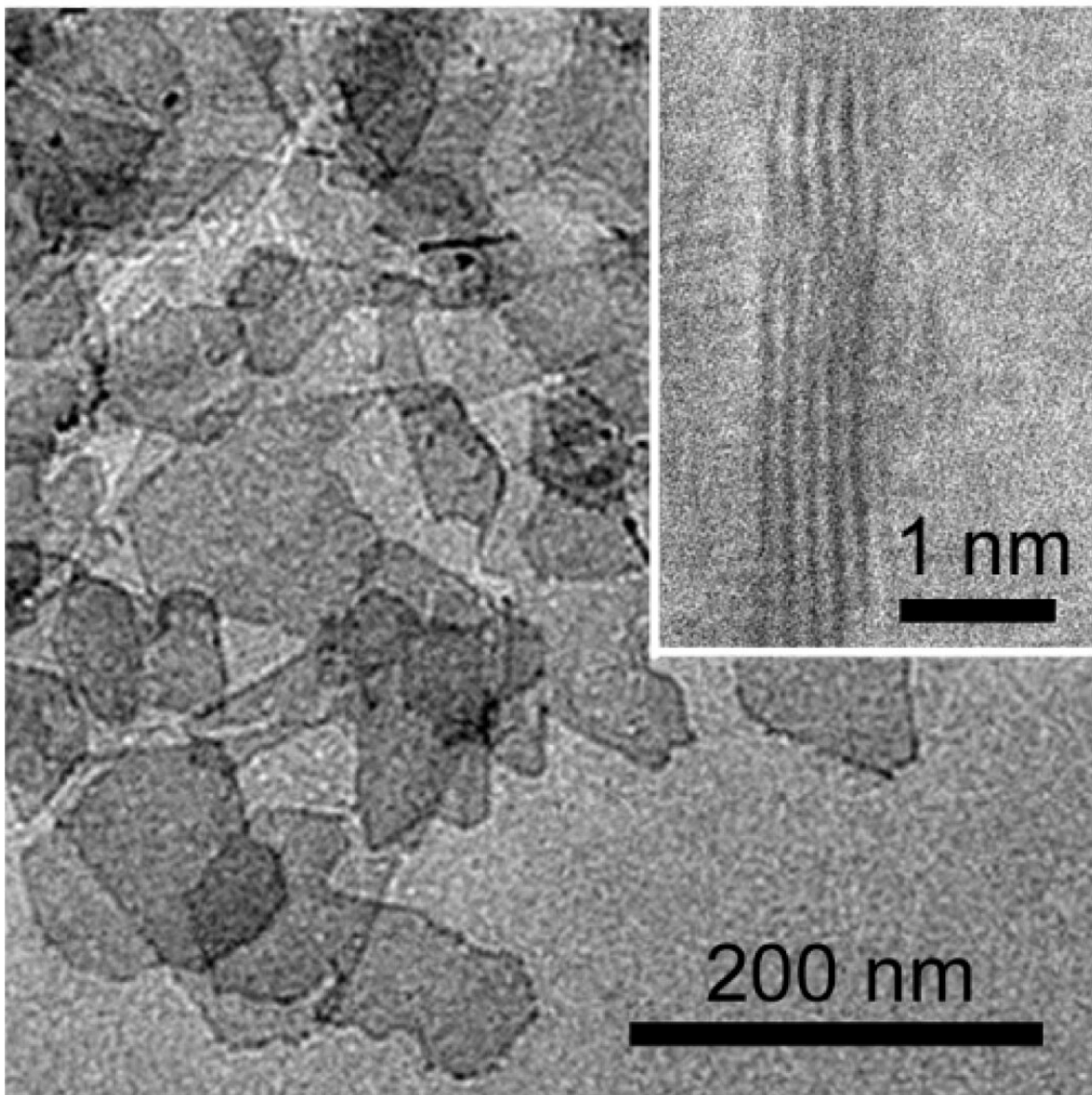


'Goldilocks' thinking to cut cost of fuel cells in electric vehicles

February 25 2019, by Kayla Wiles



A platinum-like metal only five atomic layers thick is "just right" for optimizing the performance of a fuel cell electrode. Credit: Johns Hopkins University image/Lei Wang)

The 2019 Toyota Mirai electric vehicle touts zero emissions, thanks to a fuel cell that runs on hydrogen instead of gasoline. But the Mirai has barely left California, partly because today's fuel cell electrodes are made of super expensive platinum.

Cutting down on the platinum would also cut costs, allowing more electric cars to hit the market.

A new method borrows some thinking from "Goldilocks" – just the right amount – for evaluating how much metal would be required for [fuel](#) cell electrodes. The technique uses the forces on a metal's surface to identify the ideal [electrode](#) thickness.

"There is exactly the right amount of metal that will give fuel cell electrodes the best properties," said Jeffrey Greeley, professor of chemical engineering at Purdue. "If they are too thick or too thin, the main reaction for deploying a fuel cell doesn't work as well, so there's sort of a Goldilocks principle here."

The study, to be published in the Feb. 22 issue of the journal *Science*, was a collaborative effort between Johns Hopkins University, Purdue University and the University of California at Irvine.

The researchers tested their theory on palladium, a metal very similar to platinum.

"We're essentially using force to tune the properties of thin metal sheets

that make up electrocatalysts, which are part of the electrodes of fuel cells," Greeley said. "The ultimate goal is to test this method on a variety of metals."

Fuel cells convert hydrogen, combined with some oxygen, into electricity through a so-called oxygen-reduction reaction that an electrocatalyst starts. Finding exactly the right thickness stresses the surface of the electrocatalyst and enhances how well it performs this reaction.

Researchers in the past have tried using outside forces to expand or compress an electrocatalyst's surface, but doing so risked making the electrocatalyst less stable.

Instead, Greeley's group predicted through [computer simulations](#) that the inherent force on the surface of a palladium electrocatalyst could be manipulated for the best possible properties.

According to the simulations, an electrocatalyst five layers thick, each layer as thin as an atom, would be enough to optimize performance.

"Don't fight forces, use them," said Zhenhua Zeng, a Purdue postdoctoral researcher in chemical engineering, and co-first and co-corresponding author on this paper. "This is kind of like how some structures in architecture don't need external beams or columns because tensional and compressive forces are distributed and balanced."

Experiments in Chao Wang's lab at Johns Hopkins confirmed the simulation predictions, finding that the method can increase catalyst activity by 10 to 50 times, using 90 percent less of the metal than what is currently used in [fuel cell](#) electrodes.

This is because the surface [force](#) on the atomically thin electrodes tunes

the strain, or distance between atoms, of the [metal](#) sheets, altering their catalytic properties.

"By tuning the material's thickness, we were able to create more strain. This means you have more freedom to accelerate the reaction you want on the material's [surface](#)," Wang said.

More information: Lei Wang et al. Tunable intrinsic strain in two-dimensional transition metal electrocatalysts, *Science* (2019). [DOI: 10.1126/science.aat8051](#)

Provided by Purdue University

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