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## Direct assessment of the acidity of individual surface hydroxyls on $\text{In}_2\text{O}_3(111)$

Non-contact atomic force microscopy is a versatile tool to investigate properties of individual atoms or molecules. Here we report a novel approach to determine the acidity of individual surface hydroxyls, which is directly linked to the proton affinity (PA) of the involved O atoms. The PA - the tendency to gain or lose a proton - is crucial *e.g.* in acid-base catalysis and the electro- and photocatalytic splitting of water.

The testcase of this study is the stoichiometric  $\text{In}_2\text{O}_3(111)$  surface, which has four inequivalent surface O atoms  $\text{O}_S(\alpha-\delta)$ . Water dissociation leads to a pair of OH groups: the surface  $\text{O}_S\text{H}(\beta)$  and the water  $\text{O}_W\text{H}$ . The remaining surface O atoms  $\text{O}(\alpha, \gamma, \delta)$  can be protonated via manipulation with the tip. We probe the strength of their H bond with a functionalized tip of a nc-AFM via  $F(z)$ -spectroscopy and find quantitative agreement with density-functional theory (DFT) calculations. By relating the results to known PAs of gas-phase molecules, we can calibrate our data and determine the PA of different surface sites of  $\text{In}_2\text{O}_3(111)$  with atomic precision. Measurements on hydroxylated  $\text{TiO}_2$  and zirconia extend our method to other oxides. The trends of the site-specific PA values agree well with the expectations based on area-averaging techniques.

**Primary authors:** WAGNER, Margareta; MEYER, Bernd (FAU Erlangen-Nürnberg); SETVIN, Martin (TU Wien (Vienna, AT) & Charles University (Prague, CZ)); SCHMID, Michael (Institute of Applied Physics, TU Wien); DIEBOLD, Ulrike (Institute of Applied Physics, TU Wien)

**Presenter:** WAGNER, Margareta

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