Exploring how surfaces change in contact with reactive gas phases under different conditions

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Phase diagram of the Si(100) surface in a D$_2$ gas phase by REGC method including phase boundaries (white line) and triple points as well as critical points. Credit: FHI / Y. Zhou

\[ C_{V, (T, p)} = \frac{\langle E^2 \rangle(T, p) - \langle E \rangle^2(T, p)}{k_B T^2} \]
Researchers at the NOMAD Laboratory at the Fritz Haber Institute have been engaged in describing how surfaces change in contact with reactive gas phases under different temperature and pressure conditions. For this purpose, they have developed the so-called replica exchange grand canonical method (REGC). The results were published in the journal *Physical Review Letters* on 17 June.

"Replica exchange" means that there are many replicas prepared for the silicon surface in contact with different hydrogen atmospheres. These replicas exchange with each other during the simulation. "Grand-canonical" means that the silicon surface in each replica exchanges deuterium atoms or molecules with the deuterium gas reservoir it touches, eventually reaching equilibrium with the deuterium gas reservoir.

Knowledge of the morphology and structural evolution of material surfaces in a given reactive atmosphere is a prerequisite for understanding the mechanism of heterogeneous catalysis reactions and electrocatalysis due to the structure-property-power relationship. In general, the reliable tracking of phase equilibria is of technological importance for the reasonable design of surface properties. Phase transitions are indicated by singularities of a reaction function (e.g. the heat capacity). FHI researchers have addressed this challenge by developing the Replica Exchange Grand Canonical (REGC) method in conjunction with molecular dynamics. The approach not only captures the restructuring of the studied surface under different reactive conditions, but also identifies surface phase transition lines as well as triple and critical points.

The dissociative adsorption of molecular hydrogen on the silicon surface has become a crucial criterion in the study of adsorption systems and has important applications such as surface passivation. The REGC approach is demonstrated using a silicon surface in contact with a deuterium
atmosphere. In the range of 300 to 1,000 Kelvin, the REGC approach identifies 25 different thermodynamically stable surface phases. Most of the identified phases, including some phase transitions between order and disorder, have not been observed experimentally before. It is also shown that the dynamic formation or breaking of Si-Si bonds is the driving force behind the phase transition between the experimentally confirmed adsorption patterns.

The REGC method makes it possible to combine traditional concepts of condensed matter statistical mechanics with state-of-the-art electronic structure calculations to predict stability phase diagrams of real systems. Furthermore, the approach has a significant impact on surface restructuring calculations in the field of surface science and is potentially relevant to a variety of important applications such as heterogeneous catalysis, electrocatalysis and surface segregation.


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