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# Mapping the evolution of pure CO<sub>2</sub> ices irradiated by ions, UV, and electrons using the upgraded PROCODA code (employing an effective rate constant ordering by thermochemistry data)

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## Abstract

Chemical reactions and desorption processes are being triggered by incoming ionizing radiation over astrophysical ices in cold space environments. The quantification of these processes is crucial to achieve a detailed understanding of the underlying chemistry occurring within the ice. With this goal, we have upgraded the PROCODA code (Pilling et al. 2022a) which solves a system of coupled differential equations and describes the evolution of the molecular abundances under processing by radiation, now including an effective rate constant (ERCs) ordering by employing thermochemistry data taken from literature. This methodology helps to identify the most important reactions within the reaction network and therefore decreases the degeneracy of the solutions and enhancing the accuracy of the calculations. Here, we described the chemical evolution of four irradiated pure CO<sub>2</sub> considering 11 different chemical species, 100 reaction routes and 11 radiation-induced desorption processes. The best-fit models provide the effective rate constants, several desorption parameters, as well as, the characterization of the chemical equilibrium (CE) phase. A comparison with previous code version was given and indicates that the ordering of rate constants by thermochemistry data is more important when more energy is deposited in the ice. The current work present more realistic values for the effective rate constants and a better characterization of the CE phase, such data can be used to refine astrochemical models to better describe cold space environments in the presence of incoming ionizing radiation field such molecular clouds and protoplanetary regions and the surface of comets and frozen moons and planets.

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## 1. Introduction

In cold space environments, chemical reactions and desorption processes are being triggered by incoming ionizing radiation impinging astrophysical ices (e.g. Muñoz Caro et al. 2002; Meinert et al. 2016). This radiation-induced processing leads to an increase in chemical complexity in the ices, as well as in their gaseous vicinity (e.g. van

Dishoeck 2014; Boogert et al. 2015). Among the chemical inventory in the interstellar medium, carbon dioxide (CO<sub>2</sub>), an apolar molecule with linear geometry, is ubiquitously found in the interstellar medium (e.g., de Graauw et al. 1996, Gerakines et al. 1999, Pontoppidan et al. 2008, Poteet et al. 2013, Ehrenfreund & Charnley 2000; Fraser et al. 2002, Öberg et al. 2011) and on icy moons of the solar system (Buratti et al. 2005, Cartwright et al. 2015). The irradiation of pure CO<sub>2</sub> ice by UV radiation has shown that other molecules, such as CO, O<sub>3</sub>, and CO<sub>3</sub>, can be formed in the ice (Gerakines et al. 1996). Addi-

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tionally, other experiments addressing the influence of other energy regimes have also been performed. In particular, [Mejía et al. \(2015\)](#) and [Pilling et al. \(2010a\)](#) studied the impact of cosmic ray analogs on ice chemistry, and chemical processes induced by electrons and UV photons have been addressed by [Bennett et al. \(2004\)](#) and [Martín-Doménech et al. \(2015\)](#), respectively.

The underlying chemistry occurring within the ice when exposed to ionizing radiation is not accessible from experiments. In fact, the in situ techniques used to analyze the ice such as infrared spectroscopy and mass spectrometry work as diagnostic tools to visualize the differences in the ice's chemical composition before and after irradiation. To have insights into the chemistry triggered by radiation a chemical model simulating laboratory conditions is necessary. With this goal, the PROCODA code, which stands for PROgram for solving COupled Differential equations in Astrochemistry, was developed ([Pilling et al. 2022a](#)). It solves a system of coupled differential equations and describes the evolution of the molecular abundances of ices under processing by radiation. This methodology is complementary to other attempts to understand chemical reactions in the ice carried out, for example, by [Jamieson et al. \(2006\)](#), [Bossa et al. \(2015\)](#), [Shingledecker et al. \(2019\)](#) and [Rocha et al. \(2022\)](#). Additionally, the code calculates the effective rate constants (ERCs) of the direct, bimolecular and termolecular reactions, the radiation-induced desorption effective rate constants, as well as the desorption column density for the studied species as functions of time and the summed molecular desorption yield ( $Y$ ). It is worth noting that the direct reactions mean the dissociation or destruction reactions triggered by the incoming radiation (which might also involve the action of secondary electrons). A discussion on the role of secondary electrons can be found at [Mason et al. \(2014\)](#) and [Boyer et al. \(2016\)](#).

The PROCODA code also provides abundances for both observed and non-observed (unknown) species in the IR spectra during ice chemical evolution during processing by incoming radiation, and also characterizes the chemical equilibrium (CE) phase in the ice that occurs at larger radiation fluence. Recently, the PROCODA code has been employed to map the CE phase of pure acetonitrile ( $\text{CH}_3\text{-CN}$ ) and carbon monoxide (CO) ices exposed to ionizing radiation ([Carvalho et al. 2022](#); [Pilling et al. 2022b](#)).

The current modifications in the code (the ERCs ordering hypothesis by thermochemistry data) help to identify the most important reactions within the reaction network and therefore decrease the degeneracy of the solutions and enhance the accuracy of the calculations. With the upgraded PROCODA code we map the chemical evolution of four pure  $\text{CO}_2$  ices exposed to different ionizing radiation sources (52 MeV  $^{58}\text{Ni}^{+13}$  ions, 550 MeV  $^{50}\text{Ti}^{+21}$  ions, 10 eV photons, 5 keV electrons). An extended comparison between the current code version with the previous version also applied to  $\text{CO}_2$  ices (without ordering hypothesis) described by [Pilling et al. \(2022a\)](#) for two models (Ni ions and UV photons) was given, and the main changes and

similarities were discussed. In a few words, the upgraded code version allows a decrease in the solution's degeneracy and also allows a better accuracy for the calculated rates constants, desorption yields, and molecular abundances.

It is worth noting that the current methodology highlights the employ of the term “effective” rate constant (ERC), which is a parameter with a broader concept than the “ordinary” rate constant ( $k$ ) of reaction, commonly considered in gas phase chemistry. The ERCs handle the fact that reactants within the ice are surrounded by different neighbors, which also vary with time (and also are affected by the grain's local density and temperature due to the incoming radiation). Therefore, this quantity, in some sense, represents an average value (in time and space) over the ice during this ice processing by radiation and better describes what is happening within the ice in real conditions. To provide more realistic results, the astrochemical ice models should consider this ERC instead of an “ordinary” rate constant for a given reaction (often employed the gas-phase values also for ices) since the ice composition is not strictly homogeneous and also varies with time, and are also under radiation processing.

The computational methodology in this work, which includes the hypothesis of ordering of rate constant by employing thermochemistry data and the details of the employed experimental works, is described in section 2. The results employing the upgraded PROCODA code in four pure  $\text{CO}_2$  ices exposed to ionizing radiation are shown in [Section 3](#), presenting also the characterization of the CE phase, as well as a comparison with the previous version of the code. Some astrophysical implications are provided in section 4. [Section 5](#) lists the main conclusions and presents final remarks.

## 2. Methodology

In the current work, we present the upgraded version of the PROCODA code (the original code is given by [Pilling et al. 2022a](#)) which solves a system of coupled differential equations to describe the chemical evolution, as a function of time, of typical astrophysical ices during processing by ionizing radiation. The current version considers an ordering hypothesis for the effective rate constants (ERCs) by employing thermochemistry data (gas-phase at 0 K). In the manuscript, we prefer to use the term “effective” to describe the rate constants, making explicit that this physicochemical parameter takes into account the different chemical environments in which a given reaction may occur within the ice (an average value). Details on this ordering hypothesis will be provided further.

It is worth noting that, in this manuscript, we sometimes present the ERCs as  $k$  values (e.g.,  $k_1$ ,  $k_2$ ,  $k_{\text{des}}$ ,  $k_{\text{A+B}}$ ).

### 2.1. Experimental datasets

The employed chemical network considers 100 coupled equations that describe the chemical evolution of selected

molecules formed from pure CO<sub>2</sub> ices under the presence of different types of ionizing radiation. The reaction set also takes radiation-induced molecular desorption into account. The proposed reaction network involves 11 molecular species (the parent species CO<sub>2</sub> and the newly produced species CO, O<sub>3</sub>, CO<sub>3</sub>, O, O<sub>2</sub>, C, C<sub>2</sub>, C<sub>2</sub>O, C<sub>2</sub>O<sub>2</sub>, and C<sub>2</sub>O<sub>3</sub>). Fig. 1 illustrates the Lewis structures of the species involved in the CO<sub>2</sub> ice decomposition process. These species were chosen since they present the lowest formation enthalpy energies.

Here, four experimental datasets were considered employing different ionizing sources on pure CO<sub>2</sub> ice: i) 15 K ice bombarded by 550 MeV <sup>50</sup>Ti<sup>+21</sup> ions (from Mejía et al. 2015) named as CR1 model; ii) 13 K ice bombarded by 52 MeV <sup>58</sup>Ni<sup>+13</sup> ions (from Pilling et al. 2010a) named as CR2 model; iii) 10 K ice bombarded by 5 keV electrons (from Bennett et al. 2004) named as EL model; and iv) 8 K ice irradiated by UV photons of ~ 10 eV (from Martín-Doménech et al. 2015) named as UV model. Following the authors, the experimental uncertainties were below 10–20 %. It is worth noting that, in all cases, only the four observable species (CO<sub>2</sub>, CO, O<sub>3</sub>, and CO<sub>3</sub>) were quantified in the experimental data. With the current methodology, we can map and characterize the abundances of the other seven proposed non-observable species (O, O<sub>2</sub>, C, C<sub>2</sub>, C<sub>2</sub>O, C<sub>2</sub>O<sub>2</sub>, and C<sub>2</sub>O<sub>3</sub>), expected to have formed during the irradiated CO<sub>2</sub> ice experiments.

## 2.2. The PROCODA methodology

As discussed by Pilling et al. (2022a) the typical equation in the chemical coupled system solved by PROCODA code has the following parameters:

$$\frac{dN_i}{dt} = \left[ -DES_i(t) - \sum_{d1} k_{d1} N_i(t) - \sum_{d2} k_{d2} N_i(t) N_a(t) - \sum_{d3} k_{d3} N_i(t) N_a(t) N_b(t) \right] + \left[ \sum_{p1} k_{p1} N_a(t) + \sum_{p2} k_{p2} N_a(t) N_b(t) + \sum_{p3} k_{p3} N_a(t) N_b(t) N_c(t) \right] \quad [\text{molecules cm}^{-2} \text{ s}^{-1}] \quad (1)$$

where  $dN_i/dt$  is the column density changes of a given molecule  $i$  along the time  $t$ , the  $k$  values are the ERCs for different processes, and  $DES_i(t) = k_{des,i} \Omega_i(t) N_i(t)$  is the differential column density desorption, i.e., the number of molecules (or atomic species) that desorbs from ice to gas-phase per cm<sup>2</sup> and per second due to incoming radiation which depends also on the intrinsic radiation-induced desorption rate constant ( $k_{des,i}$ ), in units of s<sup>-1</sup>, the and the dimensionless surface coverage of the species  $i$  as a function of time ( $\Omega_i(t)$ ). The desorption itself was considered a first-order type reaction. In this equation, the values  $k_{d1}$ ,  $k_{d2}$ , and  $k_{d3}$  represent the destruction processes (consumption) and the values  $k_{p1}$ ,  $k_{p2}$ , and  $k_{p3}$  represent the production processes (formation) of a given species in direct, bimolecular, and termolecular reactions, respectively (details are given at Pilling et al. 2022a). The values  $N_a$ ,  $N_b$ , and  $N_c$  indicate the concentrations of species  $a$ ,  $b$  and  $c$ , respectively, which participate in the reaction to produce or consume the respective  $i$  species.

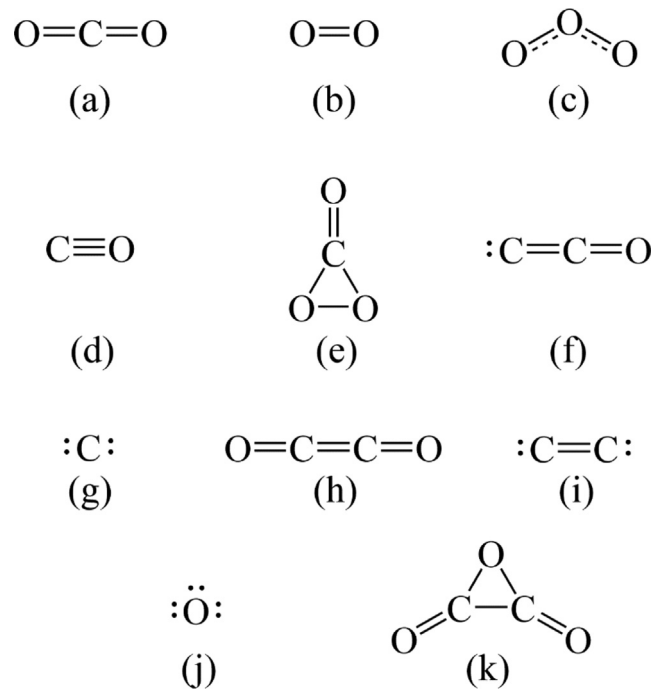


Fig. 1. Lewis structures of the eleven species formed in the CO<sub>2</sub> ice decomposition process. (a) CO<sub>2</sub>, (b) O<sub>2</sub>, (c) O<sub>3</sub>, (d) CO, (e) CO<sub>3</sub>, (f) C<sub>2</sub>O, (g) C, (h) C<sub>2</sub>O<sub>2</sub>, (i) C<sub>2</sub>, (j) O, (k) C<sub>2</sub>O<sub>3</sub>.

It is worth noting that the code applies other constraints besides the minimization of chi-square function for the data, such as, mass conservation, desorption yield and CE phase. Those constraints are employed in a minimization process of a function (named score function (SF)) that evaluates how close the system is to its minimum global solution. The score function is defined as follows

$$SF = p1 \times \sum \frac{(oCO2_{data} - oCO2_{model})^2}{oCO2_{data}} + p2 \times \sum \frac{(oCO_{data} - oCO_{model})^2}{oCO_{data}} + p3 \times \sum \frac{(oO3_{data} - oO3_{model})^2}{oO3_{data}} + p4 \times \sum \frac{(oCO3_{data} - oCO3_{model})^2}{oCO3_{data}} + p5 \times [(1 - MSC_f) + (1 - MSC_{of}) + (1 - MSC_{om})] + p6 \times (1 - DSC) + p7 \times (1 - SSC) \quad (2)$$

where the dimensionless parameters  $p1$  to  $p7$  in this equation are the weights of each term, used as a tool to search for the best solution during the computational minimization processes. In this equation, the parameter  $MSC_f$  is the model mass similarity criterion (or model mass conservation), calculated considering the similarity between the initial column mass of the modeled system and the total column mass of the model at final modeling time (at largest time) (including both masses, at the ice and desorbed),  $MSC_{of}$  and  $MSC_{om}$  are the column mass similarity criterion between the experimentally observed column mass and the observed column mass in the model at the final

modeling time and at the middle of the modeling time, respectively. From these three parameters, we calculate a parameter called mass similarity criterion  $MSC = (MSC_f \times 100\% + MSC_o_f \times 100\% + MSC_o_m \times 100\%) / 3$ , which also helps to indicate how good is the solution of the system in compared with the observed data. The parameter  $DSC$  is the desorption similarity criterion, calculated considering the similarity of the input expected desorption yield (based on the manuscripts with the experimental data employed) and the total molecular desorption yield computed by the model. The parameter  $SSC$  is the slope similarity criterion, which is related to the achievement of CE phase in the ice at larger radiation fluence, as described in Pilling et al. (2022a).

It is worth noting that the chi-squared function  $\chi^2$  (also named summed CHI2) can be obtained directly from equation [2] making  $p1 = p2 = p3 = p4 = 1$  and  $p5 = p6 = p7 = 0$ . The chi-squared function is a parameter obtained considering the experimental concentration data of the observed molecules ( $\text{CO}_2$ ,  $\text{CO}$ ,  $\text{O}_3$ , and  $\text{CO}_3$ ) and their modeled concentration.

For each model, the code was typically run three times until the best-fit solution was found. At every new run, the obtained ERCs in the previous best-fit calculations were considered as new input for the new calculations. The details of the code, including the employed equations, the coupled equation minimization algorithm (L\_BFGS-B algorithm), its explanatory flowchart, the input and output parameters, and the criteria for model convergence can be found at Pilling et al. (2022a).

### 2.3. The ordering hypothesis for the effective rate constants (code upgrade)

In this upgraded version of the code, we consider, as a first approximation, that in a given set of reactions (with the same initial reactants), the ones with smaller enthalpy variation,  $\Delta H$  (more exothermic), are expected to have a larger rate constant,  $k$  (faster reactions). This hypothesis is also supported by the Evans–Polanyi principle (also referred to as the Bell–Evans–Polanyi principle) which observes that the difference in activation energy between two reactions of the same family is proportional to the difference in their enthalpy of reaction (Bell 1936; Evans and Polanyi, 1936). The Evans–Polanyi model is a linear energy relationship that serves as an efficient way to calculate the activation energy of many reactions within a distinct family. The activation energy may be used to characterize the kinetic rate parameter of a given reaction through the application of the Arrhenius equation.

As a first approximation, the thermochemical data used in the ordering hypothesis for the ERCs, considers that all studied species are in the lowest energy level and formation enthalpies and reaction enthalpies are at gas-phase at the theoretical temperature of 0 K. It is worth noting that the hypothesis of the ordering of ERCs was implemented only for the reactions within the reaction groups. In this manu-

script, we have considered a total of 28 reaction groups (each one containing different reaction pathways or different routes inside) in the code, named from G1 to G28. Each group has its own ordering in the rate constants, and this ordering is independent of the ordering in other groups. Single reactions (the ones that do not belong to a specific reaction group) were not ranked by thermochemistry data. The rate constant ordering was introduced in the code by adding boundary constraints in the calculation of each rate of reaction inside a specific reaction group.

The values of reaction enthalpies considered in this work and the ordering of rate constants within the different reaction groups are indicated in the Table 2. For the reactions involving the species  $\text{CO}_2$ ,  $\text{CO}$ ,  $\text{O}$ ,  $\text{O}_2$ ,  $\text{O}_3$ ,  $\text{C}$ ,  $\text{C}_2$ ,  $\text{C}_2\text{O}$ ,  $\text{C}_2\text{O}_2$  it was employed the gas-phase 0 K formation enthalpies values  $\Delta_f H(\text{CO}_2) = -393.1 \text{ kJ mol}^{-1}$ ;  $\Delta_f H(\text{CO}) = -113.8 \text{ kJ mol}^{-1}$ ;  $\Delta_f H(\text{O}_3) = 144.3 \text{ kJ mol}^{-1}$ ;  $\Delta_f H(\text{O}) = 246.8 \text{ kJ mol}^{-1}$ ;  $\Delta_f H(\text{O}_2) = 0 \text{ kJ mol}^{-1}$ ;  $\Delta_f H(\text{C}) = 711.4 \text{ kJ mol}^{-1}$ ;  $\Delta_f H(\text{C}_2) = 820.2 \text{ kJ mol}^{-1}$ ;  $\Delta_f H(\text{C}_2\text{O}) = 337.1 \text{ kJ mol}^{-1}$ ;  $\Delta_f H(\text{C}_2\text{O}_2) = 14 \text{ kJ mol}^{-1}$ , obtained from the Active Thermochemical Tables version 1.122p at <https://atct.anl.gov/> (see also Ruscic et al. 2004; Ruscic et al. 2005; Ruscic & Bross 2020). For the reactions involving  $\text{CO}_3$ , it was assumed the gas-phase formation enthalpies at 0 K of  $\Delta_f H = 52.5 \text{ kJ mol}^{-1}$ , and the reactions enthalpies  $\Delta_r H$  were obtained from the NIST website (<https://cccbdb.nist.gov/>). For the reactions involving  $\text{C}_2\text{O}_3$  it was considered the 0 K formation enthalpies  $\Delta_f H(\text{C}_2\text{O}_3) = -282.2 \text{ kJ mol}^{-1}$  (from Gambi et al. (2001) together with the other formation enthalpies from ATCL at <https://atct.anl.gov/>). It is worth noting that some reactions considered in the model (see Table 2) are endothermic (at gas-phase) and might be difficult to occur in the ice at low temperatures, however since the suited ices are constantly under bombardment (with constant energy input by the incoming projectiles) such reactions may have indeed a chance to occur.

### 2.4. Limitations and advantages of the code

It is worth noting that, the current model does not consider, in an explicit way, the ionic and excited (thermally, electronically, or vibrationally states) species. However, we understand that such chemical states should belong to the reactions inventory within the ice. The ERC calculated here, for example, in the reaction  $A + \text{RAD} \rightarrow B + C$  (where  $\text{RAD}$  = any type of radiation) in some way involves the different chemical states in which  $A$ ,  $B$ , or  $C$  species appear in the ice (which includes neutral, ionic, excited and radical). The major difficulty in considering these forms separately in the models is not in writing the coupled equations, but in the way to measure them in the experimental data at IR. Future work will include explicitly introducing a few ionic species to the model (considering as one of the working hypotheses, for example, a specific ratio ion/neutral for a selected species in the ice).



Table 1

The considered reaction routes and the calculated effective rate constants (ERC) employing the upgraded PROCODA code on pure CO<sub>2</sub> irradiated ices. The reaction enthalpies considered in the ordering hypothesis for the ERC and the reaction groups (with the ordering of ERCs between parentheses) are listed. The intrinsic values for molecular desorption are listed at the bottom at the table.

Modelled reactions <sup>a</sup>	Effective rate constant (CR1 / CR2 / EL / UV)	Reaction enthalpy at 0 K <sup>b</sup> [kJ/mol]	Reaction groups ( <i>k</i> values ordered based on reaction exothermicity) <sup>c</sup>	References or comments
CO <sub>2</sub> + RAD → CO + O	k1 = (3.1e-06 / 9.4e-07 / 1.6e-06 / 1.3e-08) s <sup>-1</sup>	526.1	G1 (k1 > k3 > k5)	[3,4,5,6] gas-phase; [9,10] ice
CO + O → CO <sub>2</sub>	k2 = (3.1e-27 / 2.4e-26 / 1.4e-24 / 2.9e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-526.1	G8 (k2 > k69)	[3] gas and surface; [4] gas-phase [12] ice; [13] Desorption,
CO <sub>2</sub> + RAD → C + O <sub>2</sub>	k3 = (1.6e-8 / <1.0e-08 / 1.2e-08 / <7.6e-09) s <sup>-1</sup>	1104.5	G1 (k1 > k3 > k5)	[3] gas and surface; [4] gas-phase
C + O <sub>2</sub> → CO <sub>2</sub>	k4 = (5.3e-26 / 2.5e-25 / 3.2e-25 / 5.8e-27) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-1104.5	G9 (k4 > k70)	
CO <sub>2</sub> + RAD → C + O + O	k5 = (≤1.0e-08 / ≤1.0e-08 / ≤1.0e-08 / ≤1.0e-08) s <sup>-1</sup>	1598.2	G1 (k1 > k3 > k5)	Employing Δ <sub>r</sub> H(C <sub>2</sub> O <sub>3</sub> ) = -282.2 kJ mol <sup>-1</sup> from Gambi et al. 2001
C + O + O → CO <sub>2</sub>	k6 = (1.2e-48 / 3.4e-48 / 2.2e-48 / 1.1e-49) cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	-1598.2	Single reaction pathway	
CO <sub>2</sub> + CO <sub>2</sub> → C <sub>2</sub> O <sub>2</sub> + O <sub>2</sub>	k7 = (8.4e-31 / 4.4e-31 / 4.0e-31 / 4.1e-31) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	800.3	G2 (k13 > k9 > k11 > k7)	
C <sub>2</sub> O <sub>2</sub> + O <sub>2</sub> → CO <sub>2</sub> + CO <sub>2</sub>	k8 = (4.5e-27 / 4.6e-25 / 1.9e-26 / 6.1e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-800.3	Single reaction pathway	
CO <sub>2</sub> + CO <sub>2</sub> → CO + CO + O <sub>2</sub>	k9 = (1.0e-30 / 2.4e-28 / 1.5e-28 / 8.0e-28) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	558.6	G2 (k13 > k9 > k11 > k7)	
CO + CO + O <sub>2</sub> → CO <sub>2</sub> + CO <sub>2</sub>	k10 = (7.0e-49 / 1.2e-47 / 5.1e-48 / 1.5e-48) cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	-558.6	Single reaction pathway	
CO <sub>2</sub> + CO <sub>2</sub> → C <sub>2</sub> O <sub>3</sub> + O	k11 = (1.1e-30 / 4.2e-31 / 4.8e-31 / 3.0e-31) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	750.8	G2 (k13 > k9 > k11 > k7)	
C <sub>2</sub> O <sub>3</sub> + O → CO <sub>2</sub> + CO <sub>2</sub>	k12 = (3.0e-27 / 3.6e-25 / 4.0e-30 / 2.8e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-750.8	Single reaction pathway	
CO <sub>2</sub> + CO <sub>2</sub> → CO <sub>3</sub> + CO	k13 = (3.7e-29 / 1.9e-27 / 1.2e-28 / 1.5e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	503.7	G2 (k13 > k9 > k11 > k7)	
CO <sub>3</sub> + CO → CO <sub>2</sub> + CO <sub>2</sub>	k14 = (2.3e-27 / 2.2e-25 / 3.4e-25 / 5.4e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-503.7	Single reaction pathway	
CO <sub>2</sub> + O → CO + O <sub>2</sub>	k15 = (5.3e-28 / 1.1e-26 / 3.0e-26 / 8.0e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	32.4	G3 (k15 > k19 > k17)	
CO + O <sub>2</sub> → CO <sub>2</sub> + O	k16 = (1.9e-26 / 1.7e-25 / 2.0e-25 / 1.5e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-32.4	G19 (k77 > k16 > k75)	
CO <sub>2</sub> + O → C + O <sub>3</sub>	k17 = (2.4e-28 / 4.4e-31 / 4.9e-28 / 1.4e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	1002.0	G3 (k15 > k19 > k17)	[5] gas-phase; [10] ice; Employing enthalpies from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
C + O <sub>3</sub> → CO <sub>2</sub> + O	k18 = (2.4e-26 / 4.7e-25 / 3.9e-25 / 1.9e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-1002.0	G22 (k18 > k76)	
CO <sub>2</sub> + O → CO <sub>3</sub>	k19 = (1.1e-29 / 9.0e-27 / 3.0e-27 / 3.1e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	52.5	G3 (k15 > k19 > k17)	[5] gas-phase; [10] ice; Employing enthalpies from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO <sub>3</sub> + RAD → CO <sub>2</sub> + O	k20 = (2.6e-04 / 1.8e-02 / 2.0e-03 / ≤9.0e-09) s <sup>-1</sup>	-52.5	G12 (k20 > k78 > k87)	
CO <sub>2</sub> + O <sub>2</sub> → CO + O <sub>3</sub>	k21 = (1.2e-26 / 7.7e-25 / 1.1e-24 / 2.5e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	423.7	G4 (k21 > k25 > k23)	Employing enthalpies from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO + O <sub>3</sub> → CO <sub>2</sub> + O <sub>2</sub>	k22 = (1.1e-26 / 9.9e-26 / 2.9e-25 / 4.8e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-423.7	G10 (k22 > k79)	
CO <sub>2</sub> + O <sub>2</sub> → C + O + O <sub>3</sub>	k23 = (2.4e-27 / 1.1e-28 / 1.3e-26 / 1.7e-29) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	1495.7	G4 (k21 > k25 > k23)	
C + O + O <sub>3</sub> → CO <sub>2</sub> + O <sub>2</sub>	k24 = (4.8e-49 / 1.6e-48 / 5.3e-48 / 2.4e-49) cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	-1495.7	Single reaction pathway	
CO <sub>2</sub> + O <sub>2</sub> → CO <sub>3</sub> + O	k25 = (4.8e-27 / 3.1e-26 / 1.5e-25 / 4.7e-27) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	445.6	G4 (k21 > k25 > k23)	
CO <sub>3</sub> + O → CO <sub>2</sub> + O <sub>2</sub>	k26 = (5.3e-26 / 4.7e-24 / 7.1e-26 / 2.8e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-445.6	G28 (k26 > k80)	
CO <sub>2</sub> + C → C <sub>2</sub> + O <sub>2</sub>	k27 = (8.4e-31 / 4.8e-27 / 6.8e-29 / 2.2e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	501.9	G5 (k29 > k31 > k27)	[3] gas-phase [3,4] gas-phase
C <sub>2</sub> + O <sub>2</sub> → CO <sub>2</sub> + C	k28 = (5.0e-27 / 8.5e-26 / 8.3e-27 / 6.2e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-501.9	G24 (k62 > k28)	
CO <sub>2</sub> + C → CO + CO	k29 = (1.1e-26 / 1.3e-25 / 2.0e-25 / 1.4e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-545.8	G5 (k29 > k31 > k27)	[7,8] ice
CO + CO → CO <sub>2</sub> + C	k30 = (1.2e-26 / 1.7e-26 / 2.4e-25 / 9.1e-28) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	545.8	G11 (k59 > k30 > k63 > k61)	
CO <sub>2</sub> + C → C <sub>2</sub> O <sub>2</sub>	k31 = (4.4e-27 / 6.3e-26 / 1.4e-26 / 1.2e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-304.3	G5 (k29 > k31 > k27)	

Table 1 (continued)

Modelled reactions <sup>a</sup>	Effective rate constant (CR1 / CR2 / EL / UV)	Reaction enthalpy at 0 K <sup>b</sup> [kJ/mol]	Reaction groups ( <i>k</i> values ordered based on reaction exothermicity) <sup>c</sup>	References or comments
C <sub>2</sub> O <sub>2</sub> + RAD → CO <sub>2</sub> + C	k32 = (2.4e-03 / 2.5e-03 / 3.7e-03 / 5.4e-03) s <sup>-1</sup>	304.3	G15 (k60 > k32 > k93)	
CO <sub>2</sub> + CO → C <sub>2</sub> O <sub>3</sub>	k33 = (1.6e-28 / 4.3e-27 / 2.9e-28 / 1.6e-27) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	224.7	G6 (k33 > k35 > k37 > k39)	Employing Δ <sub>r</sub> H(C <sub>2</sub> O <sub>3</sub> ) = -282.2 kJ mol <sup>-1</sup> from Gambi et al. 2001
C <sub>2</sub> O <sub>3</sub> + RAD → CO <sub>2</sub> + CO	k34 = (1.7e-04 / 5.3e-02 / 3.1e-03 / 5.0e-03) s <sup>-1</sup>	-224.7	G16 (k34 > k95 > k97 > k99)	Employing Δ <sub>r</sub> H(C <sub>2</sub> O <sub>3</sub> ) = -282.2 kJ mol <sup>-1</sup> from Gambi et al. 2001
CO <sub>2</sub> + CO → C <sub>2</sub> O <sub>2</sub> + O	k35 = (6.6e-29 / 4.5e-28 / 4.5e-29 / 3.9e-28) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	767.8	G6 (k33 > k35 > k37 > k39)	
C <sub>2</sub> O <sub>2</sub> + O → CO <sub>2</sub> + CO	k36 = (1.7e-25 / 2.4e-25 / 3.0e-26 / 2.8e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-767.8	G17 (k36 > k98)	
CO <sub>2</sub> + CO → C <sub>2</sub> O + O <sub>2</sub>	k37 = (8.2e-29 / 2.3e-30 / 2.8e-31 / 1.4e-30) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	884.0	G6 (k33 > k35 > k37 > k39)	
C <sub>2</sub> O + O <sub>2</sub> → CO <sub>2</sub> + CO	k38 = (2.0e-26 / 4.4e-25 / 2.1e-26 / 7.7e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-884.0	G20 (k38 > k100)	
CO <sub>2</sub> + CO → C <sub>2</sub> + O <sub>3</sub>	k39 = (2.3e-28 / 4.4e-31 / 4.0e-31 / 3.4e-27) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	1471.6	G6 (k33 > k35 > k37 > k39)	
C <sub>2</sub> + O <sub>3</sub> → CO <sub>2</sub> + CO	k40 = (5.4e-27 / 1.8e-25 / 1.5e-25 / 4.4e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-1471.6	Single reaction pathway	
CO <sub>2</sub> + C <sub>2</sub> O → C <sub>2</sub> O <sub>2</sub> + CO	k41 = (2.7e-27 / 1.3e-25 / 4.4e-26 / 8.6e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-83.0	G21 (k41 > k43)	
C <sub>2</sub> O <sub>2</sub> + CO → CO <sub>2</sub> + C <sub>2</sub> O	k42 = (7.9e-27 / 5.2e-26 / 2.4e-26 / 3.7e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	83.0	Single reaction pathway	
CO <sub>2</sub> + C <sub>2</sub> O → C <sub>2</sub> O <sub>3</sub> + C	k43 = (1.1e-27 / 6.9e-26 / 1.6e-27 / 4.0e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	413.2	G21 (k41 > k43)	Employing Δ <sub>r</sub> H(C <sub>2</sub> O <sub>3</sub> ) = -282.2 kJ mol <sup>-1</sup> from Gambi et al. 2001
C <sub>2</sub> O <sub>3</sub> + C → CO <sub>2</sub> + C <sub>2</sub> O	k44 = (1.0e-26 / 3.3e-25 / 2.6e-27 / 4.4e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-413.2	Single reaction pathway	Employing Δ <sub>r</sub> H(C <sub>2</sub> O <sub>3</sub> ) = -282.2 kJ mol <sup>-1</sup> from Gambi et al. 2001
CO <sub>2</sub> + C <sub>2</sub> → CO + C <sub>2</sub> O	k45 = (2.2e-26 / 3.4e-26 / 1.4e-25 / 1.8e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-163.90	G25 (k45 > k47)	
CO + C <sub>2</sub> O → CO <sub>2</sub> + C <sub>2</sub>	k46 = (1.1e-26 / 1.1e-25 / 1.8e-26 / 1.4e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	163.90	G18 (k46 > k73)	
CO <sub>2</sub> + C <sub>2</sub> → C <sub>2</sub> O <sub>2</sub> + C	k47 = (1.8e-27 / 2.0e-26 / 1.4e-25 / 1.6e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	299.0	G25 (k45 > k47)	
C <sub>2</sub> O <sub>2</sub> + C → CO <sub>2</sub> + C <sub>2</sub>	k48 = (2.7e-27 / 1.9e-25 / 2.8e-27 / 2.4e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-299.0	G27 (k74 > k48)	
CO <sub>2</sub> + CO <sub>3</sub> → C <sub>2</sub> O + 2 O <sub>2</sub>	k49 = (1.5e-27 / 2.3e-25 / 4.6e-28 / 4.1e-31) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	858.87	G7 (k51 > k49)	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
C <sub>2</sub> O + 2 O <sub>2</sub> → CO <sub>3</sub> + CO <sub>2</sub>	k50 = (6.2e-49 / 5.1e-47 / 6.9e-50 / 1.1e-48) cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	-858.87	Single reaction pathway	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO <sub>2</sub> + CO <sub>3</sub> → CO + CO + O <sub>3</sub>	k51 = (1.5e-27 / 1.2e-24 / 1.5e-26 / 6.3e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	490.08	G7 (k51 > k49)	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO + CO + O <sub>3</sub> → CO <sub>3</sub> + CO <sub>2</sub>	k52 = (2.1e-50 / 1.0e-47 / 9.3e-48 / 6.3e-49) cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	-490.08	Single reaction pathway	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO <sub>2</sub> + O <sub>3</sub> → CO + O <sub>2</sub> + O <sub>2</sub>	k53 = (8.4e-31 / 9.8e-29 / 4.9e-29 / 2.4e-27) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	134.9	G23 (k55 > k53)	
CO + O <sub>2</sub> + O <sub>2</sub> → CO <sub>2</sub> + O <sub>3</sub>	k54 = (1.5e-48 / 6.0e-48 / 4.7e-49 / 3.4e-49) cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	-134.9	Single reaction pathway	
CO <sub>2</sub> + O <sub>3</sub> → CO <sub>3</sub> + O <sub>2</sub>	k55 = (1.2e-30 / 5.1e-27 / 2.0e-27 / 1.8e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	19.05	G23(k55 > k53)	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO <sub>3</sub> + O <sub>2</sub> → CO <sub>2</sub> + O <sub>3</sub>	k56 = (1.9e-26 / 7.9e-25 / 2.6e-26 / 3.1e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-19.05	Single reaction pathway	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO + RAD → C + O	k57 = (3.5e-06 / 1.0e-08 / 7.5e-08 / <1.0e-08) s <sup>-1</sup>	1072.05	Single reaction pathway	[3, 5] gas-phase
C + O → CO	k58 = (2.6e-26 / 7.7e-26 / 2.1e-24 / 1.1e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-1072.05	Single reaction pathway	[3] surface [11] ice and desorption
CO + CO → C <sub>2</sub> O <sub>2</sub>	k59 = (1.2e-26 / 5.2e-26 / 2.4e-25 / 1.3e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	241.6	G11 (k59 > k30 > k63 > k61)	
C <sub>2</sub> O <sub>2</sub> + RAD → CO + CO	k60 = (9.1e-03 / 5.6e-03 / 2.0e-02 / 1.1e-02) s <sup>-1</sup>	-241.6	G15 (k60 > k32 > k93)	
CO + CO → C <sub>2</sub> + O <sub>2</sub>	k61 = (8.4e-31 / 1.0e-29 / 1.1e-28 / 4.1e-31) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	1047.9	G11 (k59 > k30 > k63 > k61)	
C <sub>2</sub> + O <sub>2</sub> → CO + CO	k62 = (1.2e-26 / 8.8e-26 / 1.7e-26 / 6.2e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-1047.9	G24 (k62 > k28)	[4] gas-phase
CO + CO → C <sub>2</sub> O + O	k63 = (1.5e-28 / 7.7e-28 / 2.1e-27 / 2.8e-31) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	851.6	G11 (k59 > k30 > k63 > k61)	
C <sub>2</sub> O + O → 2 CO	k64 = (2.3e-26 / 4.3e-26 / 4.0e-26 / 4.0e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-851.6	Single reaction pathway	[3,5] gas-phase

(continued on next page)

Table 1 (continued)

Modelled reactions <sup>a</sup>	Effective rate constant (CR1 / CR2 / EL / UV)	Reaction enthalpy at 0 K <sup>b</sup> [kJ/mol]	Reaction groups ( <i>k</i> values ordered based on reaction exothermicity) <sup>c</sup>	References or comments
CO + C → C <sub>2</sub> O	k65 = (1.7e-27 / 3.7e-25 / 6.6e-28 / 1.9e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	−220.47	G26 (k65 > k67)	[5] gas-phase; [8] ice
C <sub>2</sub> O + RAD → CO + C	k66 = (2.3e-04 / 2.8e-03 / 1.8e-03 / 9.9e-04) s <sup>-1</sup>	220.47	G14 (k66 > k91)	[3, 4] gas-phase
CO + C → C <sub>2</sub> + O	k67 = (1.1e-28 / 3.4e-25 / 4.1e-28 / 5.2e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	469.58	G26 (k66 > k67)	
C <sub>2</sub> + O → CO + C	k68 = (4.7e-27 / 5.3e-26 / 7.6e-27 / 1.1e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	−469.58	Single reaction pathway	[5] gas-phase
CO + O → C + O <sub>2</sub>	k69 = (8.4e-31 / 1.8e-28 / 4.5e-27 / 2.4e-28) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	578.3	G8 (k2 > k69)	
C + O <sub>2</sub> → CO + O	k70 = (2.0e-26 / 9.9e-26 / 2.9e-25 / 1.1e-27) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	−578.3	G9 (k4 > k70)	[3] surface; [5,3] gas-phase [11] ice and desorption
CO + C <sub>2</sub> → C <sub>2</sub> O + C	k71 = (1.3e-27 / 2.5e-26 / 1.1e-27 / 1.6e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	381.9	Single reaction pathway	
C <sub>2</sub> O + C → CO + C <sub>2</sub>	k72 = (1.1e-27 / 9.6e-25 / 4.7e-26 / 1.3e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	−381.9	Single reaction pathway	[3] gas-phase
CO + C <sub>2</sub> O → C <sub>2</sub> O <sub>2</sub> + C	K73 = (4.7e-27 / 4.8e-26 / 3.9e-28 / 1.6e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	462.9	G18(k46 > k73)	
C <sub>2</sub> O <sub>2</sub> + C → CO + C <sub>2</sub> O	K74 = (2.1e-26 / 2.6e-25 / 1.4e-26 / 3.5e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	−462.9	G27 (k74 > k48)	
CO + O <sub>2</sub> → C + O <sub>3</sub>	K75 = (5.9e-27 / 4.6e-26 / 7.9e-26 / 8.4e-27) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	969.59	G19 (k77 > k16 > k75)	
C + O <sub>3</sub> → CO + O <sub>2</sub>	K76 = (7.6e-28 / 1.1e-25 / 6.9e-26 / 1.9e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	−969.59	G22 (k18 > k76)	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO + O <sub>2</sub> → CO <sub>3</sub>	K77 = (1.9e-26 / 4.3e-25 / 5.9e-25 / 1.1e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	−56.50	G19 (k77 > k16 > k75)	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO <sub>3</sub> + RAD → CO + O <sub>2</sub>	K78 = (2.5e-04 / 9.6e-03 / 2.2e-04 / 1.2e-08) s <sup>-1</sup>	56.50	G12 (k20 > k78 > k87)	[5] gas-phase Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO + O <sub>3</sub> → CO <sub>3</sub> + O	K79 = (2.1e-28 / 8.7e-26 / 2.0e-25 / 4.4e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	51.04	G10 (k22 > k79)	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO <sub>3</sub> + O → CO + O <sub>3</sub>	K80 = (1.3e-26 / 1.5e-24 / 4.3e-26 / 2.8e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	−51.04	G28 (k26 > k80)	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
O <sub>2</sub> + RAD → O + O	K81 = (4.1e-05 / 4.8e-03 / 8.0e-03 / <1.0e-08) s <sup>-1</sup>	493.68	Single reaction pathway	[3, 4] gas-phase
O + O → O <sub>2</sub>	K82 = (8.4e-31 / 1.4e-27 / 7.9e-26 / 1.3e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	−493.68	Single reaction pathway	[3] surface [4] gas-phase; [11] ice and desorption
O <sub>3</sub> + RAD → O <sub>2</sub> + O	K83 = (3.7e-05 / 1.7e-03 / 1.1e-04 / 1.2e-04) s <sup>-1</sup>	102.44	G13 (k83 > k85)	[3,5] gas-phase [10] ice
O <sub>2</sub> + O → O <sub>3</sub>	k84 = (9.4e-28 / 8.3e-27 / 1.3e-26 / 6.6e-27) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	−102.44	Single reaction pathway	[3] surface [11] ice and desorption
O <sub>3</sub> + RAD → O + O + O	k85 = (3.7e-05 / 1.6e-03 / 7.6e-05 / 2.1e-05) s <sup>-1</sup>	596.13	G13 (k83 > k85)	
O + O + O → O <sub>3</sub>	k86 = (8.5e-51 / 1.9e-49 / 2.1e-48 / 3.2e-49) cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	−596.13	Single reaction pathway	
CO <sub>3</sub> + RAD → CO + O + O	k87 = (1.5e-05 / 4.0e-03 / 1.7e-04 / <1.0e-08) s <sup>-1</sup>	573.17	G12 (k20 > k78 > k87)	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
CO + O + O → CO <sub>3</sub>	k88 = (7.0e-53 / 2.7e-49 / 8.9e-50 / 6.1e-50) cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	−573.17	Single reaction pathway	Employing Δ <sub>r</sub> H from <a href="https://cccbdb.nist.gov/">https://cccbdb.nist.gov/</a>
C <sub>2</sub> + RAD → C + C	k89 = (3.4e-05 / 2.0e-03 / 1.5e-03 / 3.2e-04) s <sup>-1</sup>	602.5	Single reaction pathway	[3, 5] gas-phase
C + C → C <sub>2</sub>	k90 = (7.6e-28 / 5.5e-26 / 1.9e-26 / 7.8e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	−602.5	Single reaction pathway	[3] surface [11] ice and desorption
C <sub>2</sub> O + RAD → C + C + O	k91 = (1.8e-04 / 1.7e-03 / 1.5e-04 / 9.3e-04) s <sup>-1</sup>	1292.52	G14 (k66 > k91)	
C + C + O → C <sub>2</sub> O	k92 = (1.3e-48 / 3.1e-48 / 2.4e-49 / 3.9e-49) cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	−1292.52	Single reaction pathway	



Table 1 (continued)

Modelled reactions <sup>a</sup>	Effective rate constant (CR1 / CR2 / EL / UV)	Reaction enthalpy at 0 K <sup>b</sup> [kJ/mol]	Reaction groups ( <i>k</i> values ordered based on reaction exothermicity) <sup>c</sup>	References or comments
C <sub>2</sub> O <sub>2</sub> + RAD → C + CO + O	k93 = (9.0e-06 / 1.0e-03 / 4.6e-04 / 2.5e-03) s <sup>-1</sup>	830.4	G15 (k60 > k32 > k93)	
C + CO + O → C <sub>2</sub> O <sub>2</sub>	k94 = (1.4e-48 / 4.5e-48 / 7.2e-50 / 2.3e-50) cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	-830.4	Single reaction pathway	
C <sub>2</sub> O <sub>3</sub> + RAD → CO + CO + O	k95 = (4.4e-05 / 2.3e-02 / 8.9e-04 / 2.2e-03) s <sup>-1</sup>	301.4	G16 (k34 > k95 > k97 > k99)	Employing Δ <sub>f</sub> H(C <sub>2</sub> O <sub>3</sub> ) = -282.2 kJ mol <sup>-1</sup> from Gambi et al. 2001
CO + CO + O → C <sub>2</sub> O <sub>3</sub>	k96 = (2.1e-49 / 2.0e-48 / 1.6e-52 / 5.2e-49) cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	-301.4	single reaction pathway	Employing Δ <sub>f</sub> H(C <sub>2</sub> O <sub>3</sub> ) = -282.2 kJ mol <sup>-1</sup> from Gambi et al. 2001
C <sub>2</sub> O <sub>3</sub> + RAD → C <sub>2</sub> O <sub>2</sub> + O	k97 = (4.4e-05 / 1.7e-02 / 1.1e-04 / 2.2e-03) s <sup>-1</sup>	543.0	G16 (k34 > k95 > k97 > k99)	Employing Δ <sub>f</sub> H(C <sub>2</sub> O <sub>3</sub> ) = -282.2 kJ mol <sup>-1</sup> from Gambi et al. 2001
C <sub>2</sub> O <sub>2</sub> + O → C <sub>2</sub> O <sub>3</sub>	k98 = (1.7e-26 / 1.3e-25 / 1.6e-27 / 1.7e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	- 543.0	G17 (k36 > k98)	Employing Δ <sub>f</sub> H(C <sub>2</sub> O <sub>3</sub> ) = -282.2 kJ mol <sup>-1</sup> from Gambi et al. 2001
C <sub>2</sub> O <sub>3</sub> + RAD → C <sub>2</sub> O + O <sub>2</sub>	k99 = (<1.0e-08 / 1.0e-03 / 1.1e-04 / 8.9e-04) s <sup>-1</sup>	619.3	G16 (k34 > k95 > k97 > k99)	Employing Δ <sub>f</sub> H(C <sub>2</sub> O <sub>3</sub> ) = -282.2 kJ mol <sup>-1</sup> from Gambi et al. 2001
C <sub>2</sub> O + O <sub>2</sub> → C <sub>2</sub> O <sub>3</sub>	k100=(5.0e-27 / 1.6e-25 / 4.3e-27 / 3.4e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	-619.3	G20 (k38 > k100)	Employing Δ <sub>f</sub> H(C <sub>2</sub> O <sub>3</sub> ) = -282.2 kJ mol <sup>-1</sup> from Gambi et al. 2001
CO <sub>2</sub> (ice) + RAD → CO <sub>2</sub> (gas-phase)	k <sub>des,CO2</sub> = (2.2e-06 / 8.1e-05 / 9.3e-08 / 5.2e-05) s <sup>-1</sup>	-	-	Intrinsic desorption rate
CO (ice) + RAD → CO (gas-phase)	k <sub>des,CO</sub> = (8.5e-07 / 7.7e-03 / 2.7e-07 / 1.6e-04) s <sup>-1</sup>	-	-	Intrinsic desorption rate
CO <sub>3</sub> (ice) + RAD → CO <sub>3</sub> (gas-phase)	k <sub>des,CO3</sub> = (2.1e-06 / 1.2e-03 / 2.0e-07 / 5.8e-04) s <sup>-1</sup>	-	-	Intrinsic desorption rate
O (ice) + RAD → O (gas-phase)	k <sub>des,O</sub> = (2.4e-05 / 1.7e-03 / 1.6e-07 / 2.7e-04) s <sup>-1</sup>	-	-	Intrinsic desorption rate
O <sub>2</sub> (ice) + RAD → O <sub>2</sub> (gas-phase)	k <sub>des,O2</sub> = (2.4e-05 / 8.8e-04 / 2.9e-08 / 4.3e-04) s <sup>-1</sup>	-	-	Intrinsic desorption rate
O <sub>3</sub> (ice) + RAD → O <sub>3</sub> (gas-phase)	k <sub>des,O3</sub> = (4.6e-06 / 2.2e-03 / 1.7e-07 / 2.8e-04) s <sup>-1</sup>	-	-	Intrinsic desorption rate
C (ice) + RAD → C (gas-phase)	k <sub>des,C</sub> = (9.8e-06 / 3.5e-03 / 9.2e-08 / 8.7e-05) s <sup>-1</sup>	-	-	Intrinsic desorption rate
C <sub>2</sub> (ice) + RAD → C <sub>2</sub> (gas-phase)	k <sub>des,C2</sub> = (1.4e-05 / 6.9e-03 / 6.1e-08 / 2.5e-04) s <sup>-1</sup>	-	-	Intrinsic desorption rate
C <sub>2</sub> O (ice) + RAD → C <sub>2</sub> O (gas-phase)	k <sub>des,C2O</sub> = (1.7e-06 / 1.6e-03 / 5.2e-08 / 5.8e-04) s <sup>-1</sup>	-	-	Intrinsic desorption rate
C <sub>2</sub> O <sub>2</sub> (ice) + RAD → C <sub>2</sub> O <sub>2</sub> (gas-phase)	k <sub>des,C2O2</sub> = (5.2e-07 / 4.6e-03 / 3.6e-08 / 2.5e-05) s <sup>-1</sup>	-	-	Intrinsic desorption rate
C <sub>2</sub> O <sub>3</sub> (ice) + RAD → C <sub>2</sub> O <sub>3</sub> (gas-phase)	k <sub>des,C2O3</sub> = (9.0e-06 / 2.0e-03 / 2.2e-07 / 3.7e-04) s <sup>-1</sup>	-	-	Intrinsic desorption rate
Molecular desorption <sup>d</sup> M (ice) + RAD → M (gas-phase)	Summed desorption rate = (4.1e + 11/ 1.3e + 14/ 1.7e + 11 / 4.8e + 12) molecules s <sup>-1</sup> Summed desorption yield = (4.9e + 03/ 8.0e-04/ 4.0e-01 / 3.0e-02) molecules projectile <sup>-1</sup>	-	-	[1,2] Summed desorption yield

<sup>a</sup>Calculated considering the total incoming projectiles and total desorbed species (employing the following experimental radiation flux:  $\phi_{CR1} = 2 \times 10^8$  Ti ions cm<sup>-2</sup> s<sup>-1</sup>,  $\phi_{CR2} = 2 \times 10^9$  Ni ions cm<sup>-2</sup> s<sup>-1</sup>,  $\phi_{EL} = 16 \times 10^{11}$  electrons cm<sup>-2</sup> s<sup>-1</sup> and  $\phi_{UV} = 2 \times 10^{14}$  photons cm<sup>-2</sup> s<sup>-1</sup>).

<sup>b</sup>For reactions enthalpy calculations it was considered only gas phase species and its lowest state energies in the calculation of the reaction enthalpy at 0 K; For the reactions involving CO<sub>2</sub>, CO, O, O<sub>2</sub>, O<sub>3</sub>, C, C<sub>2</sub>, C<sub>2</sub>O, C<sub>2</sub>O<sub>2</sub> it was employed the values Δ<sub>f</sub>H(CO<sub>2</sub>) = -393.1 kJ mol<sup>-1</sup>; Δ<sub>f</sub>H(CO) = -113.8 kJ mol<sup>-1</sup>; Δ<sub>f</sub>H(O<sub>3</sub>) = 144.3 kJ mol<sup>-1</sup>; Δ<sub>f</sub>H(O) = 246.8 kJ mol<sup>-1</sup>; Δ<sub>f</sub>H(O<sub>2</sub>) = 0 kJ mol<sup>-1</sup>; Δ<sub>f</sub>H(C) = 711.4 kJ mol<sup>-1</sup>; Δ<sub>f</sub>H(C<sub>2</sub>) = 820.2 kJ mol<sup>-1</sup>; Δ<sub>f</sub>H(C<sub>2</sub>O) = 337.1 kJ mol<sup>-1</sup>; Δ<sub>f</sub>H(C<sub>2</sub>O<sub>2</sub>) = 14 kJ mol<sup>-1</sup>, obtained from the Active Thermochemical Tables version 1.122p <https://atct.anl.gov/> (Ruscic et al. 2004; 2005; Ruscic and Bross 2020; Active Thermochemical Tables (ATcT) values based on ver. 1.122p of the Thermochemical Network (2020)); For the reactions involving CO<sub>3</sub> it was assumed the gas-phase formation enthalpies at 0 K of Δ<sub>f</sub>H = 52.5 kJ mol<sup>-1</sup>, and the reactions enthalpies Δ<sub>r</sub>H were obtained from the NIST website (<https://cccbdb.nist.gov/>); For the reactions involving C<sub>2</sub>O<sub>3</sub> it was considered the Δ<sub>f</sub>H(C<sub>2</sub>O<sub>3</sub>) = -282.2 kJ mol<sup>-1</sup> (from Gambi et al. 2001) together with the other formations enthalpies from <https://atct.anl.gov/>;<sup>c</sup> As an hypothesis, we adopt a ranking of ERC based on the thermochemical data. The lowest value of reaction enthalpy (most exothermic reaction) should have the higher ERC (faster reaction) of the reaction group considered. The assignment “single reaction pathway” in this column indicates reactions which the reactants set appeared only once in this table.<sup>d</sup>The reaction order of each equation is assumed to be equal to the number of reactants molecules (for k1 to k100). For the desorption rates, it was considered first order reaction.

[1] Pilling et al. 2010a; [2] Martín-Doménech et al. 2015; [3] KIDA database (<http://kida.astrophy.u-bordeaux.fr>); [4] UMIST database (<http://udfa.ajmarkwick.net>); [5] NIST database (<https://kinetics.nist.gov/kinetics/>); [6] Bittner et al. 2020; [7] Cottin et al. 2003; [8] Gerakines et al. 1996.; [9] Okabe 1978; [10] Moll et al. 1966; [11] Hasegawa & Herbst 1993; [12] Raut & Baragiola 2011; [13] Goumans et al. 2008; In this manuscript it was only considered non-ionic reactions.

Table 2

Comparison between selected effective rate constants (ERCs) with rate constants at solid-phase from literature.

Selected reactions <sup>a</sup>	Solid-phase effective rate constants (ERCs) in pure CO <sub>2</sub> ice (CR1 / CR2 / EL / UV) (this work)	Solid phase rate constant (literature)	Notes and references (literature)
CO <sub>2</sub> + RAD → CO + O	k <sub>1</sub> = (3.1e-06 / 9.4e-07 / 1.6e-06 / 1.3e-08) s <sup>-1</sup>	–	–
CO <sub>2</sub> + EL → CO + O	–	~ 1.7e-5 s <sup>-1b</sup>	[1]; from the irradiation of CO <sub>2</sub> : C <sup>18</sup> O <sub>2</sub> (1:2) at 10 K by 5 keV electrons.
CO <sub>2</sub> + EL → X	–	~ 4e-5 s <sup>-1b</sup>	[1]; from the irradiation of CO <sub>2</sub> : C <sup>18</sup> O <sub>2</sub> (1:2) at 10 K by 5 keV electrons.
CO <sub>2</sub> + CR → X	–	3.6e-4 s <sup>-1c</sup>	[3]; from the irradiation of CO <sub>2</sub> at 13 K by 52 MeV ions.
CO <sub>2</sub> + CR → X	–	3.4e-4 s <sup>-1c</sup>	[4]; from the irradiation of CO <sub>2</sub> at 13 K by 46 MeV Ni ions.
CO + RAD → C + O	k <sub>57</sub> = (3.5e-06 / <1.0e-08 / 7.5e-08 / <1.0e-08) s <sup>-1</sup>	–	–
CO + EL → X	–	2.9e-4 s <sup>-1 d</sup>	[2]; from the irradiation of CO ice at 15 by keV electrons;
CO + EL → X	–	~4.2e-5 s <sup>-1 e</sup>	[6] [1] from the irradiation of <sup>13</sup> CO: C <sup>18</sup> O (1:1) at 10 K by 5 keV electrons.
CO + CR → X	–	1e-4 s <sup>-1c</sup>	[5] from the irradiation of CO ice 13 K 50 MeV Ni ions.
CO + CR → X	–	3e-5 s <sup>-1c</sup>	[5] from the irradiation of CO ice at 13 K 537 MeV Ni ions.

<sup>a</sup> X indicates a unspecified product.<sup>b</sup> Rates from the average destruction of CO<sub>2</sub> and C<sup>18</sup>O<sub>2</sub>. In this model all reactions were considered of first order (distributed in one, two and three steps mechanisms).<sup>c</sup> *k* calculated from the multiplication of ion flux and dissociation cross section (determined from associative exponential function to the experimental data).<sup>d</sup> Occurs in the presence of additional CO (excited) species yielding typically CO<sub>2</sub> and C. It was considered pseudo-first-order reaction for the produced CO<sub>2</sub>. The CO<sub>2</sub> is not destroyed or does not react once produced. Other reactions in the model are all of first order.<sup>e</sup> Obtained from the average values of destruction of <sup>13</sup>CO and C<sup>18</sup>O. It was considered only first order and pseudo-first-order reactions in the chemical network.

[1] Bennett et al (2010); [2] Jamieson et al. (2006); [3] Pilling et al. (2010a). [4] Seperuelo-Duarte et al. (2009); [5] Seperuelo-Duarte et al. (2010); [6] Bennett et al (2009).

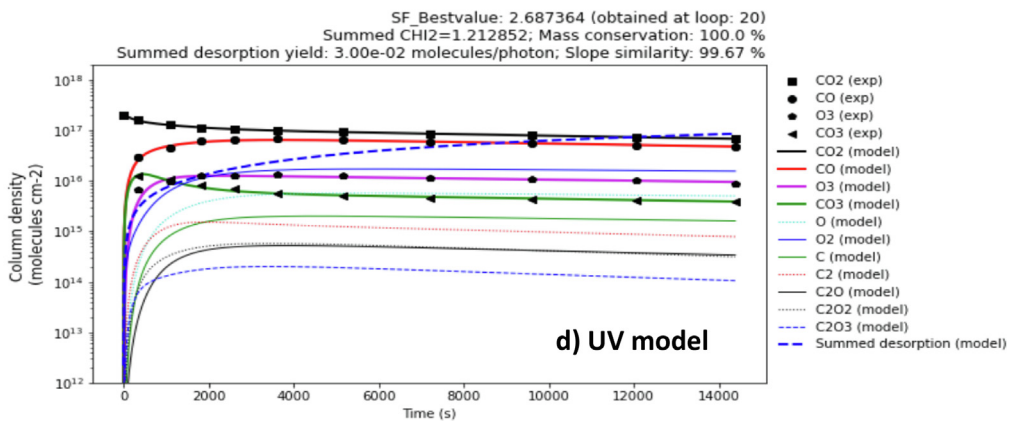
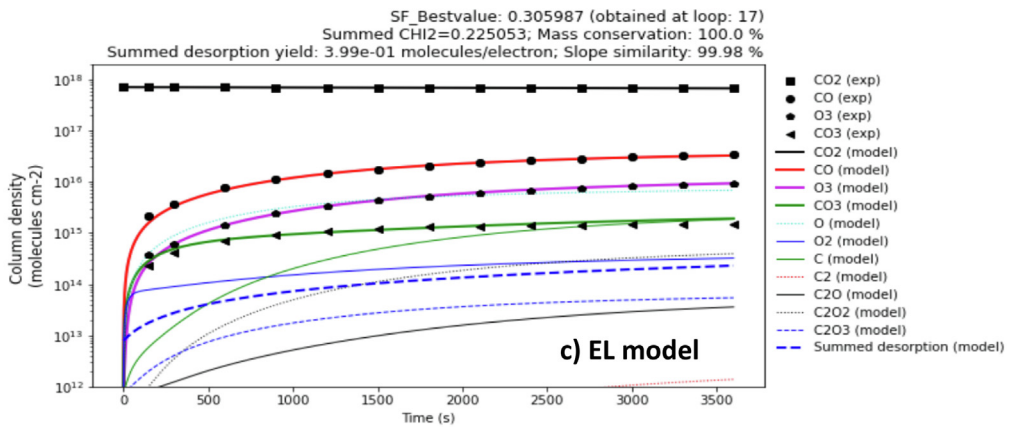
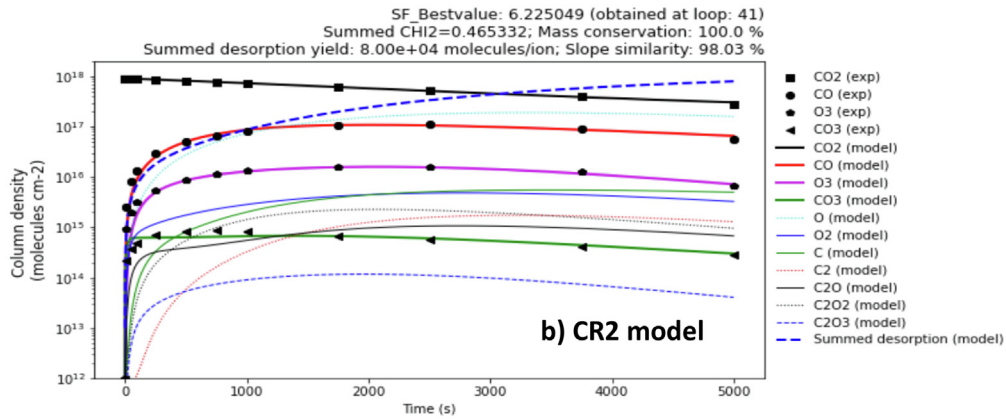
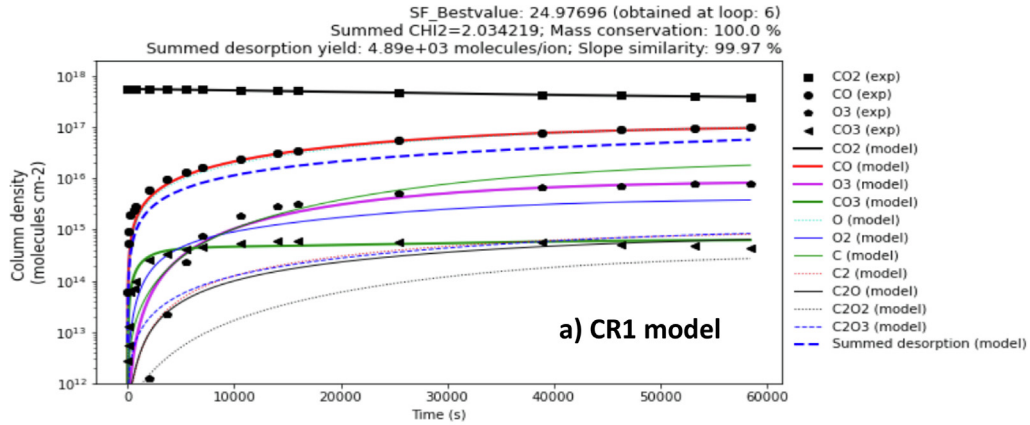
Additionally, in our methodology, we do not consider individual activation energies once such quantities in solid-phase are highly dependent on the chemical environment, local temperature, and density (see also Alves et al. 2021; Pilling and Bonfim, 2020). Since the studied CO<sub>2</sub> ices (and the astrophysical ices in general) are continuously exposed to radiation, the incoming energy (and also the distributed energy within the ice by the fast secondary electrons) would allow exceeding most of the reactions' activation barriers.

In spite of the remaining limitations of the code (e.g. limited number of species and chemical reactions), we expect the current results to be more realistic than the previous version, even understanding that solid-phase thermochemistry data (usually difficult to be determined experimentally and to be calculated) should be different from gas-phase and that some reaction might have an activation barrier. However, the issue about the presence of activation barrier can be exceeded in these irradiated ices if we consider that there is enough energy in the sample due to the incoming ionizing radiation (cosmic rays, fast electrons or UV). A comparison with previous model without the ordering of rate constants (Pilling et al. 2022a),

focused on the CR2 and UV models, is also provided and similarities and differences are discussed.

### 3. Results and discussion

The PROCODA code employed in this work is a powerful tool for the characterization of laboratory studies that perform monitoring of irradiated ices by infrared spectroscopy (e.g. Grim & d'Hendecourt 1986; Ioppolo et al. 2011; Garrod & Pauly 2011; d'Hendecourt et al. 1985; Gerakines et al. 1996; Jimenez-Escobar et al. 2016; Pilling et al. 2009; Pilling et al. 2010a,b, Öberg et al. 2009; Kaiser & Roessler 1998; Moore & Hudson 1998; Hudson & Moore 2001; Baratta et al. 2002; Gerakines et al. 2004; Bennett et al. 2006; Pilling et al. 2010a, 2010b; Ioppolo et al. 2011; Andrade et al. 2013; Almeida et al. 2014; de Barros et al. 2014; Portugal et al. 2014; Boduch et al. 2015; Martín-Doménech et al. 2015; Pilling et al. 2013; Pilling & Bergantini 2015; Suhasaria et al. 2017; Linnartz & Salama 2020; Sie et al. 2019; Tsuge et al. 2020; Mifsud et al. 2022a; Mifsud et al. 2022b; Palumbo 1997; Fillion et al. 2014).



Additionally, the code also can be very useful to help the prediction and the identification of molecules in astronomical observations (e.g. Gerakines et al. 1999; Pontoppidan et al. 2008; Evans et al. 2009, Ehrenfreund & Charnley 2000; Fraser et al. 2002; van Dishoeck 2014; Boogert et al. 2015; Tielens 2013; Kwok 2016). In addition, the ERCs calculated by the code can also be used in astrochemical models to help to understand the chemical evolution of cold space environments (e.g. Wakelam et al. 2010; Holdship et al. 2018; Shingledecker et al. 2019; Heyl et al. 2020; Rocha et al. 2022).

In this section, we will present the main results of the employ of the upgraded PROCODA code to map the chemical evolution of CO<sub>2</sub> ices exposed to ionizing radiation.

### 3.1. Evolution of molecular abundances and effective rate constants (ERCs)

Fig. 2 shows the evolution of column density, obtained for the best-fit models employing the upgraded PROCODA code in the irradiated pure CO<sub>2</sub> ices by different ionizing agents at low temperature. Panels a, b, c and d presents the different models: CR1 model (employing 550 MeV Ti ions), CR2 model (52 MeV Ni ions), EL model (5 keV electrons), and UV model (10 eV photons), respectively. In this figure the experimental data are given by black symbols, the calculated model are given by lines (with the thick lines being the model for the observed species), and the dashed thick blue line is modeled summed desorption. In all panels, important output parameters for the best-fit models are displayed in the header, including the values of chi-squared function  $\chi^2$  (sometimes also named here as summed CHI2).

The models are in a very good match with the experimental data and also reproduce the CE behavior (a slighted sloped plateau observed at large fluences). As discussed by Pilling et al. (2022a), with this methodology, we are able to quantify both the abundances of observable and non-observable species in the IR spectra irradiated ices. Such characterization helps us to better understand the underlying chemistry within the ice during the laboratory experiments and also helps to put constrain in astrochemical observations.

It is worth noting that even though each of these reactions may occur within different chemical environments (surrounded by different molecular arrangements), which

also change over time, only a single reaction rate constant is given per reaction, which does not change over time, and therefore the values in this table are actually the effective rate constants (ERCs) for the entire irradiation process. The term “effective” employed in the ERCs is also employed to take into account ices inhomogeneities in chemistry which locally affect the chemical environment and ice properties, in another word, to account in some way for an average value within the ice (see also Alves et al. 2021; Pilling and Bonfim 2020). Additional details about the model parameters can be obtained at Pilling et al. (2022a) and Carvalho et al. (2022).

Comparing the current CR2 and UV models with the its previous versions, which do not employ any thermochemistry data (Pilling et al. 2022a), showed that current models have better accuracy. This was observed in the value of chi-squared function ( $\chi^2$ ) which decreased by 22 % (from  $\chi^2 = 0.59$  to 0.46) and 36 % (from  $\chi^2 = 1.95$  to 1.25) in the current CR2 and UV model, respectively, compared with previous version. Additionally, the parameter *SSC*, employed to guarantee the CE condition at larger fluences, was closer to 100 % in the current version, which also indicates an improvement in the solution of coupled chemical system.

Table 1 lists the considered chemical reaction routes in this work and the calculated ERC obtained by employing the upgraded PROCODA code taking into account the ERC ordering hypothesis based on thermochemistry data (where reactions with smaller  $\Delta H$  (more exothermic) are expected to have larger ERCs (or *k*), being faster). The thermochemistry data employ literature 0 K gas-phase data for the considered reaction network. The employed reactions set include 11 direct reactions, 69 bimolecular reactions, 11 termolecular reactions, and 11 desorption reactions. The reaction enthalpy and the numbering of the considered reaction groups (with the ERCs ordering based on the reaction exothermicity) are also listed. The intrinsic values for molecular desorption are listed at the bottom of the table. The estimated errors for these rates are below 20 % as discussed by Pilling et al. (2022a).

From Table 1 we notice that some individual reactions are highly endothermic (at gas-phase), which would be also very difficult to occur at low temperature when considered isolated in the ice. However, due to the constant input energy in the ices (which also increase local temperature at the collision point), such endothermic reaction might have a chance to occur overpassing this endothermic

Fig. 2. The evolution of column density obtained for the best-fit models employing the upgraded PROCODA code on pure CO<sub>2</sub> ices irradiated by different ionizing agents at low temperature. Panel a, b, c and d presents the CR1 model (550 MeV Ti ions; Mejía et al. 2015), CR2 model (52 MeV Ni ions; Pilling et al. 2010a), EL model (5 keV electrons; Bennett et al. 2004), and UV (10 eV photons; Martín-Doménech et al. 2015), respectively. The black symbols represent the experimental data (observed molecular abundances for CO<sub>2</sub>, CO, O<sub>3</sub> and CO<sub>3</sub>). The bold-dashed blue line represents the summed desorption column density calculated in each model. In all panels, important output parameters for the best-fit models are displayed in the header.



threshold. Additionally, readers should have in mind that in this work, we describe a set of coupled reactions that occur simultaneously in the ice bulk under bombardment, and reactions in the solid phase behave differently than in the gas phase (they are dependent on neighborhood properties as well as the configuration of neighbor molecules (see also Alves et al. 2021; Pilling and Bonfim 2020)). This also strengthens the employ of the term ERC in this study instead of only rate constant.

Fig. 3 presents, for comparison purposes, the values of the calculated ERC obtained by the best-fit models employing the current PROCODA code (with the ordering hypothesis for ERCs based on thermochemistry data) in the pure CO<sub>2</sub> ices irradiated by different ionizing agents (CR1 model: 550 MeV Ti ions, CR2 model: 52 MeV Ni ions, EL model: 5 keV electrons; UV model: 10 eV photons). Panel a, b, c and d presents the ERC for the direct, the bimolecular, the termolecular and the intrinsic desorption reactions induced by radiation, respectively. The estimated errors for these rates are below 20 % as discussed by Pilling et al. (2022a). The numerical values of this figure are listed in Table 1.

The average values for the ERCs (with the standard deviation error) obtained with the current methodology on pure CO<sub>2</sub> ices under the presence of different ionizing agents is shown in Fig. 4. The panels a, b, c, and d present the average ERCs for the direct, the bimolecular, the termolecular and the radiation-induced intrinsic desorption reactions, respectively. The CR2 model presents the higher value for the average direct, bimolecular and termolecular ERCs and the UV model presents the lowest average values. The calculated average ERCs for the direct dissociation reaction were 6.4e-4, 7.5e-3, 2.1e-3 and 1.6e-3 s<sup>-1</sup> for the CR1, CR2, EL and UV models, respectively. For the bimolecular reactions, the calculated average ERCs were 1.0e-26, 2.4e-25, 1.3e-25 and 1.0e-25 cm<sup>3</sup> molecule<sup>-1</sup> s<sup>-1</sup>, for the CR1, CR2, EL and UV models, respectively. For the termolecular reactions, the calculated average ERCs were 6.7e-49, 8.6e-48, 2.3e-48 and 4.8e-49 cm<sup>6</sup> molecule<sup>-2</sup> s<sup>-1</sup>, for the CR1, CR2, EL and UV models, respectively. A comparison, between the ERC obtained in the current model with the previous one (Pilling et al. 2022a) showed that the average values (only for CR2 and UV) for the direct and bimolecular reactions presented roughly double the previous calculated value. Curiously, for the termolecular rates the comparison with previous calculation shows that the current CR2 model average value is roughly 8 times higher and the UV model average value is double.

The PROCODA code recently was also applied to describe the molecular evolution on pure acetonitrile (CH<sub>3</sub>CN) ice irradiated by X-rays (Carvalho et al. 2022). In this research, the authors employ 273 reaction rates involving 33 molecular species (5 species observed in the experiment and 28 non-observed or unknown) and they determined the average values for dissociation, bimolecular, and termolecular ERCs of 2.3e - 3 s<sup>-1</sup>, 9.7e - 26 cm<sup>3</sup> molecules<sup>-1</sup> s<sup>-1</sup> and 3.2e - 47 cm<sup>6</sup> molecules<sup>-2</sup> s<sup>-1</sup>, respectively.

In general, the average ERCs (with exception for the intrinsic desorption rate) for the EL model lay between the values obtained in the CR2 and UV model indicating that for these three ionizing agents the higher projectile energy the higher is the molecular ERC. Additionally, the average intrinsic desorption rate constant for the electrons was the higher among the studied ionized agents which might be a consequence of the employed experiment technique that is different from the other data (infrared data obtained in the reflection mode and not by transmission mode). A comparison between the two models employing cosmic rays shows that the CR1 model presents the lower average values for the ERC than the values found for the CR2 model, which might be attributed to the different interaction cross sections of the penetration length of projectiles (also associated with different values of stopping power; see details at Pilling et al. 2010b).

It is worth noting that the destruction cross-section for CO<sub>2</sub> obtained from the experiments employing 550 MeV Ti ions (Mejía et al. 2015) is lower than the value obtained employing 52 MeV Ni ions (Pilling et al. 2010a), 7e-14 cm<sup>-2</sup> and 1.8e-13 cm<sup>-2</sup>, respectively. This rule was true for the summed desorption yield determined experimentally in these works, which indicates that the value was lower for 550 MeV Ti ions than for 52 MeV Ni ions (around 1.3e4 and 2.2e4 molecules ion<sup>-1</sup>, respectively).

The current average ERC values for the intrinsic desorption were 8.4e-6 s<sup>-1</sup>, 2.9e-3 s<sup>-1</sup>, 1.2e-7 s<sup>-1</sup> and 2.8e-4 s<sup>-1</sup>, for the CR1, CR2, EL and UV models, respectively. Such values (in the case of Ni ions and UV photons) were around 80 % higher than the obtained by previous calculations (see Pilling et al. 2022a). This indicates that the employ of ERCs ordering hypothesis by thermochemical data induced changes also in the way intrinsic desorption was quantified since some reaction have preferences than another. Moreover, since the current model has a better chi-squared function ( $\chi^2$ ) than the previous model, we argue that current intrinsic desorption values are also more accurate.

Comparisons between selected ERCs obtained with the current methodology with some rate constants taken from literature, in both solid and gas phases, are provided in Tables 2 and 3, respectively. Table 2 compares selected ERCs (mainly CO<sub>2</sub> and CO direct reactions) with some rate constants at solid-phase taken from literature which describes a simple kinetic model to map chemical evolution of CO<sub>2</sub> (Bennett et al. 2010) and CO ices (Bennett et al. 2009; Jamieson et al. 2006), both irradiated by keV electrons. Differently to the current work, which considers, direct, bimolecular, termolecular and radiation-induced desorption reactions, the described literature models employ a number of reactions and chemical species smaller than in the current work, and additionally, all reactions were considered to be of the first order (distributed in one, two and three steps mechanisms). The ERC for the direct dissociation reaction of CO<sub>2</sub> to CO + O, obtained in the current methodology for the EL model was only



one order of magnitude lower than calculated by Bennett et al. (2010). This table also presents some rate constants determined directly from the experimental papers employ-

ing cosmic ray analogs (calculated from the experimental “effective” destruction cross-section) taken from Pilling et al. (2010) and Seperuelo-Duarte et al. (2009).

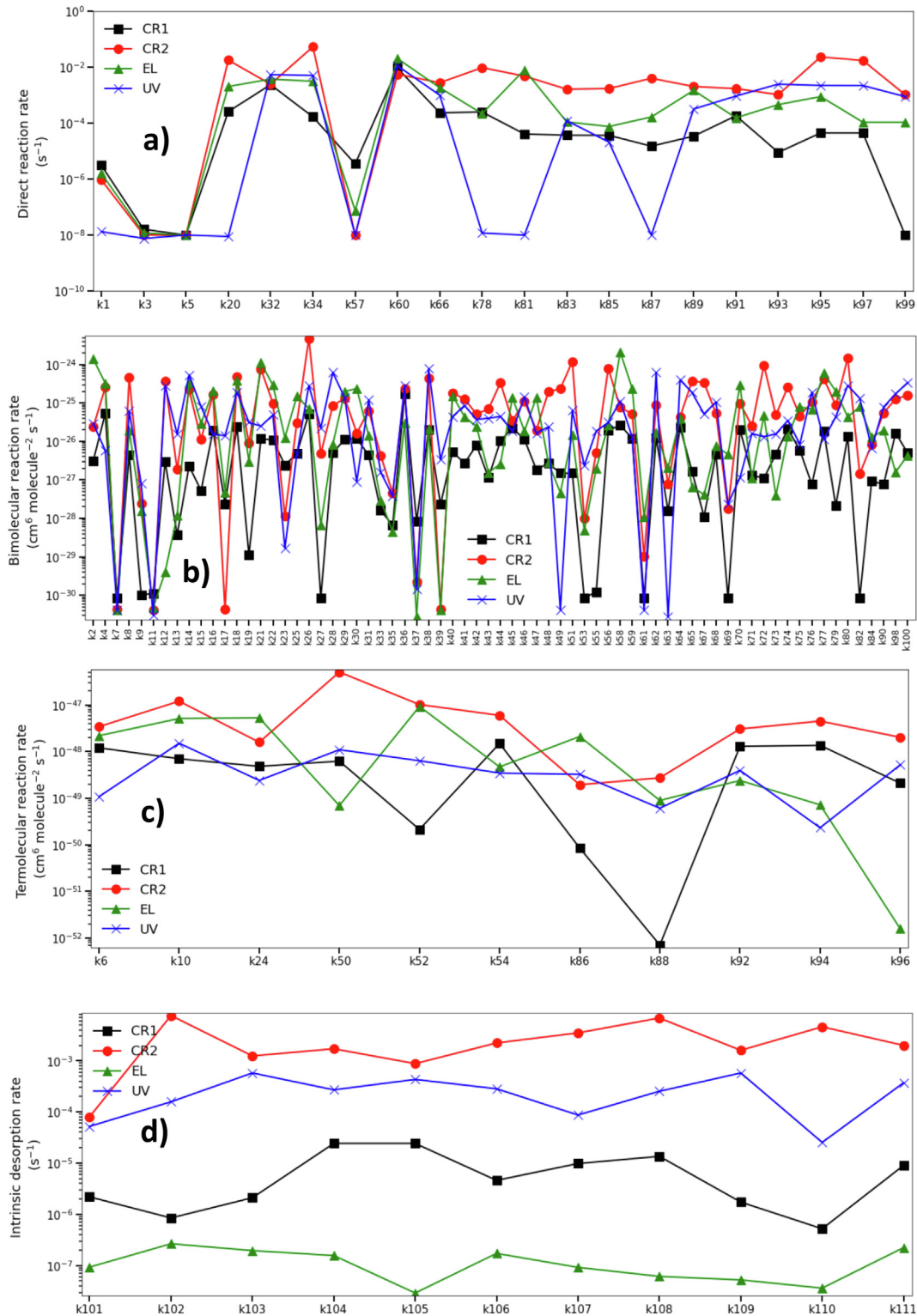


Fig. 3. Values for the effective rate constants (ERCs) obtained by the best-fit models employing current version of the PROCODA code in the irradiated pure CO<sub>2</sub> ices by different ionizing agents (CR1: 550 MeV Ti<sup>+21</sup>, CR2: 52 MeV Ni<sup>+13</sup>, EL: 5 keV electrons; UV: 10 eV photons). Panel a, b, c and d presents the ERCs values for the direct, the bimolecular, the termolecular and the radiation-induced intrinsic desorption reactions, respectively. The numerical values of this figure are listed in Table 1.

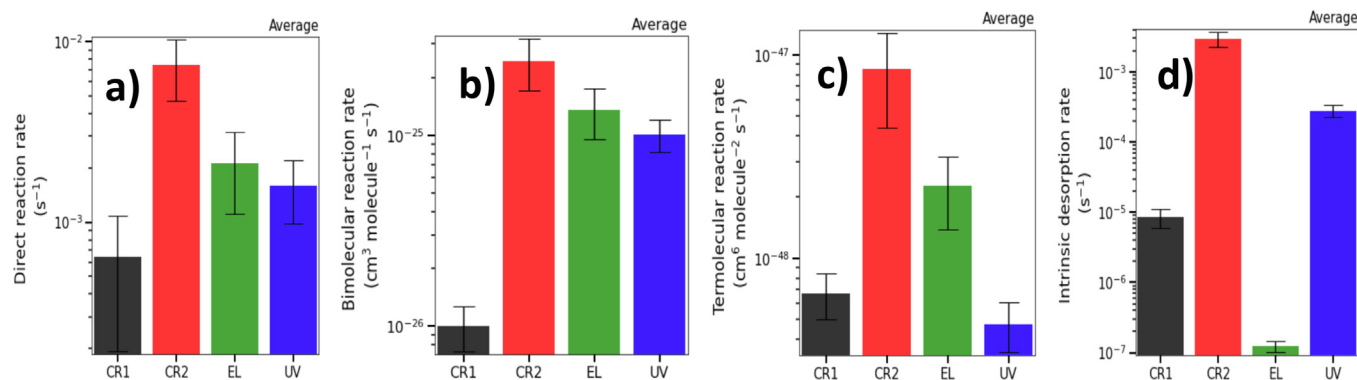


Fig. 4. Comparison between the average values of the effective rate constants (ERCs) employing the upgraded PROCODA code (with the ordering hypothesis for ERCs based on thermochemistry data) in the irradiated pure CO<sub>2</sub> ices. Panel a, b, c and d present the average values (with the standard deviation error) of the ERCs in the direct dissociation reactions, bimolecular reactions, termolecular reactions and intrinsic radiation-induced desorption reactions, respectively. (CR1 model: 550 MeV Ti ions, CR2 model: 52 MeV Ni ions, EL model: 5 keV electrons; UV model: 10 eV photons).

It is worth noting that the current ERC for direct radiation-induced reactions of a given species A (e.g.  $A + RAD \rightarrow products$ ) takes into account also excitation processes triggered by the incoming radiation itself also from the collision with secondary electrons within the ice, leading to the given product (e.g.  $A + RAD \rightarrow A^* \rightarrow products$ ). Such considerations for the excited species are also employed in the calculations of bimolecular and termolecular reactions. Details of such excitation process are described, for example, by Jamieson et al. (2006) and Bennett et al (2010).

Table 3 presents, for a comparison purpose, the ERCs for selected reactions obtained with current methodology with some rate constants at gas-phase taken from different works in the literature (e.g. Mallard et al. 1994; Gredel et al. 1989, van Dishoeck, 1988; van Dishoeck et al. 2006; Husain & Young 1975). For direct reactions, the values determined here (independently of the ionizing radiation employed) are much higher than the gas-phase values. Probably, this issue is due to the fact that ERC handle several processes at the same time, also involving ionic and excited states and also involving the collision with sec-

ondary electrons within the ice bulk, making the reaction faster than gas phase. Such phenomenon was previously discussed by Pilling et al. (2011) during a comparison of dissociation cross sections of organic species during bombardments of ices and gases by X-rays.

For bimolecular reactions, the calculated values for the ERC presented here are several orders of magnitude smaller than in the values at gas-phase, and this is probably related with the difficulty in the mobility and the diffusion of reactants within the ice bulk. The small values found, in comparison with the gas-phase, may also indicate that such collision processes should play a smaller role in ice chemistry; however it is worth noting that some new produced molecular species (e.g. C<sub>2</sub>O<sub>2</sub>) only can be explained by invoking such collision reactions within the ices.

### 3.2. Branching ratio within the considered reaction groups

As discussed in the methodology in the current manuscript we considered a rate constant ordering hypothesis based on gas-phase thermochemistry at 0 K (most exothermic reactions should be faster). In this section, we present

Table 3

Comparisons between selected effective rate constants (ERCs) in ices with rate constants at gas-phase.

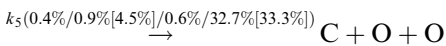
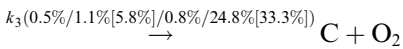
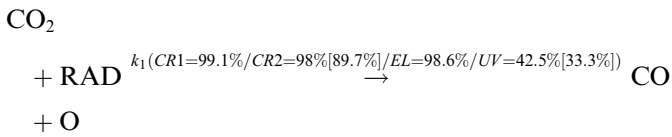
Selected reactions	Solid-phase effective rate constants (ERCs) in pure CO <sub>2</sub> ice (CR1 / CR2 / EL / UV) (this work)	Gas-phase rate constant <sup>a</sup> (literature)	Notes and references (literature)
CO + RAD → C + O	k57 = (3.5e-06 / 1.0e-08 / 7.5e-08 / <1.0e-08) s <sup>-1</sup>	1.30e-17 s <sup>-1</sup> 2.00e-10 s <sup>-1</sup>	[2] Cosmic ray [3] UV photons
CO <sub>2</sub> + RAD → CO + O	k1 = (3.1e-06 / 9.4e-07 / 1.6e-06 / 1.3e-08) s <sup>-1</sup>	1.30e-17 s <sup>-1</sup> 8.90e-10 s <sup>-1</sup>	[2] Cosmic ray [4] UV photons
O <sub>2</sub> + RAD → O + O	K81 = (4.1e-05 / 4.8e-03 / 8.0e-03 / <1.0e-08) s <sup>-1</sup>	1.30e-17 s <sup>-1</sup>	[2] Cosmic ray
CO + O <sub>2</sub> → CO <sub>2</sub> + O	k16 = (1.9e-26 / 1.7e-25 / 2.0e-25 / 1.5e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	5.99E-12 cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	[1] Theoretical
CO <sub>2</sub> + O → O <sub>2</sub> + CO	k15 = (5.3e-28 / 1.1e-26 / 3.0e-26 / 8.0e-26) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	2.46E-11 cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	[1] Theoretical
CO <sub>2</sub> + C → C <sub>2</sub> O <sub>2</sub>	k31 = (4.4e-27 / 6.3e-26 / 1.4e-26 / 1.2e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	< 1E-14 cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	[5] Under UVC photolysis
C <sub>2</sub> O + O → CO + CO	k64 = (2.3e-26 / 4.3e-26 / 4.0e-26 / 4.0e-25) cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	8.59E-11 cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	[1] Theoretical

<sup>a</sup> Employing only the  $\alpha$  term only in the Arrhenius-type formulae (see McElroy et al. 2013). [1] UMIST database; Mallard et al. 1994; [2] UMIST database, Gredel et al. 1989; [3] UMIST database; van Dishoeck, 1988; [4] UMIST database; van Dishoeck et al. 2006; [5] Husain & Young 1975.

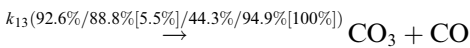
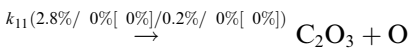
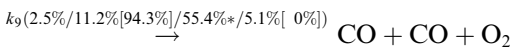
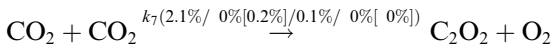
the branching ratio within the considered reaction groups (named from G1 to G28). For comparison purposes, we present in the values between brackets, the branching ratio obtained in the previous PROCODA code version without rate constant ordering hypothesis (Pilling et al. 2022a) for the CR2 and UV. It is worth noting that for some reaction groups (<2% of the calculations) the ERCs ordering hypothesis was not strictly respected during the algorithm minimization to obtain the best-fit solutions (values marked with asterisks in this section). This issue could be due to a minimization algorithm rounding procedure or a small dependence of such reactions in the entire reactions set.

### 3.2.1. Groups of reaction pathways containing CO<sub>2</sub> as one of reactants

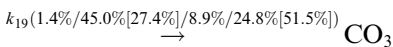
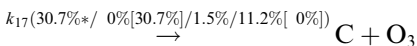
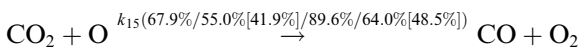
Reaction group G1 (with rate constant ordering hypothesis: k<sub>1</sub> > k<sub>3</sub> > k<sub>5</sub>)



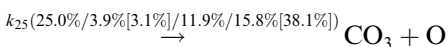
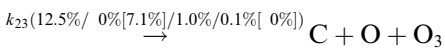
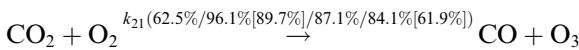
#### Reaction group G2 (k<sub>13</sub> > k<sub>9</sub> > k<sub>11</sub> > k<sub>7</sub>)



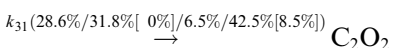
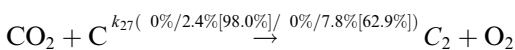
#### Reaction group G3 (k<sub>15</sub> > k<sub>19</sub> > k<sub>17</sub>)



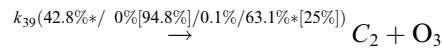
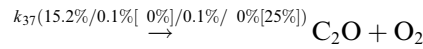
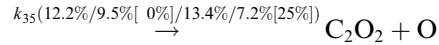
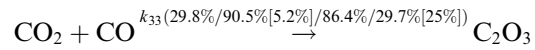
#### Reaction group G4 (k<sub>21</sub> > k<sub>25</sub> > k<sub>23</sub>)



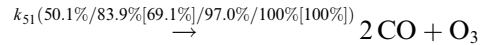
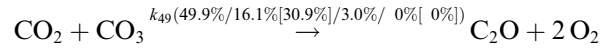
#### Reaction group G5 (k<sub>29</sub> > k<sub>31</sub> > k<sub>27</sub>)



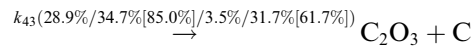
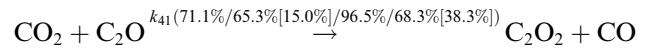
#### Reaction group G6 (k<sub>33</sub> > k<sub>35</sub> > k<sub>37</sub> > k<sub>39</sub>)



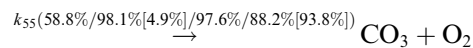
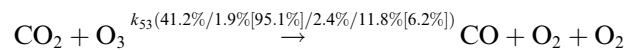
#### Reaction group G7 (k<sub>51</sub> > k<sub>49</sub>)



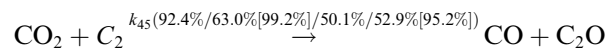
#### Reaction group G21 (k<sub>41</sub> > k<sub>43</sub>)



#### Reaction group G23 (k<sub>55</sub> > k<sub>53</sub>)



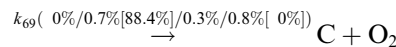
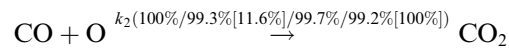
#### Reaction group G25 (k<sub>45</sub> > k<sub>47</sub>)



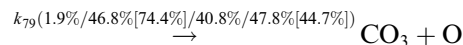
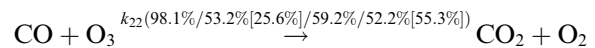
### 3.2.2. Groups of reaction pathways containing CO as one of reactants

#### Reaction group G6 (see reactions with CO<sub>2</sub>)

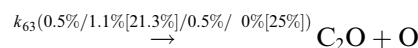
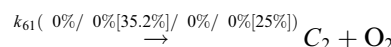
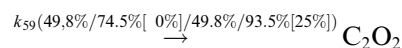
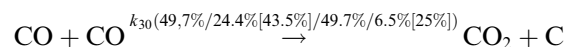
#### Reaction group G8 (k<sub>2</sub> > k<sub>69</sub>)



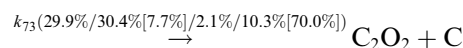
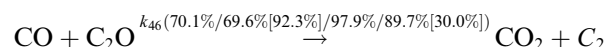
#### Reaction group G10 (k<sub>22</sub> > k<sub>79</sub>)

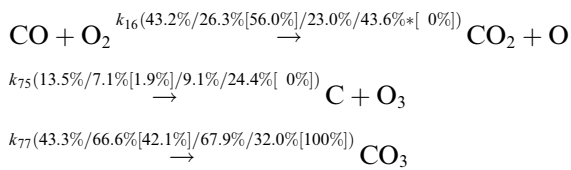
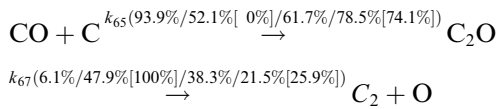


#### Reaction group G11 (k<sub>59</sub> > k<sub>30</sub> > k<sub>63</sub> > k<sub>61</sub>)

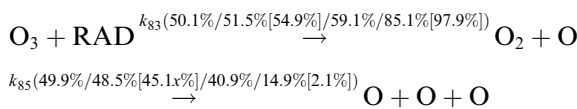
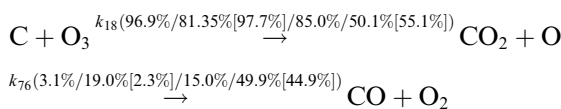
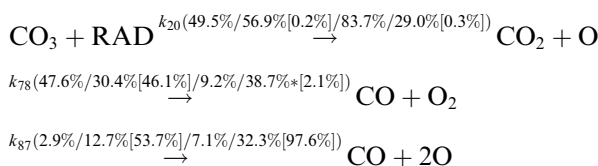
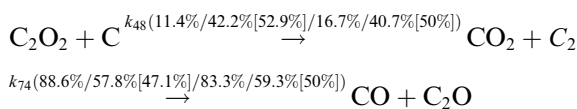
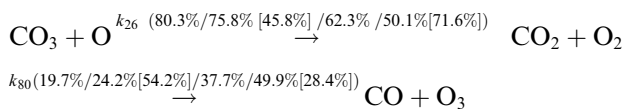


#### Reaction group G18 (k<sub>46</sub> > k<sub>73</sub>)



Reaction group G19 ( $k_{77} > k_{16} > k_{75}$ )Reaction group G26 ( $k_{65} > k_{67}$ )

## 3.2.3. Other reaction groups

Reaction group G13 ( $k_{83} > k_{85}$ )Reaction group G22 ( $k_{18} > k_{76}$ )Reaction group G12 ( $k_{20} > k_{78} > k_{87}$ )Reaction group G27 ( $k_{74} > k_{48}$ )Reaction group G28 ( $k_{26} > k_{80}$ )

A comparison between current CR2 and UV models with the previous models which do not have the hypothesis for ERCs ordering (Pilling et al. 2022a), in which the values are indicated by the values between brackets, suggests that changes in the ordering for ERC is more recurrent in case of CR2 model (observed in 46 % of the reaction groups in comparison with previous model: G2, G5, G6, G8, G10, G11, G12, G19, G21, G23, G26, G27 and G28). For the current UV model it was observed that 32 % of the reaction groups (G1, G3, G6, G11, G12, G18, G19, G21 and G27) presented changes in the ERC's ordering in comparison with the previous model. At 18 % of the reaction groups (G4, G7, G13, G22, and G25) the ordering of reactions was unaltered within the reaction groups, by comparing

current and previous code version. However, it is worth noting that even in these groups, the individual values of each ERC has changed.

A detailed physicochemical discussion about the dominant reactions in each case and why the branching ratio of specific reactions changes with the incoming radiation is out of the scope of this manuscript and should be done in future publications.

## 3.3. Radiation-induced molecular desorption

Fig. 5 presents the radiation-induced desorption column density and the intrinsic molecular desorption for the best-fit models obtained in this work. The CR2 and UV models have presented similar behavior for the desorption column density with a rapid enhancement of CO desorption after the beginning of irradiation becoming the dominant desorption species at the CE. For the CR2 model the four most largely desorbed species at the CE were CO, O, CO<sub>2</sub> and O<sub>3</sub> (in this order). For the UV model the four most important desorbed species at the CE were CO, CO<sub>2</sub>, O<sub>2</sub> and O<sub>3</sub> (in this order). For the CR1 model the four most important desorbed species at the CE were CO<sub>2</sub>, O, CO and CO<sub>3</sub> (in this order). Curiously, this indicated that depending on the type/energy of incident cosmic ray on given astrophysical ice this could yield a different molecular desorption scheme. The EL model present a desorption profile quite different than the other models being the CO<sub>2</sub> species dominant one during the entire irradiation, as well as, at the CE phase.

As observed also by Pilling et al (2022a), the ordering of the desorption column density for the considered species in the irradiated CO<sub>2</sub> ices changed as function of time during the irradiation, in which some species playing an important role in specific moments (e.g. O and CO in the CR2 model; O<sub>2</sub>, O<sub>3</sub> and CO<sub>3</sub> in the UV model). Curiously, this behavior was not seen in the EL model which might be related with low penetration of incoming electrons in the ice. The changing in the desorption column density ordering with time also indicates that gaseous atmosphere in the vicinity of icy grains in space evolved, as function of time, until the CE phase is reached. This will be investigated with details in future investigation.

A comparison with previous models without the ordering hypothesis for ERCs (Pilling et al. 2022a) shows similar behavior in the desorption column density with some punctual differences in the molecular desorption ordering at the CE phase. Changes were also observed in the intrinsic molecular desorption between these two approaches (in the UV model mainly for CO, O, C<sub>2</sub>O<sub>2</sub> and C<sub>2</sub>O<sub>3</sub>, and in the CR2 model mainly for the species CO<sub>3</sub>, O, O<sub>2</sub>, C<sub>2</sub> and C<sub>2</sub>O). The current UV model have presented the same summed desorption yield (3e-2 molecules/photons) than previous UV model. For the CR2 model we observed a small decrease (<10 %) by comparing the summed desorption yield in the current code version (8e+4 molecules/ion) with the previous version.



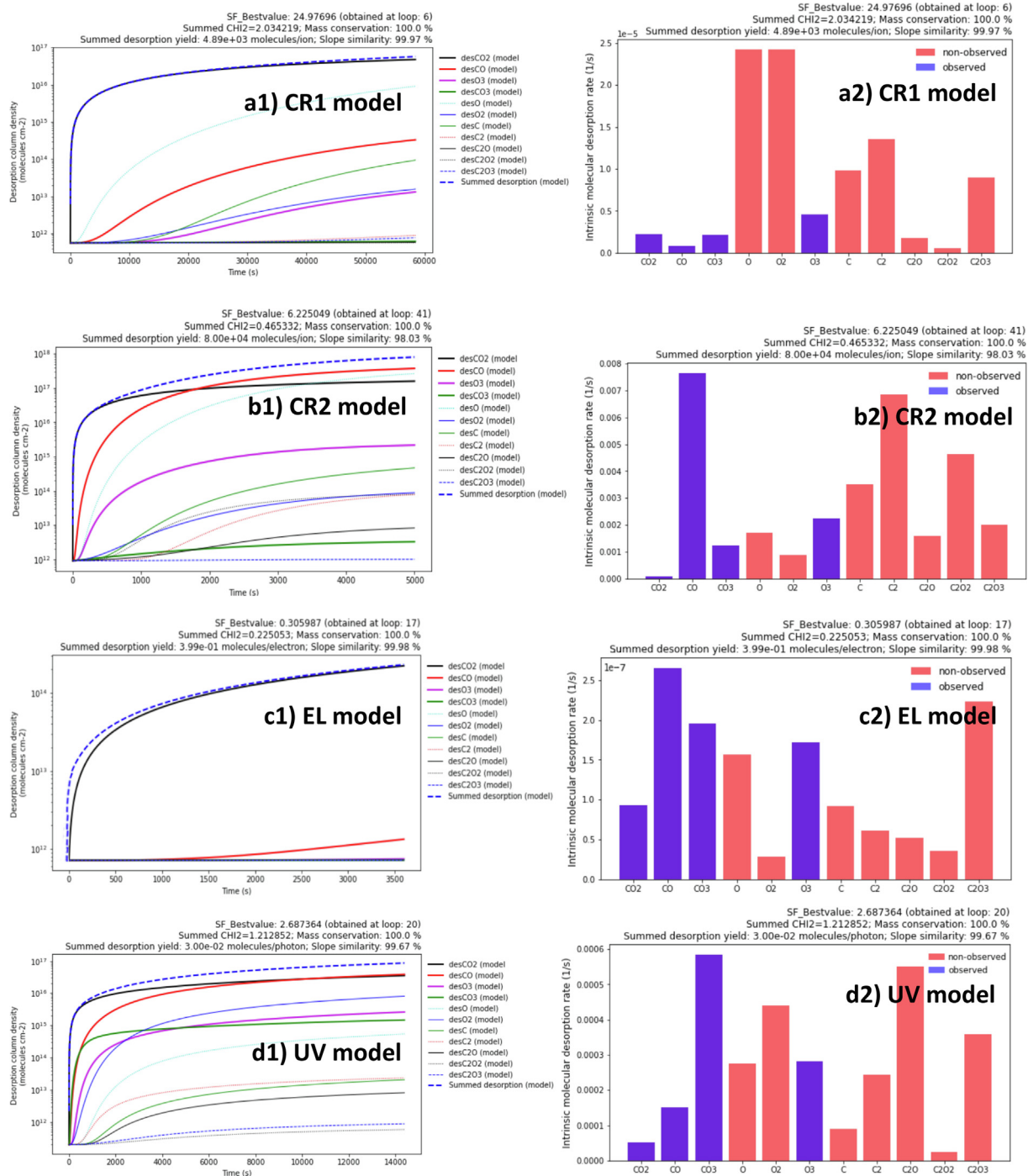


Fig. 5. Molecular desorption plots of best-fit models employing the upgraded PROCODA code. Left panels present the desorption column density for the modeled species with the summed value given by bold-dashed blue lines. Right panels present the intrinsic molecular desorption effective rate constant (DERC) for each modeled chemical species. Panels a, b, c and d presents the different models (CR1 model: 550 MeV Ti ions, CR2 model: 52 MeV Ni ions, EL model: 5 keV electrons; UV model: 10 eV photons), respectively.

Finally, it worth noting that in the current calculations we do not differentiate the desorbed species from the sputtered species (important in the case of cosmic rays). Moreover, the calculations not take into account specific desorption mechanisms, or reactive desorption species or excited states. Here, we considered only that all species that go to the gas phase (removed from the ice) are desorbed species.

### 3.4. The chemical equilibrium (CE) phase

The CE phase (or stage) of irradiated astrophysical ices occurs when the variation of summed chemical abundances is negligible, which happens at larger radiation fluences. In this situation, which requires an ice at constant temperature, the production and consumption of a given species are virtually equal, which keep chemical abundances nearly



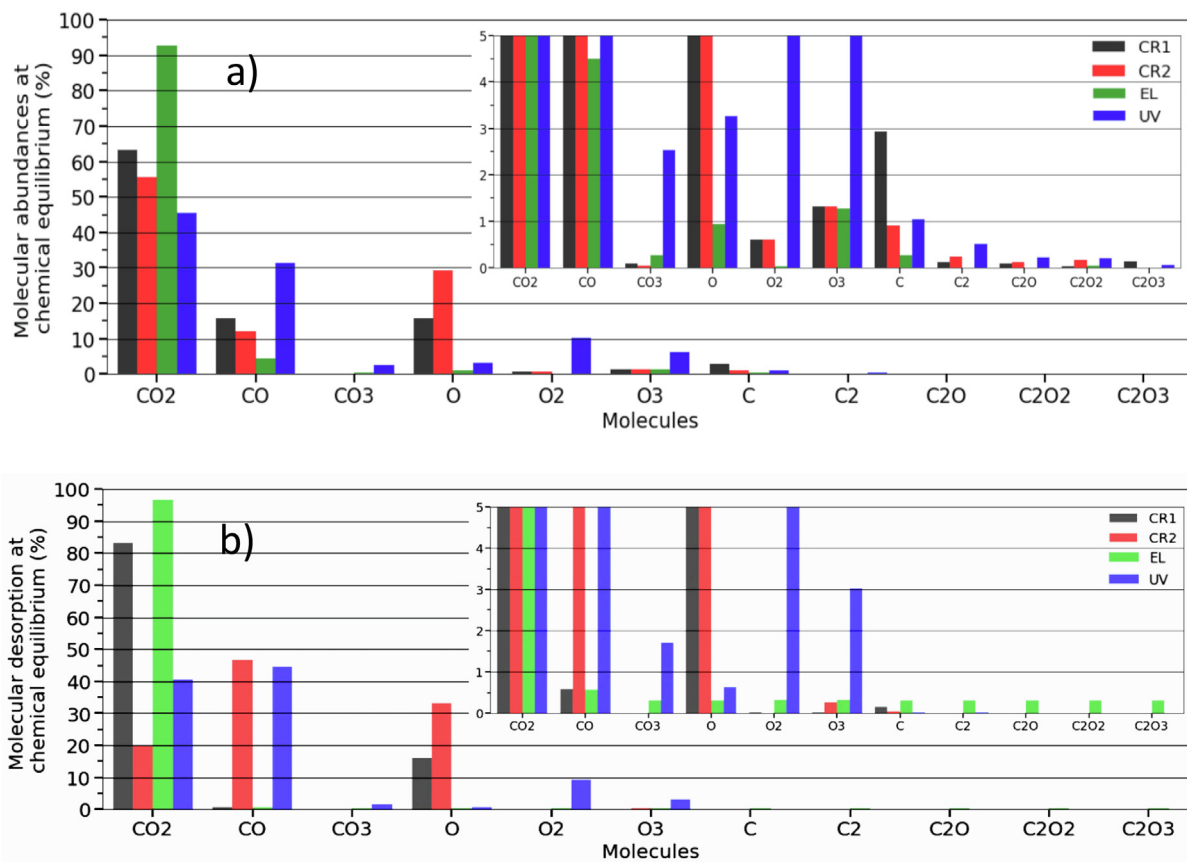


Fig. 6. Percentage values for the molecular abundances (panel a) and molecular desorption (panel b) at chemical equilibrium for the studied pure  $\text{CO}_2$  ices irradiated by different ionizing radiation, obtained the upgraded PROCODA code (with the ordering of ERC for selected reactions based on thermochemistry data). The inset panel in each figure is a zoom of the main figure for percentages below 5%. (CR1 model: black color, CR2 model: red color, EL model: green color, UV model: blue color).

constant (e.g. Almeida et al. 2017; de Souza Bonfim et al. 2017; Rachid et al. 2017; Vasconcelos et al. 2017a, 2017b, 2017c) It is worth noting that the CE phase depends on the initial molecular abundances, the ice temperature and type of incoming ionizing radiation (as is observed in this manuscript). This phase also characterizes a certain timescale in the ice's life (named timescale to reach the CE phase) where the changes in chemical abundances in an astrophysical ice remains nearly unaltered even after extended irradiation (see also Carvalho & Pilling 2020a, 2020b; Freitas & Pilling 2020, Pilling et al. 2022a).

Fig. 6 shows, in percentage, the molecular abundances (panel a) and molecular desorption (panel b) at CE phase, obtained with the upgraded version of PROCODA code. Each color indicates a specific model studied in this manuscript (CR1 model: black color, CR2 model: red color, EL model: green color, UV model: blue color). It is worth noting that the presented percentages were calculated directly from the values of molecular column densities at the CE phase. From Fig. 6a, we observe that in all models the  $\text{CO}_2$  species is the most abundant one in the CE phase, being considerably more abundant in the case of experiments employing electrons (92.6%). The second most abundant species was the CO in most experiments, except-

ing in the case of CR2 model which was the oxygen atoms (29.1%) a non-observed species in the ice. It is worth noting that the abundances at the CE phase for the species CO, O,  $\text{C}_2$  and  $\text{O}_3$ , are considerably larger in the UV model than in the other models, which can help to constrain the main radiation field in the processing of ices within different astrophysical regions. Fig. 6a also highlights an enhanced production of O atoms (and also C atoms) in the ices irradiated by CRs, which might be related with the larger destruction of eventual new-produced species such as CO (the dominant product when UV photon is applied),  $\text{O}_2$ ,  $\text{O}_3$  and  $\text{C}_2$ .

The desorption column density at the CE, in percentage, shown in Fig. 6b suggests that the experiments can be separated into two different groups: group A – large desorption of parental species (the case of CR1 and EL models) and group B – large desorption of the daughter species CO (for CR2 and UV models). Table 4 list the values of molecular abundances, in percentage, determined at the CE employing the upgraded PROCODA for the studied  $\text{CO}_2$  ices. The values between brackets, presented for comparison purposes, result for the CR2 and UV models obtained in previous calculation without the ordering hypothesis for ERCs (Pilling et al. 2022a).

A comparison between current CR2 and UV models with respective models performed with the previous code version (Pilling et al. 2022a) shows good similarities in the case of UV model and some interesting differences in the case of CR2 model (the current model yields, at the CE phase, almost the double of CO<sub>2</sub> species and a very small content of O<sub>2</sub>, C and C<sub>2</sub> with respect to the previous model). Important differences were also observed in the amount of summed observed and non-observed species in the ice in the current CR2 model with respect to the previous model. This comparison indicates, once more, that the ordering ERCs by thermochemistry data is more relevant when more energy is deposited in the sample. It is worth noting that the model that presents the largest and the lowest abundance of the parent species in the CE phase was the EL model and UV model, respectively. Both cosmic ray models presented the largest production of O atoms, being the lowest production observed in the EL model (the same behavior was also observed for C atoms). The largest production of O<sub>3</sub> was observed in the UV model, being the other models presenting curiously similar abundances in the CE phase (1.3 %). The formation of O<sub>2</sub> molecule seems to be much enhanced when UV photons is employed in respect with other ionization sources. Finally, we highlight that the differences in the abundances observed at CE phase might help to put constrain in astrophysical models depending on the dominant type of incoming radiation in a given astrophysical icy scenario.

Table 5 present the values of molecular desorption, in percentage, at the CE phase obtained in the best-fit models employing the upgraded PROCODA code at four irradiated pure CO<sub>2</sub> ices. The values between brackets, presented for comparison purposes, results for the CR2 and UV models obtained in the previous calculation without the ordering hypothesis for ERCs (Pilling et al. 2022a). A comparison between current CR2 and UV models with the previous models shows good similarities in the case of UV models and some interesting differences in the case of CR2 models (main differences occur for desorption of CO, O, and C). Important differences were also observed in the amount of summed observed and non-observed species in the ice in the current CR2 model with respect to the previous model, indicating that desorption in current model is much more ruled by observed species, an opposite behavior from previous model. This comparison indicates that the ordering of ERCs values by thermochemistry data is much more important for the ices that are exposed to higher energy delivered as the case of CR2 model in respect with the UV model.

### 3.5. Effective equilibrium constant for the irradiated ice

When a simple chemical system reaches the CE phase involving only one forward–backward reaction



where  $R_i$  and  $P_i$  indicate the different reactants and products species (with  $r_i$  and  $p_i$  its respective stoichiometric coefficient in the reaction) and  $k_{for}$  and  $k_{bck}$  being the rate constant for the forward and backward reactions, respectively, we can write a thermodynamic parameter called chemical equilibrium constant  $K$  following the equation

$$K = \frac{k_{for}}{k_{bck}} = \frac{\prod [P_i]^{p_i}}{\prod [R_i]^{r_i}} \quad (4)$$

where  $[R]^{r_i}$  and  $[P]^{p_i}$  indicate the concentration of a given reactant and product at equilibrium, in units of column density, with its respective stoichiometric coefficient in the reaction, respectively.

Depending on the value of  $K$  we can state that the CE phase is reactant-favored (for  $K < 1$ ) or product-favored (for  $K > 1$ ). The chemical equilibrium constant is related with other important thermodynamics parameters of the system such as the change in the Gibbs free energy  $\Delta G$  (“the available energy”) or the changes in the enthalpy  $\Delta H$  and entropy  $\Delta S$  by the equation

$$K = e^{\frac{-\Delta G}{RT}} = e^{\frac{-\Delta H + T\Delta S}{RT}} \quad (5)$$

where  $R$  is the perfect-gas constant (8.314 J K<sup>-1</sup> mol<sup>-1</sup>) and  $T$  is the temperature in kelvin. Additionally, since  $K$  is related with entropy changes it helps to understand also the evolution of the chemical complexity of the system (e.g. Atkins et al. 2014).

In the case of coupled and multiple equations the equilibrium constant of the ice system  $K_{ice}$  can be written as

$$\begin{aligned} K_{ice} &= \prod_{i=1}^n K_i = \prod_{i=1}^n \frac{k_{i,for}}{k_{i,bck}} \\ &= \prod_{i=1}^n \left( \frac{\prod [P_i]^{p_i}}{\prod [R_i]^{r_i}} \right) [\text{molecules cm}^{-2}]^{\Delta n} \end{aligned} \quad (6)$$

where  $[R]^{r_i}$  and  $[P]^{p_i}$  indicate the concentration of a given reactant and product at equilibrium, in units of column density, with its respective stoichiometric coefficient in the reaction, respectively (adapted from Flowers et al. 2015). In this equation  $K_i$  represents the equilibrium constant of each pair of forward–reverse reaction employing in the current model (excluding the desorption reactions). It is worth noting that the unit of the equilibrium constant is given by (molecules cm<sup>-2</sup>)<sup>Δn</sup>, where Δn = 1 is obtained by the sum of stoichiometric coefficients of products minus the sum of stoichiometric coefficients of reactants.

Considering that in the current work we employ 50 pair of forward–backward reactions (described in the Table 1 from k1 to k100; were k2 is the reverse of k1, k4 is the reverse of k3 and so on) we can write the equilibrium constant of the ice system as multiplication of 50 quotients as is illustrated shortly by

Table 4

Value of molecular abundances (in number of molecules) of frozen species, in percentage, at the CE phase obtained in the best-fit models employing the upgraded PROCODA code in the four irradiated pure CO<sub>2</sub> ices.

Molecular species	CR1 Model	CR2 Model <sup>b</sup>	EL Model	UV Model <sup>b</sup>
CO <sub>2</sub> <sup>a</sup> (parent)	63.2 %	55.6 % [33.8 %]	92.6 %	45.5 % [41.5 %]
CO <sup>a</sup>	15.6 %	11.9 % [7.9 %]	4.5 %	31.3 % [29.7 %]
CO <sub>3</sub> <sup>a</sup>	0.1 %	<0.1 % [<0.1 %]	0.3 %	2.5 % [2.2 %]
O	15.9 %	29.1 % [30.4 %]	0.9 %	3.3 % [3.8 %]
O <sub>2</sub>	0.6 %	0.6 % [3.1 %]	<0.1 %	10.2 % [14.9 %]
O <sub>3</sub> <sup>a</sup>	1.3 %	1.3 % [0.9 %]	1.3 %	6.2 % [5.8 %]
C	2.9 %	0.9 % [23.1 %]	0.3 %	1.1 % [0.8 %]
C <sub>2</sub>	0.1 %	0.2 % [0.8 %]	~0%	0.5 % [0.3 %]
C <sub>2</sub> O	0.1 %	0.1 % [0.1 %]	~0%	0.2 % [0.4 %]
C <sub>2</sub> O <sub>2</sub>	<0.1 %	0.2 % [~0%]	<0.1 %	0.2 % [0.1 %]
C <sub>2</sub> O <sub>3</sub>	0.1 %	~0% [~0%]	~0%	<0.1 % [0.5 %]
<b>Summed observed species</b>	<b>80.2 %</b>	<b>68.9 % [42.6 %]</b>	<b>98.7 %</b>	<b>85.5 % [79.2 %]</b>
<b>Summed non-observed species</b>	<b>19.8 %</b>	<b>31.1 % [57.4 %]</b>	<b>1.3 %</b>	<b>14.5 % [20.8 %]</b>

<sup>a</sup> Observed species by FTIR.

<sup>b</sup> The values between brackets, presented for comparison purpose, results for the CR2 and UV models obtained in previous calculation without the ordering hypothesis for ERCs (Pilling et al. 2022a).

Table 5

Values of molecular desorption, in percentage, at the CE phase obtained in the best-fit models employing the upgraded PROCODA code in the four irradiated pure CO<sub>2</sub> ices.

Molecular species	CR1 Model	CR2 Model <sup>b</sup>	EL Model	UV Model <sup>b</sup>
CO <sub>2</sub> <sup>a</sup> (parent)	83.2 %	19.8 % [15.6 %]	96.6 %	40.6 % [25.5 %]
CO <sup>a</sup>	0.6 %	46.7 % [12.2 %]	0.6 %	44.6 % [61.1 %]
CO <sub>3</sub> <sup>a</sup>	~0%	~0% [~0%]	0.3 %	1.7 % [0.9 %]
O	15.9 %	33.1 % [57.4 %]	0.3 %	0.6 % [0.7 %]
O <sub>2</sub>	<0.1 %	~0% [0.3 %]	0.3 %	9.4 % [10.9 %]
O <sub>3</sub> <sup>a</sup>	<0.1 %	0.3 % [0.1 %]	0.3 %	3.0 % [0.9 %]
C	0.2 %	<0.1 % [14.3 %]	0.3 %	<0.1 % [<0.1 %]
C <sub>2</sub>	~0%	~0% [<0.1 %]	0.3 %	<0.1 % [~0%]
C <sub>2</sub> O	~0%	~0% [~0%]	0.3 %	~0% [~0%]
C <sub>2</sub> O <sub>2</sub>	~0%	~0% [~0%]	0.3 %	~0% [~0%]
C <sub>2</sub> O <sub>3</sub>	~0%	~0% [~0%]	0.3 %	~0% [~0%]
<b>Summed observed species</b>	<b>83.9%</b>	<b>66.8 % [27.9%]</b>	<b>97.8 %</b>	<b>89.9 % [88.4 %]</b>
<b>Summed non-observed species</b>	<b>16.1 %</b>	<b>33.2 % [72.1 %]</b>	<b>2.2 %</b>	<b>10.1 % [11.6 %]</b>

<sup>a</sup> Observed species by FTIR.

<sup>b</sup> The values between brackets, presented for comparison purpose, results for the CR2 and UV models obtained in previous calculation without the ordering hypothesis for ERCs (Pilling et al. 2022a).

$$K_{ice} = K_1 K_2 K_3 \dots K_{50}$$

$$= \left( \frac{[CO][O]}{[CO_2]} \right) \left( \frac{[C][O_2]}{[CO_2]} \right) \left( \frac{[C][O]^2}{[CO_2]} \right) \dots \left( \frac{[C_2O][O_2]}{[C_2O_3]} \right) \quad (7)$$

However, since in the studied ices here the CO<sub>2</sub> species is the original parent species, we can write the forward–backward reaction to describe the CE phase considering the CO<sub>2</sub> the only reactant species being the other species named as the products. After making the stoichiometric balance of the combined forward–backward full reactions (employing the entire reaction set, excluding the desorption reactions) we can write an effective forward–backward reaction in the chemistry equilibrium as

$$[CO_2]^{11} \xrightleftharpoons[k_{bck}]{k_{for}} [CO][CO_3][O][O_2][O_3]^3 [C][C_2][C_2O][C_2O_2] \times [C_2O_3] \quad (8)$$

In short, eq. (8) presents the simplest effective forward–backward reaction that describes the ice system under irradiation in the chemical equilibrium phase (consider all modelled species). This procedure not considers any desorption reactions from ice to gas-phase. It is worth noting that there were other possibilities to write the effective forward–backward reaction in the chemistry equilibrium (see eq. (8)) considering 11 CO<sub>2</sub> molecules as reactants, for example, the ones obtained by changing the term [O][O<sub>2</sub>][O<sub>3</sub>]<sup>3</sup>, in the products, by [O]<sup>2</sup>[O<sub>2</sub>]<sup>2</sup>[O<sub>3</sub>]<sup>2</sup>,

$[O][O_2]^4[O_3]$ ,  $[O]^3[O_2]^3[O_3]$ ,  $[O]^5[O_2]^2[O_3]$  or  $[O]^7[O_2][O_3]$ . However such alternative combinations suggest a strongly product-favored chemical network in the equilibrium which is not observed in the models as we detailed below.

Considering eq. (8), as the representative for effective forward–backward reaction that describes the ice system under irradiation in the chemical equilibrium phase we can obtain the effective equilibrium constant of the frozen system by

$$K_{ice} = \left( \frac{[CO][CO_3][O][O_2][O_3]^3[C][C_2][C_2O][C_2O_2][C_2O_3]}{[CO_2]^{11}} \right) [\text{molecules cm}^{-2}] \quad (9)$$

where the abundances for the considered species at the CE are in units of column density. Employing the values of column density at the CH phase obtained from the models in this equation, we calculate the effective equilibrium constant  $K_{ice} = 1.7\text{e-}7$ ,  $6.4\text{e-}8$ ,  $2.0\text{e-}20$ ,  $3.2\text{e}0$  molecules  $\text{cm}^{-2}$ , for the CR1, CR2, EL, UV models, respectively. This indicates that the CE phase in the considered network is strongly reactant-favored for the CR1, CR2 and EL models (mainly this model), however it is virtually neutral (neither side favored) for the UV model (see also Silberberg, 2012).

In this work, we consider eq. (9) as the representative one for the effective forward–backward reaction in the chemistry equilibrium (the one with the lowest possible amount of products) because the effective equilibrium constant obtained employing the other discussed possibilities (different products combinations) suggest an strongly product-favored chemical network in the equilibrium which is not observed in the models (see Fig. 2).

#### 4. Astrochemical implications

As discussed previously, carbon dioxide is one of the important species in space, widely observed in molecular clouds. The quantification of its chemical pathways in the solid phase under the presence of ionizing radiation such as UV photons, swift electrons or CRs is essential to better understand the redistribution of carbon in the ISM since  $\text{CO}_2$  holds a large reservoir of depleted carbon in astrophysical ices. Additionally, the  $\text{CO}_2$  bending mode is an important diagnostic tracer of the ice structure and temperature (e.g. Chiar et al. 1995, Ehrenfreund et al. 1997; Pontoppidan et al. 2003; Baratta & Palumbo 2017).

As reported by Öberg et al. (2011),  $\text{CO}_2$  is the second most abundant frozen molecule with respect to  $\text{H}_2\text{O}$  ice as observed toward low-mass protostars (29%), and quiescent molecular clouds (38%), and it has abundance similar to CO ice toward high-mass protostars (13%). However, depending on the object, the carbon monoxide abundances may overpass the  $\text{CO}_2$  abundant in gas-phase space (see also Huang et al. 2020). The employ of PROCODA to map the chemical evolution of CO ice exposed to ionizing radiation in the laboratory (pure CO ice at 13 K irradiated by 50 MeV Ni ions) was

recently submitted for publication (Pilling et al. 2022b). This work found that the most abundant species, at the CE phase, for such experimental data were atomic oxygen (68.2%) and atomic carbon (18.2%), followed by CO (11.8%) and  $\text{CO}_2$  (1.6%).

While some computational codes simulating surface chemical reactions have been made available (e.g., Ruaud et al. 2016, Holdship et al. 2017), they mostly rely on reaction rates of gas-phase molecules, which may lead to inaccurate results. In this paper, we present new reaction rates (presented in terms of effective rate constants, ERCs) focusing on the  $\text{CO}_2$  ice chemistry triggered by ionizing radiation at different energy levels (2 different types of cosmic rays, UV and 5 keV electrons). The upgrade in the PROCODA code allows to ordering the chemical reactions using thermochemistry data, and consequently, accurate physicochemical parameters can be calculated. Additionally, the current improvements added in the code contributes to decrease the inaccuracies of chemical reactions triggered by photons, electrons or ions in frozen  $\text{CO}_2$ , as well its daughters species, in the astrochemical models. Therefore, the main astrochemical implication of current work is the claim for the employ of these calculated values of ERC in an astrochemical models. This will better describe astrochemical ices under processing by radiation and also better describe desorption of specific compounds to gas-phase induced by the incoming radiation in the ices.

Additionally, the characterization of the CE phase for the  $\text{CO}_2$  ices under the presence of different ionizing fields provide by this work show interesting aspects. For example, this species is much more consumed in the ices when 52 MeV Ni ions impinge the ices in comparison with the 550 MeV Ti ions. At this phase the production of CO is almost the same under the presence of both UV and 52 MeV Ni ions. The production of  $\text{O}_2$  and  $\text{O}_3$  is enhanced in the presence of UV photons but the production of O is enhanced in the presence of CRs. Additionally from the analysis of branching ratio of reactions within the studied reaction groups we outline that some reactions are much more important depending on the incoming radiation (See details in section 3.1). It is worth noting that combining the quantification of chemical abundances of ices taken from space observations (such the ones performed by the JWST) with the molecular abundances quantified for the CE phase (perfumed in the current work) may also help to constrain the type and the role of the incoming ionizing radiation field in a given observed astrophysical ice/region.

Among the species studied in this work, there are three of them that we would like to highlight their astrochemical relevance:  $\text{C}_2\text{O}$ ,  $\text{C}_2\text{O}_2$ , and  $\text{C}_2\text{O}_3$  (see structures at Fig. 1). The species  $\text{C}_2\text{O}$  is a linear molecule that is of interest in many areas. However, despite its simplicity, it is a very reactive species that is not encountered commonly, being detected for the first time in space environments at the cold dark molecular cloud TMC-1 (e.g. Ohishi et al. 1991). It is called an oxocarbon and can be used as a building block to synthesize oxocarbon anions (Seitz and Imming, 1992).



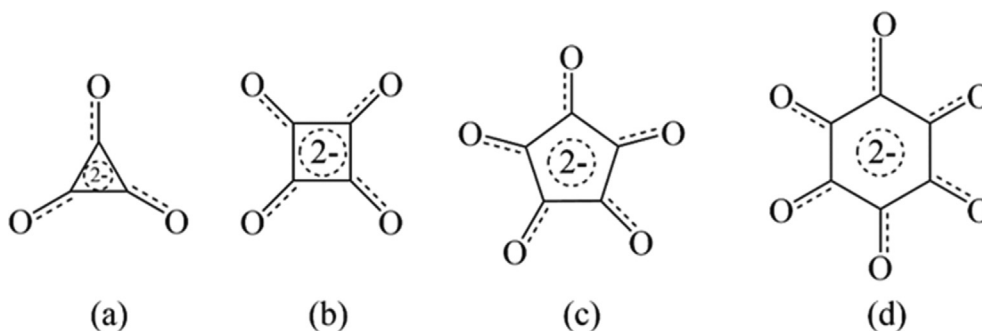


Fig. 7. Structures of oxocarbon anions. (a) deltate, (b) squarate, (c) croconate and (d) rhodizonate.

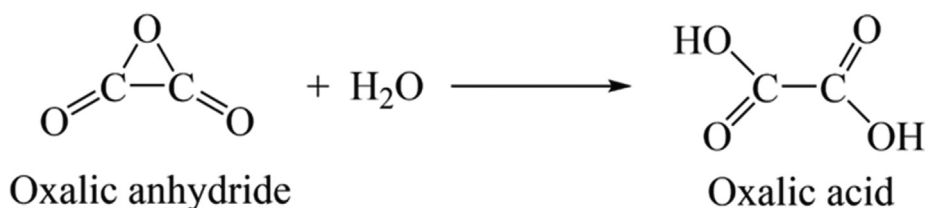


Fig. 8. Proposed hydrolysis reaction of oxalic anhydride forming oxalic acid.

Oxocarbon anions are cyclic aromatic structures that contain only oxygen and carbon atoms, such as deltate, squarate, croconate, and rhodizonate as can be seen in Fig. 7. Finally, C<sub>2</sub>O is also an important ligand used in coordination chemistry (Frenking and Tonner, 2009).

The species C<sub>2</sub>O<sub>2</sub>, named ethenedione or carbon monoxide dimer is a fascinating species for a couple of reasons. It is a pretty simple molecule that, in principle, there is no reason not to exist. However, it was a challenging task to synthesize it (Trindle, 2003; Talbi and Chandler, 2000). This species is a dimer of carbon monoxide (Fig. 1d), and new oligomers of well-known molecules have been the subject of research for a long time. (Gambi et al. 2001; Groves and Lewars, 2000) Calculations indicate that its ground state should be a triplet, such as an oxygen molecule (Trindle, 2003; Talbi and Chandler, 2000) even that it is possible to infer a closed-shell Kekulé structure for this species.

The oxalic anhydride, C<sub>2</sub>O<sub>3</sub>, is a hypothetical organic species of particular interest because it can be seen as a precursor of a supposed prebiotic species (Lavado et al. 2019) which also highlight an astrobiological implication of the current work. It is important to mention that C<sub>2</sub>O<sub>3</sub> could suffer a hydrolysis reaction to form oxalic acid, a well-known organic compound. Fig. 8 shows a proposed hydrolysis reaction of the oxalic anhydride to form oxalic acid. This reaction can easily occur since water molecules (and mainly water-rich ices) are ubiquitous structures found in the interstellar medium. (see also Perks & Liebman 2000, Talukdar 2020).

The quantification of chemical species not visible in the IR range may also contribute to a better estimate of the elemental abundance ratios in the solid-phase in astrophysical environments. For example, Sturm et al. (2022) used sub-

millimeter observations with ALMA (Atacama Large Millimeter/Submillimeter Array) to show variations in the elemental abundances of C and H in the inner and outer part of protoplanetary disks, as well as, among different protostars. The same analysis is also possible for the solid-phase molecules using IR data of JWST. Although this quantification is mostly led by the number of chemical species observed in the spectrum, the IR inactive molecules such as O<sub>2</sub> or even much less abundant (e.g., C<sub>2</sub>O and C<sub>2</sub>O<sub>3</sub>) may also be important to derive elemental abundances ratios between C and O in protoplanetary disks. The rate constants of these species derived with PROCODA and their use in gas-grain chemical models may be useful to derive reliable values in the context of molecular clouds and protostars.

Finally, we highlight that new insights on CO<sub>2</sub> ice chemistry will come out via observations with the James Webb Space Telescope (JWST), for example, with the abundance variation mapping of CO<sub>2</sub> ice in planet-forming regions. To understand the complexity of protostellar ices, chemical modeling such as the one employed by Ballering et al. (2021), which combines radiative transfer calculations with chemical data, is required. The results presented here, in special the determined ERC for reactions within the ice, the molecular abundances at CE phase, and the desorption yields, when employed in such models will help to put constraints on newly produced species (specially C<sub>2</sub>O, C<sub>2</sub>O<sub>2</sub>, and C<sub>2</sub>O<sub>3</sub>) in the astronomical observations.

## 5. Conclusions

In this manuscript, we present an upgraded version of the PROCODA code (original code at Pilling et al 2022a) which describes the chemical evolution of ices processed



by ionizing radiation considering now an ordering hypothesis for the effective rate constants (ERCs) employing thermochemistry data (taken from literature) for the reactions within the groups of reactions. The current ordering hypothesis helps to identify the most important reactions within the reaction network and decreases the degeneracy of the solutions and enhancing the accuracy of the calculations (enhances the match between model and experimental data). Here, we described the chemical evolution of four experiments employing pure CO<sub>2</sub> ices processed by radiation: two ices bombarded by cosmic ray analogues (Pilling et al. 2010a; Mejía et al. 2015), one bombarded by 5 keV electrons (Bennett et al. 2004) and another irradiated by UV photons (Martín-Doménech et al. 2015). The main conclusions of this manuscript were:

- i) The current results presented more accuracy when compared with previous calculations even understanding that solid-phase thermochemistry data is different from gas-phase and that some reaction might have activation barrier. This was observed in the value of summed chi-squared function ( $\chi^2$ ), which decreased by 22 % and 38 % in the current CR2 and UV model, respectively, compared with previous code version, as well as, in the CE criterion (slope similarity) which was closer to 100 % in the current version.
- ii) The calculated average direct ERC were 6.4e-4, 7.5e-3, 2.1e-3 and 1.6e-3 s<sup>-1</sup>, for the CR1, CR2, EL and UV models, respectively. For the bimolecular reactions the calculated average ERCs were 1.0e-26, 2.4e-25, 1.3e-25 and 1.0e-25 cm<sup>3</sup> molecule<sup>-1</sup> s<sup>-1</sup>, for the CR1, CR2, EL and UV models, respectively. A comparison rates (only for CR2 and UV), between the average ERCs obtained here with the previous code version showed that the average direct and bimolecular reactions were roughly double the previous calculated value.
- iii) Current calculated average intrinsic desorption ERCs were 8.4e-6, 2.9e-3, 1.2e-7, and 2.8e-4 s<sup>-1</sup>, for the CR1, CR2, EL and UV models, respectively. These values (only for CR2 and UV) were 80 % higher than the ones obtained previously (see Pilling et al. 2022a). The desorption column density at the CE phase suggest that the experiments can be separated in two groups: group A – large desorption of parental species (the case of CR1 and EL models) and group B – large desorption of the daughter species CO (for CR2 and UV models).
- iv) In the CE phase, the most abundant species were the CO<sub>2</sub>, being considerably more abundant in the case of experiments employing electrons (92.6 %). The second most abundant species was the CO in most experiments, except in the case of CR2 model which was the oxygen atoms (29.1 %) a non-observed species in the ice. Additionally, the abundances of CO, O, C<sub>2</sub> and O<sub>3</sub>, at the CE phase were considerably lar-

ger in the UV model than in the other models, which may help to constrain the main radiation field in the processing of ices present in different astrophysical regions.

- v) The comparison between the upgraded version of the code with the previous one indicates that the ordering of ERCs by thermochemistry data in the models is much more important for the ices that are exposed to higher energy delivered as the case of CR2 model in respect with the UV model.

The upgraded PROCODA code is not only important to map the chemical evolution of simple ices under processing of radiation in the laboratory but also important to characterize the evolution of more complex ices (e.g. H<sub>2</sub>O:CO: NH<sub>3</sub> ices, mixed ices containing water and organic species, etc.). This could allow the characterization of important organic species in the ices that were not detected by the experiments, with also important implications in the astrobiology field. We hope the current methodology helps to clarify the solid-state astrochemistry models of ices under the presence of radiation, as well as, helps to explain the observations of frozen and desorbed molecules in cold space environments.

#### Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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