A quantum-mechanical study of the adsorption of prototype dye molecules on rutile-TiO$_2$(110): a comparison between catechol and isonicotinic acid†

F. Risplendi,*ab G. Cicero,a G. Mallia,c and N. M. Harrisoncd

Received 20th June 2012, Accepted 29th October 2012
DOI: 10.1039/c2cp42078c

In this work we present a theoretical investigation of the attachment of catechol and isonicotinic acid to the rutile-TiO$_2$(110) surface. These molecules can be considered as prototypical dyes for use in Grätzel type dye sensitised solar cells (DSCs) and are often employed as anchoring groups in both organic and organo-metallic sensitisers of TiO$_2$. Our study focuses on determining the lowest energy adsorption mode and discussing the electronic properties of the resultant hybrid interface by means of density functional theory (DFT) calculations using the hybrid exchange (B3LYP) functional. We find that both molecules adsorb dissociatively at the TiO$_2$ surface giving a type II (staggered) heterojunction. Compared to isonicotinic acid, catechol, due to the greater hybridisation of its molecular orbitals with the states of the substrate, is seen to enhance performance when employed as an anchoring group in dye sensitised solar cells.

I. Introduction

Nanocrystalline dye-sensitised solar cells (DSCs), designed originally by O'Regan and M. Grätzel,1,2 have attracted great interest because of their high efficiency and thus potential application as cost-effective alternatives to present day silicon-based pn-junction photovoltaic devices. In conventional photovoltaic systems, the semiconductor assumes the tasks of both light absorption and charge-carrier transport, whereas in DSCs the two tasks are separated: a dye molecule attached to the surface of a wide band gap semiconductor (typically TiO$_2$) absorbs light, while the semiconductor is responsible for the charge-carrier transport. Charge separation occurs at the interface between the dye and the semiconductor by photo-induced electron injection from the dye to titanium dioxide (TiO$_2$). The dye is necessary, because the two commonly used TiO$_2$ phases (rutile and anatase) absorb only a small fraction of the solar spectrum; the optical band gap is 3.03 eV3,4 for the rutile phase and 3.18 eV3,5 for the anatase phase. To obtain large photoelectron injection from the dye to titanium dioxide (TiO$_2$), whose extremely high surface area facilitates high levels of dye loading and thus photoelectron generation. The structure is then immersed in an electrolyte solution containing a redox couple, such as the triiodide–iodide system, which, after electron injection, restores the original state of the dye by electron donation.

For an effective sensitisation of the substrate it is important that the dye–oxide heterostructure generates a so-called type II (or staggered) heterojunction or, in other words, that the HOMO (highest occupied molecular orbital) of the dye is in the semiconductor gap and the LUMO (lowest unoccupied molecular orbital) is above the conduction band minimum. In this arrangement the electron–hole separation is achieved through the hybrid interface either by direct electron photo-injection from the HOMO of the dye molecule to the conduction band of the oxide or, indirectly, by first excitation of an electron from the HOMO to the LUMO of the dye followed by electron injection into the semiconductor conduction band.

Recently small molecules, such as catechol, have attracted considerable attention in the realisation of DSCs.6–11 Catechol is used as an anchoring group for organic and organometallic dyes because of its efficient adsorption onto TiO$_2$ via formation of a strong adsorbate–substrate complex.6,7 Catechol has also been directly employed as a sensitisier for DSC since the TiO$_2$–catechol interface realises a type II hybrid junction.8–12 In the UV-vis absorption spectrum for catechol adsorption onto TiO$_2$ anatase nanoparticles,12 an absorption peak at ~2.88 eV (~430 nm) is observed and is interpreted as evidence of the HOMO level of the molecule in the TiO$_2$ band gap. In the case of catechol adsorbed at the rutile (110) surface absorption occurs at 2.4 eV as deduced from UV

† Electronic supplementary information (ESI) available. See DOI: 10.1039/c2cp42078c

a DISAT Department of Applied Science and Technology, Politecnico of Torino, C.so Duca degli Abruzzi 24, 10129, Turin, Italy. E-mail: francesca.risplendi@polito.it
b IIT Istituto Italiano di Tecnologia, Center for Space Human Robotics @ Polito, Corso Trento 21, 10129 Turin, Italy
c Thomas Young Centre, Department of Chemistry, Imperial College London, South Kensington, London SW7 2AZ, UK
d STFC, Daresbury Laboratory, Daresbury, Warrington, WA4 4AD, UK

This journal is © the Owner Societies 2012 Phys. Chem. Chem. Phys.
photoemission and inverse photoemission spectroscopy (UPS and IPS) measurements under ultra-high vacuum.10 Another prototypical dye often used as an anchoring group in metalorganic-dyes (e.g. N3)11 is isonicotinic acid; this presents similar properties to that of the catechol molecule in terms of anchoring/coupling to TiO2 surfaces.

Ab initio simulations have been employed to unravel the structural and electronic properties of several organic/inorganic interfaces in the case of various inorganic substrates14–16 and several theoretical methods have been exploited to investigate the electronic properties of the catechol–TiO2 heterojunction. Density functional theory (DFT) and time dependent DFT (TDDFT) methods have been applied to study the interface structure and excited states for various models of the surfaces including a TiO2-cluster8,11 and a single Ti atom;9,10 periodic DFT calculations (based on generalised gradient approximation – GGA – functionals for exchange and correlation) have been used for studying the anatase-TiO2(101)17 and rutile-TiO2(110) surfaces18 functionalised with catechol molecules. In order to classify the heterojunction, a method, which computes accurate adsorption energies and a reasonable description of the electronic structure and optical absorption spectrum, is required. Previous periodic DFT calculations, based on GGA functionals, provide a poor representation of the bulk band gap and it is unclear how well surface energy level alignments are described. Cluster models of the surface converge slowly with respect to the cluster size and so great care is required in drawing conclusions from calculations on small clusters. Concerning the isonicotinic acid–TiO2 heterojunction, there have been two previous theoretical investigations using hybrid-exchange DFT; upon adsorption of isonicotinic acid onto anatase TiO216 and onto rutile TiO2,20 the HOMO level was found to be in the energy gap while the LUMO level falls in the conduction band. Yet, it has to be noted that in the published study20 no complete geometry optimization of the combined substrate-adsorbate system was performed and that the adsorption geometry was based on a previous work on formic acid attached to the TiO2 surface.20

In order to gain deeper insight into the structural and electronic properties of the two heterostructures and to allow for a comparative study between the two heterojunctions (by considering also the relaxation effects upon adsorption), periodic hybrid-exchange DFT calculations based on the B3LYP functional are suitable. This approach provides an accurate description of the structural energetics and of the electronic structure (i.e. band gap and band offset) for periodic systems,21–34 particularly for transition metal oxides, and the implementation in the CRYSTAL code is computationally efficient for large periodic systems.35 The aim of this study is to address the binding of catechol and of isonicotinic acid to the rutile-TiO2(110) surface in terms of geometry, stability and electronic structure, using hybrid-exchange DFT. We compare in detail the interaction of these two molecules with TiO2: strong binding of the anchoring group not only serves to put the sensitiser in place, but also to control the interfacial electronic coupling between the organic molecule and the semiconductor, thus influencing solar cell efficiency. Although anatase-TiO2 is currently preferred over the rutile phase when realising DSCs, we focused our study on rutile because it presents some advantages over anatase: it is chemically more stable, can potentially be produced more cheaply and has superior light-scattering properties because of the higher refractive index. Moreover, the performance of rutile in DSCs has been shown to be comparable to that of anatase at one sun light intensity.36 In particular, the rutile-TiO2(110) face has been selected, because it is the most stable surface of rutile and thus dominates the surface area of equilibrium crystallites; it is also the best microscopically characterised surface of TiO2.18

The paper is organised as follows: in Section II the computational details are provided. In Section III the results are discussed in terms of energetics (i.e. binding energies, acidity of the functional groups, bond distances and bond natures between the adsorbate and the substrate) and in terms of electronic structure (i.e. band alignment, band gap states, electron coupling between the molecule and the surface). Conclusions are drawn in Section IV.

II. Computational details

All calculations have been performed using the CRYSTAL09 software package,37,38 based on the expansion of the crystaline orbitals as a linear combination of a local basis set (BS) consisting of atom-centred Gaussian orbitals. The titanium and oxygen atoms are described by a triple valence all-electron BS; an 86-411G** contraction (one s, four sp and two d shells) and an 8-411G* contraction (one s, three sp and one d shells), respectively;39 the most diffuse sp (d) exponents are zT1 = 0.3297(0.26) and zO = 0.1843(0.6) Bohr−2. These basis sets were developed in previous studies of the bulk and surfaces of titania in which a systematic hierarchy of all-electron basis sets was used to quantify the effects of using a finite BS.40,41 Both adsorbed molecules are described by a 6-31G** contraction,42 where the valence electrons of the carbon and nitrogen atoms are described by two sp shells and one d shell (in the case of the H atom two s shells and one p shell are used), in order to ensure a similar level of accuracy.

Electron exchange and correlation are approximated using the B3LYP hybrid exchange functional, as indicated in Section I. Matrix elements of the exchange and correlation potentials and the energy functional are integrated numerically on an atom-centred grid of points. The integration over radial and angular coordinates is performed using Gauss–Legendre and Lebedev schemes, respectively. A pruned grid consisting of 99 radial points and 5 sub-intervals with (146, 302, 1454, 1545, 590) angular points has been used for all calculations (the XXLGRID option implemented in CRYSTAL09).37 The geometry optimisation was initially performed using a smaller grid (LGRID) and then refined using the XXLGRID. This ensures numeral stability both for the bulk and the slab calculations. This grid converges the integrated charge density to an accuracy of about ×10−6 electrons per unit cell. The Coulomb and exchange series are summed directly and truncated using overlap criteria with thresholds of 10−7, 10−7, 10−7, 10−7 and 10−14 as described previously.43 Reciprocal space sampling for the bulk structure was performed on a Pack–Monkhorst net with a shrinking factor IS = 8 along each periodic direction. In the bulk, this grid corresponds to 75 k-points in the irreducible Brillouin zone of the bulk crystal and to 25 in that of the surface unit cell. The self-consistent
field procedure was converged up to a tolerance in the total energy of $\Delta E = 1 \times 10^{-2}E_h$ per unit cell.

Structural optimisation of both the bulk cell parameters and internal coordinates was performed using the Broyden–Fletcher–Goldfarb–Shanno scheme. Convergence was determined from the root-mean-square (rms) and the absolute value of the largest component of the forces. The thresholds for the maximum and the rms forces (the maximum and the rms atomic displacements) have been set to 0.00045 and 0.00030 (0.00180 and 0.0012) in atomic units. Geometry optimisation was terminated when all four conditions were satisfied simultaneously. The predicted structural parameters, with the deviation from those observed in parentheses, are: $a_{\text{bulk}} = b_{\text{bulk}} = 4.639 \text{ Å (1.1%)}$, $c_{\text{bulk}} = 2.979 \text{ Å (0.9%)}$ and $\alpha = 0.306 \text{ (0.5%)}$ where the O atom is at the fractional position $(u,u,0)$. These values are consistent with those predicted in previous calculations. With regards to the surface, a 9-atomic-layer (9AL) slab cut from the bulk has been used, since it represents a good compromise between the surface description and the computational cost. When increasing the slab thickness from 9AL to 15AL, the electronic structure is unaffected in terms of the contributions to the valence and conduction bands in the projected density of states. The corresponding lattice parameters are $a_{\text{slab}} = 2.979 \text{ Å ([001] direction of the bulk)}$ and $b_{\text{slab}} = 6.561 \text{ Å ([110] direction of the bulk)}$. In comparing the electronic structure of various adsorption modes, it is important to note that, as 2D periodic boundary conditions are used, all energies are referred to vacuum zero $V_{\text{es}}(z) = 0$ when $z = \infty$, where $V_{\text{es}}$ is the electrostatic potential and $z$ is the distance from the surface. Projection of density of states onto orbital and atomic contributions was performed using Mulliken analysis.

Adsorption has been investigated by initially placing the molecule with its symmetry plane perpendicular to the surface plane. The distance between the two anchoring oxygen atoms in both molecules is comparable to the separation between the two undercoordinated titanium atoms along $a_{\text{slab}}$, therefore the molecule is likely to be coordinated to two surface Ti atoms and thus a $3 \times 1$ supercell, containing three surface Ti atoms, has been adopted to ensure that adsorbates are separated by at least one empty surface site (see Fig. 1 and 2). In addition, it should be noted that molecules were adsorbed symmetrically on both sides of the twodimensionally periodic slab as this ensures more rapid convergence of the structure and total energy with respect to slab thickness. The binding energy (BE) per molecule of the adsorbate-substrate system was computed with respect to the isolated molecule and the clean surface. The counterpoise correction to the binding energy was applied to take into account the basis set superposition error (BSSE), details of which are documented in ref. 45 and 46. The BE reported does not take into account either the vibrational component of the free energy or London dispersion contributions to the bonding. At room temperature the former are expected to be insignificant compared to the BE reported in Table 1 (see ref. 47) and the latter are expected to be a small correction to the binding interaction which is dominated by ionic and covalent contributions.

III. Results

The binding energy of the adsorbate-substrate system with respect to the clean surface and the isolated molecule is analysed for the catechol and the isonicotinic acid along with the structural changes induced by the adsorption in Section III.A, while the electronic properties of the resulting hybrid interfaces are discussed in Section III.B. The acidity of the molecules, the ionicity of the oxygen atom and the electron conjugation between the adsorbed molecule and the substrate are used to shed light on the structural and electronic properties of the two heterojunctions in both Sections III.A and III.B.

A. Structure and energetics

The clean rutile TiO$_2$(110) surface is characterised by the presence of fivefold-coordinated titanium (Ti$_5$) and twofold-coordinated “bridging” oxygen atoms (O$_3$). The Ti ions, being undercoordinated, react with adsorbing species. There are
the outermost Ti and O atom of the surface; O$_{ap}$ is the apical oxygen atom bonded to Ti$_s$, see Fig. 1.

In the BDA case, both –OH groups of catechol deprotonate and attach to two five-coordinated Ti atoms; the two released H atoms form two hydroxyl groups with two nearest O$_s$. In the case of the isonicotinic acid C$_6$H$_5$NO$_2$ (pyridine-4-carboxylic acid according to IUPAC notation), which is characterised by the presence of a carboxyl group, two adsorption modes are considered: molecular (MA) in which the $\equiv$C–O group coordinates with a Ti$_s$ species and dissociative (DA) in which the carboxyl group releases its hydrogen atom to an O$_s$ site and the –COO$^-$ group bridges two Ti$_s$ surface ions (see Fig. 2).

Table 1, the binding energy $^1$ of the adsorbate–surface system with respect to the clean surface and the isolated molecule (BE) is given for both molecules. In both cases, the dissociative adsorption corresponds to the lowest energy configuration. The isonicotinic acid binds more strongly to the surface (BE$_{\text{DA}}$ = –1.906 eV) than the catechol (BE$_{\text{BDA}}$ = –1.756 eV). This is as one would expect from the higher acidity of the carboxyl group of the isonicotinic acid, which releases H more easily than catechol. In the case of catechol, there is a clear trend in the BE as a function of the deprotonation, the BDA being more stable than the MDA: BE$_{\text{BDA}}$ < BE$_{\text{MDA}}$ < BE$_{\text{MA}}$, in agreement with previous theoretical studies. This trend can be rationalised in terms of the bond distance between the atoms of the molecule and of the rutile surface and in terms of the displacement of the outermost surface atoms, bonded to the molecule.

The distances between the five-coordinated Ti atoms and each of the two oxygen atoms of the molecule, $d_{\text{O$_s$–Ti$_s$}}$, are reported in Table 1. In the dissociative adsorption the molecule forms shorter bonds with the surface atoms than in the molecular (MA) adsorption. In the MDA configuration of catechol, the $d_{\text{O$_s$–Ti$_s$}}$ are about 1.8 and 2.4 Å, respectively, for the deprotonated oxygen atom and for the intact hydroxyl group (both oxygen atoms belonging to the molecule), while for the BDA case both $d_{\text{O$_s$–Ti$_s$}}$ are about 1.8 Å; correspondingly, the BE increases from –1.452 to –1.756 eV. The values of $d_{\text{O$_s$–Ti$_s$}}$ and the BE for the various adsorption modes are also a consequence of the ionicity of the oxygen atom in the molecule; the more ionic the O$_s$ atom, the shorter the $d_{\text{O$_s$–Ti$_s$}}$, with the deprotonated oxygen atom more ionic (more negative) than the hydroxyl oxygen one. In the isonicotinic acid, the $\equiv$C–O group plays an important role in the interaction between the molecule and the surface, especially in the MA case. In fact, the value of $d_{\text{O$_s$–Ti$_s$}}$ = 2.184 Å for the $\equiv$C–O group is smaller than $d_{\text{O$_s$–Ti$_s$}}$ = 3.943 Å for the OH group, whose interaction with the surface is negligible. However, this value ($d_{\text{O$_s$–Ti$_s$}}$ = 2.184 Å) is larger than $d_{\text{O$_s$–H}}$.

**Table 1** Binding energy (BE) of the adsorbate–substrate complex with respect to the clean surface and the isolated molecule for the adsorption geometries considered. The bond distances between the atoms of the molecule (O$_s$ and H) and the surface atoms are reported; Ti$_s$ and O$_s$ are the outermost Ti and O atom of the surface; O$_{ap}$ is the apical oxygen atom bonded to Ti$_s$, see Fig. 1.

<table>
<thead>
<tr>
<th>Adsorption mode</th>
<th>BE/eV</th>
<th>$d_{\text{O$_s$–Ti$_s$}}$/Å</th>
<th>$d_{\text{Ti$<em>s$–O$</em>{ap}$}}$/Å</th>
<th>$d_{\text{O$_s$–H}}$/Å</th>
<th>$d_{\text{O$_s$–H}}$/Å</th>
</tr>
</thead>
<tbody>
<tr>
<td>Catechol</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MA</td>
<td>–0.359</td>
<td>2.342/2.347</td>
<td>1.907/1.893</td>
<td>0.967/0.967</td>
<td>–</td>
</tr>
<tr>
<td>MDA</td>
<td>–1.452</td>
<td>2.381/1.820</td>
<td>1.871/2.380</td>
<td>0.966</td>
<td>0.961</td>
</tr>
<tr>
<td>BDA</td>
<td>–1.756</td>
<td>1.831/1.334</td>
<td>2.202/2.249</td>
<td>–</td>
<td>0.961/0.961</td>
</tr>
<tr>
<td>Isonicotinic acid</td>
<td>MA</td>
<td>–0.457</td>
<td>3.943/2.184</td>
<td>1.815/2.014</td>
<td>0.971</td>
</tr>
<tr>
<td></td>
<td>DA</td>
<td>–1.906</td>
<td>2.033/2.059</td>
<td>2.069/2.049</td>
<td>–</td>
</tr>
</tbody>
</table>
for the deprotonated oxygen atom in both adsorbed molecules. Therefore, there is a decrease in $d_{O_{ap}}$, with increasing O ionicity, when moving from the O ion in –OH to that in $\geq C=O$, –O$^−$ of the isonicotinic acid and –O$^−$ of catechol (refer to the next subsection for a detailed analysis in terms of the electronic structure).

In order to understand the structural modifications occurring upon adsorption, it is useful to consider the relaxation of the clean surface, as a reference. With respect to a surface defined by cutting a slab from an optimised bulk geometry, the Ti$_s$ moves inwards by 0.14 Å to increase effective coordination with neighbouring O ions. In the optimised crystal bulk, the TiO$_6$ octahedron is characterised by two long apical ($d_{Ti-O_{ap}} = 2.01$ Å) and four short equatorial ($d_{Ti-O_{eq}} = 1.98$ Å) oxygen bonds. At the clean relaxed TiO$_2$ surface the distance between the outermost Ti and its neighbouring apical oxygen, $d_{Ti-O_{ap}}$, for the deprotonated oxygen atom in both adsorbed molecules. Therefore, there is a decrease in $d_{O_{ap}}$, with increasing O ionicity, when moving from the O ion in –OH to that in $\geq C=O$, –O$^−$ of the isonicotinic acid and –O$^−$ of catechol (refer to the next subsection for a detailed analysis in terms of the electronic structure).

In order to understand the structural modifications occurring upon adsorption, it is useful to consider the relaxation of the clean surface, as a reference. With respect to a surface defined by cutting a slab from an optimised bulk geometry, the Ti$_s$ moves inwards by 0.14 Å to increase effective coordination with neighbouring O ions. In the optimised crystal bulk, the TiO$_6$ octahedron is characterised by two long apical ($d_{Ti-O_{ap}} = 2.01$ Å) and four short equatorial ($d_{Ti-O_{eq}} = 1.98$ Å) oxygen bonds. At the clean relaxed TiO$_2$ surface the distance between the outermost Ti and its neighbouring apical oxygen, $d_{Ti-O_{ap}}$, for the deprotonated oxygen atom in both adsorbed molecules. Therefore, there is a decrease in $d_{O_{ap}}$, with increasing O ionicity, when moving from the O ion in –OH to that in $\geq C=O$, –O$^−$ of the isonicotinic acid and –O$^−$ of catechol (refer to the next subsection for a detailed analysis in terms of the electronic structure).

In order to understand the structural modifications occurring upon adsorption, it is useful to consider the relaxation of the clean surface, as a reference. With respect to a surface defined by cutting a slab from an optimised bulk geometry, the Ti$_s$ moves inwards by 0.14 Å to increase effective coordination with neighbouring O ions. In the optimised crystal bulk, the TiO$_6$ octahedron is characterised by two long apical ($d_{Ti-O_{ap}} = 2.01$ Å) and four short equatorial ($d_{Ti-O_{eq}} = 1.98$ Å) oxygen bonds. At the clean relaxed TiO$_2$ surface the distance between the outermost Ti and its neighbouring apical oxygen, $d_{Ti-O_{ap}}$, for the deprotonated oxygen atom in both adsorbed molecules. Therefore, there is a decrease in $d_{O_{ap}}$, with increasing O ionicity, when moving from the O ion in –OH to that in $\geq C=O$, –O$^−$ of the isonicotinic acid and –O$^−$ of catechol (refer to the next subsection for a detailed analysis in terms of the electronic structure).

In order to understand the structural modifications occurring upon adsorption, it is useful to consider the relaxation of the clean surface, as a reference. With respect to a surface defined by cutting a slab from an optimised bulk geometry, the Ti$_s$ moves inwards by 0.14 Å to increase effective coordination with neighbouring O ions. In the optimised crystal bulk, the TiO$_6$ octahedron is characterised by two long apical ($d_{Ti-O_{ap}} = 2.01$ Å) and four short equatorial ($d_{Ti-O_{eq}} = 1.98$ Å) oxygen bonds. At the clean relaxed TiO$_2$ surface the distance between the outermost Ti and its neighbouring apical oxygen, $d_{Ti-O_{ap}}$, for the deprotonated oxygen atom in both adsorbed molecules. Therefore, there is a decrease in $d_{O_{ap}}$, with increasing O ionicity, when moving from the O ion in –OH to that in $\geq C=O$, –O$^−$ of the isonicotinic acid and –O$^−$ of catechol (refer to the next subsection for a detailed analysis in terms of the electronic structure).

In order to understand the structural modifications occurring upon adsorption, it is useful to consider the relaxation of the clean surface, as a reference. With respect to a surface defined by cutting a slab from an optimised bulk geometry, the Ti$_s$ moves inwards by 0.14 Å to increase effective coordination with neighbouring O ions. In the optimised crystal bulk, the TiO$_6$ octahedron is characterised by two long apical ($d_{Ti-O_{ap}} = 2.01$ Å) and four short equatorial ($d_{Ti-O_{eq}} = 1.98$ Å) oxygen bonds. At the clean relaxed TiO$_2$ surface the distance between the outermost Ti and its neighbouring apical oxygen, $d_{Ti-O_{ap}}$, for the deprotonated oxygen atom in both adsorbed molecules. Therefore, there is a decrease in $d_{O_{ap}}$, with increasing O ionicity, when moving from the O ion in –OH to that in $\geq C=O$, –O$^−$ of the isonicotinic acid and –O$^−$ of catechol (refer to the next subsection for a detailed analysis in terms of the electronic structure).

In order to understand the structural modifications occurring upon adsorption, it is useful to consider the relaxation of the clean surface, as a reference. With respect to a surface defined by cutting a slab from an optimised bulk geometry, the Ti$_s$ moves inwards by 0.14 Å to increase effective coordination with neighbouring O ions. In the optimised crystal bulk, the TiO$_6$ octahedron is characterised by two long apical ($d_{Ti-O_{ap}} = 2.01$ Å) and four short equatorial ($d_{Ti-O_{eq}} = 1.98$ Å) oxygen bonds. At the clean relaxed TiO$_2$ surface the distance between the outermost Ti and its neighbouring apical oxygen, $d_{Ti-O_{ap}}$,
is shorter (1.853 Å), due to the under coordination of Ti, as reported in a previous work.\textsuperscript{26} When the adsorbed molecule saturates the fivefold coordinated Ti atoms (Ti\textsubscript{5}) at the rutile surface, one of the most significant effects is that Ti\textsubscript{5} moves outwards to restore a bulk-like coordination. In catechol a shorter \(d_{\text{Os}-\text{Ti}}\) \(\approx 1.8\) Å for the deprotonated oxygen atom and a consequent larger \(d_{\text{Ti}-\text{O}_{\text{ap}}}\) \(\approx 2.3\) Å can be understood in terms of the extension of the delocalisation of the \(\pi\)-electrons of the benzene ring over the Ti\textsubscript{5} atoms. For isonicotinic acid this effect does not occur and the distances, \(d_{\text{Os}-\text{Ti}}\) and \(d_{\text{Ti}-\text{O}_{\text{ap}}}\), are similar and very close to the value in the bulk (\(d_{\text{Ti}-\text{O}_{\text{ap}}} = 2.01\) Å). In Table 1, the hydrogen–oxygen distance is also reported. The distance between the H atom released by the molecule and the two-coordinated bridging oxygen atom, \(d_{\text{O}_{\text{ap}}-\text{H}}\), is close to the molecular distance, \(d_{\text{Os}-\text{H}}\).

In summary, the binding energy and the bond distance of the anchoring functional groups of a dye at the TiO\textsubscript{2} surface can be rationalised in terms of the acidity and of the ionicity of the molecular oxygen atom, and in terms of a secondary factor related to the electron conjugation with the surface. These effects are also important in their influence on the electronic structure of the heterojunctions formed.

### B. Electronic structure

In order to rationalize the behaviour of the two adsorbed molecules, the different alignment of the molecular states with respect to the semiconductor bands has to be taken into account. With this aim, the total density of states and the projections onto the atomic contribution are presented in Fig. 3 and 4.\textsuperscript{48} The lower (below \(-23\) eV) and the upper (in the range of \(-15\) eV and \(-6\) eV) valence bands have predominantly O-2s and O-2p character with some hybridisation with Ti-3d orbitals, while the lower conduction band (above \(-5\) eV) has Ti-3d character. Upon adsorption, the molecular states, attributed to the \(\pi\) orbitals in the catechol, appear in the TiO\textsubscript{2} band gap above the top of VB; while in the isonicotinic acid case a \(\sigma\) state, mainly due to carbon and nitrogen atoms, is evident at the top of VB for dissociative mode only. These abovementioned HOMO states (both \(\pi\) and \(\sigma\))
Table 2 The calculated energy difference $\Delta E$ between the highest occupied ($E_{\text{occ}}$) and the lowest unoccupied ($E_{\text{unocc}}$) states for the clean surface, the isolated molecules and the various adsorption modes. In the column $E_{\text{LUMO}} - E_{\text{HOMO}}$, the star indicates that the value of the electronic transition, HOMO $\rightarrow$ LUMO, is affected by the hybridization of LUMO with the conduction band states and by the consequent dispersion in energy of this state. In the last column, the type of electronic transition is reported: for instance, for the clean surface $\Delta E$ is the difference between the bottom of the conduction band (CB) and the top of the valence band (VB) and for the isolated molecules $\Delta E$ is the HOMO–LUMO difference. In addition, the symmetry of the state corresponding to the HOMO ($\pi$ or $\sigma$) is given in parentheses.

<table>
<thead>
<tr>
<th>Mode</th>
<th>$E_{\text{occ}}$</th>
<th>$E_{\text{unocc}}$</th>
<th>$E_{\text{LUMO}} - E_{\text{HOMO}}$</th>
<th>Transition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clean TiO$_2$</td>
<td>2.90</td>
<td>—</td>
<td>—</td>
<td>VB $\rightarrow$ CB</td>
</tr>
<tr>
<td>Isolated catechol</td>
<td>5.86</td>
<td>—</td>
<td>5ler (\pi) $\rightarrow$ LUMO</td>
<td></td>
</tr>
<tr>
<td>MA catechol</td>
<td>2.90</td>
<td>5.55*</td>
<td>HOMO(\pi) $\rightarrow$ CB</td>
<td></td>
</tr>
<tr>
<td>MDA catechol</td>
<td>2.78</td>
<td>5.24*</td>
<td>HOMO(\pi) $\rightarrow$ CB</td>
<td></td>
</tr>
<tr>
<td>BDA catechol</td>
<td>2.35</td>
<td>4.84*</td>
<td>HOMO(\pi) $\rightarrow$ LUMO</td>
<td></td>
</tr>
<tr>
<td>Isolated</td>
<td>5.21</td>
<td>5.20</td>
<td>HOMO(\pi) $\rightarrow$ LUMO</td>
<td></td>
</tr>
<tr>
<td>isonicotinic acid</td>
<td>3.07</td>
<td>4.82*</td>
<td>LUMO</td>
<td></td>
</tr>
<tr>
<td>MA isonicotinic acid</td>
<td>3.50</td>
<td>4.55*</td>
<td>HOMO(\pi) $\rightarrow$ CB</td>
<td></td>
</tr>
<tr>
<td>BDA isonicotinic acid</td>
<td>3.35</td>
<td>4.55*</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3 Wannier functions contributions $c_{\text{WF}}$ of the atoms (the two C atoms closer to the surface, the two O atoms of the molecule directly bond to them and the Ti atoms, originally fivefold coordinated) involved in the HOMO for the catechol on TiO$_2$(110) in the various adsorption modes.

<table>
<thead>
<tr>
<th>Mode</th>
<th>$c_{\text{WF}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>MA catechol</td>
<td>0.189/0.188</td>
</tr>
<tr>
<td>MDA catechol</td>
<td>0.188/0.171</td>
</tr>
<tr>
<td>BDA catechol</td>
<td>0.170/0.170</td>
</tr>
</tbody>
</table>

are useful as a reference, since their energies with respect to the vacuum level are not significantly affected by adsorption, when compared to the isolated molecule, as shown in Fig. 3 and 4. A shift of the valence band offset of the semiconductor is instead evident, especially when the DOS for the molecular mode is compared with the superposition of the DOS for the isolated molecule and the clean TiO$_2$ surface. This shift is generated by the dipole of the absorbed molecule and is significantly smaller for dissociative adsorption.

Table 4 The Mulliken populations for the atoms involved in bonding between the molecule and the surface for catechol and isonicotinic acid; the charge associated to the Ti atoms, originally fivefold coordinated, the C atoms closer to the surface, the O atoms of the hydroxyl groups and the double bond with the C atom are reported in the first column. In the last column the overlap population associated with the Ti–O bond is reported. Unit: $|$e$|$ Bohr$^{-3}$

<table>
<thead>
<tr>
<th>Mode</th>
<th>$q_{\text{C}}$</th>
<th>$q_{\text{O}}$</th>
<th>$q_{\text{Ti}}$</th>
<th>$q_{\text{Ti}--\text{O}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clean TiO$_2$(110)</td>
<td>—</td>
<td>—</td>
<td>2.130</td>
<td>—</td>
</tr>
<tr>
<td>Isolated catechol</td>
<td>0.291/0.285</td>
<td>-0.527/ -0.564</td>
<td>2.144/2.143</td>
<td>0.026/0.025</td>
</tr>
<tr>
<td>MA catechol</td>
<td>0.350/0.348</td>
<td>-0.567/ -0.562</td>
<td>2.142/2.140</td>
<td>0.024/0.015</td>
</tr>
<tr>
<td>MDA catechol</td>
<td>0.300/0.344</td>
<td>-0.578/ -0.710</td>
<td>2.142/2.146</td>
<td>0.104/0.104</td>
</tr>
<tr>
<td>BDA catechol</td>
<td>0.295/0.295</td>
<td>-0.711/ -0.710</td>
<td>2.108/2.159</td>
<td>-0.001/0.043</td>
</tr>
<tr>
<td>BDA isonicotinic acid</td>
<td>0.583</td>
<td>-0.529/ -0.405</td>
<td>2.149/2.147</td>
<td>0.066/0.070</td>
</tr>
<tr>
<td>MA isonicotinic acid</td>
<td>0.593</td>
<td>-0.581/ -0.613</td>
<td>2.149/2.147</td>
<td>0.066/0.070</td>
</tr>
</tbody>
</table>
with the oxide substrate than isonicotinic acid and thus likely to be more effective for photo-induced electron injection, when employed as a dye molecule.

IV. Conclusions

In this work different adsorption modes of the catechol and the isonicotinic acid molecules onto the rutile TiO$_2$(110) surface have been analysed in terms of geometry, binding energy and electronic structure within a hybrid-exchange DFT framework. The calculations demonstrate that for both molecules the dissociative adsorption (fully deprotonated) is the thermodynamically favoured mode of adsorption due to saturation of two fivefold coordinated Ti atoms at the surface and formation of two Ti-$\cdot$O bonds with partial covalent character. Bond formation and bond distances can be rationalised in terms of acidity and ionicity of the functional groups responsible for the molecular attachment. Upon adsorption, in the most stable configuration the HOMO of both molecules occurs in the energy gap of the TiO$_2$ substrate while the LUMO lies above the top of the conduction band, generating a type II heterojunction that makes these systems suitable for applications in DSC. In the case of catechol the HOMO state originates from a hybridisation of the delocalised $\pi$ orbitals of the benzene ring with the 3d-states of the substrate titanium atoms, while the HOMO of the isonicotinic acid corresponds to a $\sigma$ orbital of the N-C bond. In the former case the larger electron coupling between the molecule and the oxide is expected to facilitate electron injection from the dye to the TiO$_2$ and thus larger photocurrent during DSC operation. For this reason our results indicate that catechol will be a better anchoring group with respect to isonicotinic acid although being characterised by a slightly lower binding energy.

Finally, we found that the adsorption of both the organic molecules considered here gives rise to a positive shift of the TiO$_2$ bands (both VB and CB) with respect to the vacuum level (or equivalently to the hydrogen redox potential). This shift, which is mostly determined by the molecular dipole and by the type and number of bonds formed with the substrate, is important since the position of the CB determines the solar cell open circuit voltage (the higher is the positive shift the larger is $V_{oc}$). This observation highlights the need to tune the dipole of the dye (and of the co-adsorbent often used during DSC realisation to avoid dye aggregation) to molecularly engineer the electronic properties of the hybrid interface and enhance solar cell performances.

Acknowledgements

This work made use of the high performance computing facilities of Imperial College London and – via membership of the UK’s HPC Materials Chemistry Consortium funded by EPSRC (EP/F067496) – of HECTOR, the UK’s national high-performance computing service, which is provided by UoE HPCx Ltd at the University of Edinburgh, Cray Inc and NAG Ltd, and funded by the Office of Science and Technology through EPSRC’s High End Computing Programme. One of the authors, F.R., also acknowledges the TYC for providing financial assistance through the Junior Research Fellowship (JRF).

References

The DOS plots for the surfaces are plotted with respect to atomic zero and thus the variations in peak positions and Fermi energies can legitimately be interpreted in terms of changes in the surface dipole and workfunction.