The chemistry of protoplanetary fragments formed via gravitational instabilities

J. D. Ilee,1* D. H. Forgan,2,3 M. G. Evans,4 C. Hall,5,6 R. Booth,1 C. J. Clarke,1 W. K. M. Rice,5,7 A. C. Boley,8 P. Caselli,9 T. W. Hartquist4 and J. M. C. Rawlings10

1 Institute of Astronomy, University of Cambridge, Madingley Road, Cambridge CB3 0HA, UK
2 SUPA, School of Physics & Astronomy, University of St Andrews, North Haugh, St Andrews, Scotland KY16 9SS, UK
3 Centre for Exoplanet Science, University of St Andrews, St Andrews, UK
4 School of Physics and Astronomy, University of Leeds, Leeds LS2 9JT, UK
5 SUPA, Institute for Astronomy, University of Edinburgh, Blackford Hill, Edinburgh EH9 3HJ, UK
6 Department of Physics & Astronomy, University of Leicester, University Road, Leicester LE1 7RH, UK
7 Centre for Exoplanet Science, University of Edinburgh, Edinburgh, UK
8 Department of Physics and Astronomy, The University of British Columbia, 6224 Agricultural Road, Vancouver V6T 1Z4, Canada
9 Max-Planck Institute for Extraterrestrial Physics, Giessenbachstrasse 1, D-85748 Garching, Germany
10 Department of Physics and Astronomy, University College London, Gower Street, London WC1E 6BT, UK

Accepted 2017 July 28. Received 2017 July 27; in original form 2017 June 23

ABSTRACT

In this paper, we model the chemical evolution of a 0.25 M⊙ protoplanetary disc surrounding a 1 M⊙ star that undergoes fragmentation due to self-gravity. We use smoothed particle hydrodynamics including a radiative transfer scheme, along with a time-dependent chemical evolution code to follow the composition of the disc and resulting fragments over approximately 4000 yr. Initially, four quasi-stable fragments are formed, of which two are eventually disrupted by tidal torques in the disc. From the results of our chemical modelling, we identify species that are abundant in the fragments (e.g. H2O, H2S, HNO, N2, NH3, OCS, SO), species that are abundant in the spiral shocks within the disc (e.g. CO, CH4, CN, CS, H2CO) and species that are abundant in the circumfragmentary material (e.g. HCO+). Our models suggest that in some fragments it is plausible for grains to sediment to the core before releasing their volatiles into the planetary envelope, leading to changes in, e.g., the C/O ratio of the gas and ice components. We would therefore predict that the atmospheric composition of planets generated by gravitational instability should not necessarily follow the bulk chemical composition of the local disc material.

Key words: astrochemistry – hydrodynamics – planets and satellites: composition – planets and satellites: formation – protoplanetary discs.

1 INTRODUCTION

The star formation process results in the production of circumstellar discs around young protostars. These discs provide further feedstock for the star to continue accreting material, but they are also the sites of planet formation. Planet formation models can be split into two camps: the bottom-up process of core accretion (CA, Pollack et al. 1996), which relies on the coagulation of dust grains into larger bodies, or via the top down process of disc fragmentation via the gravitational instability (GI; Durisen et al. 2007; Helled et al. 2014).

At the birth of the star–disc system, it is expected that the disc mass is approximately equal to the mass of the star, and observations constrain the disc-to-star mass ratios to be no less than 0.5 (e.g. Eisner & Carpenter 2006; Joos, Hennebelle & Ciardi 2012). At such high disc-to-star mass ratios, the evolution of the system is likely to be dominated by disc self-gravity. These discs will exist in a state of marginal instability if the local Toomre Parameter (Toomre 1964),

\[ Q = \frac{c_s\kappa}{\pi G \Sigma} \sim 1, \]

where \( \kappa \) is the epicyclic frequency (representing the rotational support, and equal to the angular velocity \( \Omega \) in a Keplerian disc), \( c_s \) is the sound speed of the gas (representing the pressure support) and \( \Sigma \) is the surface density of the gas. The combined action of thermodynamics, self-gravity and rotation can result in self-regulated, quasi-steady discs where \( Q \sim 1.5\text{--}1.7 \), which is approximately the critical value for non-axisymmetric instabilities (Durisen

* E-mail: jdilee@ast.cam.ac.uk

© 2017 The Authors
Published by Oxford University Press on behalf of the Royal Astronomical Society

Downloaded from https://academic.oup.com/mnras/article-abstract/472/1/189/4079328 by University College London user on 29 January 2018
et al. 2007). Such discs possess large-scale spiral density waves, and these (combined with the gravito-turbulence produced by their non-linear coupling; Gammie 2001) are efficient outward transporters of angular momentum, and can provide a mechanism for driving mass accretion during the earliest stages of star formation. As a result, the disc material accretes rapidly on to the central star, in a process that can last for approximately $10^5$ yr (Lodato & Rice 2004; Rice, Mayo & Armitage 2010).

Self-gravitating discs may also undergo fragmentation if $Q \sim 1$ and the surface density perturbations induced by the GI are massive enough, and their cooling is efficient enough such that they become gravitationally self-bound and collapse (Gammie 2001; Rice, Lodato & Armitage 2005; Meru & Bate 2011; Paardekooper 2012; Young & Clarke 2016). The process of disc fragmentation has largely been the province of theory, but several works have shown that the Atacama Large Millimetre Array has both the high angular resolution and sensitivity to directly detect gravitationally unstable discs in the nearest star-forming regions (e.g. Cossins, Lodato & Clarke 2010; Douglas et al. 2013; Dipierro et al. 2014, 2015; Hall et al. 2016; Evans et al. 2017). Indeed, recent observational campaigns are beginning to reveal spiral arm structures at sub-millimetre wavelengths (Pérez et al. 2016), a hallmark of GI or recent fragmentation (Meru et al. 2017). Most strikingly, evidence of a fragmenting disc was gathered by Tobin et al. (2016), showing a star-disc system with two stellar mass objects in the disc connected by spiral structure. This system demonstrates that GI and disc fragmentation can form stellar mass objects. But can this process form planetary mass objects?

A standing assumption is that, in contrast to protoplanets formed via CA, a fragment formed via GI would inherit an initial composition that is likely to reflect the bulk stellar or bulk disc composition. However, Boley & Durisen (2010) and Boley, Helled & Payne (2011) discuss several situations for which this assumption may not hold. First, if a population of grains is present at the birth of the disc, then grains with stopping distances comparable to the widths of the spiral arms will be efficiently collected in the arms prior to fragmentation. Subsequent fragmentation of this arm would result in fragments with a super-stellar metallicity. Secondly, if for some reason a population of planetesimals has formed prior to fragmentation, then the gas will be inefficient at damping their motion. This effect will persist, even within a fragment, until temperatures reach the $\text{H}_2$ dissociation limit. Thus, the planetesimal concentration will be lower than the fragment concentration for some time, leading to sub-stellar metallicities. Thirdly, within a fragment itself, the initial sample of dust grains will begin to collide and grow in a very similar fashion to grain growth as predicted by CA, as well as accreting further solid material as it traverses through the circumstellar disc (Helled, Podolak & Koveç 2006). If grains can grow to sufficiently large sizes, they will sediment towards the pressure maximum at the centre of the fragment, and in the right circumstances can form a solid core. Finally, planet formation depletes metals from the gas phase. If this metal-depleted gas continues to accrete on to the central star, then the stellar atmospheric composition may diverge from the bulk composition of the initial disc, leading to sub-stellar metallicities. Such depletions are seen towards stars exhibiting the $\lambda$-Booitis phenomenon (e.g. Heiter 2002).

The combined action of these processes results in a much wider variety of objects formed via GI than was previously considered, and has resulted in a revamped model of fragmentation usually referred to as ‘tidal downsizing’ (Boley et al. 2010; Nayakshin 2010a). It has been suggested that giant planets form with a variety of core masses under this paradigm (Nayakshin & Fletcher 2015) and that the typical core mass is larger than previously predicted (e.g. Saumon & Guillot 2004). However, population synthesis models indicate that the formation of terrestrial planets via solid core formation and disruption of the outer gaseous envelope is an unlikely outcome (Forgan & Rice 2013).

The majority of GI objects appear to be giant planets and brown dwarfs, both bound to stars and free-floating (Forgan, Parker & Rice 2015). Observational constraints (Vigan et al. 2017) suggest that such objects are rare compared to GI predictions, indicating that the proportion of exoplanets that are GI objects is potentially small. However, there are examples of planetary systems that include giant objects on wide orbits (e.g. HR 8799; Marois et al. 2008), or systems which are undergoing or might have undergone fragmentation (e.g. Elias 2-27; Meru et al. 2017), suggesting it is possible for GI to form some planetary systems. In order to provide further observational predictions that can be used to examine the properties of these planets, and the discs that formed them, it is important to consider the chemical composition of both.

Chemical evolution in protoplanetary discs has been extensively studied (see the reviews by, e.g., Bergin et al. 2007; Henning & Semenov 2013; Dutrey et al. 2014). Many of the models to date have focused on immediately observable characteristics, such as species responsible for emission lines formed at high relative disc heights. However, there has recently been more interest in evaluating the chemical properties of the bulk of the planet building material in the midplane (e.g. Eistrup, Walsh & van Dishoeck 2016). Unfortunately, due to the computationally demanding nature of (i) evolving large chemical networks of many thousands of species and/or (ii) calculating the mutual influences of chemistry on the thermal evolution of the disc, these models are at most two-dimensional in nature (see the review of Haworth et al. 2016). Such a set-up limits their ability to investigate the chemical evolution of structures that are non-axisymmetric in nature. There are a handful of studies concerned with investigating chemical evolution in three-dimensional disc models – e.g. young, massive discs with non-axisymmetric structure (Ilee et al. 2011; Hincelin et al. 2013; Evans et al. 2015). However, these studies either included discs that did not fragment, or did not follow the physical evolution for a sufficiently long time to characterize any fragments that may be formed within the discs.

In this paper, we use a smoothed particle hydrodynamics (SPH) model with a hybrid radiative transfer scheme along with non-equilibrium gas-grain chemical evolution code to model the evolution of a fragmenting, self-gravitating disc. We characterize both the effect of the dynamical evolution on the chemical evolution of the disc and the chemical properties of the resulting fragments themselves. Section 2 outlines our physical and chemical modelling procedures, Section 3 presents our results, Section 4 discusses the implications of our work and finally our conclusions are presented in Section 5.

2 MODELLING APPROACH

2.1 Hydrodynamics

We use the SPH code sphNG (Bate, Bonnell & Price 1995), which was modified to include the hybrid radiative transfer formalism of Forgan et al. (2009). The gas is able to cool radiatively according to its local optical depth, which is estimated from the local gravitational potential (Stamatellos et al. 2007) and is also able to exchange energy with neighbouring fluid elements via flux-limited diffusion (see e.g. Bodenheimer et al. 1990; Whitehouse & Bate 2004; Mayer
et al. 2007). This approximation allows high-resolution simulations to be run with a minimal computational penalty. Tests show this algorithm increases runtime by only 6 per cent over non-radiative simulations, while still passing key tests of grey radiative transfer algorithms.

We simulate a 0.25 M⊙ disc, orbiting a 1 M⊙ star. The gas in the disc is represented by 4 × 10⁶ SPH particles distributed between 10 and 100 au, with a surface density profile that scales as Σ(r) ∝ r⁻¹ and a sound speed profile that scales as c_s ∝ r⁻⁰.⁵. The initial conditions are such that Q decreases with radius so that the value at the outer edge of the disc is approximately 1. Once the simulation begins, Q evolves to an approximately constant value of 1 between 50 and 90 au, before fragmentation ensues. These parameters are consistent with observations of disc-like structures in Keplerian rotation around Class 0 objects (e.g. Tobin et al. 2015), and the disc-star mass ratio is consistent with that expected after the initial rapid accretion of disc material before the envelope dissipates (Forgan et al. 2011).

The star is represented by a sink particle that can accrete gas particles if they approach sufficiently closely (within 1 au) and are gravitationally bound (Bate et al. 1995). The creation of additional sink particles is not allowed in the model. This enables us to follow the formation and evolution of any fragments, but in turn limits the maximum time for which the simulation can be run (t = 4100 yr), since beyond this point, densities within the fragments become too large to follow without an unrealistically small hydrodynamic time-step.

2.2 Chemistry

In order to calculate the chemistry of the hydrodynamic model, we adopt the approach developed in Ilee et al. (2011), including the improvements implemented in Evans et al. (2015), which we also describe here. The chemical network consists of 125 chemical species and 1334 reactions. The reactions were originally selected from the UMIST95 data base (Millar, Farquhar & Willacy 1997), but their rate coefficients were updated using data from the Kinetic Database for Astrochemistry (KIDA1; Wakelam et al. 2012). Various gas phase chemical processes are considered, including positive ion-neutral reactions, ionization by cosmic rays, charge transfer, proton transfer, hydrogen abstraction, radiative and dissociative recombination with electrons, radiative association, neutral exchange, photodissociation and photoionization.

Due to the unknown UV properties and embedded nature of young Class 0 sources, we assume that the disc is well shielded from sources of external irradiation (setting A_V = 100) but allow photochemical reactions induced by cosmic rays, the ionization rate of which is assumed to be ĉ = 10⁻¹⁷ s⁻¹ throughout the disc. In addition, several gas-grain processes are implemented, including adsorption, desorption, recombinations of ions with negative grains and simple grain surface chemistry (e.g. the rapid hydrogenation of adsorbed species). The chemical model assumes a constant gas-to-dust ratio of 100, and a uniform distribution of grains with radius 0.1 μm.

While grain surface chemistry likely plays an important role in the evolution of many chemical species, the chemical pathways are uncertain and computationally expensive to implement. In order to simulate any further potential effects of grain surface reactions, our initial gas phase species abundances are taken to be representative of observations of cometary ice abundances (see Ehrenfreund & Charnley 2000), given in Table 1. While there is some consistency between cometary and interstellar ices, it is not known whether comets undergo significant chemical processing, and thus, whether they accurately preserve the initial elemental compositions of young systems is not clear (see e.g. Caselli & Ceccarelli 2012).

All SPH particles from the full hydrodynamic simulation (4 × 10⁶ in total) are used as physical inputs for the chemical model, providing temperatures and densities as a function of space and time. The chemical model interpolates these quantities (and other parameters relevant for the full output of the physical model) on to time-scales that are more appropriate for the chemistry. This approach ensures that the chemical evolution time-steps are not fixed to the time-steps used in the physical model, allowing very rapid chemical processes to be followed with sufficient temporal resolution.

The DVODE integrator (Brown, Byrne & Hindmarsh 1989) is used to integrate the chemical rate equations, yielding fractional abundances for each species, e.g. X_i = n_i/n, where n = n_H + n_H₂ + n_CO₂. In order to improve the success of the integrations, we adopt the recursive time-step algorithm implemented in Evans et al. (2015). Integrations that are initially unsuccessful are re-attempted by continually halving the time-step until either the integration is successful or the time-step becomes so small that integration is not possible. Using this approach, we were able to achieve very high success rate, with less than 0.3 per cent of tracer particles failing to complete integration. These failed particles included very rapid jumps in temperature or density, but were evenly distributed across the simulation and so their exclusion had a negligible effect on the final chemical abundances throughout the disc. In order to visualize the results of the chemical modelling of the particles, we employ a mapping and interpolation method similar to that of Price (2007) to display abundances and column densities.

3 RESULTS

3.1 Global evolution of the disc

Fig. 1 displays several snapshots of the disc evolution. After approximately 1300 yr of evolution, corresponding to approximately

---

1 http://kida.obs.u-bordeaux1.fr/
Figure 1. The evolution of the hydrodynamic simulation. The snapshots cover approximately 4000 yr of disc evolution, through fragmentation into a number of objects, of which four are able to resist shearing and spiral arm shocks, which we label John, Paul, George and Ringo in the middle left-hand panel. John and Paul are tidally disrupted (middle right-hand and bottom left-hand panels), and a fifth fragment appears to be formed via triggering in the final snapshot (bottom left).

1.5 outer rotation periods (ORPs), the disc fragments into a large number of objects, with masses of order 0.5–4 $M_{\text{Jup}}$. Most of these objects are transient, being quickly sheared out by the disc. Of the eight or so seen in the upper left-hand panel of Fig. 1, only four remain on relatively long time-scales, which we identify using clump finding algorithms (Hall, Forgan & Rice 2017). We label them (in ascending distance from the protostar) — John, Paul, George and Ringo — in the middle left-hand panel.²

² Our simulation is identical to that of ‘Simulation 2’ described in Hall et al. (2017), in which the fragments are named Clump 5, 3, 4 and 2 in increasing radial distance from the central star, respectively.
Of these four, John and Paul migrate inwards and are tidally disrupted at \( t \approx 3100 \text{ yr} \) and \( t \approx 3500 \text{ yr} \), respectively. George and Ringo survive until the end of the simulation run at \( t = 4100 \text{ yr} \). George migrates inwards and accretes a significant amount of inner disc material, finally approaching the initial radial distance attained by Paul when the four fragments are first identified. Ringo orbits in relatively low-density surroundings, and as such only accretes material within a few au. Ringo ends the simulation at a greater distance than where it began, due to dynamical interactions with George and the intervening spiral structure.

It is important to note that the fate of George is not assured here. While this fragment has had considerably longer to become fully gravitationally bound (compared to John and Paul), it may well be the case that it still migrates inwards towards tidal disruption, perhaps achieving a closer distance to the star before being destroyed. On the other hand, Ringo appears to be safely ensconced at large distances, and it may be joined by a fifth fragment in the beginnings of formation at \( \approx 87 \text{ au} \) when the simulation is ended (see the final panel of Fig. 1). Ideally, we would continue the simulation further to track the fifth fragment, but the densities reached by the other fragments would require us to allow sink particle creation, which would obstruct our efforts to track both fragment structure and chemistry.

Fig. 2 displays the disc mid-plane temperature at \( t = 2852 \text{ yr} \) and \( t = 4100 \text{ yr} \), corresponding to the third and final panels of Fig. 1. Globally, the disc retains the initial radially decreasing temperature profile that is expected from less massive discs. However, significant deviations from this profile are seen towards the spiral arms and the fragments, where extra heating is caused by compressive forces and weak shocks.

### 3.2 Physical conditions within the fragments

We identify the population of particles that constitutes the four fragments by considering a simulation snapshot taken at \( t = 2852 \text{ yr} \) and collecting all particles that reside within 1 au of the central density peak. The initial locations of the density peak and total particle counts can be found in Table 2. The fragment locations are consistent with the results derived from more advanced clump finding algorithms that have been used on this data set in other work (see Hall et al. 2017), which we use to report the fragment masses in Table 2. Our approach is likely to be an undersampling of the material bound to each fragment – the typical Hill radii of these bodies is around 5 au or larger, and 95 per cent of the mass of George and Ringo is contained within approximately 2.5 au at the end of the simulation. However, this approach ensures that all material for which we study the chemical composition is certain to remain inside the fragment after formation, and as such will not include any material that is only incorporated into a fragment for a short time.

Table 2. The position of the fragments at \( t = 2851 \text{ yr} \), along with the number of particles \( N \) within a cylindrical radius \( < 1 \text{ au} \) of the central density peak at 2852 yr. These particles are subsequently used to sample the physical and chemical conditions of the fragments. Also given are the total masses of the fragments \( M \) at \( t = 2852 \text{ yr} \) calculated from Hall et al. (2017), and the formation \( (t_f) \) and destruction \( (t_d) \) times for each fragment.

| Position (au) | \( N \) | \( M \) (\( M_{\text{Jup}} \)) | \( t_f \) (yr) | \( t_d \) (yr) |
|--------------|---|----------------|------------|------------|
| John         | +42.9 | +10.3 | -0.2 | 27033 | 10.3 | 1935 | 3106 |
| Paul         | +22.3 | -55.0 | +0.0 | 13424 | 3.7 | 1528 | 3514 |
| George       | +90.2 | -40.6 | -0.1 | 29603 | 8.2 | 1935 | - |
| Ringo        | -99.2 | -16.4 | +0.1 | 16902 | 5.4 | 2342 | - |

### 3.3 Chemical evolution of the disc and fragments

Fig. 4 shows the column density for selected chemical species that we have modelled (plots for all species in our network can be found in Appendix B). From examination of the temporal evolution of our simulation, we split the chemical species into morphological categories – those that are primarily abundant towards the fragments (e.g. \( \text{H}_2\text{O}, \text{H}_2\text{S}, \text{HNO}, \text{N}_2, \text{NH}_3, \text{OCS}, \text{SO} \)) and those that are also abundant in the spiral shocks within the disc (e.g. \( \text{CO}, \text{CH}_4, \text{CN}, \text{CS}, \text{H}_2\text{CO} \)). Based on representative particles that reside within 1 au of the fragments, we can examine the creation and destruction routes of various species. Following the chemical evolution of an SPH particle that becomes incorporated within George at \( t = 2750 \text{ yr} \), we can see that CN is primarily formed via secondary photodissociation of HCN and HNC, and destroyed by reactions with \( \text{NH}_3 \) forming \( \text{NH}_2 \) and \( \text{HCl} \). The disc of \( \text{H}_2\text{O} \) has many formation routes, but is most significantly formed by the reaction of \( \text{H}_3\text{O}^+ \) and \( \text{NH}_3 \) and removed from the gas phase via adsorption. HCO\(^+\) is primarily created by reactions of \( \text{H}_2^+ \) with CO. It is readily destroyed when \( \text{H}_2\text{O} \) exists in significant quantities in the gas phase, forming \( \text{H}_2\text{O}^+ \) and CO. Adsorption and desorption primarily determine the gas phase abundances of \( \text{CO}, \text{CO}_2, \text{CS}, \text{HCN}, \text{H}_2\text{CO}, \text{HNO}, \text{NH}_3, \text{OCN} \) and OCS. The timescales for the adsorption and desorption range from a few seconds to fractions of a year, depending on the temperature and density of the ambient material. As in Ilee et al. (2011) and Evans et al. (2015), this is many orders of magnitude smaller than the local dynamical
Figure 2. Mid-plane temperatures of the disc simulation at $t = 2852$ (left) and 3913 yr (right), corresponding to the third and sixth panels Fig. 1. Globally, the disc exhibits an expected radially-decreasing temperature gradient, but significant increases in temperature are seen around the fragments and spiral arms.

Figure 3. Temperature (top panels) and number density (bottom panels) of the four fragments calculated from their constituent particles (see Table 2). Mean values are shown with a solid line, while the shaded region indicates the standard deviation. The dashed vertical lines indicate the formation, and where appropriate, the destruction times for each fragment. All fragments experience increasing temperatures and densities during formation, but the material that constituted John and Paul cools and rarefies rapidly after tidal disruption at approximately 3000 and 3500 yr, respectively.
Within the regions of the disc that have not undergone fragmentation, we identify two broad categories of molecules. First, there are those molecules in which the fractional abundance is primarily determined by adsorption and desorption processes (e.g. CO, CO$_2$, SO, SO$_2$, NH$_3$, H$_2$O, H$_2$CO, H$_2$S, HCN, HNC and OCS). The abundances of these molecules are therefore almost entirely set by temperature changes within the disc. Secondly, there are molecules that are significantly affected by gas-phase reactions occurring in warm, shocked regions (e.g. CS, HCS, HCS$^+$, HCO$^+$, HNO, OCN, O$_2$ and CN). The higher temperatures experienced as the shocks progress through the disc allow some reactions with high activation energies to proceed at significantly enhanced rates. Examples of molecules produced by these effects include OCN that is formed by a reaction of CO and N$_2$; HNO that is formed by NH$_2$ and O; and HCS that is formed by reactions of S and CH$_2$. We note that our choice of initial gas-phase abundances (see Table 1) likely sets the molecules whose abundance is primarily determined by adsorption and desorption, and changing these initial abundances would impact the molecules that reside within this category.

Using the collections of particles from Table 2, we are able to follow the average chemical evolution of the material that constitutes each fragment before, during and (where appropriate) after the fragment has been disrupted. Fig. 6 shows the average fractional abundance of various species (CO, H$_2$O, CN, HCO$^+$, HCS) for each of the fragments. Because of the rapid temperature increase in almost all of the material that becomes a fragment, CO is quickly liberated from ice mantles and remains in the gas phase. H$_2$O exhibits very similar evolution within John, George and Ringo – mostly located on the grains even as the fragments are fully formed initially, but then quickly liberated by thermal desorption as the fragment temperatures increase. The exception to this behaviour is Paul, which does not reach sufficiently high temperatures for the H$_2$O abundance to reach peak values before undergoing tidal disruption. After disruption in John and Paul, the H$_2$O abundance quickly drops again as the molecule freezes out. The abundance of CN is initially high in all fragments, but this quickly drops when temperatures rise above $\sim$150 K due to reactions with thermally desorbed NH$_3$ creating HCN and NH$_2$. The HCO$^+$ abundance in all fragments is low due to reactions with the ever increasing amounts of H$_2$O in the gas phase, but is boosted as John and Paul undergo disruption as material cools and the H$_2$O begins to freeze out. The abundance of HCS remains relatively constant throughout the lifetimes of the fragments, with only a small drop during the disruption for John and Paul.
The spherical averaging discussed in Section 3.2 also allows us to examine radial variations of the selected species within each fragment, shown in Appendix A (Fig. A2). At the time of classification ($t = 2852 \text{ yr}$), John and George exhibit very small radial differences in the fractional abundance of CO, H$_2$O, CN, HCO$^+$ and HCS due to their relatively high temperatures. Paul and Ringo exhibit a more varied chemical structure, in particular showing a lower peak abundance of H$_2$O, that then decreases with increasing distance from the fragment centre. Paul also shows an increasing abundance of CN at larger radii. For the fragments that survive to the end of the simulation, any initial radial variations in these species seem to have mostly been eradicated by the increasing temperatures and densities.

It is important to note that the temperatures within George rise outside the range of temperatures for which our chemical network is likely valid. Determining chemical abundances for these temperatures ($T > 300$–$500 \text{ K}$) requires dedicated high temperature reactions, which are not included within our network, and as such the specific abundances reported for George after approximately 3000 $\text{ yr}$ should be treated with caution.

4 DISCUSSION

4.1 Comparison with non-fragmenting discs

Direct comparison of our model with other, non-fragmenting discs that are subject to gravitational instabilities is challenging due to the differing assumptions between the chemical networks, reaction rates and underlying hydrodynamic models used. Nevertheless, here we outline some broad comparisons with previous work that characterizes the chemical evolution of such discs.

Perhaps the closest simulation in terms of a disc-to-star mass ratio is that of Evans et al. (2015) in which $q \sim 0.2$ (compared to our $q = 0.25$). This disc is much smaller in radial extent than the disc studied here, and therefore does not undergo fragmentation during the 2000 $\text{ yr}$ for which it is followed due to the higher ratio of cooling time to dynamical time at smaller radii. The chemical network used in Evans et al. (2015) is identical to the one used in this work, and therefore comparisons made between their results and the results presented here are free from any differing assumptions on the nature of the chemistry. When comparing CO, HCN, H$_2$O and HCO$^+$ column density maps from Fig. 4 with the results from Evans et al. (2015), it is clear that the general locations of maxima and minima are very similar if the fragments are disregarded. In particular, the CO appears to trace the spiral arms well, the HCN slightly less so, the H$_2$O in confined to the hottest regions and the HCO$^+$ is most abundant in the outer regions of both discs. Interestingly, the extent of the CO emission in our fragmenting disc model and in the model of Evans et al. (2015) is similar. This suggests that the apparent size of discs in molecular line emission may be similar even for discs that initially possess very different radial extents, due to the location of the molecular condensation front (or snow line) that is tied to the thermal structure of the disc. Evans et al. (2015) also note that the persistent shock heating can lead to permanent changes in the chemical composition of inner disc material. Species affected include HNO, CN and NH$_3$. These effects are also likely to occur in our disc, but we are unable to characterize them due to the lower temporal resolution over which we record the physical information of our hydrodynamic model. Such a lower resolution was essential in order to be able to evaluate the chemistry of the $4 \times 10^9$ particles that comprise our disc simulation.

Hincelin et al. (2013) studied the chemical evolution of a collapsing cloud that forms a gravitationally unstable disc around the first Larson core using radiation-magneto-hydrodynamics and a gas-grain chemical network. While column density plots for their model are not presented, the fractional abundances of the CO and H$_2$O ices are shown. Contrary to our results, the disc in Hincelin et al. (2013) does not seem to reach the temperatures required in the spiral arms to desorb significant amounts of either molecule into the gas phase, leading to large abundances of their ices in the spiral arms.

4.2 Molecular condensation fronts (a.k.a. ‘snow lines’)

The locations of the condensation fronts at which molecular species transition from the gas phase to ices on grain surfaces, so-called snow lines, are important because the properties of the dust grains on either side may be drastically different (see e.g. Qi et al. 2013; Banzatti et al. 2015; Cieza et al. 2016; Eistrup et al. 2016; Panić & Min 2017). Such differences lead to changes in the raw ingredients available for planet formation via core accretion and associated processes that involve the agglomeration of small particles (e.g. Öberg, Murray-Clay & Bergin 2011; Madhusudhan 2012; Booth et al. 2017). In quiescent discs, these locations are often quoted as a single radial location corresponding to where the mid-plane temperature drops below the temperature at which species begin to rapidly freeze-out (see e.g. Khajenabi, Kazrani & Shadmeheri 2017). However, in discs that undergo significant non-axisymmetric temperature and density increases, these snow lines may appear in multiple locations, generally following the regions at which the temperature corresponds to the freeze-out temperature for the molecule of interest. In particular, we note that in our model, these locations trace a three-dimensional structure across the $x$, $y$ and $z$ directions. Therefore, in reality, a more appropriate term to describe the phenomenon in such discs might be a snow bubble. However, due to the relatively low vertical resolution in our disc, we restrict our description of the condensation fronts to the radii at which the gas-phase column density of the species in question is equal to that of the column density of the corresponding ice-phase species, i.e. $N_{\text{gas}} = N_{\text{ice}}$.

Fig. 5 shows the evolution of the column density of the disc overlaid with contours showing the location of the mid-plane ‘snow lines’ for CO, CO$_2$ and H$_2$O. Early in the simulation, before the onset of global instabilities, the disc experiences the expected pattern of concentric circular snow lines. The H$_2$O snow line is closest to the central star, followed by the CO$_2$ and finally the CO snow lines – the ordering determined by their decreasing binding energy to icy grain surfaces (see Table 1 and Evans et al. 2015, their table A1). However, once the instabilities develop, the temperature structure of the disc is altered significantly by shock heating, the formation of fragments and compression under gravity. The highest temperatures are seen towards the fragments and in the spiral arms. In these regions, significant increases in the gas-phase abundances of molecules can be seen (e.g. Fig. 5, right-hand panel). These regions would, without the instabilities, lie in a temperature and density regime in which these molecules would be frozen out.

Observationally, this would manifest itself as an apparent increase in the size of the emitting region of these molecules. While in this particular disc the radial increase is relatively small (moving from approximately 40–60 au), discs which do not undergo fragmentation can exhibit increases that are much larger (e.g. Ilee et al. 2011; Evans et al. 2017). Indeed, such increases in gas-phase abundances of volatile molecules caused by dynamic effects may have already been seen towards young embedded discs in the Perseus.
molecular cloud (Frimann et al. 2017). Similar effects have been seen in other models (e.g. Cleeves, Bergin & Harries 2015); however, these have considered the temperature changes induced in the disc by the radiative output of an already-formed planet, rather than the dynamic effects occurring during the formation stages that we model here.

4.3 The C/O ratio of the dust and gas components

The total ratio of carbon to oxygen (C/O) can be an important quantity in characterizing the properties of a planetary atmosphere, in particular where in a protoplanetary disc the planet may have formed (Öberg et al. 2011; Madhusudhan 2012; Booth et al. 2017). Using
Figure 7. Average C/O ratio for each fragment (John, Paul, George, Ringo, clockwise from top left) including species in the gas phase (red) and on the grain surfaces (blue). The dashed vertical lines indicate the formation and, where appropriate, the destruction times for each fragment, and the horizontal dash-dot line indicates the initial C/O ratio of 0.26.

the collections of particles in Table 2, we are able to calculate the bulk average [C/O] for each of the fragments as a function of time for both the gas and ice phase (see Fig. 7). Globally, the general [C/O] ratio evolution can be understood as follows. With our initial chemical abundances, the initial ratio of [C/O]_gas is 0.26 and undefined for [C/O]_ice due to all species initially being in the gas phase. Particles almost immediately diverge from this initial value, with [C/O]_gas rising to unity, as CO remains in the gas phase while other species with higher binding energies freeze out very efficiently on to the surfaces of dust grains (raising the [C/O]_ice to almost the initial gaseous value). As temperatures begin to increase, CO₂ is desorbed from the grain surfaces leading to a drop in [C/O]_gas at the formation stages of each of the fragments. Further increases in temperature cause H₂O to desorb from the grain surfaces, returning the [C/O]_gas back to the initial value of 0.26, finally leaving little-to-no C or O on the grain surfaces in material that continues to experience high temperatures. For material that experiences a drop in temperatures, the reverse process occurs. Thus, the final composition of the material that goes on to make up a fragment eventually preserves the initial elemental ratio.

However, because we are able to follow the temporal evolution of the [C/O] in our model, we observe interesting transient features. During the early formation stages of each fragment, the average temperatures are not high enough to entirely desorb all species from the grain surfaces. This ‘cold start’ lasts for between 500 and 2000 yr, and as a consequence the [C/O]_gas is held at intermediate values between unity and the initial value of 0.26. In fact, Paul does not exist within the disc for a long enough period of time to reach temperatures that ensure a return of [C/O]_gas to the initial value (due to H₂O remaining frozen out) before being tidally disrupted.

This effect may have important implications for the composition of any resulting objects that are formed. It has been shown that it is possible to settle dust grains towards the centres of non-turbulent gaseous planetary cores (e.g. Boss 1998; Boley et al. 2010; Nayakshin 2010b). Therefore, the transient features we observe in [C/O]_ice are of extreme interest because, if the time-scale for dust to settle is

---

3 We note that our choice of initial chemical conditions (see Table 1) influences our initial elemental ratios of the disc material, in particular setting our initial C/O ratio to 0.26. This value is lower than measurements of the solar C/O ratio, determined to be closer to 0.5 (Allende Prieto, Lambert & Asplund 2002). We stress that changing this initial value would not alter the qualitative outcomes presented here, and would simply vertically shift the baselines plotted in Fig. 7.
comparable to the time-scale of the transient feature, then ices with non-prrimordial [C/O] ratios may be incorporated into these solid cores. From Nayakshin (2010b), the time-scale for sedimentation of dust, $t_{\text{sed}}$, is given by

$$t_{\text{sed}} \approx 5 \times 10^3 \frac{M_c}{0.01 M_\odot} (\rho_d a)^{-1} \text{yr},$$

where $M_c$ is the mass of the core, $\rho_d$ is the density of the dust grains and $a$ is the dust grain radius (where the latter two quantities are defined in cgs units). Assuming $\rho_d = 2.3 \text{g cm}^{-3}$ that would imply silicate grains, the corresponding sedimentation time-scale for centimetre-sized grains and larger would be 1000–2000 yr for all fragments, and less than 500 yr for 5 cm grains or larger. Such a time-scale is comparable to the time over which $H_2O$ can remain frozen out during the initial formation stages of the fragments (see Fig. 6). In addition, grains which are covered in water ice mantles will have a higher sticking efficiency than bare grains (e.g. Ehrenfreund et al. 2003; Wang et al. 2005). Therefore, in the envelopes of fragments that exhibit this ‘cold-start’, it may be possible to grow grains more quickly, increasing the fraction of solid material that can be sedimented towards the centre. Thus, these fragments may be able to form larger rocky cores than might be expected, even if the eventual fate of the fragment as a whole is to be tidally disrupted.

The C/O evolution discussed here not only has implications for the first few thousand years of evolution, but also for the final C/O ratio of the planet itself. If dust can be efficiently settled to the fragment core before volatiles are evaporated from their surfaces, then changes in the C/O ratio can be expected – higher values will be found in the envelopes, while lower values will be found in the core. Unfortunately, the signature of this mechanism would be indistinguishable from what was discussed for the GI formation scenarios discussed in Madhusudhan, Amin & Kennedy (2014), in which different C/O ratios correspond to differing amounts of planetesimal accretion. In all these cases, there is an inverse relation between the overall metallicity of the planetary atmospheric gas (i.e. C/H and O/H) and the resulting C/O ratio. As more ices are evaporated into the planetary atmosphere, the higher the metallicity, but the lower the C/O because the ices are rich in oxygen. Only models that include differential transport of dust grains with respect to the gas can break this degeneracy (see Booth et al. 2017). We hope to extend our work with this treatment in the near future.

### 4.4 Chemical tracers of fragments

Molecules with increased abundances towards fragments may offer a useful diagnostic of the physical conditions in these regions. Such diagnostics will be important to probe in order to examine the planet formation process in detail. Narayanan et al. (2006) have examined molecular line emission from a gravitationally unstable disc that includes a gas giant planet. Using a hydrodynamic model coupled with a non-LTE radiative transfer code and fixed molecular abundances, they determine that the gas giant planet may be observable in high-$J$ transitions of HCO$^+$ if abundances are high enough. However, in our model, we see significant depletion of HCO$^+$ due to the liberation of $H_2O$ from grain surfaces by the high temperatures towards the fragments. As such, in reality, observations of HCO$^+$ would likely not be an efficient tracer of these fragments, and if observed, would rather trace the circumfragmentary material around them instead.

Recent spatially resolved observations of the disc around d216–0939 by Factor et al. (2017) have detected blue-shifted asymmetries in the HCO$^+$ and CO emission, which they suggest could be due to either a hydrodynamic vortex or the envelope of a forming protoplanet between 1.8 and 8 $M_\text{Jup}$ located at a radius of 60 ± 20 au. While Factor et al. (2017) point out that the disc of d216–0939 is likely to be currently gravitationally stable, the detected emission would be consistent with our results for the surviving fragments George and Ringo, and as such may be evidence that the disc in d216–0939 may have undergone fragmentation in the past.

While we report here species that are increased in their fractional abundance towards the fragments, the assessment of their true observability requires molecular line radiative transfer modelling and synthetic imaging. We will be performing a detailed study on the observable continuum and line emission in this model disc in a forthcoming publication (Hall et al., in preparation).

### 5 CONCLUSIONS

In this paper, we have presented the first study of the chemical evolution in a fragmenting protoplanetary disc. Using three-dimensional hydrodynamics coupled with a non-equilibrium gas-grain chemical model, we have followed the chemical evolution of the $4 \times 10^6$ particles that comprise the simulation. We summarize our main findings as follows:

(i) Our hydrodynamic model initially fragments into a large number of objects, of which four become quasi-stable. The innermost two of these fragments migrate through the disc towards the central star and undergo tidal disruption, while the outermost two persist until the end of our simulation.

(ii) The increased temperatures and densities found towards the spiral arms and fragments cause chemical reactions that would not otherwise take place at such large radii in the disc. From these, we identify molecules that are abundant towards the fragments (e.g. $H_2O$, $H_2S$, HNO, $N_2$, NH$_3$, OCS, SO) and those which are also abundant in the spiral shocks (e.g. CO, CH$_4$, CN, CS, $H_2CO$). In particular, HCO$^+$ may be a useful tracer of the circumfragmentary disc material.

(iii) Molecular condensation fronts, or ‘snow lines’, deviate significantly from the expected concentric ring structures found in axisymmetric discs. Increases in temperature caused by passing shocks desorb material at larger radii than would normally be expected, and fragments that have formed develop snow lines themselves.

(iv) Fragments can exhibit a ‘cold start’ after formation, with temperatures not high enough to entirely desorb $H_2O$ from grain surfaces until after 500–2000 yr. In one case, sufficiently high temperatures are not reached at all before the fragment in question undergoes tidal disruption. In such cases, because the effect persists for a time comparable to the sedimentation time for dust in these fragments, grain growth in the envelopes of such fragments may be enhanced, leading to a larger proportion of solid material that can be sedimented towards the core.

(v) This ‘cold start’ phenomenon also gives rise to higher C/O ratios than would normally be expected for the envelopes of fragments formed via GI, and lower C/O ratios of any cores that may be formed towards the centre of the fragments. As such, the atmospheric composition of planets formed via GI may not necessarily follow the bulk chemical composition of the disc from which they formed.

Our results suggest that correctly accounting for the influence of the dynamical evolution on the corresponding chemical evolution of protoplanetary discs can provide useful information on the composition and properties of any resulting objects that are formed. Such effects are important not only for the formation of giant planets and
brown dwarfs but also potentially for the formation of terrestrial planets by sedimentation within the fragments. In the future, we plan to extend our modelling to account for variations in, for example, dust grain sizes and dynamics, allowing us to further probe the interlinked processes that occur as planets form via a variety of processes in young circumstellar discs.

ACKNOWLEDGEMENTS

We would like to thank the referee for suggestions that improved the clarity of the manuscript and Mikhail Kama for his reading of an early draft of this work. JDI, RB and CJC gratefully acknowledge support from the DISCSIM project, grant agreement 341137, funded by the European Research Council under ERC-2013-ADG. DHF gratefully acknowledges support from the ECOGAL project, grant agreement 291227, funded by the European Research Council under ERC-2011-ADG. DHF and WKMR also acknowledge support from STFC grant ST/J001422/1. MGE acknowledges a studentship funded by, and PC, TWH and AB acknowledge financial support from, the European Research Council (project PALs 320620). CH gratefully acknowledges funding from the European Research Council (ERC) under the European Union’s Horizon 2020 research and innovation programme (grant agreement no. 681601). Some of computational work for this paper was carried out on the joint STFC and SFC (SRIF) funded Wardlaw cluster at the University of St Andrews (Scotland, UK). This research has made use of NASA’s Astrophysics Data System Bibliographic Services and ASTROPY, a community-developed core PYTHON package for Astronomy (Astropy Collaboration et al. 2013).

REFERENCES

Allende Prieto C., Lambert D. L., Asplund M., 2002, ApJ, 573, L137
Astropy Collaboration et al., 2013, A&A, 558, A33
Banzatti A., Pinilla P., Ricci L., Pontoppidan K. M., Birnstiel T., Ciesla F., 2015, ApJ, 815, L15
Bate M. R., Bonnell I. A., Price N., 1995, MNRAS, 277, 362
Bergin E. A., Aikawa Y., Blake G. A., van Dishoeck E. F., 2007, in Reipurth B., Jewitt D., Keil K., eds, Protostars and Planets V. University of Arizona Press, Tucson, p. 751
Bodenheimer P., Yorke H. W., Rozycka M., Tohline J. E., 1990, ApJ, 355, 651
Boley A. C., Durisen R. H., 2010, ApJ, 724, 618
Boley A. C., Hayfield T., Mayer L., Durisen R. H., 2010, Icarus, 207, 509
Boley A. C., Helled R., Payne M. J., 2011, ApJ, 735, 30
Booth R. A., Clarke C. J., Madhusudhan N., Ilee J. D., 2017, MNRAS, 469, 3994
Boss A. P., 1998, ApJ, 503, 923
Brown P. N., Byrne G. D., Hindmarsh A. C., 1989, SIAM J. Sci. Comput., 10, 1038
Caselli P., Ceccarelli C., 2012, A&A, 20, 56
Cieza L. A. et al., 2016, Nature, 535, 258
Cleeves I. L., Bergin E. A., Harries T. J., 2015, ApJ, 807, 2
Cossins P., Lodato G., Clarke C. J., 2010, MNRAS, 401, 2587
Dipierro G., Lodato G., Testi L., de Gregorio Monsalvo I., 2014, MNRAS, 444, 1919
Dipierro G., Pinilla P., Lodato G., Testi L., 2015, MNRAS, 451, 974
Douglas T. A., Caselli P., Ilee J. D., Boley A. C., Hartquist T. W., Durisen R. H., Rawlings J. M. C., 2013, MNRAS, 433, 2064
Durisen R. H., Boss A. P., Mayer L., Nelson A. F., Quinn T., Rice W. K. M., 2007, in Reipurth B., Jewitt D., Keil K., eds, Protostars and Planets V. University of Arizona Press, Tucson, p. 607
Dutrey A. et al., 2014, in Beuther H., Klessen R. S., Dullemond C. P., Henning T., eds, Protostars and Planets VI. University of Arizona Press, Tucson, p. 317
Ehrenfreund P., Charnley S. B., 2000, ARA&A, 38, 427
Ehrenfreund P. et al., 2003, Planet. Space Sci., 51, 437
Eisler J. A., Carpenter J. M., 2006, ApJ, 641, 1162
Eistrup C., Walsh C., van Dishoeck E. F., 2016, A&A, 595, A83
Evans M. G., Ilee J. D., Boley A. C., Caselli P., Durisen R. H., Hartquist T. W., Rawlings J. M. C., 2015, MNRAS, 453, 1147
Evans M. G. et al., 2017, MNRAS, 470, 1828
Factor S. M. et al., 2017, AJ, 153, 233
Forgan D., Rice K., 2013, MNRAS, 432, 3168
Forgan D. H., Rice K., Stamatellos D., Whitworth A. P., 2009, MNRAS, 394, 882
Forgan D., Rice K., Cossins P., Lodato G., 2011, MNRAS, 410, 994
Forgan D., Parker R. J., Rice K., 2015, MNRAS, 447, 836
Frillmann S. et al., 2017, A&A, 602, A120
Gammie C. F., 2001, ApJ, 553, 174
Hall C., Forgan D., Rice K., Harries T. J., Klaassen P. D., Biller B., 2016, MNRAS, 458, 306
Hall C., Forgan D., Rice K., 2017, MNRAS, 470, 2517
Haworth T. J. et al., 2016, PASA, 33, e053
Heiter U., 2002, A&A, 381, 959
Helled R., Podolak M., Kovetz A., 2006, Icarus, 185, 64
Helled R. et al., 2014, in Beuther H., Klessen R. S., Dullemond C. P., Henning T., eds, Protostars and Planets VI. University of Arizona Press, Tucson, p. 643
Henning T., Semenov D., 2013, Chem. Rev., 113, 9016
Hincelin U., Wakelam V., Commorcon B., Hersant F., Guilloteau S., 2013, ApJ, 775, 44
Ilee J. D., Boley A. C., Caselli P., Durisen R. H., Hartquist T. W., Rawlings J. M. C., 2011, MNRAS, 417, 2950
Joos M., Hennebelle P., Ciardi A., 2012, A&A, 543, A128
Khajenabi F., Kazarni K., Shadmehri M., 2017, ApJ, 841, 99
Lodato G., Rice W. K. M., 2004, MNRAS, 351, 630
Madhusudhan N., 2012, MNRAS, 421, 3286
Madhusudhan N., Amin M. A., Kennedy G. M., 2014, ApJ, 794, L12
Marois C., Macintosh B., Barman T., Zuckerman B., Song I., Patience J., Lafrenière D., Doyon R., 2008, Science, 322, 1348
Mayer L., Luftin G., Quinn T., Wadsley J., 2007, ApJ, 661, L77
Meru F., Bate M. R., 2011, MNRAS, 411, L1
Meru F., Juhász A., Ilee J. D., Clarke C. J., Rosotti G. P., Booth R. A., 2017, ApJ, 839, L24
Millar T. J., Farquhar P. R. A., Willacy K., 1997, A&AS, 121
Narayanan D., Kulesa C. A., Boss A., Walker C. K., 2006, ApJ, 647, 1426
Nayakshin S., 2010a, MNRAS, 408, L36
Nayakshin S., 2010b, MNRAS, 408, 2381
Nayakshin S., Fletcher M., 2015, MNRAS, 452, 1654
Oberg K. I., Murray-Clay R., Bergin E. A., 2011, ApJ, 743, L16
Paardekooper S.-J., 2012, MNRAS, 421, 3266
Panić O., Min M., 2017, MNRAS, 467, 1175
Pérez L. M. et al., 2016, Science, 353, 1519
Pollack J. B., Hubickyj O., Bodenheimer P., Lissauer J. J., Podolak M., Greenzweig Y., 1996, Icarus, 124, 62
Price D. J., 2007, PASA, 24, 159
Qi C. et al., 2013, Science, 341, 630
Rice W. K. M., Lodato G., Armitage P. J., 2005, MNRAS, 364, L56
Rice W. K. M., Mayo J. H., Armitage P. J., 2010, MNRAS, 402, 1740
Saumon D., Guillot T., 2004, ApJ, 609, 1170
Stamatellos D., Whitworth A. P., Bate M. R., 2004, MNRAS, 353, 1078
Young M. D., Clarke C. J., 2016, MNRAS, 455, 1438
APPENDIX A: PHYSICAL AND CHEMICAL STRUCTURE OF THE FRAGMENTS

Fig. A1 shows the average physical and chemical structure of the fragments at various times throughout the simulation.

![Figure A1](image)

Figure A1. Spherically average physical conditions towards the fragments as a function of time. The upper row shows conditions at 2852 yr and the lower row at 4100 yr. Upward turns exhibited by some of the temperature profiles are due to the inclusion of particles at higher \( z \) in the disc, which experience somewhat higher temperatures.

![Figure A2](image)

Figure A2. Spherically average fractional abundances for selected species towards the fragments as a function of time. The upper row shows abundances at 2852 yr and the lower row at 4100 yr.

APPENDIX B: COLUMN DENSITY IMAGES FOR ALL SPECIES

Figs B1, B2 and B3 show the column density of the dominant neutral gas phase species, grain surface species and cations, respectively.
Figure B1. The dominant neutral gas phase species at $t = 2852$ yr. Panels span 130 au and the colour bar gives the logarithm of the column density in cm$^{-2}$. 
Figure B2. The dominant grain surface species at $t = 2852$ yr. Panels span 130 au and the colour bar gives the logarithm of the column density in cm$^{-2}$. 
Figure B3. The dominant cations in the gas phase at $t = 2852$ yr. Panels span 130 au and the colour bar gives the logarithm of the column density in cm$^{-2}$. 

This paper has been typeset from a \TeX/\LaTeX file prepared by the author.