Predicting the thermal conductivity of pyrochlore and defect fluorite nuclear waste forms using experimental data and density functional theory simulations

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Abstract:

A2B2O7 pyrochlores are considered as potential matrices for permanent immobilization of Pu and minor actinides as one potential nuclear waste form for deep geological disposal. Over the long course of disposal, the waste forms will be subjected to self-irradiation from immobilized radionuclides. The so generated heat could influence the materials by causing cracking, altering corrosion resistance, or reducing phase stability. The ability of a waste form to effectively transfer heat, i.e. thermal conductivity is therefore a key parameter impacting the performance of nuclear waste forms in a deep geological repository [1]. Furthermore, the self-irradiation of the waste form will generate defects that can lead to an order/disorder transition from a pyrochlore to a defect fluorite structure. These structural changes are also expected to affect the thermal conductivity of pyrochlore-type waste forms. We used a combination of Density Functional Theory (DFT) and the Slack Model to calculate the intrinsic thermal conductivity of a series of Nd2-xZr2+xO7±y pyrochlores and defect fluorites with various Nd/Zr ratios. We used this computed data to interpret thermal conductivity measurements obtained by Laser Flash Analysis. In particular, we were able to separate the intrinsic and extrinsic, i.e. defect-related components using an approach developed by Tian et. al [2]. Knowledge of the changes in intrinsic and extrinsic thermal conductivity of pyrochlore and defect fluorite phases provides new insight into the evolution of heat transfer capabilities of pyrochlore-type nuclear waste forms during storage, and sheds light on the effect of atomic-scale ordering/disordering in these materials. This knowledge contributes to refining the understanding of the long-term performance of Nd2Zr2O7 pyrochlore-based waste forms.

References:
