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Origin of electrolyte-dopant dependent sulfur poisoning of SOFC anodes

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The mechanisms governing the sulfur poisoning of the triple phase boundary (TPB) of Ni\textendash{}X\textsubscript{2}O\textsubscript{3} stabilized zirconia anodes have been investigated using density functional theory. Calculated sulfur adsorption energies reveal a clear correlation between the size of the cation dopant X\textsuperscript{3+} and the sulfur tolerance of the Ni\textendash{}X\textsubscript{2}O\textsubscript{3} anode; the smaller the ionic radius, the higher the sulfur tolerance. The mechanistic study shows that the size of X\textsuperscript{3+} strongly influences X\textsubscript{2}O\textsubscript{3}'s surface energy, which in turn determines the adhesion of Ni to X\textsubscript{2}O\textsubscript{3}. The Ni\textendash{}X\textsubscript{2}O\textsubscript{3} interaction has a direct impact on the Ni\textendash{}S interaction and on the relative stability of reconstructed and pristine Ni(100) facets at the TPB. Together, these two effects control the sulfur adsorption on the Ni atoms at the TPB. The established relations explain experimentally observed dopant-dependent anode performances and provide a blueprint for future search for and preparation of highly sulfur tolerant anodes.

![Figure 1 Sulfur adsorption energy on Ni\textendash{}X\textsubscript{2}O\textsubscript{3}, relative to the corresponding value on a Ni(111) terrace, plotted versus the ionic radius of X\textsuperscript{3+}. HCP11 and Hollow14 are adsorption sites on reconstructed and pristine Ni(100) facets, respectively. The dotted line indicates the ionic radius of Zr\textsuperscript{4+}.](image)

Reference