

Boosted Ethanol Electrooxidation Using Rh Single Atom Decorated Pt Nanocubes

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Direct ethanol fuel cells (DEFCs) are increasingly garnering attention as portable power sources, owing to their superior mass-energy density compared to hydrogen and lower toxicity relative to methanol. Despite these advantages, achieving full electrooxidation of ethanol to produce 12 electrons per molecule remains a significant challenge, leading to suboptimal fuel utilization efficiency. In this study, we present a novel electrocatalyst composed of unalloyed, partially oxidized Rh single atoms dispersed on the surface of Pt nanocubes. This catalyst facilitates the complete oxidation of ethanol to CO₂ at an unprecedentedly low potential of 0.35 V. Through in situ X-ray absorption fine structure (XAFS) measurements and density functional theory (DFT) calculations, we demonstrate that the Rh single-atom sites are pivotal in promoting C–C bond cleavage and efficiently removing *CO intermediates. This work not only elucidates the crucial role of unalloyed, partially oxidized single-atom catalysts (SACs) in the ethanol oxidation reaction (EOR) but also introduces a distinctive single-atom strategy that leverages low-coordination active sites on shape-controlled nanocatalysts. This approach offers a promising pathway to enhance both the activity and selectivity of complex catalytic reactions.

Paper submission Plan

No

Best Presentation

Yes

Contribution track

ICABU WG4. Applications of Particle Beams

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