Adsorption and Dissociation of Iodine Molecules on a Zr Surface

Iodine is a fission product of uranium. It can attack the inner side of zircaloy cladding in nuclear power reactors leading to cracking and fracture. Computations show that iodine molecules adsorb and dissociate on a zirconium surface without an energy barrier. The binding energy of iodine on this surface is large (nearly 300 kJ mol\(^{-1}\) per iodine atom), but the barriers for surface diffusion is only 6.8 kJ mol\(^{-1}\). This gives rise to rapid surface diffusion allowing iodine atoms to reach the crack tips faster than the propagation of cracks.

Keywords: adsorption, dissociation, iodine, zirconium, computation

1 Technological context

Iodine has been reported to increase brittle fracture of zirconium with an increase of intergranular cracking [2]. This is a safety concern in nuclear power reactors since this type of fuel-cladding interaction can cause a weakening of zircaloy thus being one of the factors which limit the life time of a fuel assembly.

2 Computed results

Computations of the adsorption and dissociation of iodine molecules on the most stable zirconium surface, namely Zr(0001) surface, reveal that this process occurs without any energy barrier as illustrated in Figure 1. The adsorption energy is large (nearly 300 kJ mol\(^{-1}\) per I atom). The iodine isotherms (Figure 2), which is obtained by computing the difference in chemical potential between iodine adsorbed on the surface and in the gas phase, shows that a large fraction of the zirconium surface is covered by iodine even at very low partial pressures. Even though iodine binds strongly to the zirconium surface, simulations show that

Figure 1: Computed energy profile of the adsorption and dissociation of an iodine molecule on a Zr(0001) surface. Note the slight elevation of the surface zirconium atoms at point (C), indicating a strong Zr-I affinity.

Figure 2: Computed adsorption isotherms for adsorption of molecular iodine on the zirconium (0001) surface plotted for different temperatures [3].


3 Significance

The quantitative understanding and description of interaction of fission products with cladding materials is of fundamental importance for the safe operation of nuclear power reactors. First-principles electronic structure calculations give detailed insight into these interactions and provide quantitative thermodynamic and kinetic data that can be used to model and predict the life-time behavior of fuel rods in nuclear power reactors. This type of simulations can help to increase the safety while potentially allowing a prolongation of the life time of the fuel elements.

MedeA modules used in this application

- **MedeA® [1] Environment** framework including crystal structure builder, surface builder, and analysis tools
- **MedeA JobServer and TaskServer**
- **MedeA Transition State Search (TSS)**
- **MedeA VASP**
- **MedeA Phonon**