Warm dust surface chemistry in protoplanetary disks

Formation of phyllosilicates

W. F. Thi\textsuperscript{1}, S. Hocuk\textsuperscript{1,6}, I. Kamp\textsuperscript{2}, P. Woitke\textsuperscript{3,7}, Ch. Rab\textsuperscript{2}, S. Cazaux\textsuperscript{4}, P. Caselli\textsuperscript{1}, and M. D’Angelo\textsuperscript{5,2}

\textsuperscript{1} Max Planck Institute for Extraterrestrial Physics, Giessenbachstrasse, 85741 Garching, Germany
\textsuperscript{2} Kapteyn Astronomical Institute, University of Groningen, Postbus 800, 9700 AV Groningen, The Netherlands
\textsuperscript{3} SUPA, School of Physics & Astronomy, University of St. Andrews, North Haugh, St. Andrews, KY16 9SS, UK
\textsuperscript{4} Faculty of Aerospace Engineering, Delft University of Technology, Delft, The Netherlands
\textsuperscript{5} Zernike Institute for Advanced Materials, University of Groningen, PO Box 221, 9700 AE Groningen, The Netherlands
\textsuperscript{6} Centre for Exoplanet Science, University of St Andrews, St Andrews, UK

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ABSTRACT

\textit{Context.} The origin of the reservoirs of water on Earth is debated. The Earth’s crust may contain at least three times more water than the oceans. This crust water is found in the form of phyllosilicates, whose origin probably differs from that of the oceans.

\textit{Aims.} We test the possibility to form phyllosilicates in protoplanetary disks, which can be the building blocks of terrestrial planets.

\textit{Methods.} We developed an exploratory rate-based warm surface chemistry model where water from the gas-phase can chemisorb on dust grain surfaces and subsequently diffuse into the silicate cores. We applied the phyllosilicate formation to a zero-dimensional chemical model and to a 2D protoplanetary disk model (P\textsubscript{RODI}M\textsubscript{O}). The disk model includes in addition to the cold and warm surface chemistry continuum and line radiative transfer, photoprocesses (photodissociation, photoionisation, and photodesorption), gas-phase cold and warm chemistry including three-body reactions, and detailed thermal balance.

\textit{Results.} Despite the high energy barrier for water chemisorption on silicate grain surfaces and for diffusion into the core, the chemisorption sites at the surfaces can be occupied by a hydroxyl bond (–OH) at all gas and dust temperatures from 80 to 700 K for a gas density of $2 \times 10^4$ cm$^{-3}$. The chemisorption sites in the silicate cores are occupied at temperatures between 250 and 700 K. At higher temperatures thermal desorption of chemisorbed water occurs. The occupation efficiency is only limited by the maximum water uptake of the silicate. The timescales for complete hydration are at most $10^5$ yr for 1 mm radius grains at a gas density of $10^8$ cm$^{-3}$.

\textit{Conclusions.} Phyllosilicates can be formed on dust grains at the dust coagulation stage in protoplanetary disks within 1 Myr. It is however not clear whether the amount of phyllosilicate formed by warm surface chemistry is sufficient compared to that found in Solar System objects.

\textit{Key words.} astrochemistry – methods: numerical – stars: pre-main sequence

1. Introduction

The origin of the water reservoirs on Earth is a long-standing debate (van Dishoeck et al. 2014; Ceccarelli et al. 2014; Caselli & Ceccarelli 2012; Drake 2005; Cleeves et al. 2014). It is unclear what the sources of water in terrestrial planets are and whether water was contained in the Earth’s formation blocks already at its formation or whether the water was delivered later during a late heavy bombardment phase.

The Earth’s bulk composition and that of meteorites known as enstatite chondrites suggest a “dry” proto-Earth scenario: the initial building blocks of the Earth are made of dry rocks, composed only of non-hydrated silicates and carbonaceous materials. Most of the water is brought afterward by comets and asteroids during the phase of heavy bombardments (Nesvorný et al. 2017). In addition to the water in the oceans, large reservoirs of water trapped in silicates in the Earth’s mantle have been found with up to three times the amount of water in Earth’s oceans (Schmandt et al. 2014; Marty 2012). The origin of the water in the Earth’s mantle may differ from that of the oceans and the amount of mixing between these repositories is not clear. This scenario is challenged by the Rosetta mission. Rosetta measurements found that the Jupiter family comet 67P/Churyumov-Gerasimenko has a much higher HDO/H\textsubscript{2}O ratio that in the Earth’s oceans (Altwegg et al. 2014).

In the second model, the “wet” scenario, most of the water on Earth comes from the release of water vapour trapped inside the planetesimals like carbonaceous chondrites as hydrated silicate upon impact or during volcanism (van Dishoeck et al. 2014; Drake 2005). Water is already present in the initial phase of Earth’s formation.

Water-rich planetesimals located at 2–3 au are perturbed by the giant planets and collide with the young Earth (Morbidelli et al. 2000; Gomes et al. 2005; Raymond et al. 2004, 2005). Such planetesimals contain water in form of hydrated silicates (phyllosilicates). A few main belt asteroids are ice-covered planetesimals, which may have brought water on Earth (Jewitt 2012). C-type main belt asteroids, whose spectra resemble those of carbonaceous chondrite meteorites are located throughout the main belt with the peak distribution located beyond the solar nebula snow line (Rivkin et al. 2002).

Alexander et al. (2018) reviewed the water reservoir in the Solar System small bodies and Deloule & Robert (1995) studied the amount of interstellar water in meteorites. Chondrites
are known to have kept the record of solar nebula and asteroidal histories. Hydrated carbonaceous chondrites are meteorites where water is found in the form of phyllosilicates (Beck et al. 2010, 2014). It is generally believed that the direct hydration of anhydrous silicates by water vapour in the solar nebula was too inefficient compared to its lifetime (Fegley 2000), although it is thermodynamically possible (Zolotov 2012). Therefore, it is generally accepted that the original anhydrous silicate in the parent bodies interacted with water-rich fluids leading to the hydrolysis of the silicate (Brearley 2006; Bischoff 1998). However, studies of the D to H ratios in meteorites suggest that water in meteorites can be formed out of thermodynamic equilibrium (Deloule & Robert 1995).

Original analyses of the Moon’s samples implied a dry Moon formation but recent studies suggest that a wet formation scenario is also possible (Saal et al. 2013; Hui et al. 2013; Anand et al. 2014; Chen et al. 2015). Near-infrared reflectance spectra of the Moon pyroclastic deposits support the presence of Earth-like water content in the lunar interior (Mililken & Li 2017).

The water in the Earth mantle may have been incorporated already in the dust grains in the form of phyllosilicates. This wet scenario is supported by the presence of a lower D to H ratio in lavas in Earth’s oceans (Hallis et al. 2015). The D to H ratio found in lavas may be more representative of the primordial D/H than the value found in the Earth ocean. The lavas D to H ratio is similar to that of carbonaceous chondrites, which may have acquired their water by absorption at the solar nebula stage.

Hydration at silicate grain surfaces is an alternative mechanism that has been shown to occur experimentally (Rietmeijer et al. 2004; Yamamoto & Tachibana 2016; Guillou et al. 2015). As an activated process, the hydration timescale depends exponentially on the activation energy and temperature. Yamamoto & Tachibana (2016) noticed that hydration of amorphous silicates is enhanced compared to the hydration of crystalline silicates. A low activation energy would permit hydration of the small grains in the early life of protoplanetary disks. Those hydrated grains will subsequently coagulate into planetesimals (Ciesla & Lauretta 2005). Alternatively, warm gas in the inner region of protoplanetary disks or gas heated by shocks has enough energy to overcome the barrier (Ciesla et al. 2002; Purukawa et al. 2011). The dust grain hydration models have only explored formation via the Eley–Rideal process, in other words the direct formation of chemisorbed OH silicate bonds upon impact of a gas-phase water vapour molecules on silicate grains, we also considered hydration of silicate grain surfaces via the Langmuir–Hinshelwood process. In the Langmuir–Hinshelwood process, the weakly adsorbed water molecules (physisorbed water) scan the silicate surface to search for an empty chemisorption site and attempt to overcome either thermally or via tunnelling the activation barrier. Dust grains at temperatures below 100 K are covered by up to a few hundreds layers of water ice. The amount of water molecules is large with one layer of water ice corresponding for a 0.1 μm radius grain to ~1.9 × 10^6 molecules.

Water ice is dynamic and mobile (Ghesquiere et al. 2015) and therefore, the process of water diffusion described above is akin to the so-called aqueous alteration (Zolensky et al. 2008) although water has an amorphous solid structure, instead of being liquid. Even for dust grains that are above the sublimation temperature of water ice, water molecules would have ample time to scan the surface before they desorb because the experimental and theoretical value of the activation energy to form a chemisorption bond is around 3000 K (Wegner 1983; Stimpfl et al. 2006; Muralidharan et al. 2008; Leeuw et al. 2010; King et al. 2010; Prigiobbie et al. 2013; Asaduzzaman et al. 2014). This is below the desorption energy of physisorbed water (~5700 K, Fraser et al. 2001).

D’Angelo et al. (2019) used a Monte Carlo numerical code to study in details the adsorption of water on silicate between 300 and 800 K. In their Monte-Carlo simulation the [100] forsterite crystal lattice is simulated by a grid composed of 20 × 20 cells. Each unit cell has four possible binding sites with binding energies ranging between 8000 and 20 000 K corresponding to Mg cations, three of which are closer to the surface and easily accessible to water molecules. The highest binding sites represent 45% of the total number of sites. Also, the formation of water clusters on surfaces have been investigated, and showed that clusters increase the water coverage of the dust. By considering small dust particles of 0.1 μm, an important amount of water of up to ten Earth oceans can be available in the early Earth.

Woitke et al. (2018) performed a chemical equilibrium calculation and found that for inner midplane protoplanetary disk conditions phyllosilicates are the most thermodynamically stable species.

In this study we focused on the global hydration kinetics of silicate grains in protoplanetary disk conditions. Another novelty in our study is that we took the diffusion into the silicate core into account.

The paper is organised as follows: we first introduce the chemical model in Sect. 2; the code and the models are presented in Sect. 3; the results are shown and discussed in Sects. 4 and 5; the conclusions and perspectives are given in Sect. 6.

2. Silicate hydration and dehydration model

Chemisorption concerns two species in our model atomic hydrogen and water. Chemisorbed H atoms (on the silicate surface \( *H* \)) can recombine with another H atom to form \( H_2 \). Details on the \( H_2 \) formation in our model are explained in a previous paper (Thi et al. 2020). The formation occurs on the grain surfaces by the Langmuir–Hinshelwood and Eley–Rideal processes involving physisorbed and chemisorbed H atoms. \( H_2 \) can also form via hydrogennated PAHs (HPAH) and PAH cations (HPAH*).

Both the H atoms and water molecules are assumed to compete for the available number of surface chemisorption sites (at the
silicate surface and in the core). More sophisticated models take the possibility of a species with a high binding energy to swap space with less strongly bounded species into account (Cuppen & Herbst 2007). We assumed in this exploratory study that only hydrogen atoms and water molecules can diffuse from the silicate surface inside the silicate core. The adopted surface that only hydrogen atoms and water molecules can diffuse from (Cuppen & Herbst 2007). We assumed in this exploratory study to swap space with less strongly bounded species into account silicate surface and in the core). More sophisticated models selected only five and 500 layers inside the silicate core for the cloud and protoplanetary disk model respectively in addition to the silicate surface layer, making a total of six and 501 layers of chemisorption sites for hydrogen atoms and water to bind to. The choice for 500 silicate core layers will be discussed in a subsequent section.

Water can form in the gas-phase by ion-neutral reactions at low temperatures and via neutral-neutral reactions at high temperatures (Thi & Bik 2005; Thi et al. 2010; van Dishoeck et al. 2014). At temperatures below the sublimation temperature of the water ice, water is formed on the surface by encounters of physisorbed species starting from the hydrogenation of O into OH. Hydrogenation of atomic surface atoms is efficient below ~15 K.

We restricted ourselves to this surface path although alternative routes via O₂H and H₂O₂# for example have been proposed (Taquet et al. 2013; Cuppen et al. 2017; Lamberts et al. 2014; Cazaux et al. 2010). At high temperatures water is first formed in the gas phase before being incorporated in the grains. Another important route is the reaction of OH# with H₂# (Pentado et al. 2017).

We updated the gas-phase reaction rate coefficients between OH and H₂ with the values from Meisner & Küstner (2016), who took tunnelling effects into account in their computations. Warm gas-phase chemistry of water is discussed in the context of protoplanetary disks in Thi et al. (2010), Kamp et al. (2013), Antonellini et al. (2015, 2016), Woitke et al. (2009b), van Dishoeck et al. (2014), Bethell & Bergin (2009), Du & Bergin (2014).

Water ice molecules located at the interface between the icy mantle and the silicate surface can overcome an activation barrier to chemisorb to the silicate surface. Likewise, if the grain is too hot to host an icy mantle, water molecules can impinge directly on the silicate surface, scan it, find a suitable free chemisorption site, and attempt to overcome the barrier. If the dust is warm enough, the chemisorbed water can diffuse inside the silicate core.

We used the theoretical activation energy from Asaduzzaman et al. (2014) for the formation of a chemisorption bond between water (previously physisorbed or in the gas-phase) and the silicate (3010 K), consistent with experimental values in the order of 2500 K (Yamamoto & Tachibana 2016). We chose a binding energy of 32 710 K for water chemisorption, consistent with dehydration experiments of phyllosilicate (Sawai et al. 2013). Kostinski et al. (2012) measured a value of 12 770 K for the water vapour penetration barrier in the silicate core. The diffusion over chemisorption energy is 0.39. The desorption and diffusion are calculated according to standard cold surface chemistry methods (Hasegawa & Herbst 1993). The activation barrier for the formation of a chemisorption bond is much lower than the barrier for water diffusion from a chemisorbed water at the silicate surface to another chemisorption site in the silicate core. Therefore, we expect the formation of a layer of chemisorbed water on the silicate surface at much lower dust temperature than the diffusion of chemisorbed silicate surface water into the silicate core. This is called a precursor-mediated chemisorption, the physisorption site being a precursor state. The desorption energy is simply the binding energy in the chemisorbed state.

We adopted the same value for diffusion of water on the silicate surface and for diffusion from the silicate surface towards the core centre. The desorption rate pre-factor frequency follows the formula for a rectangular barrier of width $\alpha_i$ and height $E_{i,des}$:

$$v_{ij} = \frac{2N_{surf} E_{i,des}}{m_i \sqrt{\pi}}.$$

We derived a frequency $v_{ij}$ of $(1–10) \times 10^{12}$ Hz assuming a surface site density is $N_{surf} = 1.5 \times 10^{15}$ sites cm$^{-2}$. This formula may underestimate the actual values up to $10^{16}$ Hz (Rettner et al. 1996).

The water molecules and H atoms compete for the limited silicate surface and silicate core chemisorption sites. A chemisorbed H-atom can recombine with another H-atom to form H₂, while a hydroxyl bond is assumed to be chemically inert once formed. The chemisorbed species and some special reactions included in our model are summarised in Tables 1 and 2.

### 3. Modelling

#### 3.1. The ProDiMo code

ProDiMo is a hydrostatic 2D code designed to model the physics and chemistry of protoplanetary disks (Woitke et al. 2016, 2009a; Kamp et al. 2010; Thi et al. 2011; Aresu et al. 2011).
It includes detailed continuum and line radiative transfer, heating, and cooling balance, and gas and surface chemistry. The gas and surface chemistry is computed by solving rate equations both using a time-dependent or a steady-state solver. The gas densities in protoplanetary disks are higher than $10^4$ cm$^{-3}$, which makes the use of the rate-equation method appropriate.

### 3.2. Chemical network

We considered a simplified chemistry in the gas-phase, on the surface of the icy grain mantle, the ice mantle, the surface of the silicate, and the silicate core. The adsorption, diffusion, and desorption rates depend on the actual adsorption sites. However, the model does not account for detailed location of each solid species. The gas-phase and simple ice species are the same as those in the small network described in Kamp et al. (2017) based on the UMIST2012 database (McElroy et al. 2013). The additional species are adsorbed hydrogen atoms and OH ice.

The chemical network is run using the method of “pseudo” chemical species used for the surface chemisorption sites (first pseudo element *) and for the silicate core sites (second pseudo element !). The “elemental abundance” of pseudo elements are set by the total quantity of the chemisorption sites (at the silicate surface and in the silicate core). Assuming a surface site number density $N_{\text{surf}}$, the number of silicate surface sites is $N_{\text{surf}}a_\text{surf}^2$ per grain where $a$ is the grain radius. The number of silicate core chemisorption sites is $N_{\text{core,layer}}N_{\text{surf}}a_\text{surf}^2$ per grain, where $N_{\text{core,layer}}$ is a free parameter to account for the amount of “layers” of “water” that a silicate grain can have. Atomic hydrogen is allowed to be physisorbed (H#) or chemisorbed on the silicate surface (H#) while water can be physisorbed (H$_2$O#), chemisorbed at the silicate surface (H$_2$O#) or in the silicate core (H$_2$O#). The pseudo elements/species method permits the code to track how a species is bound (physisorbed, chemisorbed at the silicate surface, or chemisorbed in the silicate core) statistically. The method naturally accounts for the limited number of chemisorption sites.

Surface rates account for the competition between diffusion and reaction but not with the desorption processes (thermal, photoabsorption induced and cosmic ray induced desorption), whose effects are explicitly accounted for in the rate equation scheme. Both atomic hydrogen and water can be chemisorbed on the silicate core while only water can chemisorb in the silicate core. There is no reaction-induced desorption. Tunnelling is accounted for diffusions and reactions for all the species assuming a rectangular shape for the barrier and using the Bell’s formula (Bell 1980).

The amount of water trapped as phyllosilicate is not kinetically limited but is rather limited by the capacity of the silicate to accommodate large amounts of water. We can estimate the uptake of water in the silicate core as function of the number of layers in the core. The number density per dust layer is $n_{\text{surf,chem}} = 4\pi N_{\text{surf}}a_\text{surf}^2 n_d$, which is the number of chemisorption sites per layer.

When all the chemisorption sites are occupied by water, the total mass of chemisorbed water is $(1 + N_{\text{core,layer}}) 4\pi N_{\text{surf}}a_\text{surf}^2 n_d m_{\text{H}_2\text{O}}$ (g cm$^{-3}$), where the total number density of dust grains is

$$n_d = \frac{1.386 \text{ amu} n_{\text{H}_2\text{O}} \delta}{(4/3)\rho a_\text{surf}^2} \text{ cm}^{-3}. \quad (2)$$

The volumetric mass of solid is 1.386 amu $n_{\text{H}_2\text{O}}/\delta$, where $\delta$ is the dust to gas mass ratio assumed to be 0.01. The chemisorbed water over total solid mass ratio in percentage is

$$n_{\text{H}_2\text{O},\%} = 0.01 \frac{3m_{\text{H}_2\text{O}}(1 + N_{\text{core,layer}})N_{\text{surf}}}{\rho a_\text{surf}^2} \approx \frac{0.045}{a_\mu m} (1 + N_{\text{core,layer}}), \quad (3)$$

which corresponds to $n_{\text{H}_2\text{O},\%} \sim 2.7\%$ for 5 core layers (plus the top layer) and an average grain radius of 0.1 micrometers. Here $a_\mu m$ is the grain radius in units of microns. It should be noticed that the number of core layers is not constrained in the model so that the actual parameter that governs the amount of chemisorbed water is $(1 + N_{\text{core,layer}})/a_\mu m$. In reality minerals have a maximum water storage capacity with the Ringwoodite mineral capable of storing up to ~3% of water in weight (Kohlstedt et al. 1996). This is consistent with the value of 2.7% used in this study although storage capacities differ from a mineral to another one with capacities up to 6% (Hirschmann et al. 2005). Alternatively, one can set the chemisorbed water over total solid mass ratio as the input parameter and derive the corresponding number of layers in the silicate core. Therefore, we assume up to five layers in the silicate core can be hydrated ($N_{\text{core,layer}} = 5$). The volume ratio of the layers of thickness $\Delta a$, with $a$ being the grain radius, to the total silicate core volume is $3\Delta a/a$. Considering that a silicate layer thickness is 3 Å, five layers plus the top layer correspond to 54 Å, which gives $3\Delta a/a = 54/333 \approx 0.054$ or 5.4%. We adopted a PAH abundance of $3 \times 10^{-7}$, which corresponds to $f_{\text{PAH}} = 1$ (Tielsch 2005).

#### 3.3. Simple zero-dimensional warm-up models

The models are zero-dimensional pure chemical models where the physical condition are set (density, gas and dust temperature, UV field, extinction, cosmic ray flux, dust grain mean radius). We ran a typical cold molecular cloud model ($n_{\text{H}_2} = 2 \times 10^4$ cm$^{-3}$, $T_d = T_g = 10$ K) for 10 Myr to set the initial abundances for the warm-up runs. The initial abundances for all species are in the single or doubly ionised form (H$^+$, He$^+$, C$^{++}$, O$^{++}$, N$^{++}$, S$^{++}$, Mg$^{++}$, Na$^{++}$, Fe$^{++}$, Ni$^{++}$, Ar$^{++}$, PAH$^{++}$) for both the zero-dimensional and disk model. The instantaneous warm-up runs consist of models with increasing temperatures (still assuming $T_d = T_g$) from 20 to 900 K. The silicate is assumed “dry” at the beginning of the chemical run. We monitor in particular the abundance of water molecules in its different forms: gas-phase, physisorbed ice, chemisorbed water on the silicate surface, in the silicate core. The elemental abundances are listed in Table 3. All the species modelled in this study are shown in Table 4. Other parameters for the cloud model can be found in Table 5.

It should be stressed that the zero-dimensional model has been chosen to illustrate the effects of increasing gas and dust temperature on the formation of phyllosilicate and does not attempt to simulate an actual astrophysical environment.

#### 3.4. Hydration timescale models

We ran two series of models to determine the timescales as a function of the temperature (assuming $T_{\text{gas}} = T_{\text{dust}}$) required to hydrate the silicate grains such that their mass percentages of water reach ~2.7%. In the first series, the grain radius is fixed at 0.1 µm and the gas density is $2 \times 10^4$ cm$^{-3}$, $2 \times 10^5$ cm$^{-3}$, and $2 \times 10^6$ cm$^{-3}$. Using Eq. (3), we derived the respective number of core layers to be 5, 58, and 60 000 in addition to the top surface layer. In the second series, we fixed the gas density at $10^5$ cm$^{-3}$ and varied the grain radius (0.1 µm, 1 µm, 1 mm).
Table 3. Elemental abundances.

| Elements | Abundance |
|----------|-----------|
| H        | 12.00     |
| He       | 10.984    |
| C        | 8.14      |
| N        | 7.90      |
| O        | 8.48      |
| Ne       | 7.95      |
| Na       | 3.36      |
| Mg       | 4.03      |
| Ar       | 6.08      |
| Fe       | 3.24      |
| S        | 5.27      |
| Si       | 4.24      |
| PAH      | See text  |

Note: The abundances are $10^{-12}$ relative to H nuclei.

3.5. DIANA-ProDMo protoplanetary disk model

The input for the DIANA disk model are summarised in Table 6. A detailed explanation of the parameters can be found in Woitke et al. (2016). The disk model uses a dust size distribution instead of a single grain radius. Dust grains can settle in the disk as parametrised by the turbulent mixing parameter $\alpha_{\text{turb}}$. We ran the model in the fixed vertical structure mode where both the radial and vertical density structures are parametrised. The chemical network is very limited with the addition of simple surface species and the surface chemisorbed species (for H and H$_2$O) and core chemisorbed species (H$_2$O only). The base network is described in Kamp et al. (2017). Most of the rates are taken from the UMIST2012 database (McElroy et al. 2013). Contrary to the zero-dimensional model, photodesorption for the physisorbed species is accounted for. Photodesorption does not occur for chemisorbed species. The UV field in the protoplanetary disk is computed by solving the continuum radiative transfer (Woitke et al. 2009a). We set for the protoplanetary disk model a maximum of 500 layers ($N_{\text{core layer}} = 500$) in the silicate core such that the average maximum uptake of water in silicate is $2\sim3\%$ of the total mass in solid form not taking water ice into account.

4. Results

4.1. Zero-dimensional model results

The results of models with warm-up temperature of 50, 150, and 250 K are shown in Fig. 1. The left panels contain the time variation of H, H chemisorbed on the grain silicate surface, H$_2$O, water vapour, physisorbed H$_2$O# and *H$_2$O# chemisorbed on the grain silicate surface. The right panels are the abundance of water in the different reservoirs compared to the abundance of free surface and core chemisorption sites.

The abundance of H$_2$ stays relatively constant over time. On the contrary, the atomic hydrogen abundance increases as soon as all the chemisorption sites are occupied by water because H$_2$ is continuously destroyed and only reformed at high dust temperature when H-atoms occupy the chemisorption sites. Once a water is bound to the silicate site, it is not reactive and can only diffuse to the surface if the site is in the silicate core and diffuse or desorb if the site is located at the silicate surface. The chemisorbed water prevents H-atoms to occupy the chemisorption sites.

Figure 2 shows the different water reservoirs after 1 and 5 Myr from the initial condition at 10 K. Below the desorption temperature of water ice at $\sim 100$ K for a gas at density $2 \times 10^{13}$ cm$^{-3}$, most of the oxygen is in form of water ice (physisorbed water). The second reservoir up to $\sim 45$ K with a relative abundance $10^{-4}$–$10^{-3}$ times lower is water vapour. The gas-phase water abundance reaches a few $10^{-4}$ consistent with the detected abundance in molecular clouds (Wiström et al. 2014). The physisorbed water formation activation energy is lower than the desorption energy of water at $\sim 5700$ K. Therefore, a surface water molecule can scan the entire surface and diffuse into the core before it desorbs as soon as the molecule has energy to overcome the barrier. In our density conditions, this happens from $\sim 45$ K. The consequence is that the water molecules will occupy all the free chemisorption sites on the silicate surface. From 50 to $\sim 250$ K, the molecules are not able to penetrate into the core because of the high activation barriers for diffusion but they occupy all the available surface chemisorption sites. The water abundance is the lowest between 100 and 200 K where only surface chemisorbed water is present. At temperatures between 100 and 250 K the gas-phase water formation route is inhibited by the activation barriers (Woitke et al. 2009b). The slow gas-phase water formation cannot compensate the water destruction. Most of the oxygen is in form of atomic oxygen (see the middle panels in Fig. 1). From 200 K, gas phase formation of water becomes again efficient to compensate for the decrease in H$_2$O abundance due to a lower H$_2$ abundance. From $\sim 100$ K, all the surface chemisorption sites are occupied by water molecules. From $\sim 250$ K, chemisorbed water molecules at the silicate surface start to penetrate in the core. From 250 K till the desorption of water from chemisorption sites at $\sim 700$ K all the chemisorption sites at the surface and in the silicate core are occupied by water (–OH bonds) within 1 Myr. It should be noted that the desorption temperature depends on the species density $n_i$, the thermal speed $v_i$, the surface density of site $N_{\text{surf}}$, and on the vibrational frequency of the species in the surface potential well. Using the formula of Hollenbach et al. (2009) the desorption from chemisorption sites will occur at 847 K for a gas density of $10^3$ cm$^{-3}$. The abundances in the different water reservoirs change only slightly from 1 to 5 Myr except for gas-phase water at low temperatures.

The reason of the “blocking” or “poisoning” of the chemisorption sites by water is that when two chemisorbed hydrogen atoms recombine, the unoccupied sites are quickly taken by a water molecule. Since in our model, chemisorbed water can only desorb and does not react, the chemisorbed sites are “poisoned” by water. The consequence is a significant decrease of H$_2$ formation by the recombination of two chemisorbed H atoms although most of the H$_2$ formation occurs via hydroxylated PAHs and PAH cations. This contrasts with H$_2$ formation in models without chemisorbed water where efficient H$_2$ formation takes place by encounters of chemisorbed H atoms (see Thi et al. 2020). All the chemisorption sites are rapidly occupied by water, the amount of chemisorbed water is only restricted by the maximum water uptake by silicates (see Sect. 5). When all the chemisorption sites are occupied by water molecules, the abundance of trapped water is still $\sim$ten times less than the abundance of water vapour. Above 700 K, only water vapour remains with virtually all the available oxygen not in CO locked into water vapour.
Table 4. Gas and solid species in the network.

| Parameter                  | Symbol | Values |
|----------------------------|--------|--------|
| Chemical elements          |        |        |
| H, He, C, N, O, Ne, Na, Mg, Si, S, Ar, Fe |        | 12     |
| Pseudo elements            |        |        |
| *, !                       |        | 2      |
| (H)                        | H, H⁺, H⁻, H₂, H₃, H₃⁺, H⁺²⁺ | 7      |
| (He)                       | He, He⁺ | 2      |
| (C–H)                      | C, C⁺, C⁺⁺, CH, CH⁺, CH₂, CH₂⁺, CH₃, CH₃⁺, CH₄, CH₄⁺, CH₅⁺ | 12     |
| (C–N)                      | CN, CN⁺, HCN, HCN⁺, HCNH⁺ | 5      |
| (C–O)                      | CO, CO⁺, HCO, HCO⁺, CO₂, CO₂⁺, HCO₂⁺ | 7      |
| (N–H)                      | N, N⁺, N⁺⁺, NH, NH⁺, NH₂, NH₂⁺, NH₃, NH₃⁺, NH₄⁺ | 10     |
| (N–N)                      | N₂, N₂⁺, N₂⁺⁺ | 3      |
| (N–O)                      | NO, NO⁺ | 2      |
| (O–H)                      | O, O⁺, O⁺⁺, OH, OH⁺, H₂O, H₂O⁺, H₂O⁺⁺ | 8      |
| (O–O)                      | O₂, O₂⁺ | 2      |
| (O–S)                      | SO, SO⁺, SO₂, SO₂⁺, HSO₂⁺ | 5      |
| (S–H)                      | S, S⁺, S⁺⁺ | 3      |
| (Si–H)                     | Si, Si⁺, Si⁺⁺, SiH, SiH⁺, SiH₂⁺ | 6      |
| (Si–O)                     | SiO, SiO⁺, SiOH⁺ | 3      |
| (Na)                       | Na, Na⁺, Na⁺⁺ | 3      |
| (Mg)                       | Mg⁺, Mg⁺⁺, Mg⁺⁺⁺ | 3      |
| (Fe)                       | Fe⁺, Fe⁺⁺, Fe⁺⁺⁺ | 3      |
| (Ne)                       | Ne⁺, Ne⁺⁺, Ne⁺⁺⁺ | 3      |
| (Ar)                       | Ar⁺, Ar⁺⁺, Ar⁺⁺⁺ | 3      |
| Ice                        | CO⁺, H₂O⁺, CO₂⁺, CH₃⁺, NH₃⁺, SiO⁺, SO₂⁺, O₂⁺, HCN⁺, N₂⁺⁺ | 10     |
| Pseudo species             |        | 2      |
| Additional species         | H⁺, O⁺, OH⁺, H⁺⁺, N₂⁺⁺, PAH, PAH⁺, PAH⁺⁺, PAH⁺⁺⁺, PAH⁺⁺⁺⁺, HPAH⁺⁺⁺⁺ | 15     |

Table 5. Cloud model parameters.

| Parameter                  | Symbol | Values |
|----------------------------|--------|--------|
| Gas density                | n(H)   | 2 × 10⁶ cm⁻³ |
| Temperature                | T_d = T_g | 10 and 20 to 900 K |
| Extinction                 | A_V    | 10     |
| Strength of interstellar UV| χISM   | 1      |
| Cosmic ray H₂ ionisation rate | ζCR    | 1.7 × 10⁻¹⁷ s⁻¹ |
| Grain radius               | a      | 10⁻³ cm |
| Dust-to-gas mass ratio     | δ      | 0.01   |
| PAH abundance rel. to ISM | f_PAH  | 0.01, 1 |
| Number of chemisorbed layers | N_{core,layer} | 5    |

Notes: χISM = 1 is the ISM Draine UV field. f_PAH = 1 corresponds to a PAH abundance of 3 × 10⁻⁷ (Tielens 2005).

4.2. Hydration timescales

The results of the models to determine the hydration timescales are shown in Figs. 3 and 4. The increase in gas density results in a higher desorption temperature. The hydration timescales δt are below 10⁷ yr and vary with gas density and grain radius roughly as a/n_H₂O at 450 K. The timescale dependence can be derived if one assumes that the hydration timescale is equal to the adsorption timescale

\[
4πa² \frac{2kT}{n_{H₂O} \epsilon(T)n_d} \delta t \approx 4πa²N_{core,layer}n_d. \quad (4)
\]

where \(n_{H₂O}\) is the number density of gas-phase water, \(n_d\) the number density of dust grains, \(\sqrt{\frac{2kT}{m_{H₂O}}}\) is the gas thermal speed, and \(\epsilon(T) \leq 1\) is the efficiency of the gas to overcome the adsorption and core diffusion activation barriers. Using Eq. (3), we obtain for a fixed temperature

\[
\delta t \propto n_{H₂O} \left( \frac{a}{n_{H₂O}} \right). \quad (5)
\]

At \(T > 250\) K, the water gas-phase abundance reaches a maximum steady-state value of a few 10⁻⁴, such that

\[
\delta t \propto n_{H₂O} \left( \frac{a}{n_{H₂O}} \right). \quad (6)
\]

In the high density warm regions such as the inner disk midplane of protoplanetary disks, most of the silicate grains are hydrated up to their maximum possible water intake. The timescale increases with grain size so that the hydration timescale should be compared to that of grain growth.

4.3. Disk model results

H₂ and water abundances of the disk model with 500 core layers are shown in Fig. 5. The figure shows the location of the different water reservoirs in a typical (0.01 M☉) protoplanetary disk (gas-phase, water ice, chemisorbed water at the silicate core surface, chemisorbed water in the core, aka phyllosilicate). The abundance of water in all its possible adsorbed forms is shown in the lower-right panel of Fig. 5.

Hydrogen is almost entirely in molecular form except in the hydrogen core, where it is partially ionized. This is due to the high temperature and the low hydrogen density in the core. The hydrogen abundance decreases with increasing temperature, as expected. The hydration timescale for H₂ is much shorter than that for water, indicating that H₂ is not a viable candidate for hydration.

H₂O is the most abundant molecule in the disk, followed by H₂. The abundance of H₂O increases with increasing temperature, reaching a maximum at about 100 K. The abundance of H₂ decreases with increasing temperature, as expected. The hydration timescale for H₂O is much longer than that for H₂, indicating that H₂O is a more efficient water carrier than H₂.

The abundance of water in the core is much higher than in the disk midplane, as expected. The hydration timescale for water in the core is much longer than that in the disk midplane, indicating that water is more efficiently hydrated in the core.

The abundance of H₂O in the disk midplane is much lower than in the core, as expected. The hydration timescale for H₂O in the disk midplane is much longer than that in the core, indicating that H₂O is less efficiently hydrated in the disk midplane.

The abundance of water in the disk midplane is much higher than in the disk midplane, as expected. The hydration timescale for water in the disk midplane is much longer than that in the disk midplane, indicating that water is more efficiently hydrated in the disk midplane.

4.4. Conclusion

The results of the models to determine the hydration timescales are shown in Figs. 3 and 4. The increase in gas density results in a higher desorption temperature. The hydration timescales δt are below 10⁷ yr and vary with gas density and grain radius roughly as a/n_H₂O at 450 K. The timescale dependence can be derived if one assumes that the hydration timescale is equal to the adsorption timescale

\[
4πa² \frac{2kT}{n_{H₂O} \epsilon(T)n_d} \delta t \approx 4πa²N_{core,layer}n_d. \quad (4)
\]

where \(n_{H₂O}\) is the number density of gas-phase water, \(n_d\) the number density of dust grains, \(\sqrt{\frac{2kT}{m_{H₂O}}}\) is the gas thermal speed, and \(\epsilon(T) \leq 1\) is the efficiency of the gas to overcome the adsorption and core diffusion activation barriers. Using Eq. (3), we obtain for a fixed temperature

\[
\delta t \propto n_{H₂O} \left( \frac{a}{n_{H₂O}} \right). \quad (5)
\]

At \(T > 250\) K, the water gas-phase abundance reaches a maximum steady-state value of a few 10⁻⁴, such that

\[
\delta t \propto n_{H₂O} \left( \frac{a}{n_{H₂O}} \right). \quad (6)
\]

In the high density warm regions such as the inner disk midplane of protoplanetary disks, most of the silicate grains are hydrated up to their maximum possible water intake. The timescale increases with grain size so that the hydration timescale should be compared to that of grain growth.
both water on the silicate surface and in the core are present with the water being trapped in the silicate core only when the dust temperature is high enough for the water molecules to overcome the activation energy for diffusion into the silicate core. Photodesorption is efficient in the inner disk upper atmospheres (see low-right panel of Fig. 5 when $\chi/n > 10^{-7}$) by restricting the lifetime of physisorbed water on the bare grain surface. Therefore, it also limits the efficiency of the transfer of physisorbed water to chemisorption water and subsequently influences the abundance of trapped water in the silicate core. Since the dust temperature even at 0.1 au is lower than 700 K, chemisorbed water can remain on the grains. In the protoplanetary disk mid-plane regions where $T_d$ is between the sublimation of water ice and the efficient formation of phyllosilicate (9–20 au), the amount of water both in the gas and in the solid phases is low, consistent with the zero-dimensional model results (see Fig. 2). However, the abundance of chemisorbed water can reach its maximum possible value for dust grains hotter than ~250 K (Fig. 2).

Similar to the zero-dimensional models chemisorbed water acts as a “poison” limiting the amount of chemisorbed H atoms. The consequence is a lowering of the $H_2$ formation efficiency. The lower $H_2$ formation on silicate grains is however compensated in part by an increase in $H_2$ formation via hydrogenated PAHs and PAH cations (Andrews et al. 2016; Boschman et al. 2015; Mennella et al. 2012). The efficiency of $H_2$ formation through hydrogenated PAHs depends on the actual abundance of PAHs, which is so far not well determined in protoplanetary disks. In our disk model, the PAH abundance is set to be 100 times smaller than in the general interstellar medium ($f_{PAH} = 0.01$). The occupancy of chemisorption sites by water and the drop in $H_2$ formation efficiency is a side effect of the water chemisorption model.

5. Discussion

As seen in both the zero-dimensional and the protoplanetary disk models, hydration of the anhydrous silicate into phyllosilicate is relatively efficient at temperatures above 250–300 K. Phyllosilicates are also the thermodynamically most stable forms of silicates in an oxygen-rich environment at 200–400 K (Woitke et al. 2018) and Appendix D.

Grain growth by coagulation occurs rapidly and millimetre-sized grains are detected in protoplanetary disks (Tazzari et al. 2016). The units making up the large grains can be the 0.1 micron grains considered in the zero-dimensional models. However, compact millimeter-size grains can be hydrated within $10^5$ yr if the gas density is above $10^8$ cm$^{-3}$. Our results suggest that the maximum water storage capacity of silicates is easily reached within 1 Myr for gas hotter than ~250 K for gas densities down to $2 \times 10^4$ cm$^{-3}$.

The speed of the phyllosilicate formation depends strongly on the activation energy. Previous works have used an energy of ~8420 K (Fegley 2000), which is much higher than the current accepted value around 3000 K. Interestingly such value for the activation energy is lower than the desorption energy for physisorbed water. This results in a transfer of water ice to the silicate core surface via tunnelling already at $T_d \sim 50$ K. With an activation energy higher than the desorption energy, phyllosilicate would only be formed through the Eley–Rideal process. Another difference with previous works is that at temperatures $T_d > 100$ K water can still hop a few times from a physisorption site to another physisorption site and find an unoccupied chemisorption site before it desorbs back to the

Table 6. Model parameters, and values for the reference model.

| Quantity | Symbol | Value |
|----------|--------|-------|
| Stellar mass | $M_*$ | 0.7 $M_\odot$ |
| Effective temperature | $T_e$ | 4000 K |
| Stellar luminosity | $L_*$ | 1 $L_\odot$ |
| UV excess | $f_{UV}$ | 0.01 |
| UV powerlaw index | $p_{UV}$ | 1.3 |
| X-ray luminosity | $L_X$ | $10^{30}$ erg s$^{-1}$ |
| X-ray emission temperature | $T_{X,fit}$ | $2 \times 10^7$ K |
| Strength of interstellar UV | $\chi_{ISM}$ | 1 |
| Strength of interstellar IR | $\chi_{IR}$ | 0 |
| Cosmic ray H$_2$ ionisation rate | $\zeta_{CR}$ | $1.7 \times 10^{-17}$ s$^{-1}$ |
| Disk mass | $M_{disk}$ | 0.01 $M_\odot$ |
| Dust/gas mass ratio | $\delta$ | 0.01 |
| Inner disk radius | $R_{in}$ | 0.07 AU |
| Tapering-off radius | $R_{tap}$ | 100 AU |
| Column density power index | $\alpha$ | 1 |
| Reference scale height | $H_g(100$ AU$)$ | 10 AU |
| Flaring power index | $\varepsilon$ | 1.5 |
| Minimum dust particle radius | $a_{min}$ | 0.05 $\mu$m |
| Maximum dust particle radius | $a_{max}$ | 3 mm |
| Dust size dist. power index | $\alpha_{pow}$ | 3.5 |
| Turbulent mixing parameter | $\alpha_{turb}$ | 0.01 |
| Max. hollow volume ratio | $V_{hollow}$ | 80% |
| Dust composition (1) | $M_{PAH}^{0.7}P_{PAH}^{0.3}\text{SiO}_4$ | 60% |
| (Volume fractions) | amorph. carbon | 15% |
| Porosity | 25% |
| PAH abundance rel. to ISM | $f_{PAH}$ | 0.01 |
| Chemical heating efficiency (2) | $\gamma_{chem}$ | 0.2 |
| Distance | $d$ | 140 pc |
| Disk inclination | $i$ | 45$^o$ |
| Photosorption | Yield | $10^{-2}$–$10^{-3}$ |
| Number of chemisorbed layers | $N_{core,layer}$ | 500 |

Notes: (1) The dust composition and porosity best match protoplanetary disk observations of protoplanetary disks, although the porosity of the comet 67P/Churyumov-Gerasimenko is 75–85 per cent in volume (Herique et al. 2016). (2) The chemical heating efficiency $\gamma_{chem}$ is an efficiency by which exothermic chemical reactions are assumed to heat the gas. A detailed discussion on the disk parameters and their effects on the disk thermal and chemical structure can be found in Woitke et al. (2016). The photodesorption yield depends on the species. Special rates are used for water ice (Westley et al. 1995; Öberg et al. 2009) and CO ice (Muñoz Caro et al. 2010). For the other ice species, a standard yield of $10^{-3}$ is assumed.

A detailed discussion on the $H_2$ formation processes is given in Thi et al. (2020).

At the low temperatures found in the outer disk ($T_d < 100$ K), the main water reservoir is physisorbed water (water ice). This is the so-called water ice zone. Photodesorption of physisorbed species is effective in the outer disk when $\chi/n > 10^{-7}$ ($\chi$ being the enhancement compared to the standard interstellar UV field) since we see that the water ice zone does not extend further than ~400 au while the disk outer radius is ~600 au.

Within the inner 10 au where the dust is warmer than ~150 K, water vapour is the main oxygen reservoir. Water vapour is also abundant in the “atmosphere finger” (Woitke et al. 2009b; Du & Bergin 2014). The gas-phase water chemistry can be complex and has been discussed in previous studies (Kamp et al. 2013; Thi & Bik 2005; van Dishoeck et al. 2014). In the inner disk,
Fig. 1. Species abundances as function of time at 50 (top panels), 150 (middle panels), and 250 K (bottom panels). The abundance of unoccupied silicate surface (*) and core (!) sites, of atomic hydrogen in the gas and solid phases, and of H$_2$ are shown in the left panels. Gas-phase water and water ice abundances are shown on the right panels together with gas-phase abundance of atomic oxygen and OH.
The timescales are capped at $10^{20}$ yr.

As an alternative, direct formation of phyllosilicate can also occur by implementation of energetic H$^+$ (Djouadi et al. 2011).

As phyllosilicate does not dehydrate before reaching 600–700 K (and at higher temperatures for dense gas in the inner disk regions), most of it will remain at the planet formation stage of protoplanetary disks. Figure 6 shows the ratio between water trapped into phyllosilicate form and refractory dust mass. One consequence of the fast phyllosilicate formation in contrast to formation by aqueous alteration is that phyllosilicate should be relatively uniformly abundant in partially differentiated planetesimals located in warm protoplanetary disk regions (where $T_g$ and $T_d > 300$ K), if the grain coagulation and growth of the planetesimals are slower than the uptake of water in the grains.

Phyllosilicates can be detected through their OH-stretch fundamental absorption at about 2.7 µm. The infrared signature of phyllosilicate is ubiquitous at the surface of the dwarf planet Ceres (Ammannito et al. 2016) and in a majority of other asteroids (Takir et al. 2015). Phyllosilicates also show distinctive features in the mid-infrared (Glotch et al. 2007; Pitman et al. 2010). Olivine crystal infrared features peak at 11.2 and 19.5 µm, while phyllosilicates have their maximum for the SiO$_4$ stretching at a lower wavelength (around 10 µm) in addition to bands at 8.7 µm, and 5.8 µm in addition to bands at 8.7 µm.
Fig. 5. Steady-state abundance for $H_2$ and the different $H_2O$ carriers for the standard disk model ($H_2O#$ means physisorbed water). The white contours correspond to the gas temperature in the upper right panel. $\chi$ is the strength of the UV field ($\chi = 1$ corresponds to the average local interstellar medium. The water ice location analytical model of Min et al. (2016) is shown (white dashed line). In the same panel the alternative formula from Antonellini et al. (2016) is also displayed (yellow dashed line).
Amorphous olivine peaks at $\sim 15.8$ and 22 $\mu$m due to bending vibration of the hydroxyl group as found in meteorites (Beck et al. 2014; Garenne et al. 2014). Amorphous olivine in meteorites differ in shape compared to terrestrial phyllosilicates. The zodiacal dust mid-infrared spectrum suggests the presence of phyllosilicates (Reach et al. 2003). Morris & Desch (2009) modelled the mid-IR emission of dust grains composed of anhydrous silicates and phyllosilicates in a protoplanetary disk, while Morlok et al. (2014) compared directly the IR spectra of protoplanetary disks with meteoritic samples. Min et al. (2016) modelled in detail the spectral energy distribution of the protoplanetary disk around HD 142527 between 35 and 100 microns and could constrain the amount of phyllosilicates to be lower than $\sim 47\%$ because the phyllosilicates are almost featureless in this wavelength range. Wavelength and shape variations with the level of crystallinity as well as effects of grain size render the identification of phyllosilicates in space IR spectra difficult unless high signal-to-noise ratio (S/N) spectra from the James Webb Space Telescope are available.

Phyllosilicates have been tentatively detected in a debris disk (Currie et al. 2011). Since the dust grains in a debris disk result from collisions between planetesimals, the origin of the phyllosilicates cannot be easily established. Another potential test of the proposed phyllosilicate formation model is to observe that the average abundance of phyllosilicates is anti-correlated with that of water ice (see Fig. 2 and the lower right panel of Fig. 5). The water is either in the water ice or in the silicate core. Objects with a high water ice abundance should not slow comentinually a high phyllosilicate abundance unless there is extensive radial mixing at the early stage of planetesimal formation, which will bring dry silicates and phyllosilicates in the outer disk, where they can be water ice coated (e.g. Raymond & Izidoro 2017).

Chondritic fine-grained phyllosilicates formation can be explained by our process. However, whether all observed phyllosilicate in meteorites and Solar System objects can be explained by our formation method is not clear. For instance the widespread amount found in Solar System objects and of Ceres may be due to actual aqueous alteration at some stage in the object’s geological history. Aqueous alteration should have occurred between 300 and 600 K (Brearley 2006). This temperature is reached by the heat generated by the radioactive decay of the short-lived radionuclides Aluminium-26 (Grimm & McSween 1989). The formation of phyllosilicates can occur both in the pre-dust coagulation phase and in the planetesimal formation and evolution phase.

Given the different possible sources of water on Earth, Izidoro et al. (2013) proposed a compound model for the origin of Earth’s water where they included all the possible sources. The HDO/H$_2$O ratio in Earth’s ocean water has been used as a constraint on the origin of the water on Earth (Caselli & Ceccarelli 2012; Drake 2005). At face value the D to H ratio of the water formed in our model would be below Vienna Standard Mean Ocean Water (VSMOW) and even the values measured for the deep mantle. Thus supply of high D/H material would be required. However, there are some potential mechanisms for D/H enhancements at high temperature that might be active in these zones (Thi et al. 2010) and future work will explore the HDO/H$_2$O (OD/OH) ratio of phyllosilicate grains.

6. Conclusions and perspective

We explored the kinetics of formation of phyllosilicate from anhydrous silicate grains exposed either to water vapour or in contact with a water ice mantle from 10 to 900 K using a warm gas-grain surface chemical model. The phyllosilicate formation was used in a zero-dimensional model and in a more realistic protoplanetary disk model. We showed that different reservoirs of solid water are present in protoplanetary disks. Water can be found in the core of silicate grains (phyllosilicates) within a radius of 10 au. Further away from the star, the temperatures are lower, and water can be frozen onto dust to form a thick mantle. The amount of water in phyllosilicates can reach 2–3% of the total mass of refractory material. The timescale for the formation of phyllosilicates is shorter than the typical lifetime of protoplanetary disks of 2–3 Myr. It proceeds by the adsorption to the silicate core followed by diffusion into the silicate core. The amount of water trapped as phyllosilicate can theoretically reach the maximum water storage capacity of silicates within the disks lifetime. Water can occupy the chemisorption sites required to efficiently form H$_2$. Formation of H$_2$ via hydrogenated PAHs and PAH cations becomes important depending on the abundance of PAHs in the protoplanetary disk. A search for the infrared signatures of phyllosilicates in protoplanetary disks is warranted to test the model, which predicts that a significant amount of the silicates should be hydrated in the warm region (100–700 K). Our modelling supports the scenario, in which water found on Earth may have been already trapped in the dust grains at the phase of planetesimal formation. The amount of phyllosilicates formed by this method may not be efficient enough compared to the widespread amount found in Solar System objects and subsequent aqueous alterations may be still required.

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## Appendix A: Hydrogen surface reactions

Table A.1 lists the grain surface and PAH reactions concerning atomic hydrogen.

### Table A.1. Main grain reactions involved in the formation and destruction of H$_2$.

| Reaction | Comment |
|----------|---------|
| 1 H → H# | Physisorption, barrierless |
| 2 H + * → *H# | $E^{\text{gc}}_H = E^{\text{act}}_* = 900$ K |
| 3 H# + * → *H# | $E^{\text{pc}}_H = E^{\text{act}}_*$ |
| 4 H# → H | $E^{\text{ch}}_{1H#} = 600$ K |
| 5 H# + hν → H | Photodesorption |
| 6 H# + CR → H | |
| 7 *H# → H + * | $E^{\text{ch}}_{\text{H#}} = 10000$ K |
| 8 *H# + hν → H + * | |
| 9 *H# + CR → H + * | Via CR induced UV |
| 10 H + H# → H$_2$ | Eley-Rideal (ER) mechanism, barrierless |
| 11 H# + H# → H$_2$ | $E^{\text{act}}_{\text{H#},\text{H#}} = 0$ K (Navarro-Ruiz et al. 2014) |
| 12 H# + *H# → H$_2$ + * | ER mechanism, barrierless |
| 13 H# + H# → H$_2$ + * | $E^{\text{diss}}_{\text{H#},\text{H#}} = 2 \times E^{\text{act}}_*$ |
| 14 *H# + *H# → H$_2$ + 2* | $E^{\text{diss}}_{\text{H#},\text{H#}} = 3481$ K (Diño et al. 2004) |
| 15 H$_2$ + * → *H# + H# | |
| 16 PAH → PAH# | Physisorption |
| 17 HPAH → HPAH# | |
| 18 PAH# → PAH | Desorption |
| 19 HPAH# → HPAH | |
| 20 PAH# + hν → PAH | Photodesorption |
| 21 HPAH# + hν → HPAH | |
| 22 PAH# + CR → PAH | Cosmic ray induced photodesorption |
| 23 HPAH# + CR → HPAH | |
| 24 H + PAH → HPAH | $E^{\text{act}}_{\text{PAH},\text{H}} = 692$ K (Rauls & Hornekær 2008) |
| 25 HPAH → H + PAH | Thermal H-detachment with $E^{\text{diss}}_{\text{PAH},\text{H}} = 16250$ K |
| 26 HPAH + hν → H + PAH | Photodetachment, $E(C(\text{C-H}) = 4.45$ eV (51640 K) |
| 27 HPAH + H → H$_2$ + PAH | Cross-section $\sigma = 0.06 \text{Å}^2/\text{C atom}, E^{\text{act}} = 0$ K |
| 28 H + PAH* → HPAH* | |
| 29 HPAH* + e → PAH + H | Dissociative recombination |
| 30 HPAH* + H + H$_2$ | Langevin rate (Montillaud et al. 2013) |
| 31 H$_2$ + PAH → HPAH + H | $E^{\text{diss}}_{\text{H#},\text{PAH}} = 3481$ K |
| 32 PAH + hν → PAH* + e | Photoionisation |
| 33 PAH + e → PAH* | Electron attachment |
| 34 PAH* + hν → PAH + e | Photodetachment |
| 35 PAH* + e → PAH | Electron recombination |
| 36 PAH* + X → PAH + X | Charge exchange with species X |
| 37 PAH + X → PAH* + X | |
| 38 H + e$^-$ → H$^-$ + hν | Radiative attachment |
| 39 H + H$^-$ → H$_2$ + e$^-$ | Associative detachment |
| 40 H + H → H$_2$ + H | Three-body reactions |
| 41 H + H$_2$ → H$_2$ + H$_2$ | |
| 42 H$_2$ + hν → H + H | Photodissociation including self-shielding |
| 43 H$_2$ + CR → H + H | By secondary electrons and CR-generated UV photons |

**Notes.** The energies are expressed in units of Kelvin. $E^{\text{gc}}_H$ is the energy barrier for chemisorption of a gas-phase hydrogen atom on the grain surface. $E^{\text{pc}}_H$ is the energy barrier for physisorbed H-atom to chemisorb on the grain surface. $E^{\text{act}}_*$ is the adsorption energy of a physisorbed H-atom. We assumed no activation barrier for H$_2$ recombination between two physisorbed H-atoms ($E^{\text{act}}_{\text{H#},\text{H#}} = 0$ K). $E^{\text{diss}}_{\text{PAH},\text{H}}$ and $E^{\text{act}}_{\text{H#},\text{H#}}$ are activation energy for H$_2$ recombination involving one or two chemisorbed H-atom(s). $E^{\text{diss}}_{\text{H#},\text{PAH}}$ is the H$_2$ dissociative adsorption. * and ! are pseudo-species as defined in Table 1.
Appendix B: Disk model extra results

Figure B.1 shows extra results from the disk model.

**Fig. B.1.** \textit{Top two rows:} disk distribution of various quantities: disk gas number density \((n_H)\), dust mass density \(\rho_{\text{dust}}\), UV field strength in units of \(\chi\) \((\chi = 1\) is the average local interstellar medium radiation field\) over-plotted by contours showing the dust extinction in the optical, and the dust temperature. \textit{Lower panels:} gas and dust temperature distribution in the disk.
Appendix C: Zero dimensional model at 10 K

Figure C.1 shows results from the zero dimensional model at 10 K.

\[ \text{rel. abundance } X/H \]

\[ \text{years} \]

\[ \times 10^{-2} \times 10^{0} \times 10^{-4} \times 10^{-6} \times 10^{-8} \]

\[ \times 10^{2} \times 10^{4} \times 10^{6} \times 10^{8} \]

**Fig. C.1.** Species abundances as function of time at 10 K. *Left panel:* abundance of unoccupied silicate surface (*) and core (!) sites, of atomic hydrogen in the gas and solid phases, and of H\(_2\). *Right panel:* gas-phase water and water ice abundances together with gas-phase abundance of atomic oxygen and OH.

Appendix D: Lizardite abundance from chemical equilibrium calculation

As an illustration that phyllosilicate is the most thermodynamically stable species in the inner disk midplane, we show the abundance of Lizardite Mg\(_3\)Si\(_2\)O\(_9\)H\(_4\), the hydrated form of Olivine in Fig. D.1. The abundance has been computed with the chemical equilibrium code GGCHEM, which is called by P\(_{\text{RodMo}}\) when the temperature is above 100 K (Woitke et al. 2018).

**Fig. D.1.** Lizardite Mg\(_3\)Si\(_2\)O\(_9\)H\(_4\) abundance in the inner disk as computed with the chemical equilibrium solver GGCHEM (Woitke et al. 2018).