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Neutron diffraction as a probe of liquid and glass structures under extreme conditions

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Neutrons provide a unique tool for probing the structure of liquid and glassy materials, and deliver information that cannot be obtained from other experimental techniques. Advances in neutron diffraction instrumentation and measurement protocols now make it possible to measure the structure of these disordered materials under extremes of high temperatures or high pressures. Here, we consider the use of aerodynamic levitation with laser heating to explore the structure of glass-forming oxide melts at high temperatures, and the use of a Paris-Edinburgh press to investigate the mechanisms of density-driven network collapse for glassy materials in the gigapascal (GPa) pressure regime.

Introduction
The structural complexity of liquid and glassy materials makes for a challenging field of research to which neutron diffraction has made seminal contributions, especially when it has been combined with the isotope substitution method [1, 2]. The atomic-scale disorder of these materials leads to diffuse scattering, such that experiments requiring small samples and complex sample environments must be carefully designed if the sample scattering is to be distinguished from the container plus background scattering. Aerodynamic levitation with laser heating offers a container-less method for investigating the structure of a wide variety of high-temperature liquids [3], but the small levitated sample is partially obscured from an incident neutron beam by the material of the levitator’s nozzle (Figure 1). The Paris-Edinburgh press provides a means for accessing pressures in the GPa regime [4], but a small sample must be contained within a gasket such that scattering from the gasket and anvils of the press cannot be avoided [5]. However, as will be illustrated in the following, it is possible to circumvent these experimental challenges. Indeed, it is even possible to apply the method of neutron diffraction with isotope substitution (NDIS) to provide site-specific information on a material’s structure under extreme conditions [1, 2].

Figure 1. Schematic of an aerodynamic levitation experiment as performed using the D4c diffractometer at the ILL. A small spherical sample of diameter 2–3 mm is levitated by the flow of gas through a conical nozzle, and is heated by CO₂ lasers, where the laser beams enter the diffractometer bell jar via NaCl windows. The images to the left show a molten sample as viewed from both its top and its side. In the latter, the bottom of the sample is obscured by the nozzle material. © Louis Hennet. Reproduced by permission of Louis Hennet. Permission to reuse must be obtained from the rightsholder.
Structure of high-temperature liquids by neutron diffraction with isotope substitution

CaAl$_2$O$_4$ is one of a family of glass-forming materials that are widely studied because of their geological and technological relevance. Aerodynamic levitation with laser heating has been combined with the NDIS method using Ca isotopes in order to measure the structure of a single levitated liquid droplet of CaAl$_2$O$_4$ at a temperature of 1973 K [6]. The NDIS method has also been applied to measure the glass structure at ambient temperature [6]. The experiments employed the diffractometer D4c at the Institut Laue-Langevin (ILL), and the measured real-space difference functions $\Delta G_{\text{Ca}}(r)$ and $\Delta G_{\text{M}}(r)$ are shown in Figure 2. $\Delta G_{\text{Ca}}(r)$ has contributions from only the Ca-Ca and Ca-M pair-correlation functions, where M denotes either an Al or O atom, whereas $\Delta G_{\text{M}}(r)$ has contributions from the M-M pair-correlation functions alone. The first peaks in $\Delta G_{\text{Ca}}(r)$ and $\Delta G_{\text{M}}(r)$ correspond to the Ca-O and Al-O nearest-neighbours, respectively. The diffraction results are compared to those obtained by using molecular dynamics simulations with an ionic interaction model that accounts for anion polarisation and shape deformation effects. Upon vitrification, it is found that significant structural changes occur on both the local and intermediate length scales. For instance, edge-sharing Ca-centred polyhedra occur in the liquid phase and become far more numerous as the system is quenched to form a glass (Figure 3). Thus, although glass is often used as a proxy for the corresponding liquid in order to explore possible structural configurations at extreme conditions, these configurations may not be representative of the liquid phase in so-called “fragile” glass-forming systems, a taxonomy that includes CaAl$_2$O$_4$ and a range of magma-related materials [6].

Structure of silica glass by high-pressure neutron diffraction

Silica is the canonical network glass, whose pressure-dependent behavior is often used as a reference for silicate materials that have geophysical relevance. Figure 4 shows the pressure dependence of the total structure factor $S(Q)$ as measured by neutron and x-ray diffraction, where the information provided by these techniques is complementary: The relative values of the coherent neutron scattering lengths or x-ray form factors for Si and O ensure that the neutron and x-ray $S(Q)$ functions are more sensitive to the O or Si atom pair-correlation functions, respectively [7]. For instance, the neutron $S(Q)$ function

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**Figure 2.** The difference functions (a) $\Delta G_{\text{Ca}}(r)$ and (b) $\Delta G_{\text{M}}(r)$ as (i) measured for either a single droplet of liquid CaAl$_2$O$_4$ of diameter 2.54 mm at a temperature of 1973 K or for a single bead of glassy CaAl$_2$O$_4$ of diameter 3.18 mm at room temperature (solid curves); and (ii) modelled by using molecular dynamics (chained curves). Figure has been redrawn from the data shown in [6].

Here, diffraction patterns are measured for two or more samples that are identical in every respect, except for the isotopic enrichment of one or more of the chemical species. These patterns are then subtracted, thus reducing the number of overlapping pair-correlation functions that contribute towards a single diffraction pattern.

**Figure 3.** Snapshots illustrating the largest clusters of edge-sharing Ca-centred polyhedra in molecular dynamics simulations of liquid and glassy CaAl$_2$O$_4$ at temperatures of 2500 K and 350 K, respectively [6]. © American Physical Society. Reproduced by permission of American Physical Society. Permission to reuse must be obtained from the rightsholder.
The results show that the transformation from a network of corner-sharing SiO₄ tetrahedra to a network of edge- and corner-sharing SiO₆ octahedra proceeds via the formation of five-fold coordinated Si atoms. In particular, the application of pressure leads to a reduction in size of the primitive rings that are formed by linked Si-centred motifs, a reduction that can be understood quantitatively in terms of a newly developed ‘zipper’ mechanism for ring closure [7]. The ring in Figure 5a is deemed to be ‘primitive’ because it cannot be decomposed into smaller rings. When pressure is applied, the ring closes when silicon atom Si¹ forms an additional bond with an intra-ringing oxygen atom O¹, thus increasing the Si¹ coordination number from four to five and the O¹ coordination number from two to three. With increasing density, a further ring closure event then takes place at an adjacent site to create another over-coordinated silicon atom Si² along with another over-coordinated oxygen atom O². The close proximity of these ring closure events helps to preserve locally the glass stoichiometry and, because the Si and O atoms are charged, this acts in a direction that helps to conserve local charge neutrality.
Structure of GeSe₂ glass by high-pressure neutron diffraction with isotope substitution

GeSe₂ is a prototypical chalcogenide glass-forming system in which the topological and chemical ordering contrasts to that found in typical amorphous oxides. The structural information obtained on this system from conventional neutron and x-ray diffraction experiments is almost identical because of the similarity between the coherent neutron scattering lengths of Ge and Se of natural isotopic abundance, and the similarity between the x-ray form factors for these elements [2]. It is therefore desirable to obtain site-specific structural information, where the NDIS method with Ge and/or Se isotopes offers an excellent way forward [10]. This method has now been used to explore the structure of GeSe₂ glass at pressures up to 8.2 GPa, and the resultant difference functions \( \Delta F_{\text{Ge}}(Q) \) and \( \Delta F_{\text{Se}}(Q) \) are shown in Figure 6 [11]. These functions have contributions solely from the Ge or Se atom pair-correlation functions, respectively, and were measured by using the diffractometer D4c (ILL) following the methodology developed in Ref. [12]. The structure of the glass containing Ge and Se of natural isotopic abundance was also measured at pressures up to 16.1 GPa by using the diffractometer PEARL (ISIS). The pressure-dependent structure of glassy GeSe₂ was modelled by using first-principles molecular dynamics [11].

The results give an ambient pressure network for GeSe₂ glass that is built predominantly from corner and edge sharing Ge-centred tetrahedra, where Ge and Se are four-fold and two-fold coordinated, respectively, and the chemical ordering is broken by the appearance of Ge-Ge and Se-Se homopolar bonds [10, 11]. As the pressure is increased to ~8.5 GPa, the edge-sharing tetrahedra persist in substantial numbers and the mean coordination numbers of Ge and Se remain unchanged. At larger pressures, a mediating role is found for homopolar bonds in the appearance of higher-coordinated Ge and Se atoms. The persistence of edge-sharing configurations and role played by homopolar bonds are likely to be generic features of the densification processes in other chalcogenide glasses.

Conclusions

The results presented above highlight the quality of information that can now be gained on the structure of liquid and glassy materials under extreme conditions from in situ neutron diffraction experiments. There is much to look forward to as the methodology is improved to enable the structure and properties of disordered materials to be explored over a broader range of state conditions.

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