

Understanding disorder in the $\text{Y}_2\text{Sn}_{2-x}\text{Zr}_x\text{O}_7$ pyrochlore oxides

Thursday, 12 November 2020 17:29 (1 minute)

Solid oxide fuel cells (SOFCs) offer clean alternatives to current carbon emitting energy sources; however, reducing their operation temperature is a major challenge for widespread application. Hence, the development of novel electrolyte materials with high room temperature ionic conductivity is crucial. Pyrochlore oxides, of the general chemical formula $\text{A}_2\text{B}_2\text{O}_7$, exhibit chemical and structural flexibility, resulting in a diverse range of physical properties and technological applications, such as host matrices in the immobilization of actinide-rich nuclear wastes. In particular, they have gained interest as fast-ion conductors for electrolytes in SOFCs.

The pyrochlore structure is highly ordered, which limits long-range oxygen diffusion. This can be altered via disordering, which results in the formation of oxygen Frenkel pairs that improve conductivity. The pyrochlore structure can adopt the disordered-fluorite structure via changes in composition, temperature, pressure and radiation. The disordered-fluorite exhibits lower formation energy of the Frenkel defect; however, this increase in structural disorder can increase the activation energies needed for long-range migration, which results in optimal conductivity occurring in partially disordered materials. Hence, the interplay between disorder and order in the atomic structure is key to the physical properties of these materials.

There have been many studies dedicated to understanding the structural order and disorder in pyrochlore and disordered-fluorite oxides, with a recent study claiming the local structure of the disordered-fluorite to be weberite. It was proposed that the overall structure of a disordered-fluorite consists of randomly arranged orthorhombic weberite domains that result in the long-range cubic disordered-fluorite.

In this study we use low temperature (15 K) neutron pair distribution function (PDF) and big box modelling to understand the local-scale structure of the $\text{Y}_2\text{Sn}_{2-x}\text{Zr}_x\text{O}_7$ system. We show that the local structure of the $\text{Y}_2\text{Zr}_2\text{O}_7$ disordered fluorite does not contain ordered weberite domains, which emphasizes the importance of low temperature measurements in the local structure analysis of disordered oxide materials. Our analysis of the $\text{Y}_2\text{Sn}_{2-x}\text{Zr}_x\text{O}_7$ series serves as a direct method for quantifying disorder and Frenkel defects. Understanding and quantifying these atomic-scale distortions is essential in simulations and design as it directly influences the energy landscape for anion mobility. These techniques can be used in the development and engineering of novel and advanced electrolyte materials for SOFCs.

Level of Expertise

Early Career <5 Years

Speakers Gender

Male

Do you wish to take part in the poster slam

Primary author: MARLTON, Frederick (University of Sydney)

Co-authors: KENNEDY, Brendan (The University of Sydney); LING, Chris (University of Sydney); ZHANG, Zhaoming (ANSTO)

Presenter: MARLTON, Frederick (University of Sydney)

Session Classification: Poster Session

Track Classification: Advanced Materials