarely possible to perform the integration above analytically. Instead, the BIPM recommends the use of Monte Carlo Uncertainty Propagation (MCUP) methods [49], in which we propagate representative samples of the inputs distributions, rather than the analytic distributions themselves. We generate random sets of input parameters according to their probability density function, and then, for each set, evaluate the corresponding outputs of the model, thus building a representative sample of the outputs distribution.

The MCUP method is convenient to treat models with many inputs and outputs. Its convergence does not depend on the number of input parameters. Moreover, it is directly applicable to multi-outputs models, through a single set of samples. As a result, the intrinsic correlation between the outputs is preserved and can be used for further analysis (Section 3.1.3).

However, the MCUP method is expensive in computer time (repeated runs of the model necessary to generate significant statistics). We limit our sampling to 500 runs, with random samples generated for each process type according to Carrasco et al. [34], Hébrard et al. [32], Plessis et al. [23] and Gans et al. [33].

2.4. Generation of mass spectra

We present here the procedure to generate electron impact mass spectra from the simulated neutral species concentrations. We adopt a 1 u mass resolution, for direct comparison with experimental spectra. The intensity, \( I_j \), of a mass peak at integer \( m/z \) \( j \) is modeled by

\[ I_j = \sum_j \sigma_j c_i f_i, \]  

(15)

where \( \sigma \) and \( c \) are respectively the electron impact cross-section and the concentration of the ith species, \( n \) is the maximal mass, and \( f_i \) is the fragmentation pattern of the ith species at \( m/z \) \( j \), with the constraint \( \sum_j f_i = 1 \) [50].

The electron impact cross-sections for a \( C_2H_8N_2 \) species are modeled by the formula proposed by Puccio and Miller [50]

\[ \sigma_j / A^2 = 0.460x + 1.086y + 1.348z. \]  

(16)

The fragmentation patterns for almost all stable species up to 6 heavy (C, N) atoms and some heavier species are extracted from the NIST Chemistry Webbook [51], well covering the stable species of interest in the present study, and including Earth natural isotopic abundances. However, this database provides no fragmentation data for radicals and there are only few data of this type in the literature. In consequence, we chose to exclude the radicals from the MS. Finally, we average the fragmentation patterns of isomers, if they are not distinguished in the model. For example, \( C_2H_6 \) in our model represents both propene and cyclopropane: their fragmentation patterns are averaged with equal weights.

In details, the generation of the neutral MS from a Monte Carlo sample of species concentrations proceeds according to the following steps:
1. Perform the Monte Carlo loop by calculating the kth sample of the intensities $I^{(k)}_j$ \((j = 1, 2, \ldots; n; k = 1, 2, \ldots m)\) at different m/z from the kth of m samples of the species concentrations $C^{(k)}_i$ by

$$I^{(k)}_j = \sum_{i} \sigma C^{(k)}_i f_j, \quad j = 1, 2, \ldots; n; \quad k = 1, 2, \ldots m.$$  

2. Normalize, within each set of samples, the intensities with respect to that at m/z 28 (molecular ion signal of N$_2$, the most abundant species), for comparison with other experimental mass spectra.


Note that the estimated impact cross-sections (Eq. (16)) and the NIST fragmentation patterns [51] are uncertain. However, compared to the relative uncertainties on the kinetic parameters and the species concentrations (often larger than 100%), their relative uncertainties (∼10% [50]) are treated as negligible.

There are some sources of bias to be considered when comparing the simulated MS to experimental ones: (i) the APSIS and the NIST standard MS were measured at different pressures, which may significantly influence the fragmentation patterns [28]; (ii) different mass spectrometers may overlap the major peaks (e.g. m/z 28) with neighboring peaks in different ways, interfering in the determination of the intensities of these minor peaks. These effects are difficult to quantify, and are not accounted for in the generation of the simulated MS.

3. Results and discussion

The 1-cell model has been run in reference conditions corresponding to the APSIS experiments [28]. These results are used to evaluate the model and perform uncertainty and sensitivity analysis. Additional sets of model runs have been performed for different conditions of the physical parameters (irradiation spectrum, pressure, temperature) and compared to the reference one to infer the impact of these parameters in laboratory simulations. Finally, the model has been applied to conditions similar to the experimental ones [28].

The 1-cell model has been run in reference conditions corresponding to the experimental mass spectra. The total mole fraction of ions is about $10^{-3}$. The decrease of ion concentrations with mass is less evident than for neutrals. Most of the major ions (from C1 to C10) have mole fraction between $10^{-10}$ and $10^{-15}$.

The stationary state CH$_4$ consumption ratio

$$\delta_{CH_4} = \frac{[CH_4]_0 - [CH_4]_S}{[CH_4]_0},$$

in the experiments and in the model are respectively $0.30 \pm 0.06\%$ and $0.07 \pm 0.01\%$. They are not in perfect agreement, but, considering the model simplicity and absence of adjustable parameters, a factor of 4 is tolerable. More specifically, the simplified description of advection and radiative transfer in the 1-cell model may result in a bias of the effective gas residence time and in consequence of the methane consumption.

Some species predicted by the present model were not identified in the APSIS in situ mass spectra, e.g. NH$_3$ and CH$_2$NH, because their parent MS peaks are hidden by the signals of OH$^+$ and 14$^N$NH$. But their formation is confirmed in the PAMPRE reactor in similar conditions [52], and thus likely also in the APSIS experiments. Besides, the model predicts the formation of other species with a concentration below the detection limit of the APSIS MS measurements, e.g. most C5 and C6 species, especially benzene.

The mole fractions (except for N$_2$ and CH$_4$) were summed by block. In each Cx block, the distribution of the results is expressed by a violin plot, whose widths at different mole fraction values show the corresponding probability density (Fig. 4). Experimental results from the 7 sccm APSIS experiments [28] have been overlaid for comparison. The experimental mass spectrum intensities have been divided by their relative (to N$_2$) ionization cross-sections, estimated according to Eq. (16), i.e. 3, 4 and 5 for the C2, C3 and C4 blocks, respectively. Note that the Cx (x > 2) abundances follow a powerlaw distribution, as described by Dobrijevic and Dutour [53].

The simulation predictions are in good agreement with the APSIS MS measurements (Fig. 4), except for the C2 block, for which the large measurement uncertainties of the intensities on the N$_2$ peak shoulders [28] have to be considered. Note that this fair agreement also somewhat results from the large uncertainty on the model predictions.

In the reference simulation, radicals account for about one third of the total mass of products at stationary state. Thus, the simulated mass spectra, ignoring radicals, can only be compared to the APSIS MS on a qualitative or semi-quantitative basis (see Fig. 5). Except for the C2 block, the major bands in the experimental spectrum are well covered by the 95% confidence intervals of the simulated MS (the red shaded area in Fig. 5). However, the mean of the simulated spectrum is not in satisfactory agreement with the experimental spectrum: it underestimates the production of saturated C3 and C4 MS fragments and overestimates that of unsaturated C4 fragments. In the database used to build the spectra, the major fragmentation patterns of hydrocarbons might change the number of heavy atoms, but have a strong tendency to preserve the saturation state. For instance, the main fragment of C$_3$H$_6$ is at m/z 29 (C$_2$H$_5$) vs. 30 (C$_2$H$_4$), that of C$_6$H$_{10}$ is at m/z 43 (C$_2$H$_7$) vs. 44 (C$_3$H$_8$), and for the unsaturated species, that of C$_4$H$_8$ is at m/z 52 vs. 58 (C$_5$H$_{10}$). Without considering specific details, one can conclude from this comparison that the experiment and simulation differ mostly by their production in saturated species.

The main causes of this discrepancy could be:

- incompleteness of the reaction scheme: an exhaustive search has been performed for the construction of the reaction scheme. Up to now, most C3-producing reactions have been included in
the model, while saturated C3 species are still not satisfactorily simulated. Moreover, it is unlikely that missing reactions, randomly taken for each reaction type, could result in a systematic underestimation of saturated species,

- oversimplification of the reactor geometry: we considered the effect of a dark zone (the region in the reactor not or weakly irradiated) by running the model without irradiation but with a gas inflow composition identical to that of the stationary state.

Fig. 3. Mole fractions (normalized to N2) of neutral species (left panel) and ions (right panel) at the stationary state for the 500 runs of the 1-cell model with all uncertain parameters. The species are sorted downwards by increasing mass. Only species with an average mole fraction above $10^{-15}$ are represented and CH4 has been left out of the graph for legibility. The Log10 Mole Fraction is $-0.9545 \pm 0.0001$. Cx blocks are materialized by alternating colors. Neutral species with even/odd electron counts are marked by a square/circle symbol. Error bars cover a 95% confidence intervals.
gas outflow in the reference case. The gas inflow models the non or weakly irradiated part of the reactor, where reactive species other than N2 and CH4 are supplied by transport/diffusion. The stationary state of this test case is little shifted from the reference case, demonstrating, at least, that the basic growth scheme would hardly change in the dark zone of the reactor.

- **chemistry taking place in the capillary between the reactor and the mass spectrometer:** radicals and ions (about one third of the total mass of products) pumped into the capillary cannot qualitatively change the gas composition before reaching the mass spectrometer. This was shown by another test case, where we let the stationary state reactive gas of the reference case react until equilibrium in a closed box, without irradiation. In this case, radicals and ions have infinitely long time to combine with each other and be recominated with electrons, no matter whether catalyzed by the capillary stainless steel surface or not. In these conditions, saturated species concentrations are increased by at most 40%, still far from the observed difference between the experimental and simulated spectra. However, the increase of saturated species concentration resulting from radical recombinations in this test case, points out that higher radical recombinations rates would lead to a better reproduction of saturated species.

- **wall effects:** how could radical recombinations be higher than in our reference model? It is unlikely due to homogeneous trimolecular reactions, since, although globally known with large uncertainty, their rate constants could hardly be systematically underestimated. Therefore, we may only consider heterogeneous catalysis, most probably on the reactor stainless steel walls, as detailed below.

Radicals and ions can be efficiently adsorbed or chemisorbed on stainless steel walls and undergo further chemistry. For instance, it has been observed that NH3 can be generated in this manner by successive hydrogenation of atomic nitrogen and N-containing radicals [54]. In their laboratory simulation experiments of Titan’s ionospheric chemistry Thissen et al. [26] concluded that surface effects were responsible for an excess production of NH3 and HCN. Radicals recombinations on the wall of the antechamber of the INMS mass spectrometer has been retained to explain the discrepancy between measured concentrations of C6H6 along the trajectory of the spacecraft in Titan’s atmosphere [11]. Overall, wall effects in CH4/N2 plasmas are very complex and are the subject of ongoing studies [55].

Gorodetsky et al. [56] measured a CH3 recombination coefficient of 3 × 10^-14 on stainless steel at 300 K. According to this value, the reaction rate of CH3 recombinations on stainless steel is one to two orders of magnitude below that of the homogeneous rate. They determined also the temperature dependency of the rate constant, usually more reliable than its absolute value. They showed that recombinations is accelerated about 10 times as temperature increases by 10 K. This high sensitivity may make the APSIS recombinations rate highly uncertain, since the temperature during the experiments is not regulated. In addition, the large uncertainty usually associated with measurements of heterogeneous reaction rates and the existence of multiple radical recombinations on the reactor’s wall should also be considered. The large uncertainties prevent us from making a quantitative comparison of the contributions of the homogeneous and the heterogeneous radical recombinations, but it cannot be ruled out that heterogeneous recombinations have a comparable, or even larger contribution than the homogeneous ones.

Without detailed reaction rate constants, it is difficult to model heterogeneous recombinations explicitly without introducing adjustable parameters in the model [54]. In order to account for heterogeneous recombinations in the present model, we made simulations where all the homogeneous trimolecular rate constants were multiplied by a factor 3 or 10 (cyan and dark green solid lines in Fig. 5, respectively). In the case of the factor 3, the unsaturated part of the C3 and C4 bands decreases slightly and their saturated parts increase significantly, by a factor about 3, leading to a better agreement of the C3 and C4 blocks. When trimolecular rate constants are 10 times larger than in the reference case, one gets an overestimation of the saturated part.

We observe that faster trimolecular recombinations produces more saturated species, at the expense of unsaturated species. Unsaturated species play a major role in the growth without “heterogeneous recombinations” included in the model, and their preponderance over saturated ones, not observed in the experiments, can be well compensated in the model by “heterogeneous recombinations” that consumes unsaturated species and produces saturated ones. This is a strong indication of the contribution of heterogeneous reactions in the laboratory simulation setups of atmospheric chemistry, such as the APSIS reactor.
3.1.2. Uncertainty analysis

Uncertainty analysis for simulations of N$_2$/CH$_4$ plasmas were provided for Titan's atmosphere by Hébrard et al. [38,32] and Peng et al. [39] for neutral species predicted by a neutral model and by Carrasco et al. [34,42] and Plessis et al. [25] for ions predicted in a fixed neutral bath ionospheric model. The impact on predictions uncertainty of a fully coupled ion–neutral chemistry is assessed here for the first time. A particular concern was resulting from the introduction in the model of the exhaustive database of dissociative recombination branching ratios (with large uncertainties) of Plessis et al. [23] and its strong coupling with the neutral chemistry [25], and also from the large uncertainties in the branching ratios of the photolysis of CH$_4$ at most wavelengths covered by the DISCO spectrum [33,28].

In the following, we discuss the uncertainty factors of the mole fractions in these various models. These are defined as $F_i = \exp(U_{\log y_i})$, where $U_{\log y_i}$ is the standard deviation of the logarithm of the mole fraction $y_i$ of species $i$, estimated on the Monte Carlo sample. If the distribution of $y_i$ is lognormal, then one has about 95% probability for the mole fraction to be in the interval $[y_i, y_i \times F_i]$, where $y_i = \exp(\log y_i)$ is the geometric mean. As a thumbrule, mole fractions with uncertainty factors of about 3 are known up to an order of magnitude.

Empirical Cumulated Density Functions (ECDF) of the uncertainty factors on the stationary mole fractions of neutrals and ions of the APSIS simulation are reported in Fig. 6, where we display also the ECDF for the uncertainty factors obtained in Hébrard et al. [32] for neutral species at 1200 km in Titan's ionosphere, and by Carrasco et al. [34] at the same altitude for a ionic model with fixed neutral bath. The uncertainty factors for both ions and neutrals are globally increased when compared with their uncoupled versions. The effect is stronger for ions, a direct consequence of their production from NH$_2$, itself provided for Titan's atmosphere by Hébrard et al. [38,32] and Peng et al. [39]. These authors ranked this uncertainty source as a major one in the uncertainty budget of predicted ions densities, which is confirmed by the present study. Globally however, 60% of the ions are predicted with smaller uncertainty than neutrals, an effect of the reaction rate constants of ionic processes being usually measured with better precision.

3.1.3. Sensitivity analysis

3.1.3.1. Main contributions to prediction uncertainty. To better understand the large uncertainty factors in the predictions of the present model, we analyze their variance with regard to the main types of reactions. For this, we calculate the portion of the total variance explained by each type of process: photolysis, reactions (neutral and ionic bi- and trimolecular), and dissociative recombination, by running simulations where a single type of process is treated as uncertain, while the others have fixed rate constants.

The results are summarized in Fig. 7, where we report, for each type of process, the frequency of it being the dominant contributor to the output variance of the stationary mole fraction of neutrals and ions. For reference, we plot also the occurrence frequency of each reaction type in the model (see Section 2.2). The scores for neutrals and ions are remarkably correlated to this occurrence frequency, with minor deviations for both categories. The contribution of bi- and trimolecular reactions to the mole fractions variance in the reference case is dominant, from nearly 80% for neutrals to 60% for ions. Dissociative recombination contribute significantly, with a score of about 30% for both ions and neutrals, slightly lower than the occurrence frequency of this process. Photodissociation and photoionization appear as negligible for neutral species, in accordance to the weak number of these processes in the model, but have an enhanced contribution to 15% of the cases for ions, for which photoionization is a necessary initiation pathway. It is to be noted that the highly uncertain representation of the photolysis branching ratios of methane and other photoborbers (see Section 2.2.1) does not impact significantly on the uncertainty budget of this simulation. Similarly, the contribution of dissociative recombination does not seem to be enhanced by the coupled chemical scheme. For both photolysis and dissociative recombination, the care taken in the representation of branching ratios as sum-to-one variables is certainly repaid for in the uncertainty budget.

3.1.3.2. Key reactions identification. Beyond the overall view provided by the previous variance analysis, we aim at the identification of key reactions, defined as the reactions that contribute significantly to the model output uncertainty [57]. A key reaction...
might appear for two different reasons: (i) reactions that have a strong effect on the mass flux in a reaction network; (ii) reactions with a rate constant in need of further study to improve the prediction precision of model outputs; or both. A few recent studies identified the key reactions of complex chemical networks by computing the rank correlation coefficients between rate constants and species concentrations from the Monte Carlo samples [32,39,57,58]. To select their key reactions, Hébrard et al. [32] and Peng et al. [39] retained reactions presenting correlation coefficients larger than 0.2 in absolute value with the mole fractions of more than 15 species. We adopted the same procedure, which produced 14 key reactions, as listed in Table 2 with the corresponding number \( N > \) of above-threshold species, and reaction rate uncertainty factors \( F_k \) estimated from the input Monte Carlo sample.

In order to confirm this selection, we estimated the global influence of a reaction through the sum of the squares of its correlation coefficients with the stationary mole fractions of all species. We define the score \( S_j \) of the \( j \)-th reaction as

\[
S_j = \sum_{i=1}^{N> \text{species}} r_{ij}^2
\]

where \( r_{ij} \) is the rank correlation coefficient between the \( i \)-th species and the corresponding reaction rate. The scores are reported in Table 2. All the reactions with the higher \( S \) scores are also in the top list selected by the other method, and only the scoring order is slightly perturbed.

**Table 2**

<table>
<thead>
<tr>
<th>Reaction</th>
<th>( F_k )</th>
<th>Scores</th>
</tr>
</thead>
<tbody>
<tr>
<td>( H + \text{C}_2\text{H}_3 \rightarrow \text{C}_2\text{H}_2 + \text{H}_2 )</td>
<td>3.9</td>
<td>55 6.9</td>
</tr>
<tr>
<td>( \text{H}_3 + \text{C}_2\text{H}_6 \rightarrow \text{C}_3\text{H}_5 )</td>
<td>2.0</td>
<td>48 6.6</td>
</tr>
<tr>
<td>( \text{H} + \text{CH}_3 \rightarrow \text{CH}_4 )</td>
<td>2.9</td>
<td>39 6.9</td>
</tr>
</tbody>
</table>

In order to confirm this selection, we estimated the global influence of a reaction through the sum of the squares of its correlation coefficients with the stationary mole fractions of all species. We define the score \( S_j \) of the \( j \)-th reaction as

\[
S_j = \sum_{i=1}^{N> \text{species}} r_{ij}^2
\]

where \( r_{ij} \) is the rank correlation coefficient between the \( i \)-th species and the corresponding reaction rate. The scores are reported in Table 2. All the reactions with the higher \( S \) scores are also in the top list selected by the other method, and only the scoring order is slightly perturbed.
The list of key reactions is heterogeneous in the sense that, as stated above, it contains reactions with high uncertainty factors (above 3) and reactions with small uncertainty factors (below 1.5).

The first ones (e.g. $\text{H} + \text{C}_2\text{H}_3 \rightarrow \text{C}_2\text{H}_2 + \text{H}_2; F_k = 3.9$) contribute importantly to the uncertainty budget of the simulation, and more accurate estimations of their rate constants would have a positive impact on prediction precision at a global level. In the same category, $\text{N}_2^+ + \text{CH}_4 \rightarrow \text{CH}_2^+ + \text{H}_2 + \text{N}_2 (F_k = 3.6)$ is a minor channel of a reaction with a rather accurate global rate constant ($F_k = 1.15$) which would benefit of a better evaluation of its branching ratio ($b = 0.08 \pm 0.08$; Carrasco et al. [42]). This global reaction is an important step in the formation of primary ions from $\text{CH}_4$. Another example is the dissociative recombination of $\text{CH}_2\text{NH}^+$ (the third most abundant ion in the reference simulation after $\text{HCNH}^+$ and $\text{C}_3\text{H}_7^+$, see Fig. 3), for which both the rate constant and the branching ratios are unknown. The strong impact of this process on some species has been noted in Section 3.1.2. This hampering lack of data was also mentioned by Yelle et al. [17] for a Titan ionospheric model. The branching ratios of this reaction determine the portion of $\text{CH}_2\text{NH}^+$ converted to $\text{CH}_2\text{NH}$, methanimine, a possible precursor of tholins [59].

In the second group of reactions, one has for instance the photolysis of $\text{N}_2$. This process is essential to the formation of ions and
integration of N in the system, but it is rather hopeless to expect a reduction of its already small uncertainty factor ($F_k < 1.1$).

It is striking that two trimolecular reactions appear at the top of the list. In the model of Titan’s upper atmosphere [32], most key processes were photolysis and bimolecular reactions, and only one trimolecular reaction was found to be influential, at low altitude. Indeed, trimolecular reactions are influential in the APSIS model because their rate is enhanced by the comparatively high pressure in the reactor. We note also that, despite the sizable uncertainty on its rate constant ($F_k \simeq 7$), $H + CH \rightarrow C + H_2$, the

![Heatmap of the outputs correlation matrix. Correlation coefficients from −1 to 1 are denoted by squares whose color gradually changes from blue (−1) to red (1) via white (0). The colored borders (orange, purple, green) delimit the clusters discussed in the text. The hierarchical tree is displayed at the top of the matrix, and the relevant clusters G1, G2 and G3 are underlined by color. (For interpretation of the references to colour in this figure caption, the reader is referred to the web version of this article.)](image-url)
most influential reaction in the model of Titan’s upper atmosphere reported by Hébrard et al. [32], is not identified as a key reaction for the present simulation, although we will see that it reappears in the list for simulations in Titan-like conditions (Table 4).

Two reactions involving CHCN appear in the list. This species plays a key role in the formation of C6N2 and C4N2. An issue with its chemistry has been raised by Imanaka and Smith [27], and we report the discussion to Section 3.2.3.

3.1.4. Species correlation analysis

Although we know in every detail the reaction scheme of the model, its complexity and the high uncertainty on many kinetic parameters make it delicate to predict the main pathways without a numerical search. We propose here to exploit the sample of species mole fractions generated for uncertainty/sensitivity analysis to gather information on the relationships between species through correlation analysis.

The correlation coefficient between the stationary mole fractions of two species calculated from the MCUP outputs sample contains information on the way these species respond to a global perturbation of the system (i.e. random variation of all the rate constants). Considering a scheme with a single reaction, e.g. \( R_1 = R_2 \rightarrow P_1 + P_2 \), and taking the concentrations at a given time, one expects a strong (unit) positive correlation between reactants \( r_{R_i R_j} = 1 \) and between products \( r_{P_i P_j} = 1 \), and a strong anticorrelation between reactants and products \( r_{R_i P_j} = -1 \). For more complex systems, the correlation strength depends on the structure of the reaction network between two species, and on the relative effects of various reaction rate constants on those species.

### 3.1.4.1. Inter-species correlation matrix

In a first step, we consider the inter-species correlation matrix, and more particularly examine the presence of structures and their link to the underlying chemical scheme. The rank correlation matrix of mole fractions at stationary state for the APSIS simulation is shown in Fig. 9, where the species are sorted (i) by charge; and (ii) by mass. The Cx blocks are overlined and labeled by the corresponding number of heavy atoms.

At the global scale, one observes that the neutrals and ions present rather different correlation patterns, with a massive inter-blocks correlation structure for the ions, whereas the correlation pattern seems more specific between neutral species. There is also a notable correlation pattern between neutrals and ions. The inter-ions structure is reminiscent of the inter-mass correlation matrix built recently by Westlake et al. [18] from the intensities of INMS ion data between 1200 and 2000 km, over a large sample of Cassini flybys. Westlake et al. [18] observed a block pattern in the matrix and linked it to molecular growth through the addition of C1 and C2 units.

If we enter into more details, we observe several features in the neutral and ion Cx blocks along the diagonal: some species within a Cx block form a red square, hinting to a similarity of their reactivity. The most striking example arises in the C6 neutral block (see the inset of Fig. 9), where two squares of red pixels are visible: a square in the lower left of the block, between CH3H and CH2H, relatively unsaturated species, and another in the upper right, between CH4H and CH2H4, relatively saturated hydrocarbons. Out of these two squares, there are off-diagonal light blue pixels, which indicate a negative correlation of species between the two squares. We interpret this pattern as revealing separate and competing pathways for unsaturated and saturated species in this block. This pattern is present in other blocks, although with less clarity.

Inter-blocks correlation may provide information on molecular growth. Highly positive correlation between two species with different heavy atom numbers may not only suggest a similar formation pathway, but also a direct link by a reaction realizing molecular growth, i.e. the addition of small chemical units. Positive inter-block correlations are obvious between most saturated heavy neutrals, or between most unsaturated heavy neutrals. They are more common between heavy ions, either between unsaturated ones or between saturated ones.

In general, unsaturated and saturated ions are positively correlated to their respective neutral counterparts. Saturated species have negative correlations with unsaturated ones, regardless of whether they are neutrals or ions. Given all that, saturated and unsaturated species are likely to be formed through distinct growth/formation pathways.

### 3.1.4.2. Cluster analysis

Except for simple chemical schemes, information based on inter-species correlation is too condensed to enable the reconstruction of a detailed mechanism, but we can use it as a basis for a cluster analysis, grouping together species responding similarly to the perturbation. Species within a cluster are then assumed, through their similarity, to be part of a common pathway. For instance, in the case of molecular growth which is our main target here, we expect that all species depending on a given growth unit will share a cluster, possibly with this unit.

The clustering analysis was done using the heatmap function in R [60], which builds a hierarchical clustering tree. The heatmap function plots a matrix, with rows and columns reordered to match the dendrogram plotted in the margins. The dendrogram is obtained by a hierarchical clustering algorithm hclust. The clusters are built iteratively, starting by one matrix row (column) per cluster. We used the default agglomeration method in the hclust function, the complete linkage method, which finds and groups similar clusters [61]. The correlation matrix (Fig. 9) reordered by the hierarchical clustering method is displayed in Fig. 10.

We focus on the largest clusters of positively correlated species. The three main ones (named G1, G2 and G3), totaling 133 species over the whole 244, are emphasized in the heatmap. The list of species in these clusters is provided in Table 3.

A large cluster of 74 positively correlated species (orange border) is seen at the bottom right. In this group, named G1, 77% of

### Table 3

Species lists, sorted by increasing molecular mass, from the three clusters identified in Fig. 10. The species with the highest score S (see text) in each group are reported in bold.

<table>
<thead>
<tr>
<th>Cluster name</th>
<th>Species in cluster</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1</td>
<td>CH3, CH3H, CH2CN, CNH3, NCN, C2H7, C2H5, C2H5H, C2H5H3, C2H5H4, C2H5H5, C2H5H6, C2H5H7, C2H5H8, C2H5H9, C2H5H10, C2H5H11, C2H5H12, C2H5H13, C2H5H14</td>
</tr>
<tr>
<td>G2</td>
<td>CH3, NH, CH3CN, CH3NH2, CH3NH3, CH3NH4, CH3NH5, CH3NH6, CH3NH7, CH3NH8, CH3NH9, CH3NH10, CH3NH11, CH3NH12, CH3NH13, CH3NH14</td>
</tr>
<tr>
<td>G3</td>
<td>CH3, CH3H, CH2CN, CH3CNH, CH3CNH2, CH3CNH3, CH3CNH4, CH3CNH5, CH3CNH6, CH3CNH7, CH3CNH8, CH3CNH9, CH3CNH10, CH3CNH11, CH3CNH12, CH3CNH13, CH3CNH14</td>
</tr>
</tbody>
</table>

Note: The table includes species with increasing mass from the clusters identified in Fig. 10. The species with the highest score S (see text) are in bold.
the members are ions and 47% are N-bearing species. At the top left of the heatmap, another cluster (G2 – green square) contains 29 species, and is composed for 55% of ions and 83% of N-bearing species. The third remarkable cluster, G3, is located near the middle of the heatmap (purple border) and contains 30 species. It is composed for 87% of neutral hydrocarbons.

The mass distribution and van Krevelen diagram [62,63] for the three groups are given in Fig. 11. The three clusters contain heavy species, but the G1 cluster contains the largest proportion of the heavier ones. At the compositional level, there is a neat discrimination of the three groups in the (N/C; H/C) space: the G1 cluster contains mostly H-poor (H/C≤1) species, the G3 cluster is essentially composed of rather saturated hydrocarbons (H/C>1), while the G2 cluster contains more saturated species than the first one.

As we did for key reactions, we define within each cluster a score $S$ for each species according to its cumulative squared correlation coefficient with the other species in the cluster (cf. Eq. (19)).

Cluster G1. The species with the highest score within G1 is by far C$_2$H$_2$ ($S = 15.2$).

All of the species in the cluster are rather unsaturated and most of them are ions. N-bearing species in this cluster are secondary products of the hydrocarbons discussed next. Considering these features, as well as the mechanism of molecular growth by addition of small units discussed previously, we identified a growth pathway through the recursive addition of C$_2$H$_2$ by ion–molecule reactions. A typical pathway is

$$\begin{align*}
C_2H_2 & \xrightarrow{+H} C_2H_3 \\
C_2H_3 & \xrightarrow{+H} C_2H_5 \\
C_2H_5 & \xrightarrow{+H} C_2H_7 \\
C_2H_7 & \xrightarrow{+H} C_2H_9
\end{align*}$$

(20)

In the chemical scheme, C$_2$H$_2$ can be added to most ions with a sizable rate constant, while increasing the degree of unsaturation in the products. In their synchrotron irradiation study of different N$_2$/CH$_4$/C$_2$H$_x$ gas mixtures, Thissen et al. [26] have observed that C$_2$H$_2$ is the most efficient C2 compound to promote ions growth.

On the other hand, the importance of the ion chemistry does not mean that the contribution of the neutral chemistry is negligible. A neutral growth scheme was highlighted by Peng et al. [39]:

$$\begin{align*}
C_2H & \xrightarrow{+R} C_2H_2 \\
C_2H_2 & \xrightarrow{+R} C_4H_2 \\
C_4H_2 & \xrightarrow{+R} C_6H_2
\end{align*}$$

(21)

where $R$ represents various radicals. This scheme may also be typical for the production of unsaturated species, albeit a hydrogen...
abstraction step is necessary to reactivate the intermediate. Otherwise 2 hydrocarbons of even hydrogen atom number, usually stable species, are not able to react with each other. Growth pathways by repeated addition of C2H to form polyacetylenes as proposed in Ar/C2H2 plasmas [64], or studied in molecular jets [65], are found to be negligible in APSIS conditions, where the very low concentration of C2H (Fig. 3) rapidly dampens this process.

Cluster G2. The species with the highest score within G2 is CH3NH2 (\(S = 10.9\)), with close scores for NH2 (\(S = 10.6\)) and NH3 (\(S = 10.2\)). Moreover, 83% of the species in this block are N-bearing ones. The cluster contains a major nitrogen fixation pathway [66]

\[
\text{CH}_3 + \text{N}_2 \rightarrow \text{CH}_2\text{NH} + \text{H}_2
\]

By analogy with the pathway involving C2H2, we observe also a minor growth pathway involving the repeated addition of C2H4 through ion–molecule reactions, e.g.

\[
\text{CH}_2\text{H}_4 + \text{CH}_3 \rightarrow \text{CH}_2\text{H}_2\text{NH} + \text{CH}_2\text{NH}_2
\]

and initiates the growth of hydrocarbon ions. This process has been identified as a key reaction for the APSIS simulation and its effect in promoting the growth through ion chemistry has also been highlighted in the literature [26, 69].

Irradiation at 82 nm is less efficient than at 60 nm (\(\rho_{60} = 3.3\%\)). Although at this wavelength \(N_2\) is not ionized but produces \(N\) atoms, the direct production of \(CH_2\) ion and \(CH_2\) ions [67], again at the same integrated intensity as in the reference case. Monte Carlo samples of 150 runs were performed for each case.

For comparison with the reference (DISCO) case, we calculated methane consumption ratio \(\rho_{CH_4}\) and the ratios of the mole fractions of the modified simulation over the reference simulation and reported them in Fig. 12 for neutral species. The uncertainty on the ratios is obtained by combination of variance [68].

In the 60 nm case, \(\rho_{CH_4} = 5\%\), about two orders of magnitude higher than the reference value (0.07 %, cf. Table 1). Moreover, the relative production of individual species increases with their molecular mass (Fig. 12 (top)). This increase is explained by the enhancement of ion chemistry: \(N_2\) photolysis at 60 nm essentially produces \(N_2\), which almost quantitatively ionizes methane through reactions

\[
\text{N}_2 + \text{CH}_4 \rightarrow \text{CH}_3 + \text{H}_2 + \text{N}_2
\]

and

\[
\text{N}_2 + \text{CH}_4 \rightarrow \text{CH}_3^+ + \text{H} + \text{N}_2
\]

and the ratios of the mole fractions for the 60 nm and solar irradiations are found not enable to simulate their 82.5 nm). The simulations are performed with the same integrated photon flux as the reference one. Another simulation is performed with a proxy of the solar spectrum [67], again at the same integrated intensity as in the reference case. Monte Carlo samples of 150 runs were performed for each case.

3.1.5. Impact study of physical conditions

Having highlighted the growth mechanism in the APSIS model, we explore the chemistry of the system in different physical conditions, in order to interpret other observations or experiments. To identify the effect of various physical factors, we design variants of the reference model with a single modified factor: irradiation spectrum, pressure, and temperature. The parameters for all simulations are gathered in Table 1.

3.1.5.1. Irradiation spectrum. We consider here simulations with monochromatic beams at 60 nm and 82 nm, as used by Imanaka and Smith [27] (the adopted 1 nm wavelength resolution does not enable to simulate their 82.5 nm). The simulations are performed with the same integrated photon flux as the reference one. Another simulation is performed with a proxy of the solar spectrum [67], again at the same integrated intensity as in the reference case. Monte Carlo samples of 150 runs were performed for each case.
of the branching ratios of the neutral channels of CH4 photolysis (see Section 2.2.1, Fig. 1(b)). Far from the wavelengths where measurements exist, all neutral channel branching ratios are strongly uncertain (relative uncertainties ~50–100%). Since CH4 photolysis is a key process, this large uncertainty contributes significantly to the uncertainty budget of the species concentrations. Despite this substantial uncertainty increase, the previously identified promotion of the formation of N-bearing species by the exclusive production of N atoms by N2 photolysis is significant for many species, such as N(4S); HCN, CH2NH, etc.

3.1.5.1. Pressure. In this simulation, the pressure $P$ in the reactor is lowered from the reference case by a factor $10^6$, which corresponds to the pressure around 850 km in Titan’s upper atmosphere ($\sim 5 \times 10^{-4}$ Pa). To ensure that the averaged photobereaction rates of N2 and CH4 are similar to the reference case, the reactor cell is elongated by the same factor to maintain the amount of gas ($L = 5 \times 10^5$ m).

Methane consumption is unaffected (Table 1), but the species distribution is strongly modified (Fig. 13 (top)). Some species ($C_6H, C^+, and CN^+$, for instance) are $\sim 10^8$ times more abundant than in the reference case, while others are reduced by similar amounts. We observe also a shift of the product distribution toward lower masses, signifying a limited growth. A systematic discrimination between saturated and unsaturated species is notable in each Cx block. The mole fraction ratio of a species increases with its degree of unsaturation within a block (red lines in Fig. 13 (top)). In the C5 and C6 blocks, the ratio between the extremes can reach 10 orders of magnitude.

Assuming an unchanged composition of the reactive gas, lowering the pressure by a factor of $10^6$ induces a lowering of the rate of all bimolecular reactions by a $10^{12}$ factor, while trimolecular reactions are slowed down by a factor of $10^{18}$. The latter class contains the radical recombinations forming heavy saturated species. Trimolecular reactions are important at several mbar (see Section 3.1.3.2) and contribute significantly to the production of the most saturated species. However, this contribution vanishes rapidly as the pressure decreases, resulting in very low mole fractions of the most saturated species, i.e. $C_5H_{11}, C_6H_{14}, etc.$, and less lowered mole fractions of less saturated species, such as...
C\textsubscript{4}H\textsubscript{7} and C\textsubscript{6}H\textsubscript{10}. The most unsaturated species, i.e. C\textsubscript{3}H\textsubscript{3}, C\textsubscript{3}H\textsubscript{2}… are not significantly affected.

3.1.5.2. Temperature. In this simulation, the temperature is lowered down to 150 K, close to that in Titan’s ionosphere. The relative mole fractions are shown in Fig. 13 (bottom).

Compared to the reference simulation at 300 K, methane consumption is halved (Table 1), and few stationary concentrations are significantly altered, except those of heavy saturated neutral species. Lowering the temperature is not as effective to slow down the growth as lowering the pressure, because only neutral reaction rate constants decrease with temperature: there are only marginal temperature effects on photolysis cross-sections [70,71,43], or on ion–molecule reactions.

However, we observe a discrimination between the species of various degree of unsaturation in heavy species blocks (see Fig. 13 (bottom)). Among the main growth families, only the one driven by trimolecular reaction (G3) is notably slowed down by the decrease in temperature, provoking a depletion of heavy saturated species. Moreover, dissociative recombinations are slightly accelerated by the decreased temperature of thermalized electrons (between 35 and 70%), which contributes also to increase the production of unsaturated species.

3.2. Simulation of Imanaka & Smith’s reactor

The model is now applied to Imanaka & Smith’s reactor [27] with 60 nm irradiation, in the conditions of their experiment (\(A = 1.2 \times 10^{-2} \text{ m}^2\), \(L = 0.7 \text{ m}\), \(T = 300 \text{ K}\), \(P = 13 \text{ Pa}\), \(I_0 = 10^{16} \text{ ph s}^{-1}\), \(N_2/\text{CH}_4 = 95/5\)).

Methane consumption (\(\rho_{\text{CH}_4} = 70\%\)) is about three orders of magnitude above the reference case (Table 1), and the largest in our set of simulations. This is mostly due to a higher photon flux combined with a lower CH\textsubscript{4} partial pressure. The relative charge is about \(4 \times 10^{-7}\). This charge is two orders of magnitude larger than for APSIS, because all photons at 60 nm contribute to ionization.

The experimental in situ MS and a simulated MS are compared in Fig. 14. The experimental spectrum is globally out of the 95% confidence interval of the simulated MS: the model systematically overestimates the production of unsaturated species and underestimates that of saturated species, by factors as large as 10\(^3\).

A series of factors might be responsible for this discrepancy:

- The mass spectrometer in Imanaka & Smith’s experiments was used at too high pressure to retain the linearity of its MS signals, as discussed for the APSIS experiments in Peng et al. [28]. However, the non-linearity of MS signals appears in a relatively uniform way in a band, therefore this argument cannot explain the systematic differential effects observed in all bands.

- The stable species considered in the simulated spectrum only accounts for a minor part (~10%) of total mass of products in the case of Imanaka & Smith’s reactor. The radicals, accounting for a large part of the total mass ignored in the simulated spectrum, are certainly an important source of disagreement. However, the factor 1000 between the saturated parts of measured and simulated MS bands is still not explained. Assuming that the chemical model is basically correct, the saturated part could have a concentration (not an intensity in the simulated MS) comparable to the unsaturated part in the same block, only if large radicals ignored in the simulated MS are (nearly) saturated, which cannot be the case. Even in the homogeneous model of the APSIS experiments, at a higher pressure disfavoring the growth of unsaturated species, these are still dominant in the blocks of heavy species. Therefore, the assumption of a correct chemistry is not true, i.e. the present homogeneous model is not adequate to simulate the chemistry occurring during Imanaka & Smith’s experiments.

Following the same logic as for the APSIS reactor, we address the limitations of homogeneous chemistry, by enhancing the trimolecular rate constants. In view of lower pressure and larger internal surface-to-volume ratio of Imanaka & Smith’s reactor than the APSIS reactor, both resulting in a larger relative contribution of heterogeneous chemistry, we tested three cases with 10, 30 and 100 times faster trimolecular rate constants. As for the reference case, the unsaturated part of MS bands is slightly lowered and the saturated part is significantly increased by this modification.

As trimolecular rate constants are amplified, the unsaturated part of the C3–C5 bands of the simulated MS gradually approaches the experimental results and their saturated part reaches and then overtakes the measured values.

The simulated spectrum corresponding to a factor 30 is shown in Fig. 14. Though still not satisfactory, for instance in the saturated
part of the C4–C6 band which is still overestimated, the agreement between the simulated spectrum and the experimental one is notably improved, especially for the saturated part. This may illustrate the very important effect of heterogeneous chemistry in this reactor, probably even stronger than in APSIS, because of the more favorable (tubular) geometry and lower pressure.

As discussed by Imanaka and Smith [27], the overestimation in Titan’s atmospheric models of the density of C4N2 (m/z 62 and 76) is due to a lack of loss processes for CHCN. This is verified here also for experimental laboratory conditions. In the present case, the overestimation is by more than two orders of magnitude (Fig. 14). This discrepancy is possibly higher than the ones noted for Titan modeling [27], see also Fig. 15. This would suggest that wall effects might be an efficient sink for CHCN in Imanaka & Smith setup. In any case, there is an urgent need of data on the reactivity of CHCN, either in homogeneous or heterogeneous conditions.

3.3. Simulation of a reactor in Titan’s ionospheric conditions

Combining the solar spectrum irradiation, low T and low P cases studied above, one obtains a set of conditions reminiscent of Titan’s ionosphere. Without aiming at a realistic simulation of this environment, a few adaptations of the parameters are necessary to get more representative conditions, such as adapting the photon flux and removing the carrier gas. The main difference of physical conditions between Titan atmosphere and the present case is the photon flux and removing the carrier gas. The main difference of this environment, a few adaptations of the parameters are necessary to get more representative conditions, such as adapting the photon flux and removing the carrier gas.

In Fig. 15, the simulated M5 is compared to the INMS neutral spectrum of T19 flyby between 950 and 1000 km, published in Waite et al. [9]. The agreement between the observations and the simulated spectrum for Titan is remarkably better than for either the APSIS (Fig. 5) or Imanaka & Smith’s (Fig. 14) experiments. In particular, one notes a good reproduction of the preponderance of unsaturated species within each block. This confirms the effect of low pressure and temperature underlined above (Sections 3.1.5.1.1 and 3.1.5.2) to favor the growth of unsaturated species.

The preponderance of unsaturated species at Titan indirectly supports our hypothesis that the reactor’s metallic wall plays an important role in the production of saturated species in the laboratory simulation experiments, simply because in Titan’s atmosphere, there is no such an efficient “wall” that promotes heterogeneous chemistry. Note however that wall effects on the INMS antechamber have been discussed by Vuitton et al. [11], and that heterogeneous reactions promoted by the aerosols are beginning to be studied [76].

The reproduction of the fact that C6 block’s concentration is larger than C5 block’s also validates the present reaction scheme in general terms. Though underestimated by the model, C6H6 (mainly at m/z 78) stands out as a major heavy product. Being highly unsaturated, it is efficiently formed in Titan’s atmosphere thanks to the physical conditions for its growth, i.e. low pressure and low temperature. On the other hand, overestimation of the intensities at m/z 62 and m/z 76, associated with C4N2, is due to a lack of loss pathways for CHCN, as discussed in Section 3.2.

### 3.3.1. Key reactions

For comparison with the key reactions identified in a neutral model of Titan’s chemistry by Hébrard et al. [32], we used the same procedure as in Section 3.1.3 on the present results. The list of key reactions is reported in Table 4. The intersection with Hébrard et al. [32] is reduced to two reactions N2 + hν → N(3S) + N(3P) and H + CH3CN → C + H2. The first one has already been evoked in Section 3.1.3. The second reaction has a very large uncertainty factor and has been shown to be able to induce different chemical regimes in Dobrijevic et al. [77]. A better estimation of this rate constant remains thus a priority for Titan’s chemistry modeling.

### 4. Conclusion

APSIS experiments have been studied by a 1-cell model with a fully coupled chemical scheme, i.e. involving all photolysis, bimolecular, trimolecular, ion–molecule reactions and dissociative recombinations with available kinetic data. The methane consumption is found acceptable and the production in Cx blocks is in good agreement with the experimental M5 measurements.

Ion chemistry and the full dissociative recombination scheme have been shown to be important in the model. Without them, the production of heavy species and heavy unsaturated species, respectively, are drastically lowered. By sensitivity analysis, photolysis and several reactions sensitive to pressure, mostly trimolecular reactions, appear as global key reactions. Influential photolysis reactions are consistent with the previous studies [32,39], and influential trimolecular reactions result from the relatively high pressure in the APSIS reactor.

Block patterns are observed in a species correlation matrix, indicating the importance of the addition of small units for molecular growth. Cluster analysis of the species correlation matrix reveals three major clusters of species: G1, where the growth of unsaturated species has been shown to be promoted by C2H2 through ion–molecule reactions; G2, in which the nitrogen

| Reaction | Fk | Scores
<table>
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<tbody>
<tr>
<td>N2 + hν → N(3S) + N(3P)</td>
<td>&lt; 1.1</td>
<td>23 4.6</td>
</tr>
<tr>
<td>N2 + hν → N2 + e⁻</td>
<td>&lt; 1.1</td>
<td>20 4.5</td>
</tr>
<tr>
<td>C2H2 + hν → C + H2</td>
<td>1.5</td>
<td>24 3.6</td>
</tr>
<tr>
<td>C2H2 + C2H2 → C2H4 + H</td>
<td>2.1</td>
<td>18 4.3</td>
</tr>
<tr>
<td>N2D + C2H2 → CHCN + H</td>
<td>1.3</td>
<td>18 3.1</td>
</tr>
<tr>
<td>CH3CN + hν → CH3CN + e⁻</td>
<td>1.2</td>
<td>17 4.2</td>
</tr>
<tr>
<td>CH3CN + hν → CH3CN + H + e⁻</td>
<td>1.3</td>
<td>16 3.0</td>
</tr>
<tr>
<td>CH3CN + hν → CH3CN + H + e⁻</td>
<td>1.4</td>
<td>17 3.8</td>
</tr>
</tbody>
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### Table 4

Consensus list of key reactions identified for the “Titan’s ionosphere” simulation, their rate constant uncertainty factor Fk in the conditions of the simulation, the corresponding number Ns of species used for selection, and their S score (see text). Bold characters indicate the maximum in each column.

### a key reactions in common with Hébrard et al. [32];

### b key reactions shared with the APSIS list (Table 2).
incorporation in hydrocarbons plays a central role; and G3, where the formation of heavy saturated species is ensured by trimolecular

This completes the discussion of Plessis et al. [25] about the species responsible for the promotion of the chemistry with a full dissociative recombination scheme. Plessis et al. [25] pointed out two antagonistic effects of dissociative recombination: a global chemical lysis of heavy ions and a promotion of the molecular growth due to the production of numerous small reactive species. In fact, the non H-loss dissociative recombination channels, through which many of small neutrals including C1 and C2 are produced, supply the main unsaturated reactants of the G1 cluster, but not saturated ones. In the APSIS model, these non H-loss channels increase the production rate of C2H2 by a factor of ~10, leading to a more efficient growth of unsaturated species. Therefore dissociative recombination produces small unsaturated species, especially C2H2, enhancing the most efficient growth pathways. This illustrates the promoting effects of dissociative recombination discussed by Plessis et al. [25], and makes them able to overtake the chemical lysis effect.

Titan’s ionosphere meets simultaneously the conditions favoring the growth of unsaturated species, i.e. low pressure and low temperature, leading to densities of unsaturated species, detected in an INMS neutral spectrum, in agreement with our modeling results. In contrast, the simulated MS for the APSIS and Imanaka & Smith’s experiments are not in good agreement with the experimental data in the saturated part of each Cx block. The model systematically underestimates the intensities of the saturated parts, revealing the insufficiency of homogeneous chemistry, which favors the growth of unsaturated species. If the recombinations catalyzed by the reactor’s walls are accounted for, the simulated MS are significantly improved, which strongly suggests the existence of noticeable wall effects in the laboratory simulation setups.

Based on this analysis, we have well identified competing homogeneous and heterogeneous chemistries in the reactors. The former essentially involves the growth of unsaturated species and is representative of the chemistry in Titan's upper atmosphere, and the latter promotes the production of saturated species and can be regarded as a bias of laboratory reactors. It was also shown that the unsaturated parts in the simulated MS are much less sensitive than the saturated parts to our ad hoc implementation of heterogenous chemistry. Therefore one may consider preferentially the experimental results on unsaturated species for a qualitative laboratory simulation of Titan's photochemistry.

The existence of wall effects also reveals the needs for further studies: simulation experiments at relatively low temperature (e.g. at ~0°C, leading to wall recombinations ~100–1000 times slower than at room temperature [56]) or using less catalytic wall materials to reduce these effects [78], and measurements of the kinetic parameters of involved heterogeneous reactions for their appropriate consideration in the model.

Several elementary processes have been identified at different stages of the present study as needing new or improved kinetic data, often confirming the data requests of other authors: the dissociative recombination rate and branching ratios for CH2NH [17]; the rate constant of the CH+H reactions [32,77]; new reactions consuming CHCN [27]; the photolysis branching ratios of CH4 around 80 nm, and more generally over the whole 80–140 nm range. These parameters are important sources of uncertainty and/or bias in the model predictions, and they are urgently needed to improve the detailed chemical modeling of N2/CH4 plasmas and Titan’s ionosphere.

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