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Hydration mechanisms and proton conduction in the mixed ionic-electronic conductors Ba₄Nb₂O₉ and Ba₄Ta₂O₉

Mixed conductors – materials that exhibit significant mobility of more than one type of charge carrier such as oxide ions, protons and electrons – have a range of important applications including solid oxide fuel cell membranes, electrodes, batteries and sensors. We recently studied the behaviour of hydrogen in the mixed ionic-electronic conductors γ -Ba₄Nb₂O₉ and 6H-Ba₄Ta₂O₉, using a combination of experimental (neutron diffraction and inelastic neutron scattering) and computational (*ab initio* molecular dynamics) methods. While these compounds have isostructural low-temperature polymorphs, they adopt distinct forms in the high-temperature conducting regime. We found that they also have distinct mechanisms for hydration and ionic conduction. Hydration of γ -Ba₄Nb₂O₉ is localised to 2-D layers in the structure that contain a 1:1 ratio of isolated but adjacent NbO₄ and NbO₅ polyhedra. OH⁻ and H⁺ ions combine with two polyhedra respectively to form complete layers of NbO₄OH polyhedra, giving rise to a stoichiometric hydrated form γ -III-Ba₄Nb₂O₉.1/3H₂O. Protons then diffuse through these 2-D layers by “hopping” between oxygen atoms on adjacent polyhedra. In the case of 6H-Ba₄Ta₂O₉, hydration occurs by intercalating intact water molecules into the structure up to a maximum of ~0.375 H₂O per formula unit. This explains the unusual local and long-range structural distortions in the hydrated form observed by neutron diffraction. Diffusion then occurs by water molecules moving between neighboring symmetry equivalent positions. These fundamentally different hydration and proton conduction mechanisms explain why 6H-Ba₄Ta₂O₉ has the less well-defined and higher maximum water content, while γ -Ba₄Nb₂O₉ has the higher proton conductivity.

Topic

Advanced Materials

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