

表面物理化学系列讲座 Surface Physical Chemistry Lecture Series

Single-Atom Catalysts: An Atomic-Scale View

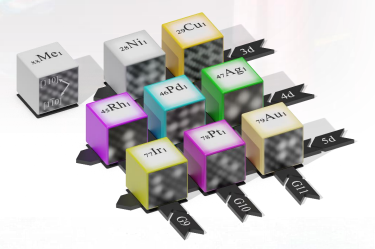
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Speaker	Prof. Gareth S. Parkinson Vienna University of Technology (TU Wien)
Date	2022.6.14, Tuesday, 15:00-17:00 (UTC+8)
Host	Prof. Kai WU 吴凯

ZOOM Meeting Number: 679 138 6278; Code: 61661

Abstract:

Understanding how the local environment of a "single-atom" catalyst affects stability and reactivity remains a significant challenge. In this talk, I will discuss under what circumstances single metal atoms can be stable on flat, well-ordered metal oxide surfaces, including examples from our work



on Fe₃O₄, Fe₂O₃, and TiO₂ single-crystal model supports. Thereafter, I will focus an in-depth study of Cu₁, Ag₁, Au₁, Ni₁, Pd₁, Pt₁, Rh₁, and Ir₁ species on Fe₃O₄(001); a model support where all metals occupy the same 2-fold coordinated adsorption site upon deposition at room temperature¹. Surface science techniques reveal that CO adsorption strength at single metal sites differs from the respective metal surfaces and supported clusters.² Charge transfer into the support modifies the d-states of the metal atom and the strength of the metal-CO bond. These effects could strengthen the bond (as for Ir₁-CO) ³ or weaken it (as for Ni₁-CO), but CO-induced structural distortions reduce adsorption energies from those expected based on electronic structure alone. The extent of the relaxations depends on the local geometry and could be predicted by analogy to coordination chemistry. In extreme cases, CO adsorption leads to sintering, and I will show that metastable (PtCO)₂ dimers are active for CO oxidation in the – nominally – Pt₁/Fe₃O₄(001) system.

References

- 1. R. Bliem et al., Science 346, 1215-1218 (2014)
- 2. J. Hulva et al., Science 371, 375 (2021)
- 3. Z. Jakub et al., Angew. Chem. Int. Ed. 58, 13961-13968 (2019).
- 4. M. Meier et al., Science Advances 8, eabn4580.(2022)