



UNIVERSITY OF LEEDS

This is a repository copy of *Formation of cometary O² ice and related ice species on grain surfaces in the midplane of the pre-solar nebula*.

White Rose Research Online URL for this paper:

<https://eprints.whiterose.ac.uk/140996/>

Version: Published Version

Article:

Eistrup, C and Walsh, C orcid.org/0000-0001-6078-786X (2019) Formation of cometary O² ice and related ice species on grain surfaces in the midplane of the pre-solar nebula. *Astronomy and Astrophysics*, 621 (A&A). A75. ISSN 0004-6361

<https://doi.org/10.1051/0004-6361/201833380>

© ESO 2019. Reproduced in accordance with the publisher's self-archiving policy. Originally published online; <https://doi.org/10.1051/0004-6361/201833380>

Reuse

Items deposited in White Rose Research Online are protected by copyright, with all rights reserved unless indicated otherwise. They may be downloaded and/or printed for private study, or other acts as permitted by national copyright laws. The publisher or other rights holders may allow further reproduction and re-use of the full text version. This is indicated by the licence information on the White Rose Research Online record for the item.

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



eprints@whiterose.ac.uk
<https://eprints.whiterose.ac.uk/>

Formation of cometary O₂ ice and related ice species on grain surfaces in the midplane of the pre-solar nebula

Christian Eistrup¹ and Catherine Walsh²

¹ Leiden Observatory, Leiden University, PO Box 9513, 2300 RA Leiden, The Netherlands
e-mail: eistrup@strw.leidenuniv.nl

² School of Physics and Astronomy, University of Leeds, Leeds LS2 9JT, UK
e-mail: c.walsh1@leeds.ac.uk

Received 7 May 2018 / Accepted 20 July 2018

ABSTRACT

Context. Detection of abundant O₂ at 1–10% relative to H₂O ice in the comae of comets 1P/Halley and 67P/Churyumov-Gerasimenko has motivated attempts to explain the origin of the high O₂ ice abundance. Recent chemical modelling of the outer, colder regions of a protoplanetary disk midplane has shown production of O₂ ice at the same abundance as that measured in the comet.

Aims. We aim to carry out a thorough investigation to constrain the conditions under which O₂ ice could have been produced through kinetic chemistry in the pre-solar nebula midplane.

Methods. We have utilised an updated chemical kinetics code to evolve chemistry under pre-solar nebula midplane conditions. Four different chemical starting conditions and the effects of various chemical parameters have been tested.

Results. Using the fiducial network, and for either reset conditions (atomic initial abundances) or atomic oxygen only conditions, the abundance level of O₂ ice measured in the comets can be reproduced at an intermediate time, after 0.1–2 Myr of evolution, depending on ionisation level. When including O₃ chemistry, the abundance of O₂ ice is much lower than the cometary abundance (by several orders of magnitude). We find that H₂O₂ and O₃ ices are abundantly produced (at around the level of O₂ ice) in disagreement with their respective abundances or upper limits from observations of comet 67P. Upon closer investigation of the parameter space, and varying parameters for grain–surface chemistry, it is found that for temperatures 15–25 K, densities of 10⁹–10¹⁰ cm⁻³, and a barrier for quantum tunnelling set to 2 Å, the measured level of O₂ ice can be reproduced with the new chemical network, including an updated binding energy for atomic oxygen (1660 K). However, the abundances of H₂O₂ and O₃ ices still disagree with the observations. A larger activation energy for the O + O₂ → O₃ reaction ($E_{\text{act}} > 1000$ K) helps to reproduce the non-detection of O₃ ice in the comet, as well as reproducing the observed abundances of H₂O₂ and O₂ ices. The only other case in which the O₂ ice matches the observed abundance, and O₃ and H₂O₂ ice are lower, is the case when starting with an appreciable amount of oxygen locked in O₂.

Conclusions. The parameter space investigation revealed a sweet spot for production of O₂ ice at an abundance matching those in 67P and 1P, and O₃ and H₂O₂ ice abundances matching those in 67P. This means that there is a radial region in the pre-solar nebula from 120–150 AU, within which O₂ could have been produced in situ via ice chemistry on grain surfaces. However, it is apparent that there is a high degree of sensitivity of the chemistry to the assumed chemical parameters (e.g. binding energy, activation barrier width, and quantum tunnelling barrier). Hence, because the more likely scenario starting with a percentage of elemental oxygen locked in O₂ also reproduces the O₂ ice abundance in 67P at early stages, this supports previous suggestions that the cometary O₂ ice could have a primordial origin.

Key words. molecular processes – methods: numerical – comets: general – comets: individual: 1P/Halley – comets: individual: 67P/Churyumov-Gerasimenko – protoplanetary disks

1. Introduction

The detection of abundant molecular oxygen at 1–10% (average $3.8 \pm 0.85\%$) relative to H₂O ice in the coma of comet 67P/Churyumov-Gerasimenko (hereinafter 67P: Bieler et al. 2015) came as a surprise, as it was the first detection of O₂ in a comet. This was not expected because O₂ ice has been found to be efficiently converted to H₂O ice in laboratory studies under interstellar conditions (e.g. Ioppolo et al. 2008). Subsequent to this detection, a re-analysis of Comet 1P/Halley data from the *Giotto* mission (Rubin et al. 2015) indicates a similar O₂/H₂O ice ratio ($3.8 \pm 1.7\%$), suggesting that indeed O₂ ice may be a common ice species in solar system comets. These detections have thus prompted speculation as to the chemical origin of the O₂ ice. Taquet et al. (2016) modelled the chemical evolution of material from the pre-stellar core stage to the midplane of the formed protoplanetary disk, and found that O₂ ice can be produced at

the early stages and survive the transport to the disk midplane. Mousis et al. (2016) found that if O₂ ice is formed from radiolysis of H₂O ice through the reaction $2\text{iH}_2\text{O} \xrightarrow{\gamma} 2\text{iH}_2 + \text{iO}_2$ (where “i” denotes a molecule in the ice form), then this likely did not happen during the pre-solar nebula (PSN) disk phase, but rather in the parent cloud, thus supporting the findings of Taquet et al. (2016). Dulieu et al. (2017) performed laboratory experiments to investigate if dismutation of H₂O₂ ice ($2\text{iH}_2\text{O}_2 \rightarrow 2\text{iH}_2\text{O} + \text{iO}_2$) on the cometary surface could be the origin of the O₂ detection. However, this explanation requires a high initial abundance of H₂O₂ ice relative to H₂O ice (twice the detected abundance of O₂, or ~7%), and a high efficiency for the conversion of H₂O₂ to O₂ in order to match the low detected level of H₂O₂ relative to O₂ of $\sim 6 \times 10^{-4}$. O₃ ice, a molecule chemically related to O₂, H₂O₂ and H₂O, was not detected in the coma of comet 67P, and has an upper limit of 10⁻⁶ with respect to H₂O ice. We note here that

molecular oxygen is also produced when CO₂ ice is exposed to far-UV radiation (see, e.g., [Martín-Doménech et al. 2015](#)). However, although this may be a viable chemical route to O₂ ice, it does not explain the strong association between the production rates for H₂O and O₂ seen for comet 67P ([Bieler et al. 2015](#)). On the other hand, experiments investigating chemistry in CO₂ ice irradiated with 5 keV ions (H⁺ and He⁺) and electrons ([Ennis et al. 2011](#); [Jones et al. 2014](#)) favour the production of ozone (O₃) over molecular oxygen (O₂). These experiments mimic the conditions that ices are exposed to in the outer solar system, and upon cosmic ray impact.

In [Eistrup et al. \(2016; hereafter Paper I\)](#), we find through chemical kinetic modelling of protoplanetary disk midplanes that O₂ ice could be produced to match the measured cometary abundance by 1 Myr, if the chemical starting conditions were purely atomised and the ionisation level was low ($\sim 10^{-19} \text{ s}^{-1}$). Purely atomised starting conditions reflect the assumption that an energetic stellar outburst or accretion heating close to the star could have fully dissociated all volatiles in the midplane, and low ionisation means that the only ionisation source in the midplane is the decay of short-lived radionuclides. This latter scenario assumes that a magnetic field could have shielded the midplane from cosmic rays (as proposed by [Cleeves et al. 2013](#)). This finding sparked interest into whether or not the chemical origin of the O₂ ice could be chemical processing of the icy material in the PSN disk midplane. Most recently, [Mousis et al. \(2018\)](#) explored the possibility that turbulent transport of icy grains between the disk midplane and the upper layers of the disk exposed the grains to a stronger cosmic-ray flux, thus chemically processing H₂O ice to produce O₂ ice via radiolysis. They find that on a 10 Myr timescale O₂ ice in the midplane remains underproduced by up to two orders of magnitude relative to the abundances observed in the comets. [Eistrup et al. \(2018\)](#) also found that this abundance for O₂ ice be reached on similarly long timescales.

Based on the promising results in Paper I, in this work we have investigated the conditions under which O₂ ice in the PSN midplane could have been chemically produced in situ to reach the observed abundance level in the comets. This work differs from that in [Mousis et al. \(2018\)](#) in that a full chemical kinetics network was used that follows the chemical connection between H₂O, O₂, and related ice species. A more suitable disk model for the PSN was used, and a more thorough investigation of the O₂ kinetic ice chemistry was conducted. Initial chemical abundances, physical conditions, parameters for grain–surface chemistry, as well as the inclusion of O₃ in the chemical network were all tested to see the effect on the O₂ ice production and abundance.

2. Methods

In order to investigate the chemical evolution in the PSN midplane, the physical disk model for the PSN from [Hayashi \(1981\)](#) was utilised. This disk model estimates the structure and mass of the PSN, assuming the total current mass of the planets in the solar system to be distributed as dust and gas in a protoplanetary disk in equilibrium. The mass of this nebula is $M_{\text{MMSN}} = 0.08 M_{\odot}$, based on integrating the PSN surface density structure from [Aikawa et al. \(1997\)¹](#) from 0.1–1000 AU. This provides temperature and density profiles for the disk midplane, as well as the surface density profile. The latter is used to calculate

¹ $\Sigma(R) = 54(R/10\text{AU})^{-3/2} \text{ g cm}^{-2}$.

the attenuation of cosmic rays (herefrom CRs) impinging on the disk, thereby estimating the contribution of CRs to the disk midplane ionisation, as was discussed in detail in Paper I. The disk structure is static in time, which was shown in [Eistrup et al. \(2018\)](#) to not cause a significantly different chemical evolution from an evolving disk structure, apart from the inward shifting of molecular icelines in response to decreasing temperature with time.

In Fig. 1a this PSN midplane temperature and density structure is plotted from 0.1–1000 AU. The temperature decreases by two orders of magnitude from $\sim 900 \text{ K}$ to 9 K from the inner to the outer disk, and the density drops from 10^{18} to 10^6 cm^{-3} . The dynamic ranges, especially in density, are thus larger than what was used in Paper I. However, this disk also extends out to 1000 AU, instead of the 30 AU seen in Paper I. This is because this PSN disk midplane is significantly warmer and more massive than the disk structure in Paper I (0.08 versus 0.01 M_{\odot}), thus the relevant temperature regime for O₂ ice chemistry (10–30 K) is found outside 100 AU here.

2.1. Ionisation levels

In Paper I, we confirm that ionisation is an important driver of chemistry in disk midplanes (see, e.g., [Aikawa et al. 1997](#); [Walsh et al. 2015](#)). It is also found that abundant O₂ ice was produced at low ionisation (only radionuclide decay, and no contribution from cosmic rays). Therefore, three different levels of ionisation are explored here: a low level (Level 1, SLRs only), a high level (Level 2, the fiducial level which includes galactic cosmic rays), and an extra high level (Level 3, which includes enhanced cosmic rays), see Fig. 1b. The two former levels assume the same contributions to the midplane ionisation as was the case for the low and high levels in Paper I: ionisation Level 1 includes only the contribution from short-lived radionuclides (SLRs) in the midplane. Ionisation Level 2 includes in addition CRs impinging on the disk, using the canonical cosmic ray ionisation rate for the local ISM ($\zeta_{\text{H}_2} = 10^{-17} \text{ s}^{-1}$). Ionisation Level 3 assumes a CR ionisation contribution 10 times higher than for ionisation Level 2, based on estimates from e.g. [van Dishoeck & Black \(1986\)](#), [Dalgarno \(2006\)](#), and [Indriolo et al. \(2015\)](#) that the galactic cosmic ray rate ζ_{GCR} could be in the range $\zeta_{\text{H}} = 10^{-17} - 10^{-16} \text{ s}^{-1}$ in diffuse clouds. We explore this enhanced CR scenario because O₂ ice has been found to be efficiently synthesised in experiments studying H₂O ice radiolysis in relation to solar system icy bodies (see, e.g., lab work by [Teolis et al. 2017](#), and others). For Levels 1 and 2, the ionisation ranges between 10^{-18} and 10^{-17} s^{-1} throughout the disk, whereas for Level 3, $\sim 10^{-16} \text{ s}^{-1}$ is reached at radii larger than 10 AU. In the inner disk inside $\sim 2 \text{ AU}$, all three ionisation levels are the same, because the surface densities here ($>600 \text{ g cm}^{-2}$) attenuate the impinging CRs, so that only the SLRs contribute. In the outer, more diffuse disk, the contribution from CRs impinging on the disk becomes more dominant, thus leading to three markedly different ionisation levels.

2.2. Chemical network

The chemical model used here includes gas-phase chemistry, gas–grain interactions and grain–surface chemistry. The gas-phase chemistry is from the latest release of the UMIST Database for Astrochemistry ([McElroy et al. 2013](#)) termed RATE12. The rates for gas–grain interactions and grain–surface chemistry are calculated as described in [Walsh et al. \(2015, and references therein\)](#). A gas to dust mass ratio of 100 is adopted.

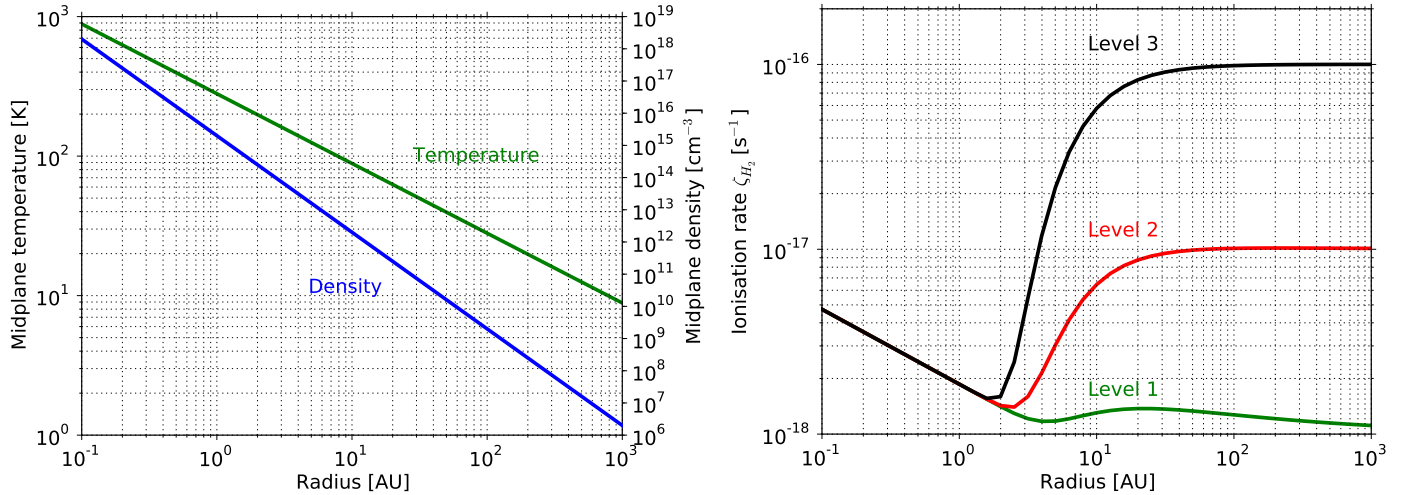


Fig. 1. Panel a: midplane temperature and density profiles for the pre-solar nebula. Panel b: ionisation levels utilised in the models. Level 1: SLRs only. Level 2: SLRs + fiducial CRs. Level 3: SLRs + enhanced CRs.

Table 1. Binding energies E_{bin} [K] for relevant species.

Species	UDfA (used here)	UDfA reference	Penteado et al. (2017)
O	800	Tielens & Allamandola (1987)	1660 ± 60
OH	2850	Garrod & Herbst (2006)	3210 ± 1550
O ₂ H	3650	Garrod & Herbst (2006)	800
H ₂ O ₂	5700	Garrod & Herbst (2006)	6000 ± 100
O ₂	1000	Garrod & Herbst (2006)	898 ± 30
O ₃	1800	Garrod & Herbst (2006)	2100 ± 100
H ₂ O	5700	Brown & Bolina (2007)	4800 ± 100
H	600	Cazaux & Tielens (2002)	650 ± 100
H ₂	430	Garrod & Herbst (2006)	500 ± 100

This setup is similar to the “full chemistry” setup from Paper I. The adopted grain size is 0.1 μm .

Four scenarios of initial chemical abundances are explored: the first two are cloud inheritance (hereafter the “inheritance” scenario) and chemical reset (hereafter the “reset” scenario). In the inheritance scenario the initial abundances are molecular (but no O₂ initially), and assumed inherited from the parent molecular cloud. In the reset scenario, the initial abundances are atomic, because all molecules are assumed dissociated prior to arrival in the disk midplane, due to exposure to a sufficiently strong accretion shock en route into the disk, or accretion heating very close to the star leading to high temperatures. The reset scenario is of particular interest for this paper because in Paper I, we see that for this scenario that the O₂ ice to H₂O ice ratio was similar to that found in comets 67P and 1P (Bieler et al. 2015; Rubin et al. 2015). Table A.1 lists the initial atomic and molecular abundances used in these two scenarios. Tables 1 and 2 list the relevant binding energies for species, and activation barriers for reactions applicable to the O₂ chemistry, respectively. Radiolysis of H₂O ice through the reaction $2\text{iH}_2\text{O} \xrightarrow{\gamma} 2\text{iH}_2 + \text{iO}_2$ and dismutation of H₂O₂ ice through the reaction $2\text{iH}_2\text{O}_2 \rightarrow 2\text{iH}_2\text{O} + \text{iO}_2$ are not explicitly included in the network. For the inheritance scenario, in this chemical network, atomic oxygen can be produced in situ in the ice mantles via the cosmic-ray induced photodissociation of H₂O ice and other abundant oxygen-bearing molecules therein (e.g. CO₂ ice).

The third and fourth scenarios for the initial chemical abundances are more simple: first, the “water”-scenario, which assumes initial abundances of gas-phase H, He, H₂ and H₂O. The intention with this scenario is to investigate if O₂ ice can be produced through processing of H₂O ice. Second, the “atomic oxygen”-scenario, which assumes gas-phase H, He, H₂, and O as initial abundances. Here it is investigated whether or not O₂ ice production from atoms depends on the presence of elements other than oxygen. Eistrup et al. (2018) showed that H₂S ice on the grain surfaces acts as a catalyst for the conversion of adsorbed oxygen atoms into O₂ ice. Whether or not the sulphur-bearing species are essential for the production of O₂ ice can be tested in these two scenarios, because they exclude sulphur, and only include a source of oxygen in the form of oxygen atoms or H₂O.

Lastly, we have explored whether the observed cometary O₂ ice abundance can be maintained if the PSN started out with a percentage (5%) of elemental oxygen locked up in primordial O₂, as suggested by Taquet et al. (2016).

3. Results

In this section the chemical evolution of various species in the PSN are presented in a number of figures. Due to the focus on O₂ ice in this paper, the attention will be on species that are chemically related to O₂ ice (a schematic overview of these species can be found in Cuppen et al. 2010).

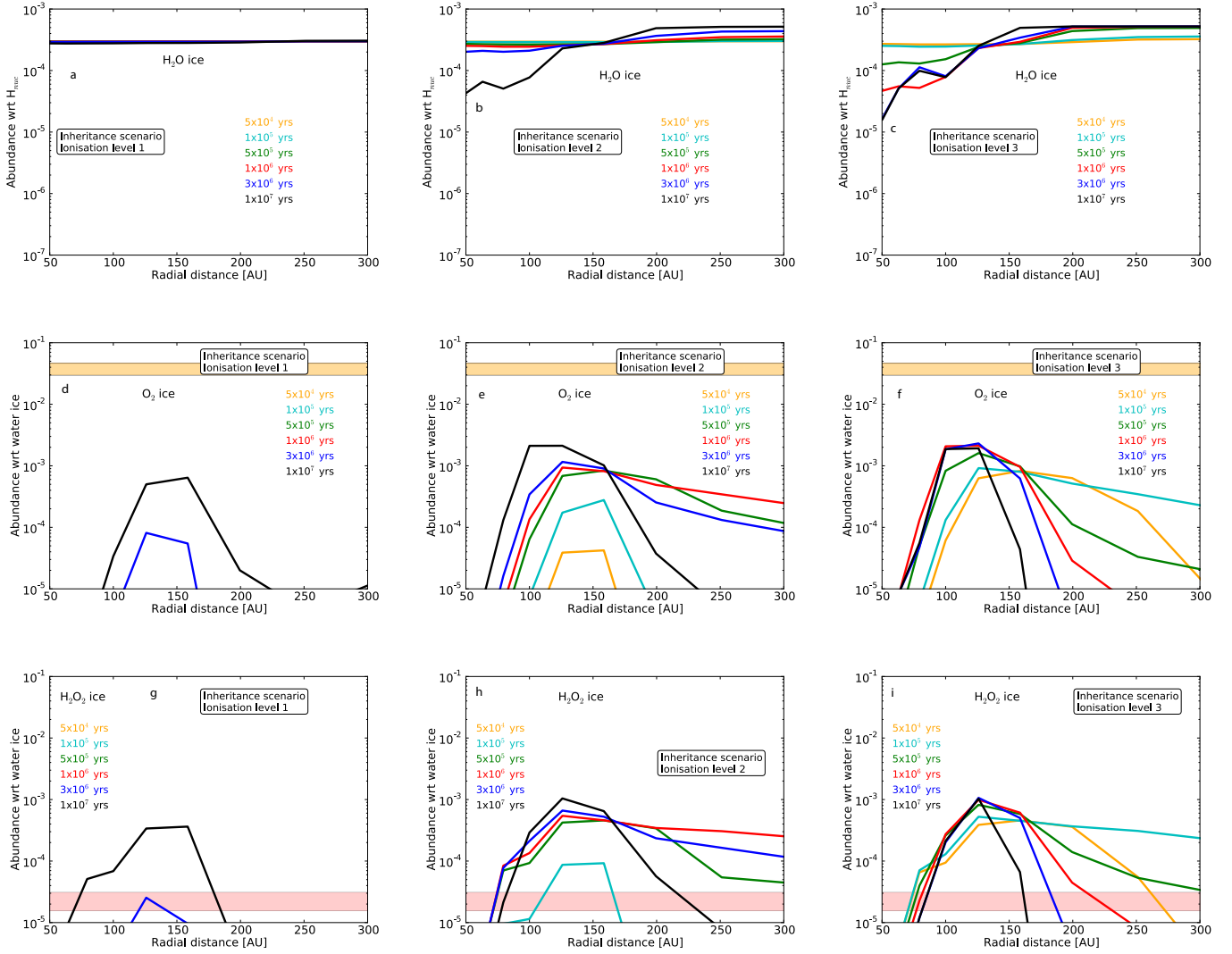


Fig. 2. Radial abundance profiles for the inheritance scenario at multiple evolutionary time steps. *Top to bottom panels:* H₂O, O₂, and H₂O₂. *Left to right panels:* increasing ionisation levels. For O₂ and H₂O₂, the limits detected in comet 67P are indicated as yellow and red shaded areas, respectively. The chemical network utilised does not include O₃ chemistry.

Table 2. Activation energies E_{act} [K] for relevant grain–surface reactions.

Reaction	E_{act} [K]
$\text{OH} + \text{H} \rightarrow \text{H}_2\text{O}$	0
$\text{OH} + \text{H}_2 \rightarrow \text{H}_2\text{O} + \text{H}$	2100
$\text{O}_3 + \text{H} \rightarrow \text{O}_2 + \text{OH}$	0
$\text{O} + \text{O} \rightarrow \text{O}_2$	0
$\text{O} + \text{O}_2 \rightarrow \text{O}_3$	500
$\text{OH} + \text{O} \rightarrow \text{O}_2\text{H}$	0
$\text{O}_2\text{H} + \text{H}_2 \rightarrow \text{H}_2\text{O}_2 + \text{H}$	5000
$\text{H} + \text{H}_2\text{O}_2 \rightarrow \text{H}_2\text{O} + \text{OH}$	2000

3.1. PSN abundance evolution

3.1.1. Inheritance scenario

Figure 2 features abundances as a function of midplane radius for different evolutionary times up to 10 Myr for the inheritance scenario. H₂O ice is plotted with respect to H_{nuc} abundance, whereas O₂ and H₂O₂ ice are plotted with respect to H₂O

ice. This is done to match the convention used in the reporting of cometary abundances. The model details are listed in each panel, along with colour-coding indicating evolution times. In the middle and lower panels, the respective observed limits for O₂ and H₂O₂ ices are marked with orange and red shaded regions, respectively. The orange shading for O₂ ice marks the limits to the mean O₂ ice abundance observed in comet 67P ($3.8 \pm 0.85 \times 10^{-2}$ with respect to H₂O ice), and the red shading marks the limits for H₂O₂ ice ($6 \pm 0.7 \times 10^{-4}$ wrt O₂ ice, thus $2.34 \pm 0.8 \times 10^{-5}$ wrt H₂O ice; Bieler et al. 2015). We note that in 67P variations in O₂ over H₂O ice were seen spanning 1–10%.

An evolution time of 10 Myr is chosen because this is assumed to be the maximum lifetime of a gaseous protoplanetary disk, and based on the results from Eistrup et al. (2018), the outer icy disk midplane should have reached steady state by this time. Since the focus in this work is the outer, icy PSN disk midplane, the radial range starts at 50 AU, which lies inside the O₂ iceline (~120 AU at ~29 K).

For ionisation Level 1 in Fig. 2a, H₂O ice stays abundant throughout the evolution. For higher ionisations, the H₂O ice abundance decreases with time inside, and increases with time outside ~120 AU. This radius marks the O₂

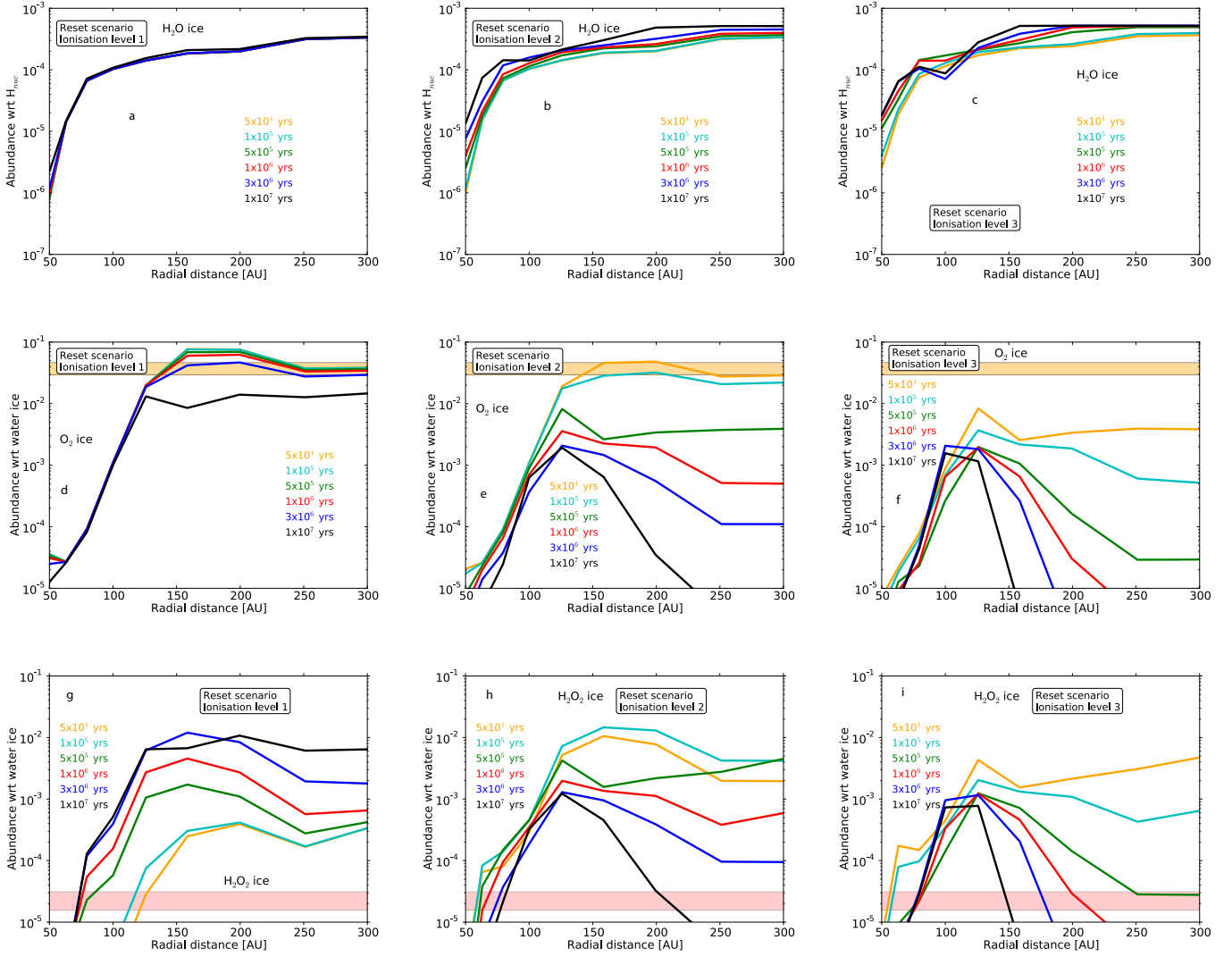


Fig. 3. Radial abundance profiles for the reset scenario at multiple evolutionary time steps. *Top to bottom panels* are H₂O, O₂ and H₂O₂ ices. *Left to right panels* are increasing ionisation levels. For O₂ and H₂O₂ the limits detected in comet 67P are indicated as yellow and red shaded areas, respectively. The chemical network utilised does not include O₃ chemistry.

Panels d–f show the evolution of O₂ ice for the different ionisation levels. The profiles for O₂ ice for ionisation Levels 1 and 2 peak at similar abundance levels of $2\text{--}3 \times 10^{-3}$ with respect to H₂O ice, which is an order of magnitude lower than the observed abundance. We see that for higher levels of ionisation, this abundance is reached faster, with ionisation Level 1 producing of order 10^{-3} with respect to H₂O ice by only 10 Myr.

We see that for H₂O₂ ice abundance profiles featured in panels g to i they evolve in a similar way to O₂ (we note the same y -axis range as for O₂ ice), and reach peak abundances at ~ 120 AU by 10 Myr, that are a factor of between two and three lower than those for O₂ ice. This abundance level is 30–100 times higher than that observed for H₂O₂ ice in comet 67P. For all ionisation levels, there are narrow radial regions where the H₂O₂ ice abundance matches the observed levels over a time range defined by the ionisation level (e.g. between 70 and 80 AU ionisation Level 3 at all times). However, at these points the O₂ ice abundance does not match that observed. For the inheritance scenario, it is thus seen that O₂ ice is underproduced by at least an order of magnitude, and H₂O₂ is overproduced by up to two orders of magnitude.

3.1.2. Reset scenario

The abundance evolution profiles for the reset scenario are shown in Fig. 3. For O₂ ice in panels d to f we see that a high abundance ($10^{-2}\text{--}10^{-1}$ with respect to H₂O ice) is reached for both ionisation Levels 1 and 2, with each reproducing the observed abundance for a large radial range covering 150–300 AU. For ionisation Level 1 the O₂ ice abundance matching the observation is maintained up to 2 Myr. Until 1 Myr between 150–220 AU, the modelled abundance lies above the observations ($>5 \times 10^{-2}$ with respect to H₂O ice). For Level 2 the abundance level matching the observed values happens between $5 \times 10^4\text{--}10^5$ yr in the range 150–200 AU, but subsequently drops to reach $2\text{--}3 \times 10^{-3}$ with respect to H₂O ice by 10 Myr. This was also the abundance level reached for O₂ ice by 10 Myr for the inheritance scenario. For ionisation Level 3, the abundance is below the observed level by 5×10^4 yr of evolution already, and by 10 Myr it also has reached the level of $2\text{--}3 \times 10^{-3}$ with respect to H₂O ice. For ionisation Level 1, the abundance remains higher than $2\text{--}3 \times 10^{-3}$ with respect to H₂O ice throughout the evolution, indicating that, for both inheritance and reset, at this ionisation level, 10 Myr of chemical evolution is not

enough to bring the O_2 ice abundance to what appears to be its steady-state level according to the results from the inheritance scenario in Fig. 2 (for a description of disk midplane chemical steady state, see Eistrup et al. 2018).

We see H_2O_2 ice in Fig. 3, panels g–i, to evolve significantly over time. For all ionisation levels, H_2O_2 ice is produced over time with the timescale dependent on ionisation level. The fastest production is seen for Level 3, where the peak abundance of $>10^{-2}$ with respect to H_2O ice is reached already by 10^5 yr. On the other hand, it takes 0.5 Myr and 3 Myr to reach a similar abundance for Levels 2 and 1, respectively. The H_2O_2 ice is subsequently destroyed after reaching its peak. For ionisation Levels 2 and 3, a steady-state abundance of order 10^{-3} with respect to H_2O ice is reached at radii ~ 100 – 120 AU. Beyond this radius it has disappeared by 10 Myr.

3.1.3. Water and atomic oxygen starting conditions

Models were also run starting with simpler sets of initial abundances, either with all oxygen already present in H_2O (water scenario) or all free (atomic oxygen scenario), in addition to H and He. This was to test the formation of O_2 ice solely from processing of H_2O ice, and the formation via assembly from free oxygen atoms, in the absence of other chemistry. It was found that the water scenario did not, at any point in time or radius, reproduce O_2 ice to the abundance seen in the comets. The maximum abundance was $<10^{-7}$ with respect to H_2O ice, regardless of ionisation level. This is shown in Fig. A.1, where the gas and ice abundances of O, O_2 , H_2O , and H_2O_2 are plotted as a function of radius by 10 Myr of evolution.

The atomic oxygen scenario, on the other hand, can reproduce the observed abundance of O_2 ice by 10 Myr, as is seen in panel a of Fig. 4. This panel of Fig. 4 shows the abundances of H_2O , O, O_3 and H_2O_2 with respect to H_{nuc} as a function of radius, by 10 Myr of evolution. The radial range (80–300 AU) for this reproduction is similar to the range in the reset scenario, but in the atomic oxygen scenario H_2O_2 ice is found to be more efficiently formed than O_2 ice. This scenario is revisited later in the paper.

3.2. Including ozone ice chemistry

The chemical network utilised so far has not included O_3 . This is because it has not been considered to be an important player in ISM chemistry, as it is formed primarily in the gas phase via a three-body mechanism. However, data exist on O_3 ice chemistry, as outlined in, for example, Cuppen et al. (2010) and Lamberts et al. (2013). The production pathway to O_3 ice is



with a fiducial barrier of 500 K, and the destruction pathway is



which has no barrier. Hence, it is expected that the pathway to O_3 ice via O_2 ice may impact on the O_2 ice abundance, and will be dependent on the both the availability of free oxygen atoms on the ice, and the efficiency of hydrogenation.

Figure 5 presents the evolving abundances of H_2O ice, O_2 ice, H_2O_2 ice, and O_3 ice as a function of radius for the inheritance scenario, thus now including O_3 ice chemistry. These panels are directly comparable with those in Fig. 2. The abundance evolution of O_3 ice is shown in the bottom row. The

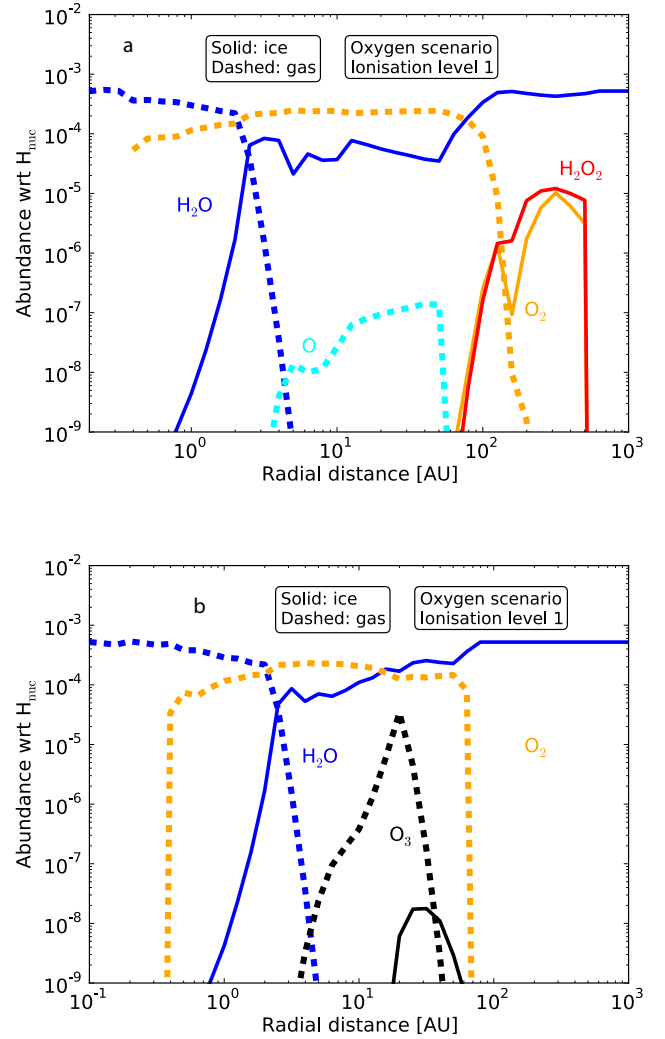


Fig. 4. Abundances by 10 Myr evolution in the oxygen-only scenario, for ionisation Level 1. *Panel a:* without O_3 chemistry included. *Panel b:* with O_3 chemistry included.

abundance evolution seen for O_2 ice in panels d–f and for H_2O_2 ice in panels g–i are similar to the trends in Fig. 2 for the chemistry without O_3 . However, the abundances reached by 10 Myr with O_3 chemistry are lower than those without: both O_2 ice and H_2O_2 ice abundances peak at $\sim 5 \times 10^{-4}$ with respect to H_2O ice, thus a factor of two lower for H_2O_2 ice and a factor of five lower for O_2 ice when compared with the scenario that does not include O_3 chemistry. The peak O_2 ice abundance is almost two orders of magnitude lower than the observed value, and the peak of the H_2O_2 ice is about 20 times higher than the observed value.

For O_3 ice in panels j–l, we see a steady build-up over time. We note that the range on the y -axis range is different for O_3 ice than for H_2O_2 and O_2 ices. The O_3 ice reaches maximum abundance fastest at ionisation Level 3 ($\sim 10^{-4}$ with respect to H_2O ice), thus a factor of five lower than that reached for H_2O_2 ice and O_2 ice, yet two orders of magnitude higher than the observed upper limit at 10^{-6} with respect to H_2O ice for O_3 ice in comet 67P. Only at larger radii (between 150 and 270 AU, depending on evolution time) do the O_3 and H_2O_2 ice abundances fall within the observed limits, although O_2 ice does not. All three ice species thus show similarities in their evolution, and order-of-magnitude similarities in their peak abundances. The observed large differences between their respective abundances

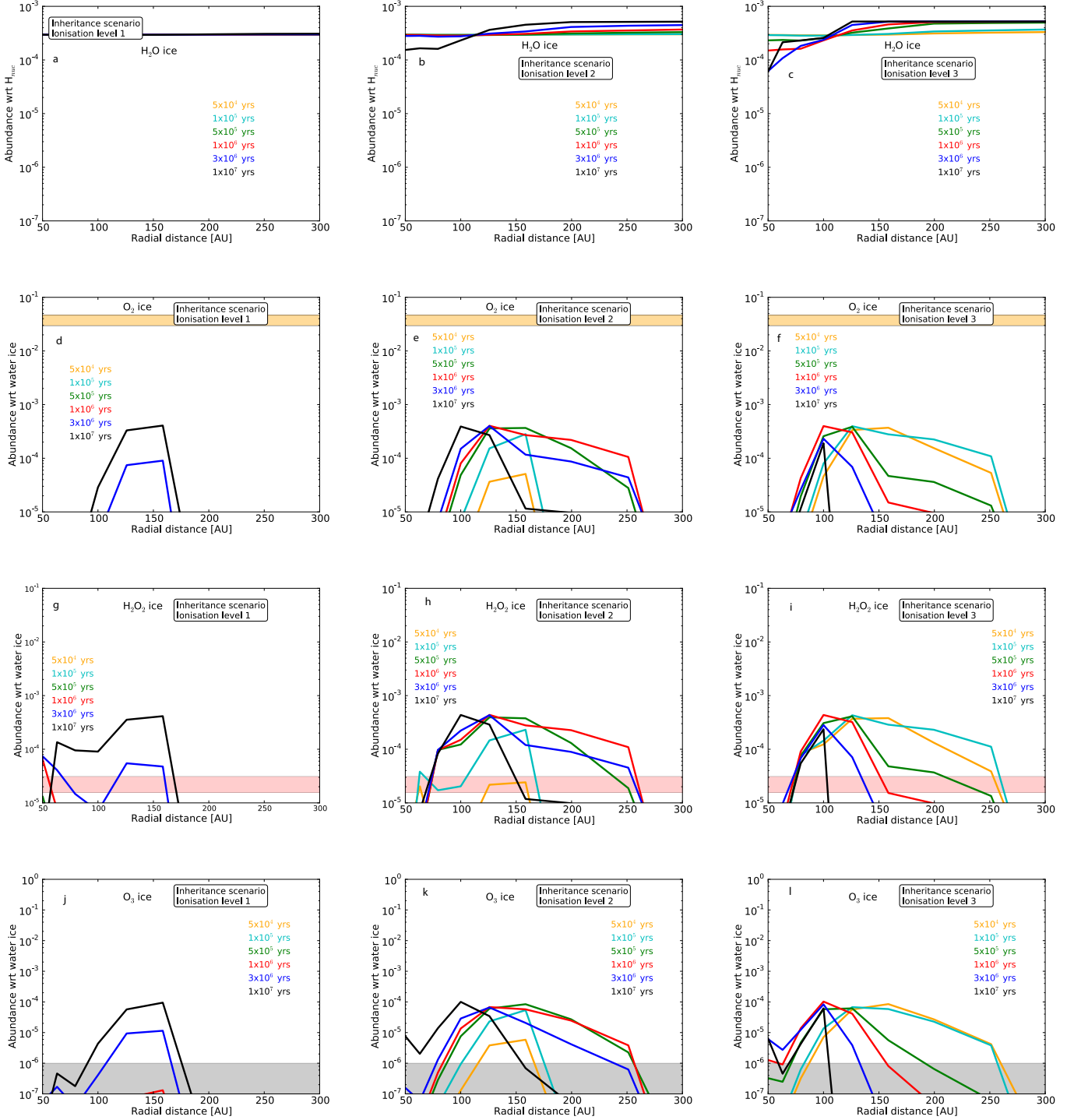


Fig. 5. Radial abundance profiles for the inheritance scenario at multiple evolutionary time steps. *Top to bottom panels:* H₂O, O₂, H₂O₂, and O₃ ices. *Left to right panels:* increasing ionisation levels. For O₂, H₂O₂, and O₃ the limits, and upper limit, detected in comet 67P, are indicated as yellow, red and grey shaded areas, respectively. The chemical network utilised includes O₃ chemistry.

(O₃:H₂O₂:O₂ \lesssim 1:23.5:39 200) are not reproduced by these models.

For the reset scenario, including O₃ chemistry, Fig. 6 shows a higher abundance level of O₂, H₂O₂ and O₃ ices than for the inheritance scenario. At ionisation Level 1, O₂ and H₂O₂ ices peak at $\sim 10^{-2}$ with respect to H₂O ice at ~ 150 AU between 0.5–1 Myr evolution, but subsequently drop by about an order of magnitude by 10 Myr. For ionisation Levels 2 and 3, abundance levels for both species of $\sim 10^{-2}$ with respect to

H₂O ice are only achieved by 5×10^4 yr, and the peak abundances by 10 Myr are at $1\text{--}2 \times 10^{-4}$ with respect to H₂O ice, thus neither matching the observed values of H₂O₂ ice nor of O₂ ice. At no point in time or radius does the reset scenario including O₃ chemistry reproduce the observed mean abundance of O₂ ice. However, for ionisation Levels 1 and 2 the abundance between 150–200 AU reaches above 1% of H₂O ice, which is within the observed range in comet 67P (1–10%).

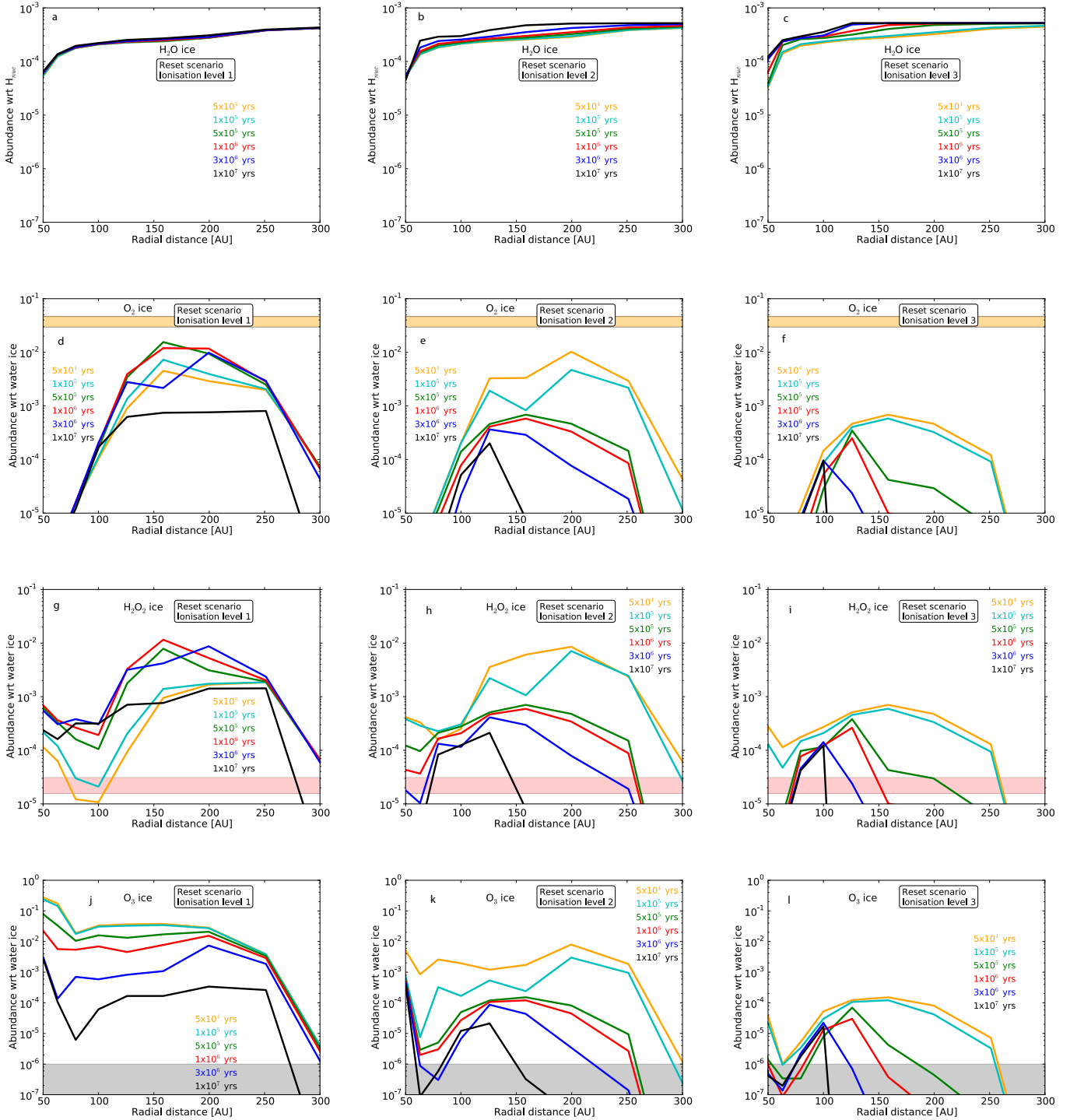


Fig. 6. Radial abundance profiles for the reset scenario at multiple evolutionary time steps. *Top to bottom panels:* H₂O, O₂ and H₂O₂ and O₃ ices. *Left to right panels:* increasing ionisation levels. For O₂, H₂O₂, and O₃ limits, and upper limit, detected in comet 67P, are indicated as yellow, red, and grey shaded areas, respectively. The chemical network utilised includes O₃ chemistry.

The O₃ abundance for the reset scenario at ionisation Level 1 starts out high at $1\text{--}3 \times 10^{-2}$ with respect to H₂O ice between 100 and 200 AU up to $5 \times 10^5 \text{ yr} \cdot \text{s}$ seen in Fig. 6. We note that the higher ratios reached within 70 AU is due to generally low absolute ice abundances (see panels a–c in Fig. 6). For ionisation Levels 2 and 3, these high abundances reduce over time leading to a peak of at a few times 10^{-5} with respect to H₂O ice by 10 Myr between 100–120 AU. The maximum abundance achieved by 10 Myr is an order of magnitude higher than the

observed upper limit. Only in the wings of the abundance profiles do the models reproduce this upper limit for the entire radial range, depending on evolution time and ionisation level.

The scenarios that begin with only H₂O or elemental oxygen were also investigated after the inclusion of O₃ chemistry. However, neither of them reproduced a significant amount of O₂ ice. The abundances by 10 Myr are shown for the oxygen-only scenario in Fig. 4b. Upon inclusion of O₃ chemistry, the peak abundance of O₂ ice from panel a (without O₃) drops by at

least four orders of magnitude to $<10^{-9}$ with respect to H₂O ice in panel b. Starting with atomic oxygen only therefore does not reproduce the observed abundance. For the water scenario, there is also no O₂ ice. An overview of gas and ice abundances of H₂O, O₂, O₃, and H₂O₂ as a function of radius by 10 Myr of evolution for the inheritance, reset, water and oxygen scenarios is shown in Fig. A.2. In none of the tested cases including O₃ chemistry is the O₂ ice abundance reproduced to match the levels observed in the comets.

3.3. Exploring the sensitivity of the abundances to assumed grain–surface parameters

The underproduction of O₂ ice and overproduction of H₂O₂ and O₃ ices may be a result of the adsorption and reaction parameters assumed for the network of reactions involving oxygen and hydrogen. If, for example, the reaction $i\text{O} \xrightarrow{i\text{O}} i\text{O}_3$ is too efficient on the grain surfaces, this could lead to over- and under-production of O₃ and O₂ ices, respectively. This reaction is dependent on the adsorption of atomic oxygen, for which the utilised binding energy here is $E_{\text{bin}} = 800$ K. This is based on estimates from Tielens & Allamandola (1987). However, recent experimental work by He et al. (2015) experimentally determined a binding energy of $E_{\text{bin}} = 1660$ K, thus more than doubling the former value. This higher binding energy should act to keep oxygen atoms from desorbing at higher temperatures, thereby facilitating *each* step in the reaction pathway leading from $i\text{O} \xrightarrow{i\text{O}} i\text{O}_2 \xrightarrow{i\text{O}} i\text{O}_3$. On the other hand, at low temperature the mobility of atomic oxygen will be reduced. This latter case could possibly lead to a decreased production of O₃ ice.

In this subsection, the sensitivity of the chemistry to several parameters is explored for single-point models (i.e. single n , T) which cover the temperature and density ranges in the PSN midplane where O₂ ice production is expected, based on the model results described thus far. The first parameter explored is the barrier width for quantum tunnelling b_{qt} which is set to either 1 or 2 Å representing the limits to the range of values usually assumed for b_{qt} when modelling grain–surface chemistry, see Cuppen et al. (2017), and references therein. The second parameter is the ratio of the diffusion energy to the molecular binding energy $E_{\text{diff}}/E_{\text{bin}}$, which is taken to be either 0.3 or 0.5 (see Ruffle & Herbst 2000; Garrod & Herbst 2006, and references therein). This amounts to four combinations of the two reaction parameters. The values of each parameter associated with a given model setup is given in each panel. The values for the grain–surface chemistry used in Eistrup et al. (2016, 2018), are $b_{\text{qt}} = 1$ Å and $E_{\text{diff}}/E_{\text{bin}} = 0.5$. Only the reset scenario is studied here because this is the case for which maximal O₂ ice formation is seen.

3.3.1. Abundance mosaics

Figures A.3, 7, A.4, and A.5 show the abundances of H₂O ice with respect to H_{nuc} , and O₂ ice, O₃ ice and H₂O₂ ice with respect to H₂O ice, again, maintaining the convention used by the comet community. From top to bottom span evolution times from 5×10^4 yr up to 10^7 yr. In each panel in the figures, a mosaic of colour-indicated abundances shows the results across different combinations of midplane temperature (ranging from 10 to 30 K on the y -axis), and densities (ranging from 10^8 to 10^{10} cm⁻³ on the x -axis). The ionisation level is kept constant at $\zeta = 10^{-17}$ s⁻¹, approximately the same as ionisation Level 2 from Fig. 1 in the

outer disk midplane. This ionisation level was seen in Fig. 3 (model without O₃) to match the observed O₂ ice abundance early in the evolution (up to 0.1 Myr).

From left to right in each of Figs. A.3 and A.4, two grain–surface reactions parameters are changed. The first one is the barrier width for quantum tunnelling b_{qt} which is either 1 Å (Columns one and two, in each figure) or 2 Å (Columns three and four, in each figure). The second parameter is the ratio of diffusion energy to molecular binding energy $E_{\text{diff}}/E_{\text{bin}}$, which can be either 0.3 (Columns one and three, in each figure) or 0.5 (Columns two and four, in each figure). This amount to four combinations of the two reaction parameters, thus the four columns of mosaics in each figure. The values of each parameter associated with a given model setup is given in each panel.

The range of evolution time steps is chosen to cover both the evolutionary stage at which O₂ ice was abundant in Fig. 3 (5×10^4 yr, for ionisation Level 2, $\sim 10^{-17}$ s⁻¹), as well as the later stages of disk evolution. The colours in the mosaic are chosen to enable distinguishing between abundance levels that change per order-of-magnitude, with dark red (top colour in colourbar) representing the highest abundance, and dark blue (bottom colour in colourbar) representing the lowest. The range of abundance levels in each figure is chosen to cover the full range of abundances produced for each species in the models. By showing the abundance evolutions in this way model results matching the observed abundances of, for example, O₂ ice (1–10% of H₂O ice) shows as orange (second highest colour in colourbar) in Fig. 7.

For H₂O ice in Fig. A.3 it is seen that for Columns one and two ($b_{\text{qt}} = 1$ Å) the abundance level largely remains within the initial order of magnitude of 10^{-4} with respect to H₂O ice. For Columns three and four ($b_{\text{qt}} = 2$ Å), some change is seen: especially for the third column ($E_{\text{diff}}/E_{\text{bin}} = 0.3$), the abundance is an order of magnitude lower for temperatures 25–30 K than for 10–20 K, and the abundance increases with time at 20 K. Turning to Fig. 7 for O₂ ice, limited production is seen for Columns one and two, with an early peak abundance between 10^{-4} – 10^{-3} with respect to H₂O ice at 25 K for $n = 10^8$ – 10^9 cm⁻³ by 5×10^4 yr. More interesting still are Columns three and four: until 0.5 Myr, abundances of 10^{-2} –1 with respect to H₂O ice are reached for temperatures 15–25 K for all densities. Thus, the O₂ ice abundance lies above or reproduces the observed values. For these cases, by >1 Myr, the produced O₂ ice is destroyed and reaches 10^{-5} – 10^{-3} with respect to H₂O ice by 10 Myr.

Having narrowed in on this parameter range (early evolutionary times, and $b_{\text{qt}} = 2$ Å), it can now be checked if H₂O₂ and O₃ ices can match the observed levels for the same model parameters. Figure A.4 shows the same suite of plots for O₃ ice. The results in Columns three and four up to 0.5 Myr evolution show that O₃ ice is abundantly produced, at a similar or higher level than O₂ ice, thus not matching the observed upper limit (darkest shade of blue, $<10^{-6}$ with respect to H₂O ice). The subsequent destruction by >1 Myr does not bring the abundance level much further down, and the O₃ ice abundance is generally similar to that for O₂ ice. O₃ ice only matches the observed upper limit for $T = 10$ K.

In Fig. A.5, H₂O₂ ice is abundantly produced for $b_{\text{qt}} = 2$ Å and $E_{\text{diff}}/E_{\text{bin}} = 0.3$. For 25 K it is consistently at a level of 10^{-1} –1 with respect to H₂O ice during the entire evolution. For 20 K it is also abundant, although 1–2 orders of magnitude lower than that found at 25 K, and it is less abundant than O₂ ice until ~ 0.5 Myr by between one and two orders of magnitude. At later times, for 20–25 K, H₂O₂ ice becomes 1–3 orders of magnitude

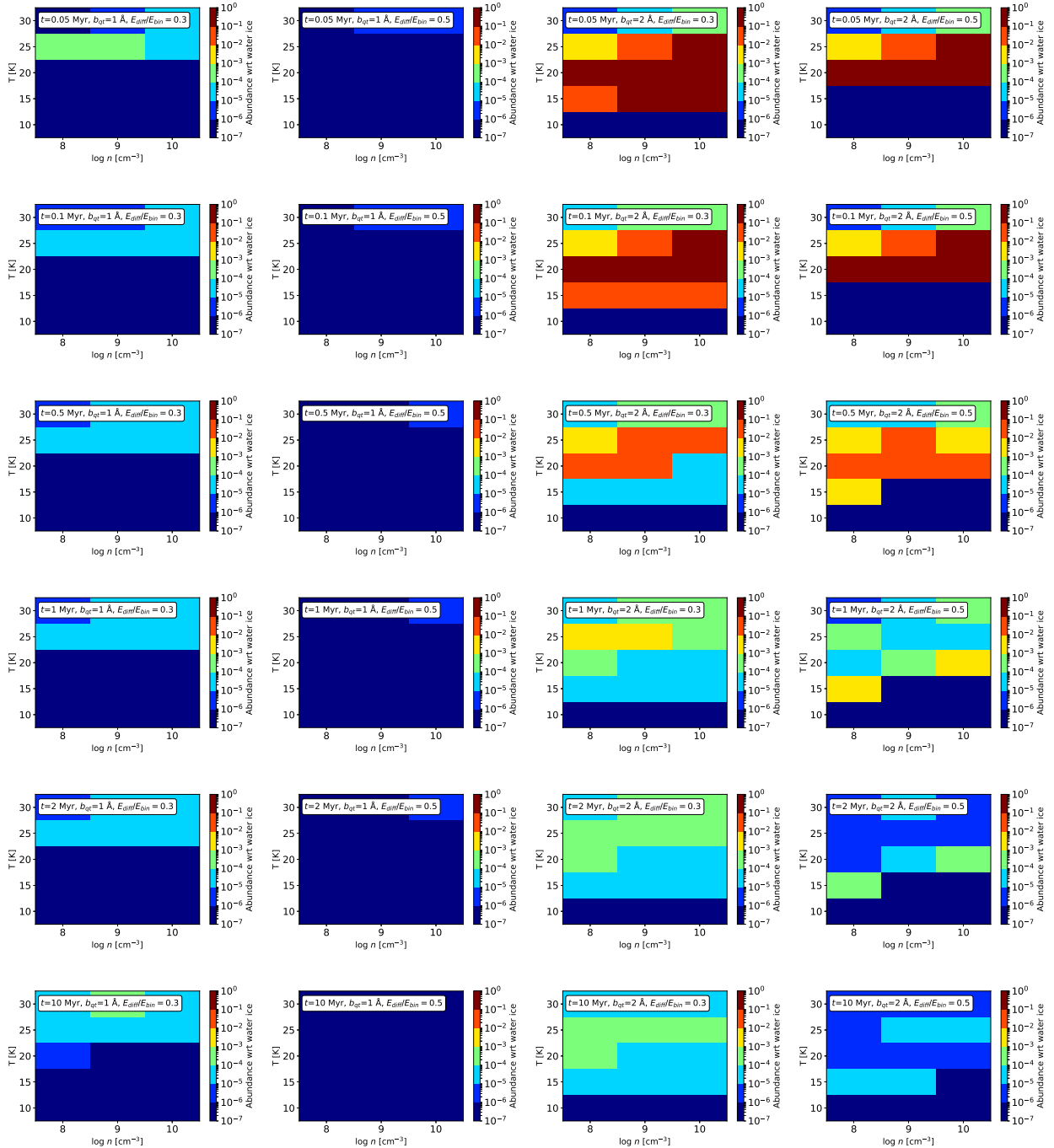


Fig. 7. Abundances (given as colours) for O_2 ice as function of midplane density (x -axes), and temperature (y -axes) at different evolutionary steps for the reset scenario. From left to right panels: abundances from model runs with different parameters for grain–surface reactions: Cols. 1 and 2: $b_{qt} = 1 \text{ \AA}$, Cols. 3 and 4: $b_{qt} = 2 \text{ \AA}$, Cols. 1 and 3: with $E_{\text{diff}}/E_{\text{bin}} = 0.3$, and Cols. 2 and 4: with $E_{\text{diff}}/E_{\text{bin}} = 0.5$. From top to bottom panels: different evolutionary times, from 0.05 Myr (top) to 10 Myr (bottom). To the right of each plot is a colourbar, indicating the abundance level with respect to H_2O ice for each colour. The chemical network utilised includes O_3 chemistry. Orange matches cometary O_2 abundances.

more abundant than O_2 ice. For $E_{\text{diff}}/E_{\text{bin}} = 0.5$ it is produced at levels 10^{-6} – 10^{-3} with respect to H_2O ice, depending on density.

3.3.2. Abundance evolutions for selected parameter sets

In Sect. 3.3.1 the colour mosaics revealed a promising set of physical and chemical parameters that supported the production of O_2 ice in situ under conditions suitable for the PSN: $b_{qt} = 2 \text{ \AA}$

(rather than the fiducial value of 1 \AA), $T = 15$ – 25 K , $n = 10^9$ – 10^{10} cm^{-3} and evolution times up to 0.5 Myr. The ratio of diffusion-to-binding energy for ice species appeared to play a minor role, with both values of 0.3 and 0.5 facilitating O_2 ice production.

In Fig. 8 evolving abundances are plotted for O_2 , O_3 , H_2O_2 , and O_2H ices with respect to H_2O ice, for the three different temperatures and two densities outlined above, and with $E_{\text{diff}}/E_{\text{bin}} = 0.3$. Details about each plot can be found in the labels. Each panel

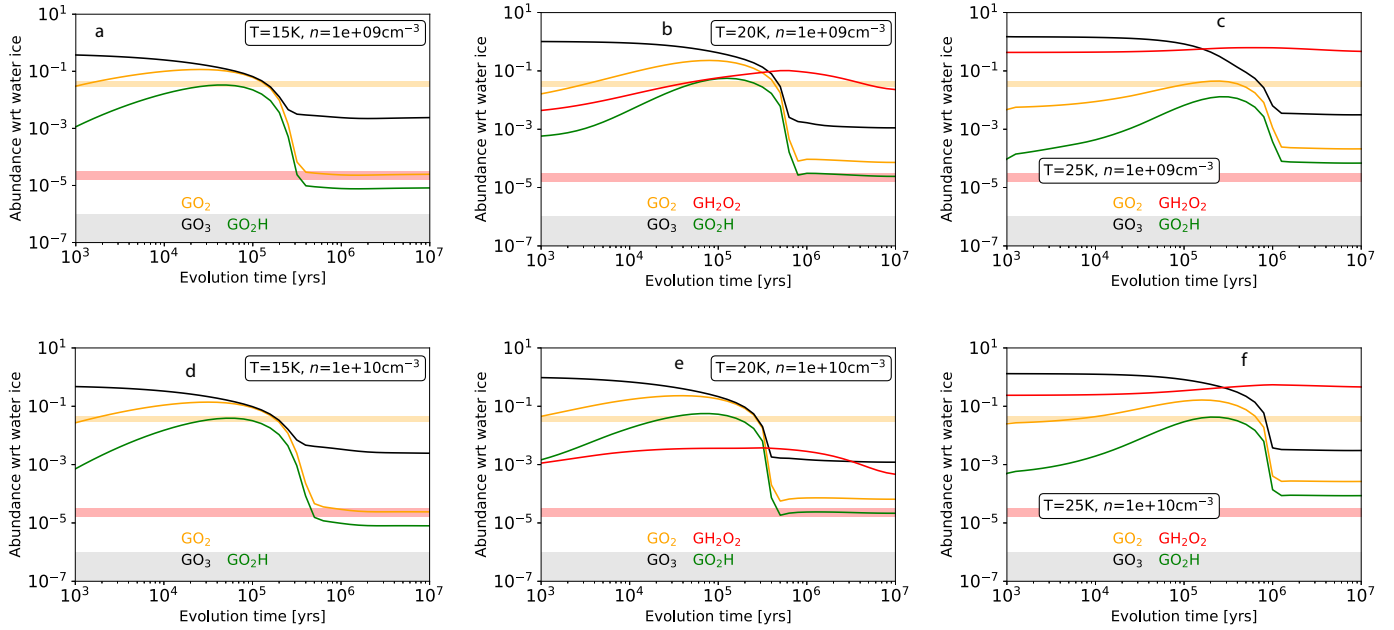


Fig. 8. Evolving abundances as function of time for the reset scenario for six different combinations of temperature (15, 20, or 25 K), density (10^9 or 10^{10} cm^{-3}) and molecular diffusion-to-binding energy ratio of 0.3. The barrier width for quantum tunnelling is $b_{qt} = 2$ Å. The orange shaded regions indicate the limits to the measured abundance of O₂ ice in the coma of comet 67P. The red shaded regions indicate the limits to the measured abundance of H₂O₂, and the top of the grey shaded region marks the upper limit to the measured abundance of O₃ ice in comet 67P. The chemical network utilised includes O₃ chemistry.

includes shaded regions indicating the observed abundances for O₂ ice (yellow), H₂O₂ ice (red), and upper limit for O₃ ice (grey).

For all six panels, it is seen that O₂ ice is within the measured abundance values, at least for a period during evolution. In all cases an initial production is seen, with the abundances of O₂ ice across model setups peaking at 30–40% with respect to H₂O ice between $\sim 5 \times 10^4$ – 5×10^5 yr, with the highest abundances reached at 20 K. For temperatures at 15–20 K, the O₂ ice abundances are generally higher than for $T = 25$ K. By $\sim 3 \times 10^5$ – 10^6 yr, depending on model setup, the O₂ ice abundance drops at least two orders of magnitude below maximum. Hence, the observed abundance of O₂ ice can be reproduced in all six setups, but only at early times.

Regarding ice species besides O₂, Fig. 8 shows that O₃ is the most abundant of the plotted species, at least until 10^5 yr. At early evolutionary stages, O₃ ice is even more abundant than H₂O ice. This is because the calculations are started with free oxygen atoms (reset scenario). This high level is not seen in the measurements of comet 67P, in which an upper limit of 10^{-6} with respect to H₂O ice was reported.

Likewise for H₂O₂ ice, it is mostly produced at a high abundance ($>10^{-3}$ with respect to H₂O ice) in these models. At 25 K after ~ 0.5 Myr of evolution, H₂O₂ ice is the most abundant of the plotted species. At no point in time in any of the plots does the abundance of H₂O₂ ice match the levels observed (red shaded region): for 15 K it is at least an order of magnitude too low, and between 20 and 25 K it is overproduced by between a factor of three and two orders of magnitude.

For a ratio of diffusion-to-binding energy of 0.5, similar plots are shown in Fig. 9. Between 20 and 25 K there are evolutionary times when O₂ ice is reproduced to match the observations. However the highest abundance reached is at 20 K. For $T = 15$ K, no reproduction of the observed abundances of neither O₂ ice nor H₂O₂ ice is seen, and O₃ ice is (next to H₂O ice) the dominant carrier of elemental oxygen. At $T = 25$ K for $n = 10^9$ cm^{-3} ,

the abundance of H₂O₂ ice matches that measured in the comet both by ~ 0.03 Myr and by ~ 3 Myr, and by 0.03 Myr the O₂ ice abundance is only ~ 2 times lower than the observed abundance. Along the same lines, for $T = 25$ K and $n = 10^9$ cm^{-3} by ~ 0.4 Myr O₂ ice matches the observed abundance, and H₂O₂ ice is only approximately twice as high as the observed level. This indicates that something close to a match with the observed abundances between both O₂ and H₂O₂ ice abundances with respect to H₂O ice is reached at 25 K. However, all the models still vastly overproduce the O₃ ice abundance compared with the measurements.

This parameter space investigation has shown that there are sweet spots, both for the physical and chemical setup, where O₂ ice can be produced to match the measurements on comet 67P, even after expanding the grain–surface chemical network to include O₃ ice chemistry. However, the low measured abundance of O₃ ice remains unexplained by the models.

A possible way to adjust the models to lower the O₃ ice production could be to increase the reaction barrier for the $i\text{O}_2 \xrightarrow{i\text{O}} i\text{O}_3$ reaction, where the default barrier is at $E_{\text{act}} = 500$ K. Because the abundance levels of O₂ ice and H₂O₂ ice in panel c of Fig. 9 ($T = 25$ K, $E_{\text{diff}}/E_{\text{bin}} = 0.5$, $n = 10^{-9}$ cm^{-3} , $b_{qt} = 2$ Å and $\zeta = 10^{-17}$ s^{-1}) featured evolutionary stages when they were simultaneously in proximity to the observed levels, this model setup is now tested with three different values for the activation energy for the $i\text{O}_2 \xrightarrow{i\text{O}} i\text{O}_3$ reaction. In addition to the fiducial 500 K, also 1000 K and 2000 K activation energies are tested. We note that these are likely too high relative to what is known from laboratory work where O₃ is seen to form readily at low temperatures (Lamberts et al. 2013). However, given that O₃ ice is not seen to be efficiently produced in space including comets, the hypothetical situation with a higher barrier is explored, as a means to mitigate as of yet unknown chemical pathways away from O₃ ice. To compare the effects of these changes, the same

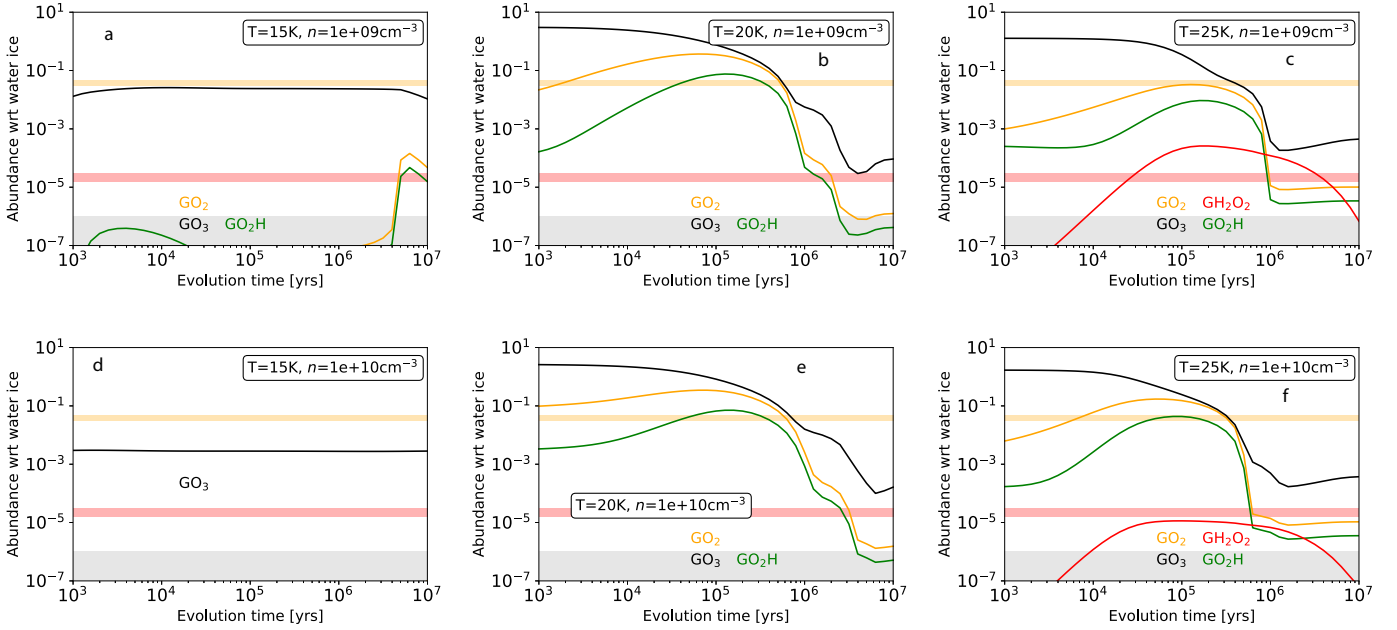


Fig. 9. Evolving abundances as function of time for the reset scenario for six different combinations of temperature (15, 20, or 25 K), density (10^9 or 10^{10} cm^{-3}) and molecular diffusion-to-binding energy ratio of 0.5. The barrier width for quantum tunnelling is $b_{qt} = 2$ Å. The orange shaded regions indicate the limits to the measured abundance of O_2 ice in the coma of comet 67P. The red shaded regions indicate the limits to the measured abundance of H_2O_2 , and the top of the grey shaded region marks the upper limit to the measured abundance of O_3 ice in comet 67P. The chemical network utilised includes O_3 chemistry.

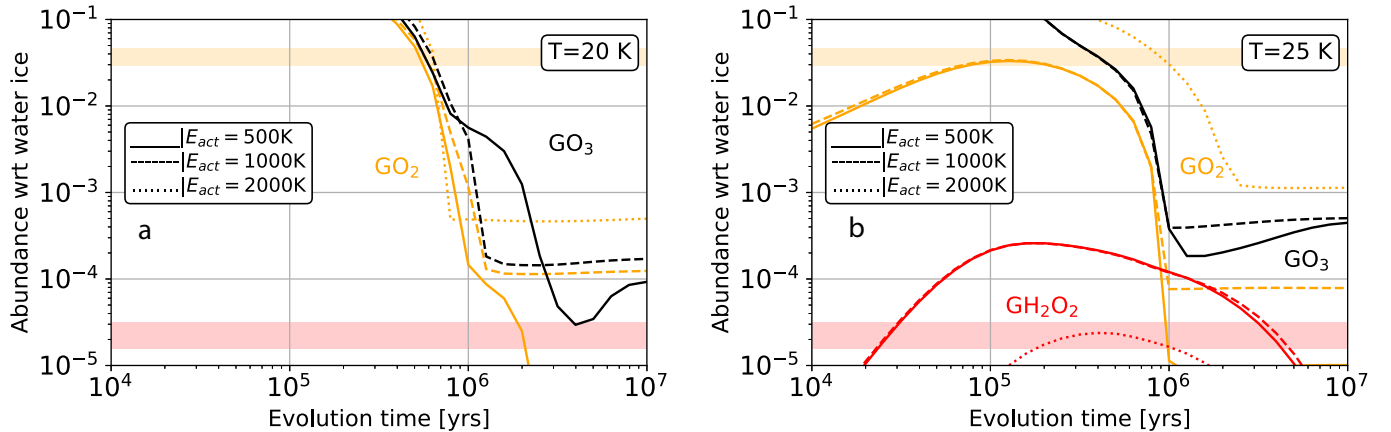


Fig. 10. Evolving abundances for O_2 ice, H_2O_2 ice and O_3 ice for three different activation energies for the reaction $i\text{O} + i\text{O}_2$. Panel a is for $T = 20$ K. Panel b is for $T = 25$ K. The barrier width for quantum tunnelling is $b_{qt} = 2$ Å, and the molecular diffusion-to-binding energy is 0.5.

tests of activation energies were performed for the setup in panel b in the same figure ($T = 20$ K, keeping all other parameters the same).

In panels a and b of Fig. 10, abundances for O_2 , O_3 , and H_2O_2 ices are plotted as a function of time, with solid, dashed and dotted profiles representing activation energies for the O_3 ice production reaction of 500 K, 1000 K and 2000 K, respectively. In panel a only the abundance of O_2 ice is found to match the observed level. However, in panel b, at 25 K, it is seen that for at $E_{\text{act}} = 2000$ K, O_2 ice and H_2O_2 ice are both reproduced to within the observed values between 0.8–1 Myr evolution. For the same evolutionary timescale, O_3 ice is much lower in abundance ($\sim 10^{-16}$ with respect to H_2O ice), thus also agreeing with the upper limit for the cometary abundance. This is therefore a sweet spot in the physical and chemical parameter space, in which the observed abundances of all three species are reproduced. We

note that the ionisation level for this sweet spot includes the contributions from both SLRs and CRs, whereas it was found in Paper I that O_2 ice be reproduced only without CRs. However, the framework here is different from that in Paper I, as O_3 chemistry and the updated binding energy for atomic oxygen have been included. We note that the anomalously high activation required for $\text{O} + \text{O}_2$ is likely masking as of yet unknown routes in the O_3 chemistry.

3.4. Including a primordial source of O_2 ice

Here, the results from the scenario starting with 5% of atomic oxygen locked in O_2 is tested, exploring the theory for the primordial origin of O_2 ice, as suggested by Taquet et al. (2016) and Mousis et al. (2016). This is a modification of the inheritance scenario so that remaining oxygen is already locked in molecules

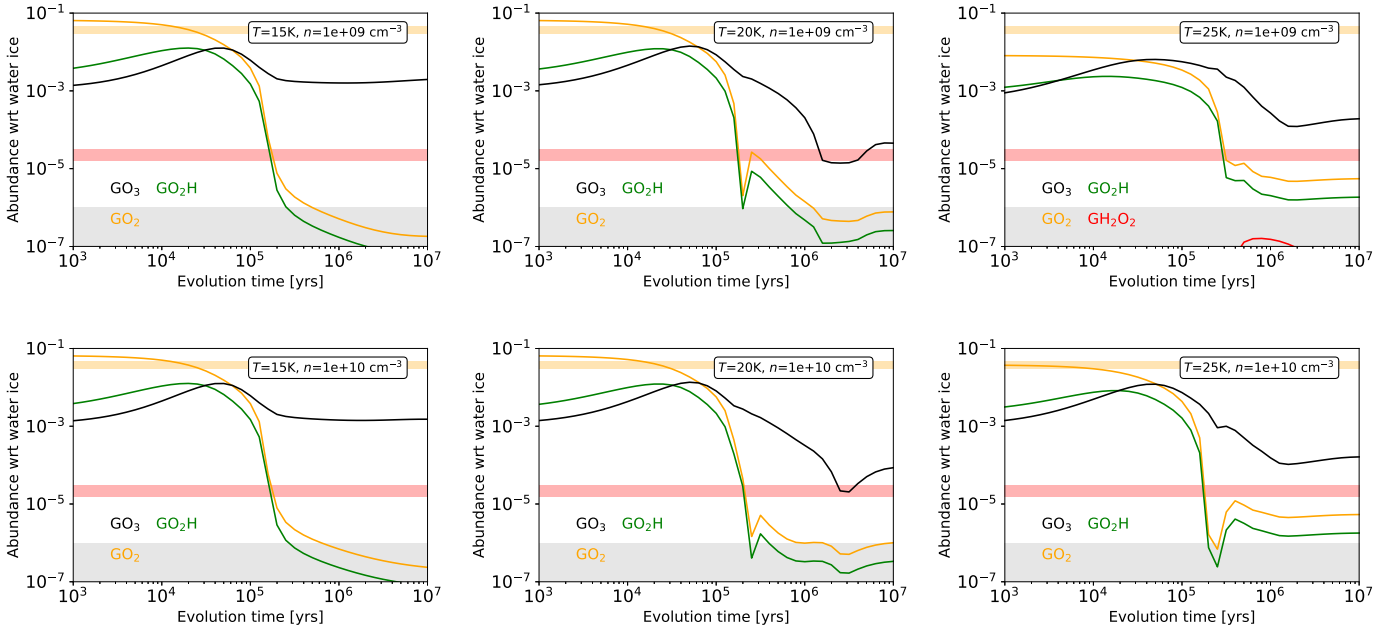


Fig. 11. Evolving abundances as function of time for the inheritance scenario, plus 5% extra elemental oxygen as O₂, for six different combinations of temperature (15, 20, or 25 K), density (10⁹ or 10¹⁰ cm⁻³), and molecular diffusion-to-binding energy ratio of 0.5. The barrier width for quantum tunnelling is $b_{qt} = 2 \text{ \AA}$. The orange shaded regions indicate the limits to the measured abundance of O₂ ice in the coma of comet 67P. The red shaded regions indicate the limits to the measured abundance of H₂O₂, and the top of the grey shaded region marks the upper limit to the measured abundance of O₃ ice in comet 67P.

(mostly H₂O, CO, and CO₂, see the inheritance scenario abundances in Table A.1). In Fig. 11 evolving abundance profiles for this scenario are shown, for temperatures between 15 and 25 K from left to right, and densities between 10⁹ and 10¹⁰ cm⁻³ from top to bottom. The ionisation rate is at $\zeta = 10^{-17} \text{ s}^{-1}$, the ratio of diffusion-to-binding energy is 0.5, and the barrier width for quantum tunnelling is 2 Å. The activation energy for the O + O₂ reaction is the fiducial value of 500 K.

We see in panels a, b, d, and e, of Fig. 11, that an early abundance level of O₂ ice at or above the observed mean is maintained until a few times 10⁴ yr for temperatures between 15 and 20 K. At 25 K, the abundance of O₂ ice is lower early on, and in particular for $n = 10^9 \text{ cm}^{-3}$ the abundance is $< 10^{-2}$ with respect to H₂O ice, which is outside the observed range. Simultaneously with O₂ ice matching the observed abundance in some cases, the abundance of O₃ ice is one to two orders of magnitude lower than O₂ ice by 3×10^3 yr for all cases except in panel c. Between a few times 10⁴ yr and 10 Myr, the O₂ ice abundance decreases by more than two orders of magnitude compared with the initial abundance, and from $\sim 10^5$ yr, O₃ ice is the dominant oxygen carrier of the plotted species. Hence, this scenario can reproduce the observed abundance of O₂ at early stages, and with lower abundances of H₂O₂ and O₃ ices, although abundances for the latter two species are still higher than measured.

4. Discussion

The abundance of O₂ ice in comet 67P has motivated much work as to its origin. Rubin et al. (2015), Mousis et al. (2016, 2018), Taquet et al. (2016), and Dulieu et al. (2017) have all attempted to explain the O₂ level from a point of view of possible chemical origins. Mousis et al. (2016) and Rubin et al. (2015) approached the problem by investigating chemical processing of either H₂O ice or H₂O₂ ice into O₂ ice. Mousis

et al. (2018) attempted an explanation with a scenario in which ice-covered grains were transported from the midplane to the upper layers of the disk. Here, the grains undergo photochemical processing producing O₂ ice, which is then cycled back to the disk midplane. It is noted that Mousis et al. (2018) only consider the O₂ ice abundances, not H₂O₂ or O₃ ices. On the other hand, Taquet et al. (2016) approached it from a broader point of view considering primordial production pathways in the parent cloud, and utilising extensive chemical networks and subsequently tracking the primordially produced O₂ ice to the disk midplane. While those works all conclude that a primordial origin and subsequent retention of O₂ ice was the most likely, Paper I concluded that starting from a fully atomised disk midplane, there is an evolutionary phase during which a range of radii in the outer icy midplane (outside the O₂ ice-line) will reasonably reproduce the measured O₂/H₂O ice ratio level.

The investigation here has taken the step further from Paper I: exploring the physical and chemical parameter space that may facilitate the production of O₂ ice in situ in the PSN, and simultaneously tracing the abundances of the chemically related species H₂O₂, O₃, and O₂H, as well as using a more massive disk model more appropriate for the PSN.

4.1. Chemical starting conditions

The chemical starting conditions' effects on the O₂ ice production were explored. It is clear that only an atomised start can facilitate the production, although models starting with a percentage of elemental oxygen in O₂ did retain this O₂ ice for a short period of evolution. The presence of atoms other than oxygen (such as sulphur), however, has no large impact on the production of O₂ ice. That means that oxygen-only and hydrogenated oxygen species lock up the majority of the available elemental oxygen.

4.2. Dependence on ionisation levels

The ionisation levels have been shown to facilitate different chemical evolution of abundances. The chemical timescales are shorter for higher ionisation levels. Ionisation Level 1 only reproduced the observed O_2 abundance for models without O_3 , and ionisation Levels 2 and 3 cause similar evolutionary trends, meaning that Level 3 does not facilitate any more O_2 ice production than Level 2. We note that the sweet spot for reproduction of O_2 , H_2O_2 and O_3 ices to within the observed abundances was found for a ionisation level of $\zeta = 10^{-17} \text{ s}^{-1}$, which is the local ISM value for dense clouds. This ionisation resembles ionisation Level 2 at 100–130 AU in the PSN disk midplane (see Fig. 1, panel b), which is also the radial range covering the temperature and density of the sweet spot, which is just outside the O_2 iceline.

4.3. Changing E_{bin} for atomic oxygen, and inclusion of O_3

We introduced O_3 and included it in the chemical network with production through $iO_2 \xrightarrow{iO} iO_3$ (activation energy of 500 K), and destruction through barrierless hydrogenation. After this change, the binding energy E_{bin} for oxygen atoms was increased from 800 to 1660 K, as measured by He et al. (2015). This change was expected to make oxygen atoms less volatile, thus having them reside on the grain surfaces at higher temperatures than before. This was expected to aid the production of O_2 and O_3 ices through the pathway $iO \xrightarrow{iO} iO_2 \xrightarrow{iO} iO_3$.

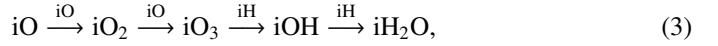
The expansion to include O_3 chemistry induces very different behaviours of O_2 , O_3 , and the chemically related H_2O_2 . Still assuming the PSN midplane, and ionisation Level 1, O_2 ice is not reproduced to match the observed cometary abundances.

The updated binding energy for atomic oxygen also had an effect, as the parameter space investigation revealed O_2 ice production matching the observed value, even when including O_3 chemistry. This change made the oxygen atoms less volatile, thus adsorbing to the grain at higher temperatures.

4.4. Narrowing down on O_2 ice production

With the lower produced level of O_2 ice after inclusion of O_3 chemistry the question remained if O_2 ice, in any case, can be produced to within the measured levels. Besides updating E_{bin} for atomic oxygen, two parameters for grain–surface chemistry, which had been kept constant thus far were now varied: the barrier width for quantum tunnelling, and the ratio of diffusion-to-binding energy of icy molecules. Adjusting these parameters, in particular the barrier width for quantum tunnelling, appears to have important impacts on the production level of O_2 ice. Increasing the barrier for quantum tunnelling from 1 to 2 Å, for temperatures ranging between 15–25 K, and densities 10^9 – 10^{10} cm^{-3} , results in O_2 ice produced to within the mean measured level in comet 67P. Increasing the barrier width from 1 to 2 Å lowers the mobility of H and H_2 on the grain surfaces. This means that O+O reactions proceed more efficiently than hydrogenation of O, O_2 , and O_3 ice, which in turn leads to higher abundances of O_2 and O_3 ices for a b_{qt} of 2 Å. Generally, with this wider tunnelling barrier, the measured mean abundance of O_2 ice can be reproduced relatively early in the evolution, from 0.05–0.5 Myr. At 25 K the corresponding level of H_2O_2 ice was close to its observed value. O_3 ice, however, remains more abundant than O_2 ice. Early in the evolution, O_3 ice is even more abundant than H_2O ice, but this is explained by O_3 ice being a

precursor to H_2O ice in the reaction pathway:



due to the availability of free oxygen. Indeed, in Fig. 8, showing the time evolution of species, O_3 is the dominant oxygen carrier at early times, and H_2O at later times. Exceptions to this are the cases of a low ratio of diffusion-to-binding energy of 0.3, and temperatures of 25 K (panels c and f in Fig. 8), where H_2O_2 dominates at later times. In those cases, a lower diffusion energy and higher temperature means a higher mobility of icy molecules on the grain surface. This leads to OH (the product of $O_3 + H$) reacting with O_2 ice (precursor to O_3 ice and product of the hydrogenation of O_3 ice), thereby lowering the abundance of O_3 ice, and increasing the abundance of first O_2H ice, and subsequently H_2O_2 ice via the reaction pathway $iO_2 \xrightarrow{iH} iO_2H \xrightarrow{iH} iH_2O_2$.

4.5. Activation energy for O_3 ice production pathway

A crucial element in modelling chemical evolution is the assumed activation energies for reactions. Especially for grain–surface chemistry, these energies remain somewhat uncertain, due to the difficulty of estimating them through production rates in the lab. In this work, the fiducial activation energy for the reaction $iO + iO_2 \rightarrow iO_3$ was $E_{\text{act}} = 500 \text{ K}$ (Lamberts et al. 2013), resulting in model O_3 ice at a higher abundance than its parent species, O_2 ice. This means that the modelled abundance of O_3 ice is several orders of magnitude higher than the upper limit determined in comet 67P (see Figs. 8 and 9).

Testing out higher activation energies for the production of O_3 ice (thereby impeding the $O + O_2$ reaction) revealed that for $E_{\text{act}} = 2000 \text{ K}$, the production of O_3 ice was sufficiently impeded to bring the abundance below the limit observed in the comet. This test simultaneously showed a match between both the abundances of O_2 ice and H_2O_2 ice and the cometary abundances at 25 K between 0.8 and 1 Myr evolution for an ionisation level of $\zeta = 10^{-17} \text{ s}^{-1}$, starting from the reset scenario. Thus, a sweet spot in the parameter space, where all three species match observed abundances, was found. However, it should be emphasised that such a high barrier is not supported in previous analyses of laboratory experiments. An upper limit for the activation energy of $O + O_2$ of $E_{\text{act}} = 150 \text{ K}$ on amorphous silicate surfaces was derived from experimental results by Minissale et al. (2014). Lamberts et al. (2013) required an activation energy of $E_{\text{act}} = 500 \text{ K}$ in order to reproduce experimental results from Ioppolo et al. (2010) of O_3 production on thick ices, and in agreement with this result, Taquet et al. (2016) were able to reproduce the low abundances O_3 and O_2H on ice mantles in dark clouds using an activation energy of $E_{\text{act}} = 300 \text{ K}$.

In this work O_3 remains overproduced for these levels of activation energies, even when increasing it to $E_{\text{act}} = 1000 \text{ K}$. Only using an artificially high energy of $E_{\text{act}} = 2000 \text{ K}$ for the $O + O_2$ reaction lowered the abundance of O_3 ice to the observed level. This could suggest that as-yet unknown reactions may be missing from the network: reactions involving either O_2 or O that do not lead to O_3 (so O and O_2 get locked up in other molecules than O_3) and/or new routes for destruction of O_3 ice.

4.6. Location of O_2 ice production sweet spot in PSN disk midplane

Taking the sweet spot for matching abundances of O_2 ice, H_2O_2 ice, and O_3 ice, it is interesting to look at where in the PSN disk

midplane the physical conditions for this match may be satisfied. A close look at Fig. 1a shows that indeed in the radial range ~100–130 AU are found a temperature of ~25 K and a density of 10^9 – 10^{10} cm⁻³. Thus, for the sweet-spot O₂ ice production scenario, this would be the predicted radial range for formation of both comets 1P and 67P, and a formation timescale by 0.8–1 Myr, given the model assumptions. This radial range is two to three times larger than the orbit of the present day Kuiper belt (30–50 AU).

4.7. Primordial origin of O₂ ice

One scenario does, however, predict early presence of O₂ ice to within the observed level, and H₂O₂ ice and O₃ ice orders of magnitude lower than that of O₂ ice. This is the scenario starting with an initial abundance of O₂ of ~5% with respect to the total elemental oxygen abundance, which was inspired by the results of Taquet et al. (2016) assuming that the O₂ is inherited from the parent cloud. Given early formation of the comets (by thousands to tens of thousands of years after formation of the PSN midplane), this scenario remains likely for the formation of the ices, and the fiducial values of $b_{\text{qt}} = 1 \text{ \AA}$ and $E_{\text{act}} = 500 \text{ K}$ for the O + O₂ reaction in the ice. This scenario agrees with the findings of Taquet et al. (2016).

5. Conclusion

Since the somewhat unexpected detection of abundant O₂ ice in the coma of comet 67P, several studies have attempted to explain the origin of the O₂. Building on the results from Paper I, this work has investigated the possibility of in situ formation of O₂ ice on grains in the midplane of the PSN disk midplane.

While a high abundance of O₂ ice, matching that observed in comet 67P, is reproduced outside the O₂ ice in the PSN disk midplane at intermediate evolutionary stages when assuming the initial chemistry to be reset, the same production is not seen after including O₃ ice chemistry into the chemical network. For the fiducial choice of parameters for grain–surface chemistry, and an activation energy of 500 K for the O + O₂ → O₃ reaction on the grains, O₃ ice is, in most cases, found to be the dominant oxygen-carrier next to H₂O ice, and O₂ ice was orders of magnitude too low in abundance compared to the abundance observed in comet 67P.

In order to test the sensitivity of the production of O₂, O₃, and H₂O₂ ices to the assumed parameters for grain–surface chemistry, in particular the barrier width for quantum tunnelling b_{qt} and the ratio of diffusion-to-binding energy of ice molecules $E_{\text{diff}}/E_{\text{bin}}$, a parameter space investigation was conducted. Several temperatures and densities were also tested, and the reset scenario assumed for initial abundances. This led to a sweet-spot set of parameters being revealed: $b_{\text{qt}} = 2 \text{ \AA}$, $E_{\text{diff}}/E_{\text{bin}} = 0.3$ – 0.5 , $T = 15$ – 25 K and $n = 10^9$ – 10^{10} cm⁻³ which facilitated O₂ reproduction matching the observed level. However, the abundances of O₃ and H₂O₂ ices were still in disagreement with the observed values by orders of magnitude.

As a last adjustment of the chemistry intended to lower the O₃ ice abundance, the activation energy for production of O₃ ice from the association of O and O₂ ices was increased in order to mitigate possible unknown chemical pathways away from O₃. For an activation energy of $E_{\text{act}} = 2000 \text{ K}$, $E_{\text{diff}}/E_{\text{bin}} = 0.5$, and the remaining physical and chemical conditions as given above, the abundances of O₂ ice, H₂O₂ ice, and the upper limit for O₃ ice in the comet were all reproduced. This matches a formation

location in the PSN disk midplane between 120 and 150 AU, just outside the O₂ iceline. However, this high activation energy for O₃ ice production is not supported by laboratory estimates, and thus more laboratory work is needed to determine potential missing chemical pathways for O₃ ice chemistry.

A model that starts out with a percentage of elemental oxygen locked in O₂ and thus assuming a primordial origin of O₂, also reproduced the observed abundance of O₂ ice at early stages of evolution, without increasing E_{act} for the O + O₂ reaction. Here, the O₃ and H₂O₂ ice abundances were below the O₂ ice abundance, but not matching the observed abundance levels. However, since the observed abundances of all three ices species are only reproduced in this work in the case for a set of rather extreme choices for the chemical parameters, the most plausible explanation for the origin of the cometary O₂ ice remains the primordial, as originally proposed by Taquet et al. (2016).

Acknowledgements. The authors thank Ewine van Dishoeck and Arthur Bosman for many useful discussions and comments that helped the investigation and improved the quality of the manuscript. The authors also thank an anonymous referee for their review of this work. Astrochemistry in Leiden is supported by the European Union A-ERC grant 291141 CHEMPLAN and the Netherlands Research School for Astronomy (NOVA). C.W. also acknowledges the Netherlands Organisation for Scientific Research (NWO, grant 639.041.335) and the University of Leeds for financial support.

References

- Aikawa, Y., Umebayashi, T., Nakano, T., & Miyama, S. M. 1997, *ApJ*, **486**, L51
 Bieler, A., Altwegg, K., Balsiger, H., et al. 2015, *Nature*, **526**, 678
 Brown, W. A., & Bolina, A. S. 2007, *MNRAS*, **374**, 1006
 Cazaux, S., & Tielens, A. G. G. M. 2002, *ApJ*, **575**, L29
 Cleaves, L. I., Adams, F. C., & Bergin, E. A. 2013, *ApJ*, **772**, 5
 Cuppen, H. M., Ioppolo, S., Romanzin, C., & Linnartz, H. 2010, *Phys. Chem. Chem. Phys.*, **12**, 12077
 Cuppen, H. M., Walsh, C., Lamberts, T., et al. 2017, *Space Sci. Rev.*, **212**, 1
 Dalgarno, A. 2006, *Proc. Natl. Acad. Sci. USA*, **103**, 12269
 Dulieu, F., Minissale, M., & Bockelée-Morvan, D. 2017, *A&A*, **597**, A56
 Eistrup, C., Walsh, C., & van Dishoeck E. F. 2016, *A&A*, **595**, A83
 Eistrup, C., Walsh, C., & van Dishoeck E. F. 2018, *A&A*, **613**, A14
 Ennis, C. P., Bennett, C. J., & Kaiser, R. I. 2011, *Phys. Chem. Chem. Phys.*, **13**, 9469
 Garrod, R. T., & Herbst, E. 2006, *A&A*, **457**, 927
 Hayashi, C. 1981, *Prog. Theor. Phys. Suppl.*, **70**, 35
 He, J., Shi, J., Hopkins, T., Vidali, G., & Kaufman, M. J. 2015, *ApJ*, **801**, 120
 Indriolo, N., Neufeld, D. A., Gerin, M., et al. 2015, *ApJ*, **800**, 40
 Ioppolo, S., Cuppen, H. M., Romanzin, C., van Dishoeck, E. F., & Linnartz, H. 2008, *ApJ*, **686**, 1474
 Ioppolo, S., Cuppen, H. M., Romanzin, C., van Dishoeck, E. F., & Linnartz, H. 2010, *Phys. Chem. Chem. Phys.*, **12**, 12065
 Jones, B. M., Kaiser, R. I., & Strazzulla, G. 2014, *ApJ*, **781**, 85
 Lamberts, T., Cuppen, H. M., Ioppolo, S., & Linnartz, H. 2013, *Phys. Chem. Chem. Phys.*, **15**, 8287
 Martín-Doménech, R., Manzano-Santamaría, J., Muñoz Caro, G. M., et al. 2015, *A&A*, **584**, A14
 McElroy, D., Walsh, C., Markwick, A. J., et al. 2013, *A&A*, **550**, A36
 Minissale, M., Congiu, E., & Dulieu, F. 2014, *J. Chem. Phys.*, **140**, 074705
 Mousis, O., Ronnet, T., Brügger, B., et al. 2016, *ApJ*, **823**, L41
 Mousis, O., Ronnet, T., Lunine, J. I., et al. 2018, *ApJ*, **858**, 66
 Penteado, E. M., Walsh, C., & Cuppen, H. M. 2017, *ApJ*, **844**, 71
 Rubin, M., Altwegg, K., van Dishoeck, E. F., & Schwehm, G. 2015, *ApJ*, **815**, L11
 Ruffle, D. P., & Herbst, E. 2000, *MNRAS*, **319**, 837
 Taquet, V., Furuya, K., Walsh, C., & van Dishoeck E. F. 2016, *MNRAS*, **462**, S99
 Teolis, B. D., Plainaki, C., Cassidy, T. A., & Raut, U. 2017, *J. Geophys. Res.*, **122**, 1996
 Tielens, A. G. G. M., & Allamandola, L. J. 1987, *Astrophys. Space Sci. Lib.*, **134**, 397
 van Dishoeck, E. F., & Black, J. H. 1986, in *Interstellar Processes: Abstracts of Contributed Papers*, eds. D. J. Hollenbach & H. A. Thronson, Jr.
 Walsh, C., Nomura, H., & van Dishoeck E. 2015, *A&A*, **582**, A88

Appendix A: Additional table and figures

Figure A.1 shows the gas and ice abundances of O, O₂, H₂O, and H₂O₂ as a function of radius by 10 Myr of evolution, for the chemical network without O₃ chemistry. Figure A.2 shows the gas and ice abundances of O₂, H₂O and H₂O₂ and O₃ as a function of radius by 10 Myr of evolution, for the chemical network including O₃.

Table A.1. Initial abundances (with respect to H_{nuc}) for atomic (reset scenario) and molecular (inheritance scenario) setups.

Species	Atomic	Molecular
H	9.1×10^{-5}	5.0×10^{-5}
He	9.8×10^{-2}	9.8×10^{-2}
H ₂	5.0×10^{-1}	5.0×10^{-1}
N	6.2×10^{-5}	
O	5.2×10^{-4}	
C	1.8×10^{-4}	
S	6.0×10^{-6}	
H ₂ O		3.0×10^{-4}
CO		6.0×10^{-5}
CO ₂		6.0×10^{-5}
CH ₄		1.8×10^{-5}
N ₂		2.1×10^{-5}
NH ₃		2.1×10^{-5}
CH ₃ OH		4.5×10^{-5}
H ₂ S		6.0×10^{-6}
O ₂		0

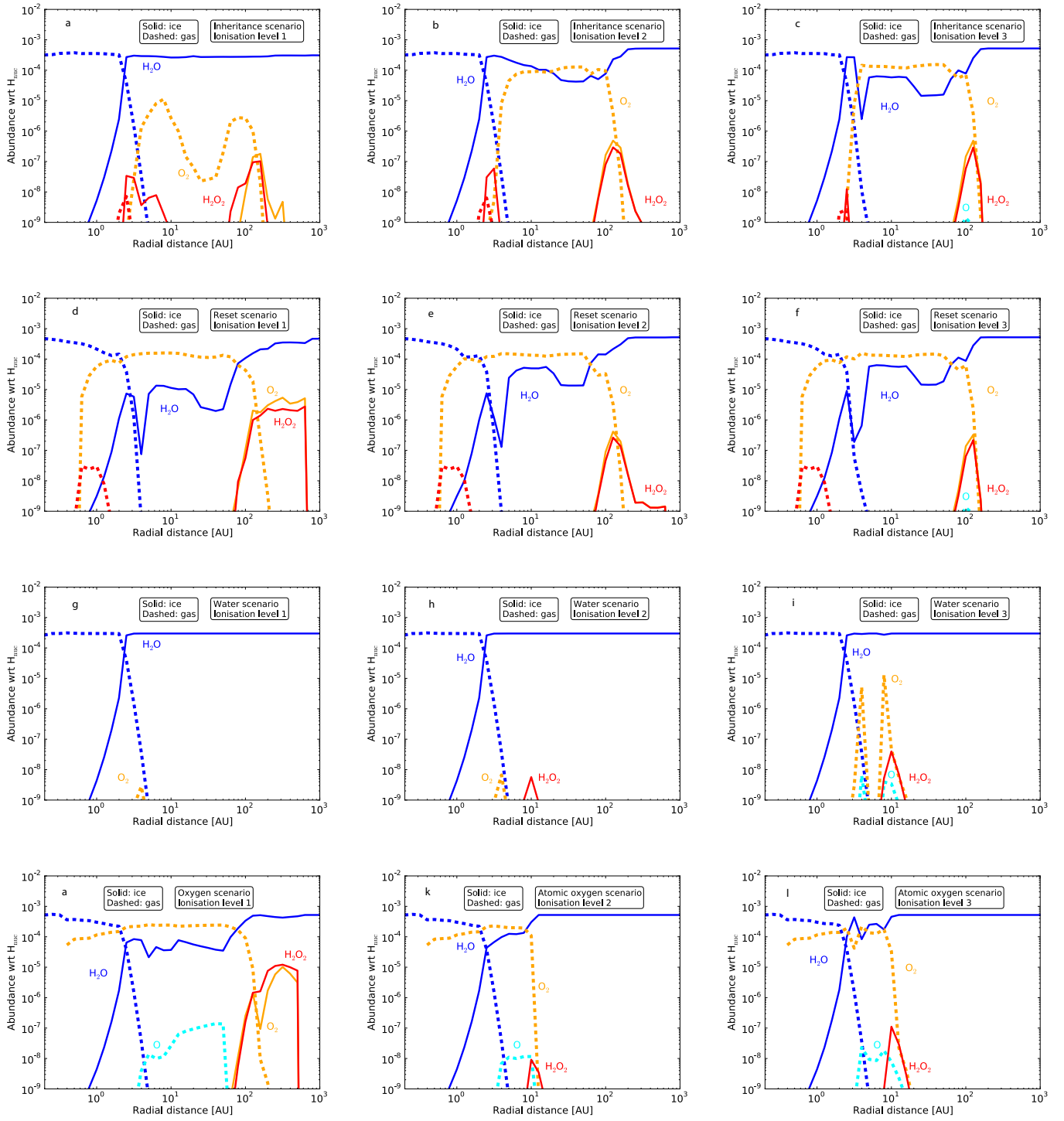


Fig. A.1. Abundances by 10 Myr evolution. *From top to bottom panels:* inheritance scenario, reset scenario, water scenario, and oxygen scenario. *From left to right panels:* changing ionisation levels. The chemical network utilised does not include O₃ chemistry.

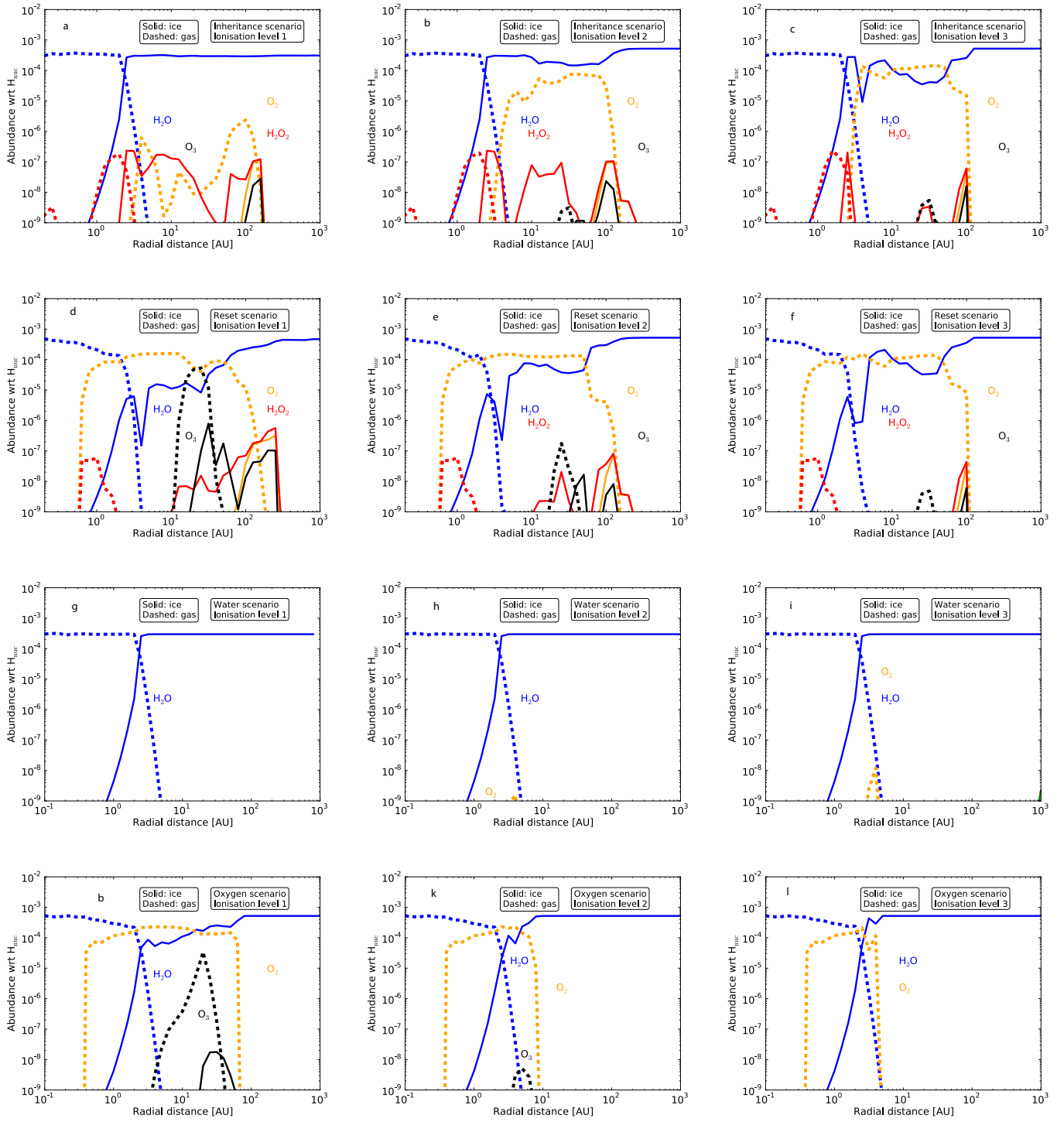


Fig. A.2. Abundances by 10 Myr evolution. *From top to bottom panels:* inheritance scenario, reset scenario, water scenario, and oxygen scenario. *From left to right panels:* changing ionisation levels. The chemical network utilised includes O_3 chemistry.

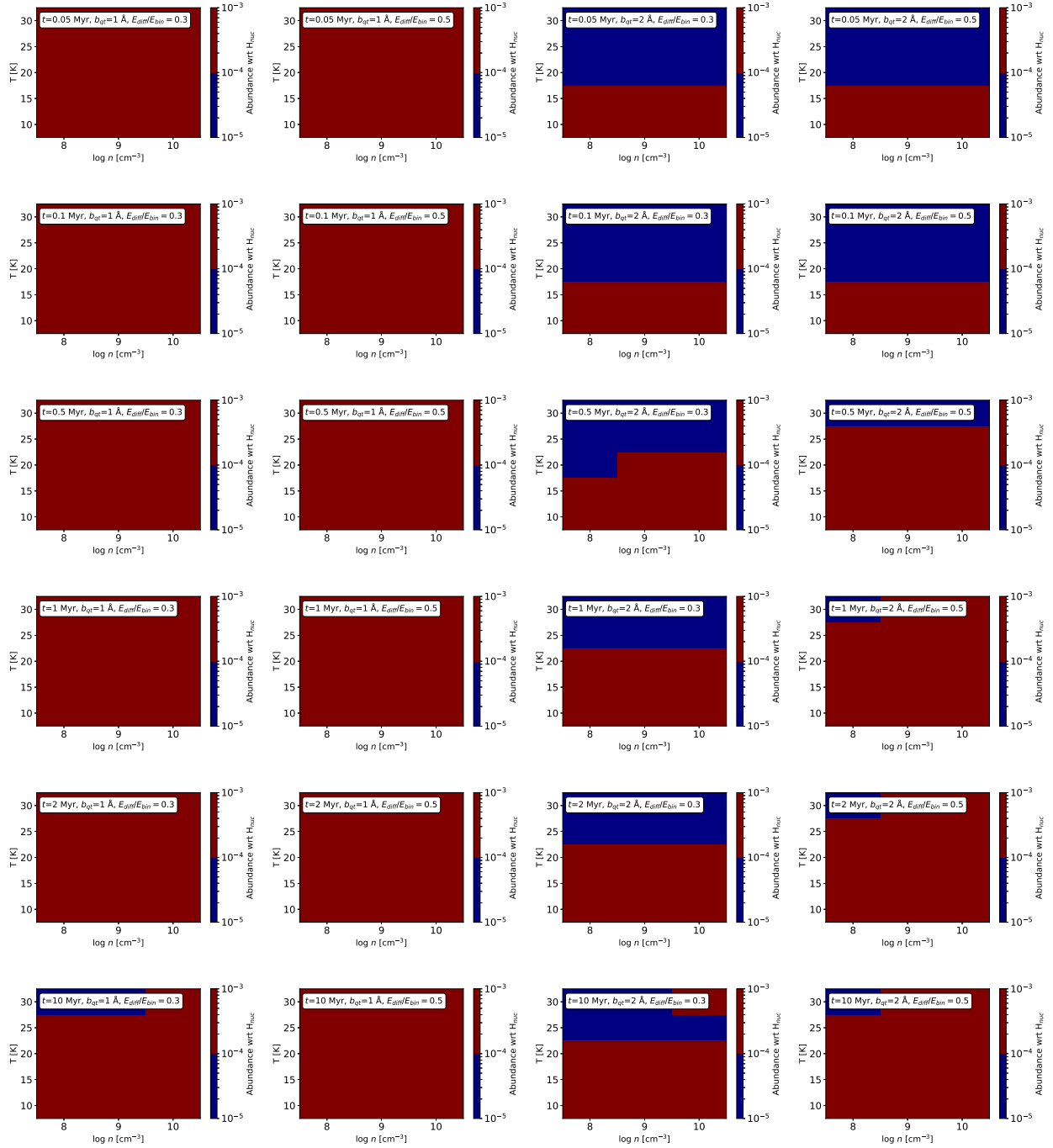


Fig. A.3. Abundances (given as colours) for H₂O ice as function of midplane density (x -axes), and temperature (y -axes) at different evolutionary steps for the reset scenario. *From left to right panels:* abundances from model runs with different parameters for grain–surface reactions: *Cols. 1 and 2:* $b_{gr} = 1 \text{ \AA}$, *Cols. 3 and 4:* $b_{gr} = 2 \text{ \AA}$, *Cols. 1 and 3:* with $E_{diff}/E_{bin} = 0.3$, and *Cols. 2 and 4:* with $E_{diff}/E_{bin} = 0.5$. *From top to bottom panels:* different evolutionary times, from 0.05 Myr (*top*) to 10 Myr (*bottom*). To the right of each plot is a colourbar, indicating the abundance level with respect to H_{nuc} for each colour. The chemical network utilised includes O₃ chemistry.

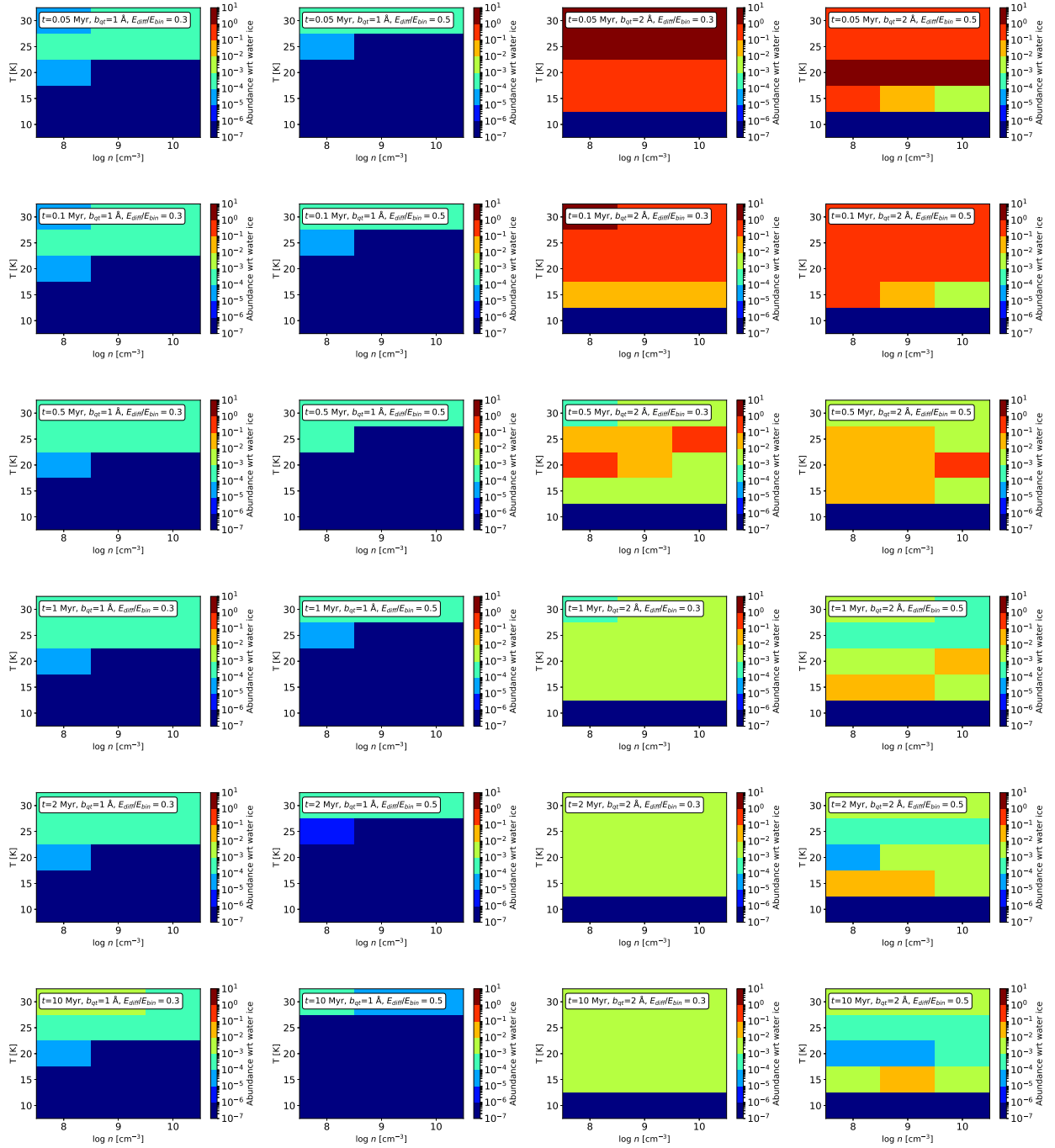


Fig. A.4. Abundances (given as colours) for O₃ ice as function of midplane density (x -axes), and temperature (y -axes) at different evolutionary steps for the reset scenario. *From left to right panels:* abundances from model runs with different parameters for grain–surface reactions: *Cols. 1 and 2:* $b_{qt} = 1 \text{ \AA}$, *Cols. 3 and 4:* $b_{qt} = 2 \text{ \AA}$, *Cols. 1 and 3:* with $E_{\text{diff}}/E_{\text{bin}} = 0.3$, and *Cols. 2 and 4:* with $E_{\text{diff}}/E_{\text{bin}} = 0.5$. *From top to bottom panels:* different evolutionary times, from 0.05 Myr (*top*) to 10 Myr (*bottom*). To the right of each plot is a colourbar, indicating the abundance level with respect to H₂O ice for each colour. The chemical network utilised includes O₃ chemistry. Darkest hue of blue matches the upper limit for cometary O₃ abundances.

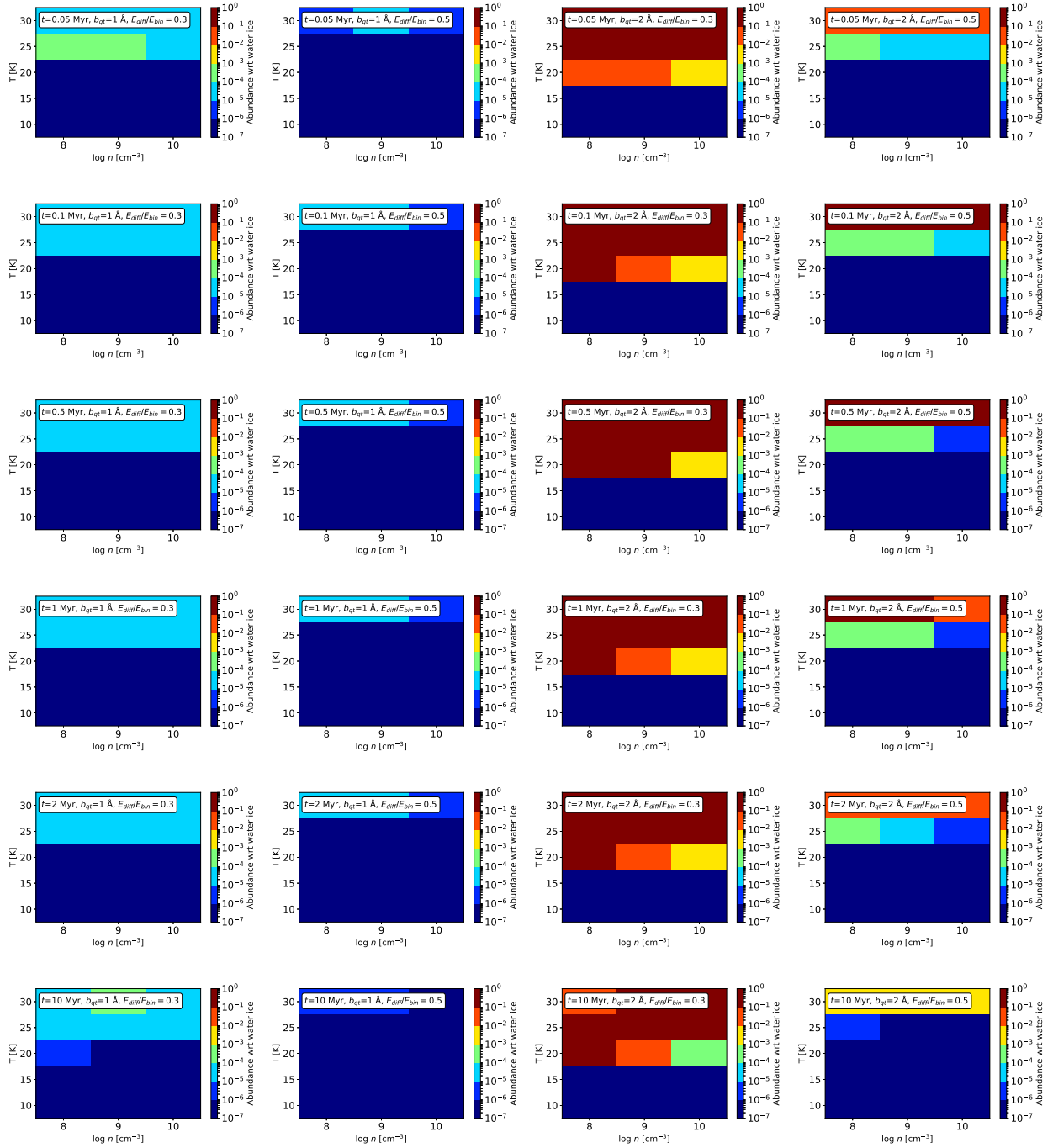


Fig. A.5. Abundances (given as colours) for H₂O₂ ice as function of midplane density (x -axes), and temperature (y -axes) at different evolutionary steps for the reset scenario. *From left to right panels:* abundances from model runs with different parameters for grain–surface reactions: *Cols. 1 and 2:* $b_{qt} = 1 \text{ \AA}$, *Cols. 3 and 4:* $b_{qt} = 2 \text{ \AA}$, *Cols. 1 and 3:* with $E_{\text{diff}}/E_{\text{bin}} = 0.3$, and *Cols. 2 and 4:* with $E_{\text{diff}}/E_{\text{bin}} = 0.5$. *From top to bottom panels:* different evolutionary times, from 0.05 Myr (*top*) to 10 Myr (*bottom*). To the right of each plot is a colourbar, indicating the abundance level with respect to H₂O ice for each colour. The chemical network utilised includes O₃ chemistry. Lightest hue of blue matches the cometary H₂O₂ abundances.