Molecules and dust in Cassiopeia A

I. Synthesis in the supernova phase and processing by the reverse shock in the clumpy remnant

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ABSTRACT

Aims. We study the chemistry of the type IIb supernova ejecta, which led to the Cas A supernova remnant, to assess the chemical type and quantity of dust that forms and evolves in the remnant phase. We later model a dense oxygen-rich ejecta knot, which is crossed by the reverse shock in Cas A, to study the evolution of the clump gas phase and the possibility to reform dust clouds in the post-reverse shock gas.

Methods. The chemistry is described by a chemical network that includes all possible processes that are efficient at high gas temperatures and densities. The formation of key bimolecular species (e.g., CO and SiO) and dust clusters of silicates, alumina, silica, metal carbides and sulphides, pure metals, and amorphous carbon is considered. A set of stiff, coupled ordinary differential equations is solved for the conditions pertaining to both the SN ejecta and the post-reverse shock gas.

Results. We find that the ejecta of type IIb SNe are unable to form large amounts of molecules and molecular clusters that are precursors to dust grains, when compared to their type II-P counterparts, because of their diffuse ejecta. The ejecta gas density needs to be increased by several orders of magnitude to allow for the formation of dust clusters. We show that the chemical composition of the dust clusters that form changes drastically and gains in chemical complexity with increasing gas density. Hence, the ejecta of the Cas A supernova progenitor must have been in the form of dense clumps to account for the dust chemical composition and masses that have been inferred from infrared observations of Cas A. As for the impact of the reverse shock on dense ejecta clumps, we show that the ejecta molecules that are destroyed by the shock reform in the post-reverse shock gas with lower abundances than those of the initial ejecta clump, except for SiO. These molecules include CO, SiS, and O$_2$. On the other hand, dust clusters are destroyed by the reverse shock and do not reform in the post-reverse shock gas, even for the highest gas density. These results indicate that the synthesis of dust grains out of the gas phase in the dense knots of Cas A and in other supernova remnants is unlikely.

Key words. astrochemistry – molecular processes – ISM: supernova remnants – circumstellar matter – supernovae: general – dust, extinction

1. Introduction

Large amounts of cosmic dust at high redshift are inferred from the reddening of background quasars and damped Ly$_\alpha$ systems (Pei et al. 1991; Pettini et al. 1994) and pose the problem of identifying the providers of dust grains in primeval galaxies and at cosmic time less than one billion years. Type II supernovae (SNe) are strong contenders as dust factories at high-redshift because of the short evolution time-scale of their massive stellar progenitors (~10$^6$ yrs). Furthermore, dust and molecules are formed by the reverse shock and do not reform in the post-reverse shock gas, even for the highest gas density. These results indicate that the synthesis of dust grains out of the gas phase in the dense knots of Cas A and in other supernova remnants is unlikely.

Owing to its proximity (3.4 kpc) and angular size (~5'), the SN remnant Cas A has been extensively studied observationally. This remnant results from the explosion of a SN of type IIb 330 years ago. The progenitor was a massive supergiant star having lost most of its hydrogen envelope with a ZAMS mass that comprises between 15 and 25 M$_\odot$ (Krause et al. 2008). The explosion blast wave travelled in a diffuse medium, leading to a low density ejecta as compared to regular Type II SNe (Filipenko 1988). The Cas A ejecta is now crossed by the RS and warm dust has been detected in the IR. Dust grains have been detected with the submillimetre (submm) Herschel telescope in several SN remnants, several decades or centuries after the SN explosion. The derived masses are larger than the masses observed in the IR at early times and vary between ~0.08 M$_\odot$ for the Cas A remnant (Barlow et al. 2010) and ~0.7 M$_\odot$ for the young remnant SN1987A (Matsuura et al. 2011). These large values are surprising because the physical conditions are harsh in SN remnants. The SN ejecta gas is re-processed by a reverse shock (RS) that is created when the mass of gas swept up by the explosion blast wave exceeds the ejecta mass (Chevalier 1977). The RS travels inwards and, therefore, induces the partial destruction of the molecules and dust synthesised and expelled in the SN ejecta phase.
an estimated mass of \( \sim 0.075 \, M_\odot \). Of prime importance is the detection of molecules in the remnant. Warm carbon monoxide, CO, has been observed at the RS position with Spitzer (Rho et al. 2009, 2012) and in one dense clump with Herschel (Wallström et al. 2013). In the young remnant SN1987A, cool CO has been recently observed with ALMA along with the partial spectroscopic detection of SiO (Kamenetzky et al. 2013). This detection points to chemical species, CO and SiO, formed in the ejecta after the SN explosion. As in other evolved circumstellar environments, molecules appear to be strongly coupled to the formation of dust grains in SNe (Cherchneff 2010). Once more, the presence of dust and molecules in Cas A and other remnants confirms this inherent aspect of the synthesis of cosmic dust and indicates a strong chemical reprocessing of the material ejected during the SN phase by the RS.

Theoretical studies on both the formation of dust in SNe and the reprocessing by the RS have been carried out. Models of dust production in SN ejecta lead to contradictory results on the dust quantity produced after the SN explosion. Early studies based on classical nucleation theory (CNT) indicate large dust masses (Kozasa et al. 1989; Todini & Ferrara 2001; Nozawa et al. 2003) that cannot be reconciled with IR observations of SNe. Models based on chemical kinetics and that consider the coupling between molecules and dust grains result in a lower amount of dust formed in the nebular phase (Cherchneff & Lilly 2008; Cherchneff & Dwek 2010). More recently, Sarangi & Cherchneff (2013, hereafter SC13) show that the molecular clusters precursors to dust (hereafter, dust clusters) gradually grow from low to high masses on a time-span of a few years after the explosion and provide a genuine explanation to the discrepancy on dust mass derived from IR data of SNe and submm data of SN remnants. The reprocessing of dust by the RS in SN remnants has been modelled by assuming pre-shock dust distributions derived from CNT and in the context of a homogeneous SN ejecta (Nozawa et al. 2010; Silvia et al. 2012). None of these studies addresses the chemistry of the RS, the survival of molecules, and the possibility to reform dust after the passage of the RS. Therefore, whether SNe and SN remnants are dust makers and/or destroyers is still unclear on the basis of both available theoretical predictions and observational data.

To advance our understanding of the net dust budget produced by SNe, we carry out a global study of the synthesis of molecules and dust in SN ejecta and their reprocessing by the RS in the remnant phase by focusing on Cas A. In this paper (Paper I), we study the formation of molecules and dust clusters in the type IIb SN that led to Cas A and the later processing of the produced gas-phase material by the RS. In a forthcoming publication (Paper II), we will address the condensation of dust clusters and derive grain size distributions for the type IIb SN. We will then study the dust reprocessing in clumps and the inter-clump medium for various RS models. Presently, we model the non-equilibrium chemistry of the SN ejecta and that of the post-RS gas, by assuming that the processed ejecta material resides in dense, fast-moving knots, as observed by Fesen et al. (2001, 2006) and Docenko & Sunyaev (2010, hereafter DS10).

### Table 1. Thermal and non-thermal processes and chemical reaction types included in the chemical model for both the type IIb SN ejecta and the post-RS gas.

<table>
<thead>
<tr>
<th>Reaction description</th>
<th>Gas regime/location</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Thermal</strong></td>
<td></td>
</tr>
<tr>
<td>Unimolecular</td>
<td></td>
</tr>
<tr>
<td>AB ( \rightarrow ) A + B</td>
<td>Neutral exchange</td>
</tr>
<tr>
<td>A + B ( \rightarrow ) AB + hv</td>
<td>Radiative association</td>
</tr>
<tr>
<td>AB + M ( \rightarrow ) A + B + M</td>
<td>Collision dissociation</td>
</tr>
<tr>
<td>AB(^+) + C ( \rightarrow ) BC(^+) + A</td>
<td>Ion–Molecule</td>
</tr>
<tr>
<td>AB(^+) + C ( \rightarrow ) AB + C(^+)</td>
<td>Charge exchange</td>
</tr>
<tr>
<td>A(^+) + e(^-) ( \rightarrow ) A + hv</td>
<td>Radiative recombination</td>
</tr>
<tr>
<td>AB(^+) + e(^-) ( \rightarrow ) A + B</td>
<td>Dissociative recombination</td>
</tr>
<tr>
<td><strong>Non-Thermal</strong></td>
<td></td>
</tr>
<tr>
<td>A + CE ( \rightarrow ) A(^+) + e(^-) + CE</td>
<td>Ionisation by Compton e(^-)</td>
</tr>
<tr>
<td>AB + CE ( \rightarrow ) A + B + CE</td>
<td>Dissociation by Compton e(^-)</td>
</tr>
<tr>
<td>AB + hv ( \rightarrow ) A + B</td>
<td>Photo-dissociation</td>
</tr>
<tr>
<td>A + hv ( \rightarrow ) A(^+) + e(^-)</td>
<td>Photo-ionisation</td>
</tr>
</tbody>
</table>

### 2. Chemical model

The synthesis of dust in SN environments stems from the chemical ability of the elements produced via nucleosynthesis during the stellar evolution of the massive progenitor and the SN explosion to assemble into molecules and dust molecular clusters. Supernova ejecta are characterised by extreme physical conditions, including high gas temperature and velocity in the back of the explosion blast wave, radioactivity induced by the decay of \( ^{56}\text{Ni} \), and \( \gamma \)-rays and ultraviolet (UV) fields. For both environments studied in this paper, we use similar processes, as in the study of dust formation in primeval and type II-P SNe by Cherchneff & Dwek (2009, 2010) and SC13, respectively. A chemical kinetic approach is based on a chemical reaction network that includes the thermal and non-thermal processes summarised in Table 1. Thermal processes include unimolecular, bimolecular, and termolecular reactions. Because of the high gas temperatures present a few months after the SN explosion, neutral exchange reactions, which often have activation energy barriers reflecting the energy required to break and re-arrange chemical bonds, are prevalent formation processes in the build-up of chemical complexity and the formation of dust clusters (Cherchneff & Dwek 2009, 2010, SC13).

In addition to these reactions, we consider non-thermal processes that include the destruction of molecules and ionisation...
of atoms by energetic Compton electrons created when the γ-ray photons generated by the decay of $^{56}$Ni and $^{56}$Co degrade via collisions with electrons. Rates are estimated following Cherchneff & Dwek (2009), and we assume an average γ-ray optical depth for the ejecta with a 19 $M_{⊙}$ stellar progenitor. As shown by SC13, the use of a γ-ray optical depth for each ejecta zone as calculated by Kozma & Fransson (1992) for a 20 $M_{⊙}$ progenitor has little impact on the trends and results derived for an average γ-ray optical depth. Photo-ionisation and -dissociation by UV radiation are considered in the chemistry of the shocked clump.

The various atoms, molecules, and ions considered to participate in the SN ejecta chemistry are summarised in Table 2. Small molecular clusters form according to the processes described in SC13, where the description of the growth pathways of small silicate clusters, namely forsterite dimers (Mg$_2$SiO$_4$, Mg$_5$Si$_2$O$_7$) and enstatite dimers (Mg$_2$Si$_2$O$_5$), is based on the work by Goumans & Bromley (2012). This study indicates possible chemical routes to the formation of the silicate dimers, which involve the formation of the SiO dimer ring, (SiO)$_2$, and its growth to Si$_2$O$_3$ through the reaction with O$_2$ and SO. The subsequent pathway involves the addition of a magnesium atom to the Si$_2$O$_3$ structure. The later growth of clusters is described by one O-addition step followed by one Mg inclusion as a recurrent growth scenario. We have also considered a new pathway for the formation of the alumina dimer, (Al$_2$O$_3$)$_2$, that was not considered in SC13. Because the structure of the molecule Al$_2$O$_3$ is akin to that of Si$_2$O$_3$, we have considered a similar growth route starting from the formation of AIO and followed by its dimerisation to Al$_2$O$_3$ and the formation of Al$_2$O$_3$ via oxidation. The alumina dimers are then formed from the recombination of two Al$_2$O$_3$ molecules. Oxidising agents for the formation of silicates and alumina include atomic O, O$_2$, AIO, and SO, but reactions with O$_2$ and SO are prevalent for silicate clusters while reaction with AIO dominates in the case of alumina.

The chemical modelling of the type IIb SN that led to Cas A includes two temperature regimes and chemical networks because of the initial gas temperature and density quickly drop with time from 100 days to 3000 days (~8 years) post-explosion (see next Section and Table 4). These values are lower than in the case of type II-P SNe. The ejecta quickly expands at high velocities owing to the lack of a progenitor circumstellar wind that would slow down the explosion blast wave through interaction. We thus consider a high temperature reaction network from day 100 (~7000 K) to day 1000 (300 K) and a low temperature network from day 1001 until day 3000 (~60 K), for which the reaction rates of relevant reactions have been adjusted according to available, low-temperature data.

The chemical kinetic description is applied to the gas parameters that characterise the SN ejecta and shock models (see Section Sect. 3). The variation of the number density of the chemical species $i$ with time is described by the following rate equation:

$$\frac{dn_i}{dt} = P_i - L_i = \sum_{j} k_{i \rightarrow j} n_j - \sum_{k} k_{k \rightarrow i} n_k,$$  \hspace{1cm} (1)

where $P_i$ and $L_i$ are the total chemical production and loss processes for species $i$, $n_i$ is the number density for species $i$, and $k_{ij}$ the rate for the reaction of $i$ with $j$. This rate is expressed in Arrhenius form as

$$k_{ij}(T) = A_{ij} \times \left( \frac{T}{300} \right)^{\nu} \times \exp(-E_{ij}/T),$$  \hspace{1cm} (2)

where $T$ is the gas temperature in Kelvin, $\nu$ the temperature dependence exponent, and $E_{ij}$ the activation energy barrier in Kelvin. The coefficient $A_{ij}$ has the units of s$^{-1}$ for unimolecular processes, cm$^3$ s$^{-1}$ for bimolecular reactions, and cm$^6$ s$^{-1}$ for termolecular reactions.

The system includes 93 species, 412 chemical reactions, and a set of 93 stiff, coupled ordinary differential equations is solved using a Gear method (Hindmarsh 1983).

### 3. Physical models

We study the chemistry of two distinct environments: 1) the ejecta of the SN type IIb that led to the remnant Cas A; and 2) one oxygen-rich knot in Cas A that is crossed by the RS. The aim is to derive a comprehensive picture of the production and processing of molecules and dust in Cas A and to assess whether the chemical species synthesised in SNe survive RS processing in the remnant phase. First, we present our physical model for the ejected material of a type IIb SN, and second, we describe our model of a shocked clump or knot in Cas A.

#### 3.1. The type IIb SN ejecta

Stars with masses in the range 8–30 $M_{⊙}$ explode as core-collapse SNe of type II, showing strong hydrogen lines in their spectrum. Type IIb SNe show a weak hydrogen line, which subsequently disappears with time, and are characterised by higher expansion
where \( T(r, t) \) is the gas temperature at 100 days and \( \gamma \) is the adiabatic index. Values are \( T_0 = 6664 \, \text{K} \) and \( \gamma = 1.435 \). A flat density profile was assumed across the He-core with a value at 100 days equal to \( \rho(100) = 5.1 \times 10^{-15} \, \text{g cm}^{-3} \). All values were taken from Nozawa et al. (2010). The SN type IIb parameters for the Cas A progenitor and the values for the ejecta gas number density and temperature as a function of time are summarised in Table 4.

3.2. The reverse shock models

The Cas A remnant shows evidence for several high-density knots or clumps embedded in a rarefied inter-clump medium. Fesen et al. (2001) have observed fast-moving knots (FMKs) in the optical, while Rho et al. (2009, 2012) have detected ro-vibrational transitions of CO in emission in several small knots located at the RS position. High energy line detection of shocked CO with Herschel confirmed the existence of high-density gas regions in the remnant (Wallström et al. 2013). In Cas A, the RS velocity relative to the ejecta is \( \approx 2000 \, \text{km s}^{-1} \) (Morse et al. 2004). When encountering a dense clump with a density contrast with respect to the inter-clump medium \( \chi = n_c/n_{ic} \) (\( n_c \) and \( n_{ic} \) are the number density of the clump and the interclump medium, respectively), the RS velocity decreases by a factor \( \sqrt{\chi} \) owing to the conservation of energy. For clump density contrast ranging between 100 and 1000 and RS velocities of 1000–2000 km s\(^{-1}\), the RS velocity in the clump spans 30–200 km s\(^{-1}\). Such velocities have been confirmed by DS10 from their analysis of fine-structure far-IR atomic lines coming from an oxygen-rich FMK in Cas A.

In the present study, we want to assess the fate of molecules and dust clusters within ejecta clumps that are processed by the RS. We thus investigate various RS velocities, derive the

**Table 3.** Gas mean molecular weight \( \mu_{\text{gas}} \), C/O ratio, and initial elemental mass yields as a function of ejecta zone for the 19 \( M_\odot \) progenitor of Rauscher et al. (2002).

<table>
<thead>
<tr>
<th>Zone</th>
<th>( \mu_{\text{gas}} )</th>
<th>C/O</th>
<th>He</th>
<th>C</th>
<th>O</th>
<th>Ne</th>
<th>Mg</th>
<th>Al</th>
<th>Si</th>
<th>S</th>
<th>Ar</th>
<th>Fe</th>
<th>Ni</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a</td>
<td>42.3</td>
<td>0.08</td>
<td>0</td>
<td>1.0(–7)</td>
<td>1.3(–6)</td>
<td>6.4(–7)</td>
<td>1.8(–5)</td>
<td>2.7(–5)</td>
<td>3.9(–2)</td>
<td>2.3(–2)</td>
<td>4.7(–3)</td>
<td>5.8(–3)</td>
<td>1.1(–1)</td>
</tr>
<tr>
<td>1b</td>
<td>22.5</td>
<td>1(–3)</td>
<td>0</td>
<td>1.1(–4)</td>
<td>1.2(–1)</td>
<td>1.0(–4)</td>
<td>8.7(–4)</td>
<td>2.2(–4)</td>
<td>9.8(–2)</td>
<td>5.6(–2)</td>
<td>1.5(–2)</td>
<td>2.7(–3)</td>
<td>6.1(–4)</td>
</tr>
<tr>
<td>2</td>
<td>17.0</td>
<td>0.049</td>
<td>1.1</td>
<td>3.3(–1)</td>
<td>8.4(–2)</td>
<td>9.1(–3)</td>
<td>1.5(–2)</td>
<td>1.3(–1)</td>
<td>1.2(–4)</td>
<td>7.5(–4)</td>
<td>2.2(–4)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3a</td>
<td>15.1</td>
<td>0.27</td>
<td>0</td>
<td>2.9(–2)</td>
<td>1.1(–1)</td>
<td>2.8(–3)</td>
<td>2.0(–3)</td>
<td>3.0(–5)</td>
<td>1.1(–4)</td>
<td>3.2(–5)</td>
<td>8.9(–6)</td>
<td>7.8(–5)</td>
<td>2.8(–5)</td>
</tr>
<tr>
<td>3b</td>
<td>10.3</td>
<td>0.48</td>
<td>7.4(–2)</td>
<td>1.2(–1)</td>
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<td>2.8(–5)</td>
</tr>
<tr>
<td>4a</td>
<td>4.1</td>
<td>3.0</td>
<td>7.2(–1)</td>
<td>1.2(–2)</td>
<td>4.2(–3)</td>
<td>1.0(–2)</td>
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<tr>
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<td>4.3(–4)</td>
<td>2.4(–5)</td>
</tr>
</tbody>
</table>

Notes: The total yield is the elemental mass yield for the total ejected material. Mass yields are in \( M_\odot \) and the gas mean molecular weight \( \mu_{\text{gas}} \) is in g cm\(^{-3}\). Nitrogen, N, is only present in Zone 4b with a yield of \( 4 \times 10^{-7} \). All values were taken from Nozawa et al. (2010). The SN type IIb parameters for the Cas A progenitor and the values for the ejecta gas number density and temperature as a function of time are summarised in Table 4.

<table>
<thead>
<tr>
<th>Zone 1a</th>
<th>Zone 1b</th>
<th>Zone 2</th>
<th>Zone 3a</th>
<th>Zone 3b</th>
<th>Zone 4a</th>
<th>Zone 4b</th>
</tr>
</thead>
<tbody>
<tr>
<td>( 1.68–1.88 , M_\odot )</td>
<td>( 1.89–2.18 , M_\odot )</td>
<td>( 2.19–3.86 , M_\odot )</td>
<td>( 3.87–4.02 , M_\odot )</td>
<td>( 4.03–4.49 , M_\odot )</td>
<td>( 4.5–5.26 , M_\odot )</td>
<td>( 5.27–5.62 , M_\odot )</td>
</tr>
<tr>
<td>( T_{\text{gas}} )</td>
<td>( n_{\text{gas}} )</td>
<td>( n_{\text{gas}} )</td>
<td>( n_{\text{gas}} )</td>
<td>( n_{\text{gas}} )</td>
<td>( n_{\text{gas}} )</td>
<td>( n_{\text{gas}} )</td>
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<td>1.8(8)</td>
<td>2.0(8)</td>
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<tr>
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<td>9.0(6)</td>
<td>1.7(7)</td>
<td>2.3(7)</td>
<td>2.5(7)</td>
<td>3.7(7)</td>
</tr>
<tr>
<td>300</td>
<td>1600</td>
<td>2.7(6)</td>
<td>5.0(6)</td>
<td>6.6(6)</td>
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<td>5.8(5)</td>
<td>1.1(6)</td>
<td>1.4(6)</td>
<td>1.6(6)</td>
<td>2.3(6)</td>
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<tr>
<td>1000</td>
<td>330</td>
<td>7.2(4)</td>
<td>1.4(5)</td>
<td>1.8(5)</td>
<td>2.0(5)</td>
<td>3.0(5)</td>
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<tr>
<td>1500</td>
<td>220</td>
<td>2.2(4)</td>
<td>4.0(4)</td>
<td>5.3(4)</td>
<td>6.0(4)</td>
<td>8.8(4)</td>
</tr>
<tr>
<td>2000</td>
<td>140</td>
<td>9.0(3)</td>
<td>1.7(4)</td>
<td>2.3(4)</td>
<td>2.5(4)</td>
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<td>2500</td>
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<td>4.6(3)</td>
<td>8.7(3)</td>
<td>1.1(4)</td>
<td>1.3(4)</td>
<td>1.9(4)</td>
</tr>
<tr>
<td>3000</td>
<td>80</td>
<td>2.7(3)</td>
<td>5.0(3)</td>
<td>6.6(3)</td>
<td>7.5(3)</td>
<td>1.1(4)</td>
</tr>
</tbody>
</table>

Notes: Time is in days, \( T_{\text{gas}} \) in Kelvin and \( n_{\text{gas}} \) in cm\(^{-3}\). (a) The type IIb SN ejecta is characterised by the following parameters: progenitor mass = \( 19 \, M_\odot \); explosion energy \( E_{\text{kin}} = 1 \times 10^{51} \) erg; He-core mass = \( 5.62 \, M_\odot \); ejecta velocity = \( 4000 \, \text{km s}^{-1} \); and \( \gamma \)-ray Optical depth \( \tau_\gamma = 15 \).
time-variation of the post-RS gas parameters, based on existing shock models, and apply our chemical kinetic formalism to study the post-RS chemistry. Borkowski & Shull (1990; hereafter BS90) model steady state radiative shocks in a pure oxygen gas with velocities that range between 35 km s\(^{-1}\) and 170 km s\(^{-1}\). This study provides the adequate range of conditions for the case of the RS impacting a dense knot in Cas A.

Considering a knot of gas with an initial density contrast with respect to the inter-clump medium, the parameters pertaining to a description of the RS, as illustrated in Fig. 1, are the shock velocity relative to the knot velocity \(V_s\), the pre-shock number density \(n_0\) and temperature \(T_0\), the gas density \(n_1\) and temperature \(T_1\) in the post-shock hot region (HR), and those for the photoionised region (PIR) \((n_2\) and \(T_2\), respectively), and the time length of the PIR, \(t_{\text{PIR}}\). In the HR, the compression induces an equilibration between ions and electrons, which draw their temperature through Coulomb collisions with ions when they cool via inelastic collisions with ions. Because the cooling time is shorter than the recombination time for ions, a constant ionisation state prevails in the HR until the gas quickly cools down and recombines. Ionisation is still sustained in the PIR by the UV flux coming from the HR so that thermal equilibrium is reached over a time \(t_{\text{PIR}}\) (or length \(L_{\text{PIR}}\), which is defined by the depletion of the HR UV flux.

From our SN ejecta model, we can constrain the knot physical properties as follows: considering the oxygen-rich zones of the ejecta (e.g. zone 2) and assuming the ejecta expansion velocity of Table 4, the volume of the homogenous ejected material at time \(t\) (in days) after explosion assuming a spherical expansion is

\[
V(t) = \frac{4}{3}\pi \left( v_\text{ej} \times t \times 8.64 \times 10^3 \right)^3, \tag{5}
\]

where \(v_\text{ej}\) is the ejecta velocity. For our chosen values at day 100, \(V(100) = 1.7 \times 10^7\) cm\(^3\). Values for the clump volume filling factor, \(f_c\), derived from radiative transfer studies of the dust emission in SN ejecta typically range from 0.05 to 0.2 (Gallagher et al. 2012). Assuming \(f_c = 0.05\), we derive that the total volume in the form of clumps in the SN ejecta is \(V_{\text{tot},(100)} = f_c \times V(100) = 8.6 \times 10^5\) cm\(^3\). For the ejecta mass given in Table 4, we derive a typical clump mass ranging from \(10^{-3} M_\odot\) to \(3 \times 10^{-3} M_\odot\), where the number of clumps \(N_c\) is 1000 and 3000, respectively. These \(N_c\) values are usually used in radiative transfer models. Our derived clump mass is also consistent with values obtained from 3D hydrodynamic models of SN explosion (Hammer et al. 2010). The number density of one oxygen-rich clump at day 100 post-explosion is then given by

\[
n_c(100) = \frac{M_c}{V_{\text{tot},(100)} N_c}, \tag{6}
\]

with \(n_c(100) = 3.4 \times 10^{10}\) cm\(^{-3}\) for \(N_c = 1000\). This gas number density is roughly a factor of 200 larger than the values listed in Table 4 for the O-rich zones. Using Eq. (3), we recover a typical oxygen-rich clump number density 330 years after explosion of \(\sim 20\) cm\(^{-3}\), which is consistent with the value of 100 cm\(^{-3}\) inferred by DS10 for the clump density not yet shocked by the RS. If we assume an inter-clump medium number density of 1 cm\(^{-3}\) and 0.1 cm\(^{-3}\) (Nozawa et al. 2003, BS90), we recover a clump/inter-clump density contrast \(\gamma\) value of between \(\sim 20\) and 200, respectively. The density contrast used by DS10 is 100, while the value assumed by Sylvia et al. (2012) is 1000. For a range of initial unattenuated RS shock speeds of 1000–2000 km s\(^{-1}\), the RS velocity through the clump ranges from \(\sim 35\) km s\(^{-1}\) to 200 km s\(^{-1}\). These RS velocities have been modelled by BS90, and we use their results as a basis for our RS analytical models.

The RS parameters are listed in Table 5. We assume five velocities for the shock relative to the clump, an initial pre-shock density in the clump of \(n_0 = 100\) cm\(^{-3}\) and a PIR length \(L_{\text{PIR}} = 2 \times 10^{13}\) cm for all models, as derived by DS10. In the PIR, we assume the oxygen column densities listed in Table 10 of BS90 for the various shock speeds listed in Table 5. The time length of the PIR is derived by assuming the conservation of momentum through the shock front and is given by

\[
t_{\text{PIR}} = \frac{N(O)}{n_2 \times v_2} \times \frac{N(O)}{n_0 \times V_s}, \tag{7}
\]

where \(N(O)\) is the oxygen column density and \(v_2\) the gas velocity in the PIR. The density in the PIR \(n_2\) is derived from the definition of the PIR length \(L_{\text{PIR}}\) given by

\[
L_{\text{PIR}} = v_2 \times t_{\text{PIR}} = \frac{n_0}{n_2} \times V_s \times t_{\text{PIR}}. \tag{8}
\]

The temperature in the PIR is that derived by DS10 for the high velocity shocks and is fixed to 1500 K. We also test the impact on the PIR chemistry of the higher PIR temperature derived...
by BS90 ($T_2 = 4500$ K). For the lowest shock, we take the PIR temperature derived for that shock speed by BS90.

Non-thermal processes include the penetration of the PIR by UV photons coming from the HR. These UV photons partially sustain the level of ionisation in the PIR via the photoionisation of oxygen. The photon fluxes emergent from the HR were estimated as a function of the shock velocity and energy band by BS90. We consider the ionisation of OI in various levels of OII that involve UV photons with energy between 13.62 and 35.12 eV as the main source of electrons in the PIR. The UV flux values are listed in Table 5 of BS90 for the shock speeds considered in this study. The unattenuated photoionisation rate for oxygen $\zeta_{0,v}$ at frequency $v$ is given by

$$\zeta_{0,v} = \int_v \sigma_{v} \times 4\pi \frac{J_v}{h v} \times d v,$$

where $4\pi J_v$ is the radiation intensity averaged over the solid angle and $\sigma_v$ is the photoionisation cross-section of oxygen taken from Angel & Samson (1988) at the frequency $v$. The attenuated rate due to extinction in the post-shock gas is calculated as follows: we omit potential attenuation from dust grains present in the PIR (this will be considered in Paper II). We estimate the unattenuated rate for the various frequency bands $k$ and the integrated fluxes given in Tables 3 and 5 of BS90, respectively, and calculate the frequency-dependent optical depth $\tau_k$ for each frequency band $k$, which is given by

$$\tau_k(t) = \sigma_k \times n_2 \times t \times n_2.$$

Here $t$ is the time in the PIR. The total attenuated photoionisation rate at time $t$ in the PIR is then given by

$$\zeta(t) = \sum_k \zeta_{0,k} \times \exp(-\tau_k(t)),$$

where $\zeta_{0,k}$ is the unattenuated photoionisation rate in the frequency band $k$. The unattenuated photoionisation of oxygen is then included as a non-thermal process in the chemical scheme as described in Sect. 2, with the rate calculated from Eq. (9) for the frequency bands $k$, and the attenuated rate is calculated at each integration time step according to Eq. (11).

### 4. Results

We study the variation of the abundances and masses of chemical species and dust clusters that form in the ejecta of the Cas A supernova precursor and present the results in Sect. 4.1. We then focus on an oxygen clump in Cas A, whose chemical composition has been derived in Sect. 4.1, and study the impact of the RS on such a dense clump. Various RS velocities that correspond to clump/interclump density enhancement of 100–1000 are considered, and results are presented in Sect. 4.2.

#### 4.1. Molecules and dust clusters in the Cas A supernova progenitor

Type IIb SNe have low-density ejecta compared to their type II-P counterparts. In Sect. 4.1.1, we present the results for a stratified ejecta whose parameters are given in Table 4. We define this low-density type IIb SN ejecta as our “standard case”. Results on the impact of increasing the gas density on the chemical composition of the gas and the dust are presented in Sect. 4.1.2.

![Fig. 2. Masses of molecules and dust clusters in the ejecta as a function of post-explosion time for the type IIb SN with 19 $M_\odot$ progenitor that led to Cas A. The ejecta parameters are given in Table 4.](image)

**4.1.1. Low-density type IIb ejecta**

The masses of the prevalent molecules and dust clusters that form in all the ejecta zones as a function of post-explosion time are shown in Fig. 2 for the “standard case”. The dominant species is SiS, which efficiently forms in zones 1A/B of the ejecta from the radiative association reaction

$$\text{Si} + \text{S} \rightarrow \text{SiS} + h\nu.$$  \hspace{1cm} (12)

This reaction has already been identified as the prevalent formation process for SiS in the innermost zones of the denser ejecta of type II-P SNe (SC13). This reaction is temperature-independent, and it can therefore proceed at late post-outburst time and ensure the growth of SiS mass with time. The SiS destruction is provided by the reaction with Ar$^+$. Carbon monoxide forms in the oxygen-rich zones 3A/B and 2 via the radiative association reaction

$$\text{C} + \text{O} \rightarrow \text{CO} + h\nu;$$  \hspace{1cm} (13)

and the reaction with $\text{O}_2$

$$\text{C} + \text{O}_2 \rightarrow \text{CO} + \text{O}.$$  \hspace{1cm} (14)

The CO destruction stems from charge-exchange reaction with $\text{O}^+$ and with reaction with Ne$^+$. Silicon monoxide primarily forms in zones 1B and 2 with a small contribution from zones 3A/B from the following radiative association process

$$\text{Si} + \text{O} \rightarrow \text{SiO} + h\nu;$$  \hspace{1cm} (15)

while SiO is destroyed by the reaction with Ne$^+$. Its abundance increases until ~1300 when SiO starts to be depleted in SiO dimers, Si$_2$O$_2$, and trimers. Finally, SO and $\text{O}_2$ also form from radiative association reactions in zone 1A and 1B/2/3, respectively. Inert gas ions do not efficiently recombine at an early time because of the low ejecta gas density, and their abundance gradually decreases owing to the decrease in $^{56}\text{Ni}$ mass and $\gamma$-ray photons with time. The formation of the prevalent molecules is then postponed to $t > 1000$ days, in contrast with type II-P SNe (SC13).

The masses of the most abundant synthesised dust clusters are also shown in Fig. 2. A small amount of Si$_5$O$_5$ is formed in the O-rich zones 1B, 2, and 3A/B out of SiO and its polymerisation. The dimerisation of SiO is possible after day 1000. Further growth is inhibited owing to the very low ejecta gas density. No significant amount of silicate dust clusters is formed in the O-rich zones of the ejecta because the first chemical steps in
the synthesis of forsterite and enstatite dimers involve neutral-neutral processes with a moderate activation barrier (Goumans & Bromley 2012, SC13), and higher gas temperatures than those found at day $\sim 1000$ in the ejecta are thus required. The low-gas density at day $>1000$ also hampers the efficient nucleation of silicate clusters. Some SiC dimers form in the carbon-rich outermost zones 4A/B after day 1800. As for SiO polymers, their later growth is hampered by the low gas density at late times. The masses of molecules and dust clusters at 3000 days after explosion and for the standard case are summarised in Table 6. We see that the mass fraction of the ejected material in the form of molecules only amounts to 1.2% and that dust clusters form in small amount. Therefore, the low-density, homogeneous ejecta of a type I Ib SN seems to be inefficient at forming significant quantities of dust clusters and grains.

These results are in contrast with the theoretical study of dust formation in the SN progenitor of Cas A presented by Nozawa et al. (2010), where the authors derive a total dust mass of $0.167 \, M_\odot$ for similar ejecta conditions and a complex chemical composition for the dust, including silicates, carbon, and metal oxides. Their study of dust synthesis is based on a CNT formalism and assumes that all available carbon and silicon atoms are locked up in CO and SiO when the C and Si elemental abundances are less than that of oxygen. The formalism fails to include the chemistry of the ejecta gas phase, and the formation and destruction of molecules and dust clusters from chemical kinetics. We find that only $\sim 1 \times 10^{-3} \, M_\odot$ of silicon carbide, silica, and alumina dust clusters can form for the standard case. Our results point to the importance of the nucleation phase, which describes the formation of molecules and dust clusters out of the elements comprised in the ejecta zones, as a bottleneck to dust condensation. This issue has already been addressed by Cherchneff & Lilly (2008), Cherchneff & Dwel (2009, 2010), and SC13 and will be discussed in Sect. 5. On the observational front, the Spitzer data of Cas A reveal that the IR excess due to dust in the remnant can be reproduced by an ensemble of dust grains of various composition, including silicates, metal oxides, metal sulphides, pure metal grains, and some amorphous carbon grains (Rhio et al. 2008; Arendt et al. 2014). Dust is clearly present in the diffuse remnant phase, implying that it could form in the denser ejecta of the supernova. This is in contrast with our present findings and forces us to reconsider the impact of the gas density on the nucleation of dust.

### Table 6. Masses of molecules and dust clusters (in $M_\odot$) at 3000 days post-explosion for the type I Ib SN with a 19 $M_\odot$ progenitor that led to Cas A.

<table>
<thead>
<tr>
<th>Density case</th>
<th>Standard</th>
<th>×200</th>
<th>×2000</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Molecules</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SiS</td>
<td>$4.1 \times 10^{-2}$</td>
<td>$1.3 \times 10^{-1}$</td>
<td>$5.7 \times 10^{-2}$</td>
</tr>
<tr>
<td>CO</td>
<td>$4.8 \times 10^{-3}$</td>
<td>$4.9 \times 10^{-1}$</td>
<td>$4.9 \times 10^{-1}$</td>
</tr>
<tr>
<td>SiO</td>
<td>$6.7 \times 10^{-10}$</td>
<td>$8.7 \times 10^{-12}$</td>
<td>$1.8 \times 10^{-12}$</td>
</tr>
<tr>
<td>O$_2$</td>
<td>$9.5 \times 10^{-9}$</td>
<td>$5.8 \times 10^{-1}$</td>
<td>$1.07$</td>
</tr>
<tr>
<td>SO</td>
<td>$2.9 \times 10^{-7}$</td>
<td>$2.6 \times 10^{-2}$</td>
<td>$7.7 \times 10^{-2}$</td>
</tr>
<tr>
<td>Total Mass</td>
<td>$4.6 \times 10^{-2}$</td>
<td>$1.22$</td>
<td>$1.69$</td>
</tr>
<tr>
<td>Efficiency</td>
<td>$1.2%$</td>
<td>$31%$</td>
<td>$43%$</td>
</tr>
<tr>
<td><strong>Clusters</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Silicates$^a$</td>
<td>–</td>
<td>$3.2 \times 10^{-3}$</td>
<td>$1.3 \times 10^{-2}$</td>
</tr>
<tr>
<td>SiC</td>
<td>$7.9 \times 10^{-6}$</td>
<td>$3.5 \times 10^{-4}$</td>
<td>$1.3 \times 10^{-4}$</td>
</tr>
<tr>
<td>Silica$^a$</td>
<td>$2.7 \times 10^{-7}$</td>
<td>$1.4 \times 10^{-9}$</td>
<td>$8.0 \times 10^{-15}$</td>
</tr>
<tr>
<td>Alumina</td>
<td>$1.7 \times 10^{-8}$</td>
<td>$1.7 \times 10^{-2}$</td>
<td>$1.4 \times 10^{-2}$</td>
</tr>
<tr>
<td>Pure metals$^a$</td>
<td>$4.5 \times 10^{-10}$</td>
<td>$8.8 \times 10^{-5}$</td>
<td>$8.0 \times 10^{-5}$</td>
</tr>
<tr>
<td>Carbon</td>
<td>–</td>
<td>$7.5 \times 10^{-12}$</td>
<td>$9.4 \times 10^{-3}$</td>
</tr>
<tr>
<td>Iron sulphide</td>
<td>–</td>
<td>$6.6 \times 10^{-10}$</td>
<td>$6.2 \times 10^{-7}$</td>
</tr>
<tr>
<td>Total Mass</td>
<td>$8.2 \times 10^{-6}$</td>
<td>$0.021$</td>
<td>$0.045$</td>
</tr>
<tr>
<td>Efficiency</td>
<td>$0.0002%$</td>
<td>$0.52%$</td>
<td>$1.1%$</td>
</tr>
</tbody>
</table>

Notes. The gas density cases correspond to the “standard case” and increase of 200 and 2000 in the gas number density, as discussed in the text. Efficiencies are the mass fraction of the ejected material in the form of molecules or dust clusters. $^a$Silicates correspond to forsterite clusters, pure metals to Fe, Si, and Mg clusters, and silica to polymers of SiO.

![Fig. 3. Masses of molecules and dust clusters in the ejecta as a function of post-explosion time for the type I Ib SN with 19 $M_\odot$ progenitor that led to Cas A, assuming a number density increase of 200 with respect to the “standard case”. Top: masses of molecules. Bottom: masses of dust clusters.](image-url)
in the previous section because of the high gas density. While radiative association reactions contribute to the synthesis of all molecules in the ejecta, bimolecular processes, including reactions with O$_2$, contribute to forming CO, SO, and AlO in the various zones. The species SiS is essentially formed in the innermost zones 1A/B. As shown by SC13 for type II-P SNe, CO is mostly synthesised in the O-rich zones 2 and 3A/3B and does not trace the formation of carbon dust clusters. The molecule SiO primarily forms in zone 1B and 2 and gets quickly depleting in silicate clusters after day $\sim$600. The dust clusters start to form as early as 400 days post-explosion for the pure metal clusters, and significant masses of alumina and forsterite clusters form in the O-rich zones after day 800.

The masses of dust clusters at day 3000 are shown in Fig. 4 as a function of gas number density increase in the ejecta. The build-up of the chemical complexity of the dust in the ejecta is clearly seen. As seen in the previous section, small amounts of SiC, silica, and alumina dust clusters form for the "standard case" given by the explosion model of a type IIf SN with 19 $M_\odot$ progenitor that led to Cas A. The x axis represents the increment of density with respect to our "standard case" (which corresponds to $x = 1$).

Fig. 4. Masses of dust clusters in the ejecta at 3000 days post-explosion as a function of density increase for the type IIf SN with 19 $M_\odot$ progenitor that led to Cas A. The x axis represents the increment of density with respect to our “standard case” (which corresponds to $x = 1$).

The masses of molecules and dust for the cases $x = 200$ and $x = 2000$ are summarised in Table 6 at day 3000 after explosion. We clearly see that the efficiency at forming molecules and dust clusters increases with gas density enhancement, but species are affected in different ways. For example, O$_2$ and SO are more responsive to a rise in gas density owing to the larger number of chemical reactions involved in their formation and since SO forms from O$_2$ at high density (SC13). More generally, we conclude that a low-density stratified ejecta as derived from non-clumpy explosion models of type IIf SNe are not conducive to the synthesis of large amounts of molecules and are almost dust-free. High-density clumps thus need to be considered as the main sources of molecules and dust in the SN ejecta to explain the presence of dust observed at IR wavelengths in Cas A. These clumps may retain the chemical composition of the ejecta zones from where they originate and evolve and are shocked by the RS in the remnant phase.

4.2. Molecules and dust clusters in the reverse shock

The FMKs have been observed in Cas A and interpreted as dense ejecta clumps that experience the crossing of the RS (Chevalier & Kirshner 1978). Their chemical composition includes a large fraction of oxygen atoms, along with neon, silicon, sulphur, argon, and iron. This specific composition coincides with that of the oxygen-rich zones of the SN ejecta. Radiative shock theoretical models are able to reproduce optical and IR lines observed in these knots and provide information on the physical conditions at the RS (Itoh 1981, BS90; Sutherland & Dopita 1995, DS10). We want to assess the impact on the RS on a FMK in terms of molecule and dust cluster reprocessing and thus, consider an oxygen-rich clump whose chemical composition is derived from our SN ejecta study presented in Sect. 4.1.2. We follow the chemistry in the post-shock gas using the analytical model for the RS, as illustrated in Fig. 1 and described in Sect. 3.2. We consider the five shock velocities of Table 5. According to BS90, oxygen atoms are fully ionised to different ionisation states in the HR. The high HR temperature ($\sim 10^5$ K) ensures that all molecule and dust clusters present in the pre-shock clump are destroyed by the RS and atoms are ionised in the HR. We then model the chemistry in the PIR, assuming all atoms are singly ionised as initial conditions, and study the chemistry, including the potential reformation of small dust clusters.

Abundances for the prevalent atoms, ions, and molecules in the PIR are shown in Fig. 5 for the 200 km s$^{-1}$ shock. This shock model corresponds to the PIR conditions derived by DS10 with a different initial chemical composition. The ionisation of O atoms by the UV field coming from the HR is effective at maintaining a high ionisation fraction until $\sim$150 days, despite the rapid recombination to O atoms before $\sim$25 days. The O$^+$ ions then recombine and exchange charge with other atoms to sustain an ionisation fraction of $\sim$0.03, which is specifically through charge exchange reaction with Mg. Molecules chiefly reform from the gas phase in the PIR and CO has the largest abundance, followed by SiO, O$_2$, and SiS. The formation processes involved are radiative association reactions, such as Reactions (12), (13), or (15). Other bimolecular processes that are active in the dense ejecta of type II-P SNe are not efficient at the low-gas number density of the PIR (in this case $n_{\text{gas}} = 10^6$ cm$^{-3}$), e.g., Reaction (14). The CO and O$_2$ reform with lower abundances by a factor $\geq$1000 than their initial values in the clump before the passage of the RS, while SiS reforms with a similar abundance. The SiO is an exception because the reformed gas-phase abundance is higher than its pre-shock value. As seen in Sect. 4.1.2, SiO is quickly
because of the low-gas density involved. We explore the possibility of synthesising dust clusters by artificially increasing the gas number density in the PIR for the 140 km s$^{-1}$ RS. We consider two PIR enhanced densities, $10^7$ cm$^{-3}$ and $10^9$ cm$^{-3}$, and results are presented in Fig. 7. We see that even the largest gas density fails at building up significant amounts of dust cluster, with a maximum abundance of $\sim 10^{-7}$ for forsterite and aluminas dimers and (SiO) trimers. The PIR corresponds to a region, where both the gas temperature and number density are high compared with values that characterise the pre-shock gas in the clump and non-shocked gas in the remnant. This environment is thus more conducive to dust cluster formation from the gas phase. If dust clusters are unable to form from the gas phase in this region, we can then anticipate that the dust grains destroyed by the RS crossing dense gas clumps are unable to reform in the remnant from the gas phase.

We have so far considered the effects of the PIR region after the passage of the RS. Since molecules reform but dust clusters do not for the standard physical conditions of the PIR region given by Table 5, we want to investigate whether molecules could keep forming over a time span $t \gg \tau_{PIR}$. Following Sutherland & Dopita (1995), the clump-crushing time is given by $\tau_{cc} = \chi^{1/2}/v_0$, where $v_0$ is the pre-RS velocity and $\chi$ is the density contrast defined in Sect. 3.2. According to Silvia et al. (2010), typical clump-crushing times for the $\chi$ factors and pre-RS velocities considered in this study range from 6 to 100 years. We thus consider the RS shock model with $V_s = 200$ km s$^{-1}$, keep the PIR number density constant over a period of ~15 years, and decrease the gas temperature to 300 K to account for radiative cooling. The abundances of the prevalent molecules are shown in Fig. 8. Most molecules slowly keep forming from radiative association processes, while dust clusters form with negligible abundances. The final abundance values for molecules at $\tau_{cc}$ remain all below $10^{-3}$ for $6 < \tau_{cc} < 100$ years. We can thus conclude that molecules quickly reform in the dense and warm PIR region and continue forming in the post-RS gas until $t = \tau_{cc}$. We also confirm that dust clusters are unable to form from the gas phase in the PIR and at later times in the shocked clump.

5. Discussion

We have investigated the synthesis of molecules and dust clusters in the stratified ejecta of the Cas A type IIn SN progenitor. Focussing on the oxygen-rich material formed in the oxygen-rich zones of the ejecta, we have studied their processing by the RS in a dense clump of the Cas A remnant. Our results highlight the following points:

- The stratified, homogeneous ejecta of a type IIn SN is too diffuse to form large amounts of molecules and dust. For normal ejecta conditions, only $\sim 10^{-2} M_{\odot}$ of molecules and $\sim 10^{-5} M_{\odot}$ of silicon carbide and silica dust clusters form. When the gas density is raised, the masses of molecules and clusters also increase, and most importantly, the chemical complexity of the dust composition is enhanced.
- In the remnant, molecules and dust clusters in the gas phase of oxygen-rich clumps are destroyed by the crossing of the RS but chemical species, including CO, SiO, SiS, and O$_2$, reform in the post-shock gas. Stronger shocks lead to denser post-shock, PIRs and thus higher abundances of reformed molecules.
- No clusters of silicate, silica, silicon carbide, amorphous carbon, or pure metal grains can reform in the back of the shock.
The main results of this study point to the importance of density in the formation processes of molecules and dust in SNe and their remnants. In the homogeneous ejecta of a type IIb SN, we find that no molecular cluster precursors to dust could form at the low-gas density involved. This points to the complexity of the dust formation processes at a microscopic scale and that dust nucleation is driven by non-equilibrium chemistry in the gas-phase. Existing studies based on the CNT formalism ignore the dust nucleation bottleneck and indicate fairly large masses of grains formed in type IIb SNe (e.g., Nozawa et al. 2010). Our present results are in contrast with these studies and show that the formation of dust is extremely gas-density dependent. This

because the post-shock gas densities are too low to permit the chemical build-up of dust clusters and the formation of dust grains.

Fig. 6. Abundances of molecules in the PIR (with respect to total gas number density) for various shock velocities. Top left: 140 km s\(^{-1}\). Top right: 100 km s\(^{-1}\). Bottom left: 70 km s\(^{-1}\). Bottom right: 35 km s\(^{-1}\). All shock models are from BS90.

Fig. 7. Abundances of dust clusters in the PIR (with respect to total gas number density) for the 140 km s\(^{-1}\) reverse shock. Dashed lines and full lines are for a gas density of 10\(^7\) cm\(^{-3}\) and 10\(^9\) cm\(^{-3}\), respectively.

Fig. 8. Abundances of molecules (with respect to total gas number density) as a function of time in the post-shock region of the 200 km s\(^{-1}\) RS. The vertical dotted line indicates \(t = t_{\text{PIR}}\), and results for \(t < t_{\text{PIR}}\) are similar to those of Fig. 5.
dependency does not pertain to the condensation phase of dust clusters but to their formation phase, or nucleation out of the gas phase. Increase in gas density by a factor of 200–2000 permits an efficient nucleation of dust grains and indicates the need for non-homogeneous, type Ibb SN ejecta in the form of clumps or knots.

The chemical complexity of the dust grains that form in SN ejecta is also density dependent and grows with increasing gas density. In particular, carbon dust only forms at high gas number densities and traces the densest SN ejecta conditions. In Cas A, the IR spectra measured by Spitzer indicate a variety of spectral signatures ascribed to warm dust, which vary according to position and observed atomic lines in the remnant (Rho et al. 2008; Arendt et al. 2014). These grains include silicates, aluminas, pure metals, carbides, iron sulphide, and amorphous carbon and have been processed by the RS, but they originate from the SN ejecta that led to Cas A. According to our results, the SN ejecta gas must be in the form of dense clumps with a density contrast of at least 200 compared to the homogeneous type Ibb SN ejecta to properly account for the dust chemical diversity in Cas A that is inferred from the IR data. More generally, the chemical type of dust detected in SNe and SNRs can be used as a tracer of the physical conditions of the dust birthplace, which is a denser gas leading to dust grains of greater chemical variety.

Recent observations of high excitation energy rotational lines of CO in one region of Cas A with Herschel, where CO was already detected at the RS position with Spitzer, have brought evidence of hot and dense gas in the remnant (Wallström et al. 2013). According to the line analysis, this O/CO-rich region is characterised by warm and dense gas ($n_{\text{gas}} \sim 10^6 \text{ cm}^{-3}$) and probably corresponds to post-RS gas. Such conditions are found in the wake of our 200 and 140 km s$^{-1}$ RS crossing a dense clump, where CO reforms on a time scale of $\sim 100$ days. These observations therefore support the present model results, which indicate that molecules, including CO and SiO, reform after the RS passage in sufficient amount to warrant detection. The molecular content of Cas A should then include two components reflecting various evolutionary phases of the remnant. A cold component would comprise species formed in the SN clumpy ejecta (e.g., SiS, CO, O$_2$, and SO), which have not yet been processed by the RS and have large abundances. Such a cool component has been confirmed in SN1987A by the detection of low excitation rotational lines of CO with ALMA (Kamenetzky et al. 2013). A warm/hot component would coincide with shocked clumps and newly reformed molecules (e.g., CO and SiO) in the post-RS gas, as seen in Cas A with Herschel. The warm CO component should have a lower abundance than its cold counterpart. The amount of reformed molecules depends on the shock strength: the faster shocks produce larger quantities of chemical species in the post-shock gas. For all shock speeds, dust clusters are unable to reform in sufficient quantities in the post-shock gas to guarantee a viable dust formation pathway from the gas phase in SN remnants.

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References

Cherchneff, I. 2010, ASP Conf. Ser., 425, 237
Danziger, I. J., Lucy, L. B., Bouchet, P., & Gouiffes, C. 1991, in Supernovae, the tenth Santa Cruz Workshop in Astronomy and Astrophysics, ed. S. E. Woosley (New York: Springer-Verlag), 69
Itoh, H. 1981, PASJ, 33, 1