

CARBONATE FORMATION IN NON-AQUEOUS ENVIRONMENTS BY SOLID-GAS CARBONATION OF SILICATES

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Abstract. We have produced synthetic analogues of cosmic silicates using the Sol Gel method, producing amorphous silicates of composition $\text{Mg}_{(x)}\text{Ca}_{(1-x)}\text{SiO}_3$. Using synchrotron X-ray powder diffraction on Beamline I11 at the Diamond Light Source, together with a newly-commissioned gas cell, real-time powder diffraction scans have been taken of a range of silicates exposed to CO_2 under non-ambient conditions. The SXPDP is complemented by other techniques including Raman and Infrared Spectroscopy and SEM imaging.

1 Introduction

Dust is present throughout the universe, existing in a wide variety of astrophysical environments and playing a major role in star and planet formation. Throughout their lifetime, dust grains are exposed to high levels of radiation and extreme temperatures, leading to the evolution of the amorphous, structurally disordered grains, formed within the atmospheres of evolved stars (Gail & Sedlmayr 1998), to the highly processed materials that were present in the early solar nebula. We are investigating this processing and mineralization of grains using laboratory produced analogues of cosmic dust. In-situ synchrotron X-ray powder diffraction (SXPDP) is conducted on Beamline I11 at Diamond Light Source (Thompson *et al.* 2009, 2011) and is complemented by SEM imaging, Raman and FTIR spectroscopy. This has relevance to the existence of carbonates in circumstellar environments that are believed to form through solid-gas interaction of cosmic silicates with gaseous CO_2 at high temperatures (Rietmeijer *et al.* 2008). We describe preliminary proof-of-concept work which demonstrates that this process can indeed proceed.

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2 Experimental

Beamline I11 is equipped to simulate extreme temperature and gas pressure conditions ($20 < T < 1000^\circ\text{C}$, 1–100 bar gas pressure), through the use of a cyberstar hot air blower and gas cell capillary holder (Parker *et al.* 2012; Thompson *et al.* these proceedings). Amorphous magnesium and calcium silicates of composition $\text{Mg}_x\text{Ca}_{1-x}\text{SiO}_3$, where $0 < x < 1$, were produced in the laboratory as analogues of cosmic dust, using an adapted sol gel method (Thompson *et al.* 2012). To inhibit the direct formation of carbonate, the gels were dried *in vacuo* and the dried silicates stored under argon. Powdered samples were loaded into quartz capillaries in a gas cell and mounted onto the sample stage of the diffractometer. The heating and dosing of samples with CO_2 was controlled remotely allowing real time, in-situ measurements to be taken. SXPD data were collected using a fast Position Sensitive Detector (Thompson *et al.* 2011), specifically designed for fast data collection. Data were collected at regular intervals as the samples were put under vacuum and then exposed to CO_2 up to a pressure of 1 bar. While this is clearly higher than in circumstellar environments (so the reaction rate is correspondingly higher), the nature of the experimental setup requires such pressure in order to provide adequate diffusion of CO_2 through the high packing density of the material in the capillary. After a number of scans at room temperature the hot air blower was introduced and the temperature was increased in steps of 50°C , over a period of ~ 1 hour, up to a maximum of 900°C ; two 4-s scans were taken at each temperature.

3 Characterisation

3.1 Raman spectroscopy

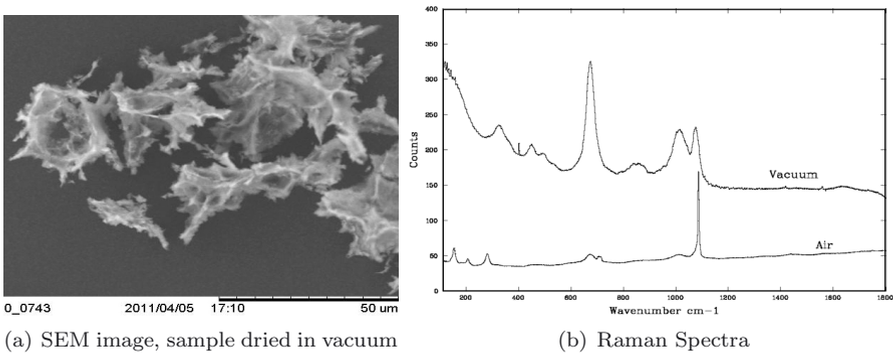


Fig. 1. SEM image of vacuum dried sample and a comparison of Raman Spectra of MgCaSiO_3 grains, dried in air and under vacuum.

The sol gels were dried in air and *in vacuo*, producing noticeably different end products. The gel dried in air produced solid, angular grains of a few mm to a few

cm in size, varying from clear to white in colour depending on composition. On the other hand, gels dried *in vacuo* produce a very fine white powder, with SEM images showing a branch-like network structure with individual grains no larger than a few μm in size (see Fig. 1a).

Raman spectra show that there is also a compositional difference between the samples dried in air and those dried *in vacuo* (see Fig. 1b). The air-dried samples exhibit features of crystalline calcium carbonate (calcite) exhibiting characteristic features at 712 cm^{-1} and 1088 cm^{-1} , while samples dried *in vacuo* show a predominately silicate composition with prominent features in the 670 cm^{-1} and 1000 cm^{-1} bands. This implies that the carbonation of the samples is due to the reaction of Ca with atmospheric CO_2 during the drying phase.

3.2 FTIR

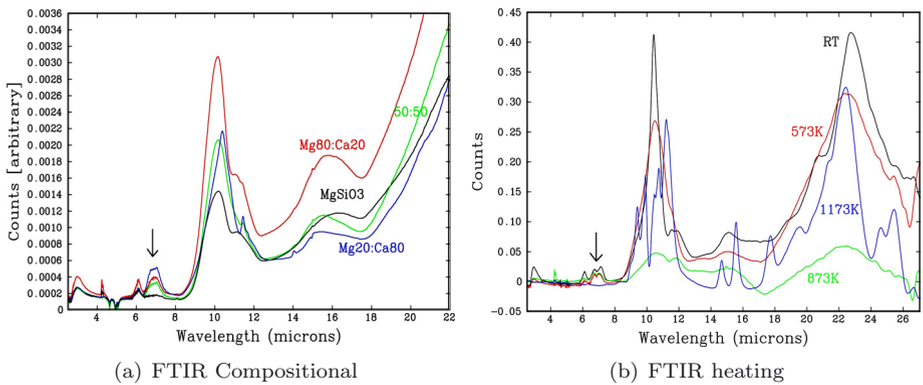


Fig. 2. Comparison of FTIR Spectra of MgCaSiO_3 grains, of varying composition and temperature.

FTIR spectra of the samples were taken for a range of compositions and annealing temperatures. Figure 2a shows the comparison of four compositions, $x = 0, 0.2, 0.8$ and 1 . A peak (arrowed) can be clearly seen at $7\ \mu\text{m}$ strengthening with increasing Ca content. Also with increasing Ca content, the broad silicate features around $11\ \mu\text{m}$ and $16\ \mu\text{m}$ seem to weaken. Figure 2b shows the spectra of a CaSiO_3 sample annealed at four different temperatures. Few differences are observed at temperatures of up to 873 K , but note the disappearance of the $7\ \mu\text{m}$ feature and the rise of crystalline silicate features at temperatures greater than 873 K . This indicates that any carbonate phase present within the sample becomes unstable at this temperature and breaks down.

3.3 Synchrotron X-ray powder diffraction

Figure 3a shows a sequence of *in-situ* SXP patterns of $\text{Mg}_{0.5}\text{Ca}_{0.5}\text{SiO}_3$ exposed to CO_2 at 1 bar pressure and heated to 1272 K over a period of a few hours. A feature

indicative of calcite is present at low temperatures but disappears above 873 K, again suggesting that the calcite phase is not stable at higher temperatures. The sample then begins to crystallize to diopside at just below 1173 K. Reitveld refinement of the powder diffraction patterns, using the TOPAS-Academic software package (Coelho 2007), has provided information about the relative weight percentages of the diopside and calcite phases within the sample as it is being heated. This shows an inverse correlation of the two phases (see Fig. 3b).

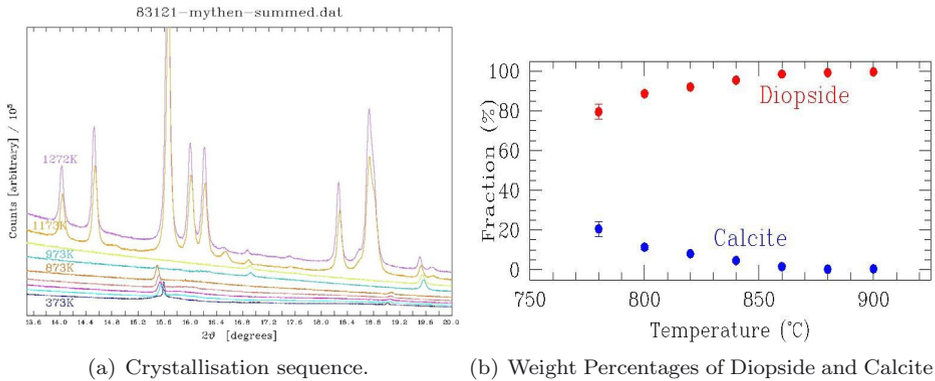


Fig. 3. Sequence of powder diffraction patterns showing the structural evolution of a silicate sample and derived weight percentages of diopside and calcite phases.

4 Results and conclusions

Analysis of these data is still ongoing and the crystallization temperatures and effect of CO₂ on the samples will be much better constrained through further experiments and analysis. At this stage we provide proof of physical principle regarding the formation of carbonates rather than a direct simulation of any astrophysical environment. However by knowing that the carbonate phase becomes unstable above 873 K, we can begin to constrain the astrophysical environments in which carbonates would be able to form and survive.

References

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