Wettability of fully hydroxylated and alkylated (001) α-quartz surface in carbon dioxide atmosphere

Aleksandr Abramov  
*Edith Cowan University*

Alireza Keshavarz  
*Edith Cowan University*

Stefan Iglauer  
*Edith Cowan University*

Follow this and additional works at: [https://ro.ecu.edu.au/ecuworkspost2013](https://ro.ecu.edu.au/ecuworkspost2013)
Wettability of Fully Hydroxylated and Alkylated (001) alpha-Quartz Surface in Carbon Dioxide Atmosphere

Aleksandr Abramov*, Stefan Iglauer, Alireza Keshavarz, School of Engineering, Edith Cowan University, 270 Joondalup Drive, Joondalup, WA 6027, Western Australia, Australia

*aabramov@our.ecu.edu.au

Abstract

Wettability of alkylated quartz surface is of primary importance in several technological applications including the development of oil and gas reservoirs and carbon geo-sequestration. It is intuitively understood and experimentally confirmed that hydroxylated quartz surface is hydrophilic. By gradually saturating hydroxylated (001) alpha-quartz surface with pentyl groups we show using molecular dynamics simulations that the surface can also exhibit extreme hydrophobicity. Within a range of surface pentyl group density from 0.29 to 3.18 per nanometre squared the contact angle of a water droplet under 10 MPa pressure of carbon dioxide at 300 K changes from 10–20 to 180 degrees. This study has shown that a complete description of wettability of alkylated quartz surface requires three contact angles, one at the tip level of pentyl groups, and two at the level of quartz surface. The latter two are the contact angle of the spherical droplet and the hidden contact angle of a water "skirt" formed between the tip level of pentyl groups and the quartz surface. Analysis of the hidden contact angle unveils a binary wettability, where the surface relatively abruptly transforms from hydrophilic (the contact angle is less than 90 degrees) to hydrophobic (the contact angle is 180 degrees) with an increase in surface pentyl group concentration.

Introduction

In oil and gas reservoirs and in saline aquifers mineral surfaces are exposed to water and hydrocarbons over geological time scales 1-2. Hydroxylation and chemisorption of hydrocarbons with following change in wettability is thus expected and was demonstrated experimentally 3. Because of natural abundance of quartz its wettability is of primary importance for carbon geo-sequestration and recovery of hydrocarbons.

Properties of quartz crystal and its surfaces are of key importance in many areas, including the construction industry, piezoelectric devices 4, biotoxicity 5, astrochemistry 6 and the geology of the solid earth 7. Depending on conditions of pressure and temperature, crystalline silica (SiO₂) can form different polymorphs, some of them are α- and β-quartz, α- and β-cristobalite and coesite 8; structures and properties of these polymorphs are known and understood. Low temperature polymorphs, such as α-quartz, exist at ambient conditions. Above approximately 3 GPa and 1200°C α-quartz becomes unstable and with increase in pressure transforms to coesite, and with increase in temperature transforms to β-quartz 9.

The (001) surface of α-quartz is regarded as the most stable surface of pristine 10 and hydroxylated α-quartz 11-12. Thus, numerous research studies have been dedicated to understanding the properties of these quartz surfaces using computational chemistry and materials science methods 10-17.

Contemporary (001) α-quartz surface models consider two extremes in terms of the degree of surface hydroxylation, namely (i) the pristine (no hydroxyl groups, stable Si-O-Si bridges) and (ii) the hydroxylated model. The ideal fully hydroxylated theoretical silica surface is covered with geminal silanol groups, which results in OH group (or single silanol group) concentration of 9.4 per nm² 18 (slightly depends on calculated/measured lattice constants). However, any real hydroxylated silica
surface (excluding possible contaminations and defects) is a combination of isolated silanols, vicinal silanols, geminal silanols and Si-O-Si bridges. Average OH group (or single silanol group) concentration on real hydroxylated surfaces under ambient conditions amounts to 4.5-6.2 per nm². This silanol density aligns with data reported in a comprehensive review paper, where the average concentration of OH groups is said to be a physicochemical constant of 4.6 or 4.9 groups per nm², depending on the averaging technique, the least square or the arithmetical mean, respectively.

In recent years one specific factor - the wettability of quartz surfaces, has been intensively studied because of its implications in carbon capture and storage technology where it determines reliability, security and capacity of the structural and the residual trapping mechanisms. Quartz abundance in the Earth’s continental crust (the second most abundant mineral after feldspars with weight concentration of 12% and in caprocks (more than 50% by weight) makes it crucial for characterisation of the overall wettability of real multi-mineral caprocks. In addition, sandstone reservoirs, mainly made up of quartz, are prime targets for CO2 storage.

Experimental wettabiltiy studies of quartz surfaces have explored different gaseous atmospheres including air, hydrogen sulphide and carbon dioxide. Various water solutions have been tested ranging from distilled water to brines of different salinities. The diversity of surface-liquid-gas combinations examined in the experiments is enriched by the multiplicity of contact angle measurement methods. Out of nine methodologies offered in 33, researchers commonly use the sessile drop method, the captive bubble method, the capillary rise method and direct measurement with atomic force microscopy.

Experimental results have answered many questions on quantification of quartz wettabiltiy in CO2 atmosphere with respect to the number of independent parameters such as pressure, temperature, salinity and salt type. When these parameters, for example the degree of hydroxylation, are difficult or even impossible to control experimentally because of chemical side reactions or contaminations, theoretical methods complement obtained data and provide deeper theoretical understanding. There are number of studies employing computational methods to simulate wetting behaviour of water on surfaces of minerals including quartz, in particular in presence of carbon dioxide and often in the context of carbon geo-sequestration. Theoretical studies have explored possible surface structures and predicted contact angles in CO2 environment on pristine and hydroxylated surfaces, as well as the dependence of the contact angle on the degree of hydroxylation. Chen studied the effect of CO2 contamination with CH4, Ar and H2S on the water-CO2 interfacial tension and on the contact angle of a water droplet in CO2 atmosphere on a silica surface. In the computational work the (001) surface of α-quartz was reconstructed. In studies reported previously developed force field and a surface model database for silica were used. In the surface models were derived from the (101) plane of α-cristobalite, which is stable above 1470°C and metastable at lower temperatures, and from the (100) plane of α-quartz. In the (202) plane of α-cristobalite was used to study how ion type and salinity affect wettabiltiy of quartz in presence of CO2.

In practical realizations of carbon capture and storage technology, quartz surfaces are far from the studied ideal ones. Deep saline aquifers, coal beds and depleted oil or gas reservoirs are considered as possible storage media for carbon dioxide. Within these formations mineral surfaces can be alkylated surfaces with chemisorpt organics, or are almost completely composed of organic matter (coal). Experimental studies indicate that these surfaces are intermediate-wet to strongly CO2-wet. To understand wetting of these alkylated surfaces and to explore wettabiltiy beyond the single contact angle we use fully hydroxylated (001) α-quartz surfaces with the highest theoretically possible concentration of hydroxyl groups. Note that the concentration of OH groups on
this surface is twice the average concentration observed in nature 4.6 OH/nm². This surface thus has the highest theoretically possible hydrophilicity. We demonstrate that this extreme hydrophilicity can be dramatically altered (lowered) even by minute concentrations of chemisorption organics, consistent with experimental data, which has serious implications for the carbon geo-storage capacity and security.

**Computational Details**

A general purpose parallel molecular dynamics simulation package DL_POLY 4.08 was applied to perform classical molecular mechanics computations. Ab-initio Density Functional Theory (DFT) calculations were performed with the version 6.3 of the Quantum ESPRESSO suite of computer codes, specifically PWscf and PostProc. The Bader charge analysis was performed with the version 1.03 of "Code: Bader Charge Analysis". Results of the simulations were visualized with VMD and VESTA software, the latter package was also used for some structural manipulations. The DL_POLY Graphical User Interface was used to prepare a model C₅H₁₁ group.

In all classical molecular dynamics simulations integration of the equations of motion was done with the velocity Verlet algorithm. The time step in all simulations was set to 2 fs. In the NPT simulations the Nose-Hoover thermostat and barostat with the relaxation constants 0.05 and 0.5 ps, respectively, were used. In the NVT simulations the Nose-Hoover thermostat with the relaxation constant 0.05 ps was used. Interactions cutoff distance was set to 17 Å in all calculations. Electrostatics were computed with the smoothed particle mesh Ewald summation.

**Force Fields**

A combination of force fields was employed in this study. The quartz crystal, the hydroxylated quartz surface and the SPC water were modelled with parameters of the CLAYFF force field. Carbon dioxide was represented by the EPM2 model. Chemisorption pentyl groups were simulated with the DREIDING force field. Ab-initio DFT calculations were used to rationalize parameters applied for coupling the CLAYFF and the DREIDING force fields, specifically the charges on atoms of the OC bond connecting the hydrocarbons to quartz.

Water and carbon dioxide molecules in all simulations, as well as the OH groups in the initial preparation of the hydroxylated quartz surface were treated as rigid bodies. Geometrical details of the rigid bodies were taken from the corresponding molecular models and force fields: CO distance (EPM2 CO₂) 1.149 Å; OH distance (SPC H₂O) 1 Å; HOH angle (SPC H₂O) 109.47°; OH distance (hydroxyl) 1 Å. To get convenient for pentyl groups integration onto the surface orientation of the OH groups involved in the weak hydrogen bonds (pointing slightly upwards, see 11, 13), and to highlight and isolate the weak hydrogen bonds - the place of attachment of the pentyl groups, only non-bond CLAYFF force field terms were used in the initial preparation of the hydroxylated quartz surface. All OH bond and Si-O-H angular potentials of the silanol groups were accounted for in the equilibration and the production runs of the fully assembled systems (alkylated quartz surface, water droplet, molecules of the CO₂ phase).

Parameters of potentials used to model the van der Waals interactions between all atomic species of studied in this work systems are listed in Table 1.

<table>
<thead>
<tr>
<th>Atomic species</th>
<th>A, A₁²-kcal/mol</th>
<th>B, A⁵-kcal/mol</th>
<th>q, e</th>
<th>m, amu</th>
</tr>
</thead>
<tbody>
<tr>
<td>C (CO₂)</td>
<td>4.31282·10⁴</td>
<td>9.82063·10¹</td>
<td>0.6512</td>
<td>12.0107</td>
</tr>
<tr>
<td>O (CO₂)</td>
<td>3.87857·10⁵</td>
<td>4.98236·10²</td>
<td>-0.3256</td>
<td>15.9994</td>
</tr>
<tr>
<td>Atomic species</td>
<td>A, A^{12}-kcal/mol</td>
<td>B, A^{6}-kcal/mol</td>
<td>q, e</td>
<td>m, amu</td>
</tr>
<tr>
<td>----------------</td>
<td>---------------------</td>
<td>-------------------</td>
<td>------</td>
<td>-------</td>
</tr>
<tr>
<td>O (H\textsubscript{2}O)</td>
<td>6.29342 x 10\textsuperscript{5}</td>
<td>6.25459 x 10\textsuperscript{2}</td>
<td>-0.82</td>
<td>15.9994</td>
</tr>
<tr>
<td>H (H\textsubscript{2}O)</td>
<td>0</td>
<td>0</td>
<td>0.41</td>
<td>1.0079</td>
</tr>
<tr>
<td>Si (SiO\textsubscript{2})</td>
<td>1.23698 x 10\textsuperscript{1}</td>
<td>9.54288 x 10\textsuperscript{-3}</td>
<td>2.1</td>
<td>28.0855</td>
</tr>
<tr>
<td>O (SiO\textsubscript{2} bridging)</td>
<td>6.29342 x 10\textsuperscript{2}</td>
<td>6.25459 x 10\textsuperscript{2}</td>
<td>-1.05</td>
<td>15.9994</td>
</tr>
<tr>
<td>O (SiO\textsubscript{2} hydroxyl)</td>
<td>6.29342 x 10\textsuperscript{2}</td>
<td>6.25459 x 10\textsuperscript{2}</td>
<td>-0.95</td>
<td>15.9994</td>
</tr>
<tr>
<td>H (SiO\textsubscript{2} hydroxyl)</td>
<td>0</td>
<td>0</td>
<td>0.425</td>
<td>1.0079</td>
</tr>
<tr>
<td>C (O-C\textsubscript{5}H\textsubscript{11})</td>
<td>1117341.71</td>
<td>667.516584</td>
<td>0 (0.338301)*</td>
<td>12.0107</td>
</tr>
<tr>
<td>H (O-C\textsubscript{5}H\textsubscript{11})</td>
<td>17198.6333</td>
<td>32.3369279</td>
<td>0</td>
<td>1.0079</td>
</tr>
<tr>
<td>O (O-C\textsubscript{5}H\textsubscript{11})</td>
<td>232115.998</td>
<td>298.083888</td>
<td>-0.863301**</td>
<td>15.9994</td>
</tr>
</tbody>
</table>

*charge on carbon of the OC bond is shown in brackets (estimated on basis of DFT calculations); **estimated on basis of DFT calculations.

Parameters in Table 1 correspond to the following analytical expression for the potential \( U \): \( U(r) = \frac{A}{r^{12}} - \frac{B}{r^{6}} \), where \( r \) is the interatomic distance.

The following formulae were used to convert original CLAYFF parameters \((D_0, R_0)\) to parameters representing the 12-6 potential:

\[
A = D_0 R_0^{12}, \\
B = 2 D_0 R_0^6. 
\]

The EPM2 potential parameters were converted to 12-6 potential parameters using:

\[
A = 4 \epsilon \sigma^{12}, \\
B = 4 \epsilon \sigma^6. 
\]

The interaction parameters between unlike atoms were obtained as the geometric mean for both A and B, Lorentz-Berthelot rules:

\[
A_{ij} = \sqrt{A_i A_j}, \\
B_{ij} = \sqrt{B_i B_j},
\]

where \( i \) and \( j \) are indices of the unlike atoms.

Apart from the rigid bodies all bonds and angles were treated explicitly using bond and angle potentials. The energy of every bond stretch was described by a harmonic relationship:

\[
U(r_{ij}) = \frac{k_1}{2} (r_{ij} - r_0)^2,
\]

where \( k_1 \) is the force constant, and \( r_0 \) represents the equilibrium bond length, see Table 2.

<table>
<thead>
<tr>
<th>Species ( i )</th>
<th>Species ( j )</th>
<th>( k_1 ), kcal/mol/A\textsuperscript{2}</th>
<th>( r_0 ), A</th>
</tr>
</thead>
<tbody>
<tr>
<td>O (SiO\textsubscript{2} hydroxyl)</td>
<td>H (SiO\textsubscript{2} hydroxyl)</td>
<td>1108.2698</td>
<td>1.0</td>
</tr>
<tr>
<td>C (O-C\textsubscript{5}H\textsubscript{11})</td>
<td>H (O-C\textsubscript{5}H\textsubscript{11})</td>
<td>700.0</td>
<td>1.09</td>
</tr>
<tr>
<td>Species i</td>
<td>Species j</td>
<td>$k_1$, kcal/mol/A²</td>
<td>$r_{0}$, A</td>
</tr>
<tr>
<td>-------------------</td>
<td>--------------------</td>
<td>-------------------</td>
<td>------------</td>
</tr>
<tr>
<td>C (O-C₅H₁₁)</td>
<td>C (O-C₅H₁₁)</td>
<td>700.0</td>
<td>1.53</td>
</tr>
<tr>
<td>C (O-C₅H₁₁)</td>
<td>O (O-C₅H₁₁)</td>
<td>700.0</td>
<td>1.288</td>
</tr>
<tr>
<td>Si (SiO₂)</td>
<td>O (O-C₅H₁₁)</td>
<td>700.0</td>
<td>1.455</td>
</tr>
</tbody>
</table>

The Si-O-H angles ($\theta$) of attached to the quartz surface hydroxyl groups were restrained by the harmonic potential ⁷⁴:

$$U(\theta) = \frac{k_2}{2} (\theta - \theta_0)^2,$$

where $k_2 = 60.0$ kcal/mol/rad² is the force constant, and $\theta_0 = 109.47^\circ$ is the equilibrium bond angle ⁷¹.

Bending of the Si-O-C₅H₁₁ angles of attached to the quartz surface O-C₅H₁₁ groups were modelled and parametrized according to the cosine potential ⁷⁴:

$$U(\theta) = A_1 (1 + \cos \theta),$$

where $A_1 = 100.0$ kcal/mol ⁷³.

Bending of all intramolecular angles of the O-C₅H₁₁ groups were modelled and parametrized using the harmonic cosine potential ⁷⁴:

$$U(\theta) = \frac{k_3}{2} (\cos \theta - \cos \theta_0)^2,$$

where $k_3 = 112.499693$ kcal/mol is the force constant, and $\theta_0 = 109.471^\circ$ is the equilibrium bond angle ⁷³.

Torsion of all intramolecular dihedral angles ($\varphi$) of the O-C₅H₁₁ groups were modelled and parametrized according to the cosine potential ⁷⁴:

$$U(\varphi) = A_2 (1 + \cos 3\varphi),$$

where $A_2 = 0.11111111$ kcal/mol ⁷³.

Modelling of the Oxygen-Carbon Bond

Utilized in this work force fields are designed and well suited for modelling minerals (the CLAYFF force field ⁷¹) and for modelling of organic molecules (the DREIDING force field ⁷³). However, specific for the purpose of this study description of a polar OC bond connecting pentyl groups to the quartz surface is not provided by either the CLAYFF or the DREIDING force fields. Thus, to extend capabilities of the force field pair a physically reasonable estimation of charges on oxygen and carbon atoms of the OC bond is necessary. This estimation was obtained from the DFT calculations which were performed as follows. Out of the orthorhombic cell of quartz crystal ⁷⁷ a super cell 2x1x1 was constructed (overall 36 atoms, 12 silicon atoms and 24 oxygen atoms). Using this cell three stages of computations were then performed. In the first stage the lattice vectors of the crystal were allowed to be changed and optimized during relaxation of ionic positions of the cell. In the second stage a three-layer slab (three heights of the orthorhombic cell) was constructed with a fully hydroxylated surface and integrated pentyl group, 30 Å of vacuum space was provided above the surface, see Figure 1. Only ions of this system were then relaxed with atoms of the bottom layer (one height of the orthorhombic unit cell) of the slab being fixed. In the third stage distribution of the charge density of the relaxed slab was determined with a single point energy calculation. In the third stage of these computations the kinetic energy cutoffs for plane waves and for charge density were set to 40 and 1500 Ry, respectively. The first Brillouin zone in the reciprocal space was sampled with the Monkhorst-Pack mesh ⁷⁸ 5x5x1. We
used the PAW (Projector Augmented Wave method) potentials H.pbe-kjpaw_psl.0.1.UPF, C.pbe-n-kjpaw_psl.0.1.UPF, O.pbe-n-kjpaw_psl.0.1.UPF and Si.pbe-n-kjpaw_psl.0.1.UPF from. The electron exchange and correlation energy was approximated by the PBE (Perdew, Burke and Ernzerhof) functional.

Figure 1. Optimized at the DFT level of theory three-layer quartz slab with integrated pentyl group. White balls - hydrogen atoms, light blue balls - carbon atoms, red balls - oxygen atoms, yellow balls - silicon atoms. Blue lines demark the unit cell of the simulated system, images of which are shown with lines instead of the balls.

The Bader charge analysis performed on the DFT computed electronic density resulted in a charge of -1.311489 e on the oxygen atom, and a charge of 0.338301 e on the carbon atom. This positive carbon charge corresponds to 0.796 of the charge on the hydroxyl hydrogen of the CLAYFF force field (the hydrogen atom which is to be removed and replaced by the carbon atom of the pentyl group). We assign this fraction of the CLAYFF hydroxyl hydrogen charge to the carbon of the OC bond. The rest of the hydroxyl hydrogen charge assigned to the oxygen of the OC bond makes the atom approximately 90% as negative (-0.863301 e) as the hydroxyl oxygen (-0.95 e) of the CLAYFF force field. This 90% proportion aligns well with results of the Bader charge analysis where the charges on these two oxygen atoms of the same geminal silanol group are -1.311489 and -1.477656 e, respectively. With these operations we thus resolve two problems at the same time. We replace the hydroxyl hydrogen with carbon of the pentyl group, and we redistribute the charge of the replaced hydrogen atom between the carbon and the oxygen atoms of the OC bond in a manner consistent with the DFT results.
Preparation of Alkylated Quartz Surfaces

Construction and modelling of modified (contaminated with hydrocarbons) quartz surfaces is hampered by the fact that structures of surfaces of pure quartz polymorphs are not well characterised even in 2018\textsuperscript{16-17}. On the basis of previous reconstructions of the hydroxylated (001) \(\alpha\)-quartz surface\textsuperscript{11-13} we suggest several model quartz surfaces with integrated pentyl groups. These models are then used to quantify the effect of quartz surface contamination with hydrocarbons on its wettability.

Preparation of the modified surfaces proceeded as follows: out of the primitive unit cell of \(\alpha\)-quartz\textsuperscript{82} its orthorhombic cell was created according to\textsuperscript{77}. The latter cell was then used to construct a super cell 2x1x1 (overall 36 atoms, 12 silicon atoms and 24 oxygen atoms). Out of this cell a super cell 4x5x8 was created to equilibrate the crystal at 1 atm pressure and 300 K temperature over 10\textsuperscript{5} steps. The system was simulated in the NPT ensemble. Used throughout this study, lattice vectors of the equilibrated orthorhombic super cell 2x1x1 amounted to the following values 9.989x8.651x5.492 Å, only 1.6\% larger than the crystallographic values obtained from\textsuperscript{82}. Four of these 2x1x1 cells (two in x direction and two in y direction) were then used to prepare a unit cell of fully hydroxylated quartz slab. The unit cell of the top layer of the 5x6x4 slab is shown in Figure 2. To obtain an ordered surface structure this four-layer (a layer in terms of the unit cell height) slab was relaxed at 10 K for 10\textsuperscript{5} steps. The system was simulated in the NVT ensemble. Atoms of the lowest laying layer (one unit cell height) of the slab were fixed to emulate the bulk crystal.
To explore the influence of different alkyl group surface concentrations on the contact angle, a uniform alkylation of the hydroxylated quartz surface was applied according to the cumulative sequence illustrated in Figure 2. The hydrogen atom of the geminal silanol group, which is not involved in formation of the strong hydrogen bond, was replaced with a pentyl (C₅H₁₁) group. The resulting surfaces with C₅H₁₁ surface concentrations ranging from 0.29 to 4.63 groups/nm² (in 0.29 groups/nm² steps) are demonstrated in Figure 3. The fully pentylated surface is depicted in Figure 4.
Figure 3. Top views of the top layer unit cells. 16 quartz surfaces with surface pentyl concentrations ranging from 0.289 to 4.629 groups/nm² (surfaces 1 to 16) are shown, unit cells are ordered from left to right and from bottom to top. White balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms, yellow balls - silicon atoms.
Figure 4. Top view (left) and views along x (right top) and y (right bottom) directions of fully pentylated quartz surface (surface 16). White balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms, yellow balls - silicon atoms. Top SiO₄HC₅H₁₁ groups are shown with balls, lower atoms are shown with lines of the same colour. Only top layer unit cell of four-layer slab is demonstrated.

It is worth noting that even a fully pentylated quartz surface still has a very high concentration of hydroxyl groups. Thus, with an increase in pentyl group concentration from 0 to 4.63 groups/nm² the concentration of hydroxyl groups simultaneously decreases linearly from 9.26 to 4.63 groups/nm², see the supporting information.

Simulation Setups

Using these alkylated quartz surfaces, 16 models overall containing a water droplet and CO₂ gas were prepared. Simulations were performed at 300 K and 10 MPa CO₂ pressure. The number of CO₂ molecules corresponding to this PT-condition in every simulation box was determined according to and amounted to 7188, 7137, 7086, 7035, 6984, 6933, 6881, 6830, 6779, 6728, 6677, 6625, 6574, 6523, 6472, 6421 for surfaces 1 through 16, respectively. When estimating the space available for CO₂ in the simulation boxes, gas volumes of the system with hydroxylated surface (no hydrocarbons) and of the system with fully pentylated surface were calculated as free space from the surface up to the CO₂ constraining potential. The volume of the water droplet was subtracted from the latter values, see Figure 5. The space available for CO₂ for the systems with intermediate pentyl group concentrations were obtained by interpolation.
Uniformly distributed carbon dioxide and water molecules were pre-equilibrated at 300 K in 100x104x60 and in 50x50x50 Å boxes, respectively. Water density was taken to be 1000 kg/m$^3$. Dynamics of these two systems were modelled in the NVT ensemble. Overall $5 \cdot 10^5$ steps were performed for CO$_2$ and $10^5$ steps for H$_2$O.

A half sphere of radius 26 Å atop of the C$_5$H$_{11}$ groups was filled with water molecules from the above described box. 946 H$_2$O molecules fit into the "water droplet". These "water droplets" were placed in the centre of the slabs. The rest of the space up to 60 Å above the C$_5$H$_{11}$ groups was filled with pre-equilibrated carbon dioxide. Vacuum space of 70 Å was provided on top of the 60 Å above the C$_5$H$_{11}$ groups. All simulations were performed with periodic boundary conditions in x and y directions. In z direction a repulsive force was set at 60 Å above the tip level of C$_5$H$_{11}$ groups:

$$F = k(z_0 - z), z > z_0,$$

where $k=1$ kcal/mol - is the force constant, and $z_0$ is the position of the repulsive potential.

The distance to the repulsive potential (60 Å above the C$_5$H$_{11}$ groups) ensures that the contact angle at the surface is not affected by perturbations in the CO$_2$ density caused by the wall.

Dynamics of the quartz-C$_5$H$_{11}$-H$_2$O-CO$_2$ systems were then simulated in the NVT ensemble. Overall $1.5 \cdot 10^6$ production steps (3 ns simulation time) were performed for every system preceded by $3 \cdot 10^5$ equilibration steps. Atoms of the lowest laying layer (one unit cell height) of the four-layer quartz slab were fixed in all simulations to emulate the bulk crystal. Dimensions of the simulation boxes were 99.89x103.81x159.17 Å. Figure 6 demonstrates the initial simulation setup for the surface with 1.16 C$_5$H$_{12}$/nm$^2$ surface concentration (four C$_5$H$_{11}$ groups in the unit cell, surface 4).
Determination of the Contact Angle

After the production runs an additional $5 \times 10^4$ simulation steps were performed at the same conditions. Five simulation snapshots separated by $10^4$ steps were taken from these final runs for the analysis. The contact angle was calculated using the water iso-density charts. To construct the charts the axis of a sufficiently large cylinder was aligned with the vertical axis passing through the centre of mass of all water molecules of a snapshot. The cylinder was then divided into thin disks $dH$, see Figure 7. Starting from the maximum radius $R_{\text{max}}$ and proceeding towards the central vertical axis, water density in every ring $dR$ of thickness $dH$ was calculated and compared with the half of water normal density (i.e. $0.033/2$ water molecules per cubic angstrom). In every disk a position $(H+dH/2, R+dR/2)$, where the water density first exceeds, if it exceeds at all, the half of the normal water density was recorded as the point of the iso-density.
The choice of \( dH \) and \( dR \) is a trade-off between two factors. Very small values result in only few molecules present within the ring's volume, which is also too small. The small number in the denominator in the density calculations in turn results in data scattering and significant errors. Very large values on the other hand smooth the density too much, which removes details and reduces accuracy. The following protocol was applied to substantiate the used values. Starting with 0.25 Å for the \( dH \) and \( dR \) they were then increased one after another in 0.25 Å steps until the algorithm became insensitive to molecules which escaped to the gas phase from the water droplet. Thus \( dH \) and \( dR \) amounted to 1.5 and 1.25 Å, respectively.

Constructed in the described way iso-density charts were then used to find the radius and position of a circle which best approximates the iso-density. The circle's position was limited to the vertical axis of the iso-density plot, which is the vertical axis passing through the centre of mass of all water molecules of the simulation snapshot. The BFGS algorithm\(^{84-87}\) was applied to find the minimum of a sum of absolute differences between the data points and the circle's profile. The theoretical and the apparent contact angles (for the definitions see the qualitative analysis of the results below) were then calculated as the angles of the tangential line to the circle's contour at its interception point with the surface. The quartz surface was set at the average vertical position of the hydrogen atoms of the hydroxyl groups, and the tip level of the pentyl groups was found as the average position of the top three hydrogen atoms of all \( \text{C}_5\text{H}_{11} \) groups in the snapshot. Averaged over the five simulation snapshots iso-density charts were used to calculate the hidden contact angle.
Results and Discussion

Qualitative Analysis of the Results

As an example, water iso-density chart with close to the average parameters (the contact angle, radius and vertical position of the spherical water droplet) and the circle fitted to the data points for quartz surface with pentyl concentration 1.157 C₅H₁₁/nm² (surfaces 4) is shown in Figure 8. The same iso-density charts for quartz surfaces with pentyl concentrations 0.289 through 3.182 groups/nm² (surfaces 1 through 11) are collected in the supporting information. All averaged contours of the water droplets for surfaces 1-11 gathered in one image are shown in Figure 9.

Figure 8. Water iso-density chart for quartz surface (surface 4) with concentration of pentyl groups 1.157 C₅H₁₁ per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line, the dotted line illustrates the planar projection of the water “skirt”.

Figure 9. All averaged contours of the water droplets for surfaces 1-11.
Examination of the water iso-density contours in the supporting information and Figure 9 reveals that the spherical water droplet loses contact with the quartz surface and the tip level of the pentyl groups when their concentration is higher than 2.604 C₅H₁₁ per square nm. The surface 10 does not have any connection with the droplet at all, even via a "tail" of water molecules. These observations drawn from the iso-densities are also supported by visual inspection of corresponding simulation snapshots with removed CO₂ molecules shown in the supporting information and exemplified here for the surface 10 in Figure 10. Above a 2.893 groups/nm² pentyl surface concentration (surface 10), the water droplet floats freely in the CO₂ phase. For concentrations higher than 3.182 groups/nm² (surface number higher than 11), the dimensions of the simulation box become too small for the droplets' motion and its distance travelled over the simulation period, the droplets interact with the repulsive wall, and cross borders of the simulation box.
Figure 10. Simulation snapshot of the quartz surface with pentyl concentration 2.893 groups per square nm (surface 10). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.

Additionally, the water iso-density charts indicate that a water "skirt" is formed between the quartz surface and the tip level of the pentyl groups. The water "skirt" is envisaged by authors as a conical frustum or a truncated cone. In the iso-density chart in Figure 8, and in the charts provided in the supporting information, planar projections of the "skirts" are shown with the dotted lines. These lines represent the fits to the first closest to the quartz surface peak in the water iso-density and the next highest data point of the iso-density. Formation of the "skirt" can also be supported by a simple visual examination of the side views of corresponding simulation snapshots with removed CO₂ molecules and C₅H₁₁ groups, see Figure 11 and the supporting information. The "skirt" is particularly clearly visible in the snapshot for the surface 4, view along y axis; in the snapshot for the surface 5, view along x axis; in the snapshot for the surface 6, view along y axis; in the snapshot for the surface 7, view along x axis; in the snapshot for the surface 8, view along x axis; in the snapshot for the surface 9, view along x axis. At low pentyl concentration, surface 1, the planar projection of the "skirt" is almost a tangential line to the droplet's profile. With increasing pentyl group concentration, the planar projection of the "skirt" starts to intersect the droplet's profile more explicitly, surface 2 and following. With further increase in concentration of C₅H₁₁ groups radius of the "skirt" decreases, and the "skirt" is pulled towards the centre of the rising droplet, surfaces 8 and 9.
This qualitative review, observations and inspections of the simulation results demonstrate that the wettability of alkylated quartz surface is a complex phenomenon. There are at least two planes (levels) which can be called surfaces: the surface of quartz itself and the tip level of pentyl groups; position of the latter also depends on the concentration of C5H11 groups. There is also a water "skirt" which is formed in between these two planes (levels).

To account for mentioned effects in quantitative description of the wettability of alkylated quartz surfaces we introduce the following parameters, see Figure 12. The theoretical contact angle ($\theta_T$), the contact angle of a spherical water droplet (spherical cap) at the level of quartz surface. The apparent contact angle ($\theta_A$), the contact angle of a spherical water droplet (spherical cap) at the tip level of pentyl groups; this contact angle is most likely to be measured in an experiment, with the difference between $\theta_T$ and $\theta_A$ being large and more noticeable for longer alkyl groups. The hidden contact angle ($\theta_H$), the contact angle of a water "skirt" at the level of the quartz surface.
Quantitative Analysis of the Results

Circles fitted to the iso-density charts and corresponding circumscribed spheres shrink with increasing pentyl group concentration. When the pentyl concentration exceeds 1.5 C₅H₁₁/nm² (surface number 6 and higher) the radius stabilizes at 18.4 Å, see Figure 13. At this point almost all water molecules become part of a spherical droplet.

![Graph showing dependence of circumscribed sphere radius on concentration of pentyl groups.](image)

*Figure 13. Dependence of the circumscribed sphere (circle) radius on concentration of pentyl groups.*

With increasing pentyl concentration the droplet moves upwards, see Figure 14, and loses contact with the tip level of pentyl groups at 2.604 C₅H₁₁/nm² (surface 9), see Figure 15, although the water "skirt" is still present at this concentration, see the supporting information, but disappears at 2.893 C₅H₁₁/nm² (surface 10). Spherical part of the droplet loses contact with the quartz surface at 2.025 C₅H₁₁/nm² pentyl surface concentration (surface 7), see Figure 15. At this pentyl concentration the "skirt" is still in place.
Figure 14. Dependence of the vertical position of the circumscribed sphere (circle) centre on concentration of pentyl groups. Position of the quartz surface is at zero.

Figure 15. Circumscribed sphere (circle) or spherical droplet contact with the quartz surface (height zero) and the tip level of pentyl groups as a function of \(\text{C}_5\text{H}_{11}\) concentration.

Dependences of the theoretical and the apparent contact angles on surface pentyl group concentration are shown in Figure 16 and Figure 17, respectively. Both contact angles increase with rising pentyl concentration, while the apparent contact angle is generally smaller and increases less sharply. It is reasonable to expect that for longer alkyl groups the difference between the theoretical and the apparent contact angles will be higher because of higher tip level of the alkyl groups.
Four wettability zones can be identified in Figure 16 and Figure 17, see Table 3. The first one is the water-wet zone, where pentyl concentrations range from 0.289 to 0.579 $\text{C}_5\text{H}_{11}/\text{nm}^2$, surfaces 1 and 2. The second zone is intermediate-wet, where pentyl concentrations range from 0.868 to 1.447 $\text{C}_5\text{H}_{11}/\text{nm}^2$, surfaces 3, 4 and 5. The third zone is CO$_2$-wet, where pentyl concentrations range from 1.736 to 2.314 $\text{C}_5\text{H}_{11}/\text{nm}^2$, surfaces 6, 7 and 8.

For higher concentrations in the zone four, starting from 2.604 $\text{C}_5\text{H}_{11}/\text{nm}^2$, surface 9, the surfaces are completely hydrophobic (the contact angle is 180°) according to both the theoretical and the apparent contact angles.

<table>
<thead>
<tr>
<th>Wettability</th>
<th>Zone 1</th>
<th>Zone 2</th>
<th>Zone 3</th>
<th>Zone 4</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Water wet</td>
<td>Intermediate</td>
<td>CO$_2$</td>
<td>Extremely hydrophobic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>wet</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3. Wettability zones according to the theoretical and the apparent contact angles.
<table>
<thead>
<tr>
<th>Pentyl concentration, groups/nm²</th>
<th>Zone 1</th>
<th>Zone 2</th>
<th>Zone 3</th>
<th>Zone 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface, number</td>
<td>1-2</td>
<td>3-5</td>
<td>6-8</td>
<td>9 and higher</td>
</tr>
<tr>
<td>Theoretical contact angle, degree</td>
<td>22.01-40.33</td>
<td>89.90-122.37</td>
<td>173.98-180</td>
<td>180</td>
</tr>
<tr>
<td>Apparent contact angle, degree</td>
<td>15.40-31.54</td>
<td>77.30-105.49</td>
<td>143.13-177.88</td>
<td>180</td>
</tr>
</tbody>
</table>

The numerical values of the theoretical and apparent contact angles predicted here follow the same trend as observed experimentally in comparable systems obtained for calcite surfaces exposed to stearic acid, where a calcite surface was aged in highly diluted stearic acid solutions (concentrations ranged from $10^{-10}$ to $10^{-2}$M). For these concentrations, the contact angle of a water droplet in CO$_2$ atmosphere measured at $10$ MPa and $323$ K increased from $43$ to $126^\circ$. The same trend is observed in this study, where the contact angle rapidly increased in the range of small pentyl concentrations ($0.289-2.314$ C$_5$H$_{11}$/nm$^2$) from $15$ to $180^\circ$.

An elaborate analysis of the hidden contact angle reveals an even more detailed picture of the processes occurring on the quartz surface. The hidden contact angle clearly demonstrates that surface 9 and the corresponding pentyl concentration ($2.604$ C$_5$H$_{11}$/nm$^2$) represent a transition concentration at which hydrophobicity takes over, see Figure 18. All surfaces with pentyl concentrations $2.893$ C$_5$H$_{11}$/nm$^2$ or higher (surface number 10 or higher), have less affinity to water. Indeed, surface 10 is the surface where the water "skirt" disappears completely, see Figure 10.

![Figure 18. Dependence of the hidden contact angle on concentration of pentyl groups.](image)

With increasing surface pentyl group concentration the hidden contact angle behaves differently in comparison with the theoretical and the apparent contact angles. Noticeable increase in the value of the hidden contact angle is observed at concentrations higher than $2$ C$_5$H$_{11}$/nm$^2$. At concentrations $2.025$ and $2.314$ C$_5$H$_{11}$/nm$^2$ (surfaces 7 and 8) the hidden contact angle is almost $30^\circ$. It increases further to slightly more than $60^\circ$ at $2.604$ C$_5$H$_{11}$/nm$^2$ concentration (surface 9). Interestingly, the value of the hidden contact angle at a concentration of $1.736$ C$_5$H$_{11}$/nm$^2$ (surface 6) and lower is almost constant and on average amounts to slightly less than $10^\circ$. 
The hidden contact angle is thus a useful characteristic which shows that the alkylated quartz surface exhibits close to a binary behaviour in terms of its wettability. It is water wet and hydrophilic when concentrations of pentyl groups are less than $2.604 \, \text{C}_5\text{H}_{11}/\text{nm}^2$ (surface 9), and it is extremely hydrophobic at higher pentyl concentrations, Figure 18. Very sharp rise of the contact angle over a small range of pentyl concentrations practically makes the surface either hydrophilic or hydrophobic. We attribute this close to the binary wettability to the steric effects preventing water molecules from reaching the quartz surface, compare number of water molecules at the surface 9 (and lower) and at the surface 10 (and higher) in the supporting information.

In addition to determined contact angles, it is possible to estimate when the wettability regime of alkylated quartz surfaces changes from homogenous to the heterogeneous. To do so let us first consider the basic equation of the Cassie-Baxter model 88:

$$\cos \theta_{CB} = f_1 \cos \theta_1 + f_2 \cos \theta_2,$$

where $\theta_{CB}$ is the Cassie-Baxter contact angle; $f_1$ and $f_2$ are the area fractions for surface components 1 and 2, for which the contact angles are $\theta_1$ and $\theta_2$, respectively.

The first surface component in our case is the fully hydroxylated quartz surface for which we approximate the contact angle with the average hidden contact angle ($9.81^\circ$) at low pentyl concentrations $1.736 \, \text{C}_5\text{H}_{11}/\text{nm}^2$ (surface 6) and smaller. The second surface component is the alkylated quartz surface with minimal pentyl concentration for which the contact angle becomes equal to $180^\circ$. The Cassie-Baxter contact angle is the apparent contact angle characterised in this work. Taking into account the relationship between the area fractions ($f_2=1-f_1$), the following expression can then be obtained:

$$f_1 = \frac{\cos \theta_1 + 1}{\cos \theta_2 + 1}.$$

Using the last formula we calculate the surface area per one $\text{C}_5\text{H}_{11}$ group and plot it as a function of $\text{C}_5\text{H}_{11}$ concentration, Figure 19. At concentrations $0.868 \, \text{C}_5\text{H}_{11}/\text{nm}^2$ (surface 3) and higher, the surface area per one pentyl group stabilizes at $0.46 \, \text{nm}^2$ per group. The concentration at which this surface area becomes constant is the concentration when the wetting regime becomes heterogeneous. In this regime every pentyl group added to the surface unit cell increases the surface fraction $f_1$ by the same amount. When there are eight groups within the unit cell (surface 8) the fraction becomes approximately equal to one. For all concentrations higher than $2.314 \, \text{C}_5\text{H}_{11}/\text{nm}^2$ (surface 8) the apparent contact angle remains at $180^\circ$ and the wetting regime again becomes homogenous, but now on the alkylated surface. It should be noted that the range of surface pentyl concentrations of the heterogeneous wetting regime coincides with the range of surface pentyl concentrations of above mentioned wettability zones 2 and 3.
Figure 19. Surface area per one C₅H₁₁ group as a function of its surface concentration. Dotted line averages values at the plateau.

In atomistic simulations analysed in this study, surfaces are smooth from a macroscopic perspective, and it has been shown previously experimentally that a minor surface roughness has no significant influence on the water/CO₂ contact angle of silica systems 32.

Conclusions

A systematic computational study of wettability of fully hydroxylated and alkylated (001) α-quartz surface under 10 MPa pressure of carbon dioxide at 300 K with respect to surface concentration of pentyl groups is performed. Considered concentrations of the alkyl groups vary from 0.29 to 4.63 C₅H₁₁/nm². It is shown that when the surface density of pentyl groups is increased from 0.29 to 3.18 C₅H₁₁/nm² the contact angle of a water droplet changes from 10‐20° to 180°.

A complete description of wettability of alkylated quartz surface requires three parameters: the theoretical contact angle, the apparent contact angle and the hidden contact angle. These contact angles are determined at the tip level of pentyl groups and the level of the quartz surface. The hidden contact angle is calculated as the angle of a water "skirt", which is formed between the level of the quartz surface and the tip level of pentyl groups. The hidden contact angle remains relatively small (less than 90°) up to the threshold pentyl density of 2.604 C₅H₁₁/nm², after which it rather abruptly increases to 180°. We call such a behaviour the binary wettability, when the surface is either hydrophilic or extremely hydrophobic, with intermediate wetting regime observed only in a very narrow range of the surface pentyl density of 0.29 C₅H₁₁/nm². We conclude that this binary wettability of alkylated quartz surface originates from the steric effects preventing water molecules from reaching the quartz surface.

The theoretical and the apparent contact angles are determined as the contact angles of the spherical droplet at the level of the quartz surface and at the tip level of pentyl groups, respectively. The apparent contact angle is the angle which is most likely to be measured in an experiment and it will deviate more from the theoretical one with longer chains of chemisorpt hydrocarbons. On basis of the theoretical and the apparent contact angles four wettability zones can be identified. In the zone 1 (water wet zone) the pentyl density is in range 0.289-0.579 C₅H₁₁/nm², and the contact angle is in range 15-40°. In the zone 2 (intermediate wet zone) the pentyl density is in range 0.868-1.447 C₅H₁₁/nm², and the contact angle is in range 77-122°. In the zone 3 (CO₂ wet zone) the pentyl
density is in range 1.736-2.314 C₅H₁₁/nm², and the contact angle is in range 143-180°. In the zone 4 (extremely hydrophobic zone) the pentyl density is 2.604 C₅H₁₁/nm² or higher, and the contact angle is 180°. Additionally, the wettability regime in zones 2 and 3 can be regarded as heterogeneous.

A possible future research direction is the modification of the quartz surface with varied alkyl chain length. It is reasonable to expect that in this case the contact angle will be mostly determined by the parameters for the hydrocarbons, thus making the surface even more hydrophobic.

Supporting Information
Surface hydroxyl group concentration vs surface pentyl group concentration, water iso-density charts for surfaces 1 through 11, simulation snapshots for surfaces 1 through 11, simulation snapshots with removed pentyl groups for surfaces 1 through 11.

Acknowledgments
This work was supported by resources provided by the Pawsey Supercomputing Centre with funding from the Australian Government and the Government of Western Australia.

References


24. IPCC *Carbon Dioxide Capture and Storage*; Intergovernmental Panel on Climate Change; 2005, 2005.


Wettability of Fully Hydroxylated and Alkylated (001) alpha-Quartz Surface in Carbon Dioxide Atmosphere

Aleksandr Abramov*, Stefan Iglauer, Alireza Keshavarz, School of Engineering, Edith Cowan University, 270 Joondalup Drive, Joondalup, WA 6027, Western Australia, Australia

*aabramov@our.ecu.edu.au

Surface Hydroxyl Group Concentration vs Surface Pentyl Group Concentration

Figure S1. Surface hydroxyl group concentration as a function of surface pentyl group concentration.
Figure S2. Water iso-density chart for quartz surface (surface 1) with concentration of pentyl groups 0.289 \( \text{C}_5\text{H}_{11} \) per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line, the dotted line illustrates the planar projection of the water "skirt".
Figure S3. Water iso-density chart for quartz surface (surface 2) with concentration of pentyl groups 0.579 \( \text{C}_5\text{H}_{11} \) per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line, the dotted line illustrates the planar projection of the water "skirt".
Figure S4. Water iso-density chart for quartz surface (surface 3) with concentration of pentyl groups 0.868 C₅H₁₁ per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line, the dotted line illustrates the planar projection of the water "skirt".
Figure S5. Water iso-density chart for quartz surface (surface 4) with concentration of pentyl groups 1.157 \( \text{C}_5\text{H}_{11} \) per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line, the dotted line illustrates the planar projection of the water "skirt".
Figure S6. Water iso-density chart for quartz surface (surface 5) with concentration of pentyl groups 1.447 C\textsubscript{5}H\textsubscript{11} per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line, the dotted line illustrates the planar projection of the water "skirt".
Figure S7. Water iso-density chart for quartz surface (surface 6) with concentration of pentyl groups 1.736 C₅H₁₁ per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line, the dotted line illustrates the planar projection of the water “skirt”.
Figure S8. Water iso-density chart for quartz surface (surface 7) with concentration of pentyl groups 2.025 C$_{5}$H$_{11}$/per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line, the dotted line illustrates the planar projection of the water "skirt".
Figure S9. Water iso-density chart for quartz surface (surface 8) with concentration of pentyl groups 2.314 $\text{C}_5\text{H}_{11}$ per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line, the dotted line illustrates the planar projection of the water "skirt".
Figure S10. Water iso-density chart for quartz surface (surface 9) with concentration of pentyl groups 2.604 C₅H₁₁ per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line, the dotted line illustrates the planar projection of the water "skirt".
Figure S11. Water iso-density chart for quartz surface (surface 10) with concentration of pentyl groups 2.893 \( \text{CsH}_{11} \) per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line.
Figure S12. Water iso-density chart for quartz surface (surface 11) with concentration of pentyl groups 3.182 C\textsubscript{5}H\textsubscript{11} per square nm. The iso-density data points are shown with blue dots, fitted circle is shown with blue line, the tip level of the pentyl groups is shown with the dashed line.
Simulation Snapshots for Surfaces 1 Through 11

Figure S13. Simulation snapshot of the quartz surface with pentyl concentration 0.289 groups per square nm (surface 1). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.
Figure S14. Simulation snapshot of the quartz surface with pentyl concentration 0.579 groups per square nm (surface 2). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.
Figure S15. Simulation snapshot of the quartz surface with pentyl concentration 0.868 groups per square nm (surface 3). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.
Figure S16. Simulation snapshot of the quartz surface with pentyl concentration 1.157 groups per square nm (surface 4). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.
Figure S17. Simulation snapshot of the quartz surface with pentyl concentration 1.447 groups per square nm (surface 5). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.
Figure S18. Simulation snapshot of the quartz surface with pentyl concentration 1.736 groups per square nm (surface 6). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.
Figure S19. Simulation snapshot of the quartz surface with pentyl concentration 2.025 groups per square nm (surface 7). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.
Figure S20. Simulation snapshot of the quartz surface with pentyl concentration 2.314 groups per square nm (surface 8). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.
Figure S21. Simulation snapshot of the quartz surface with pentyl concentration 2.604 groups per square nm (surface 9). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.
Figure S22. Simulation snapshot of the quartz surface with pentyl concentration 2.893 groups per square nm (surface 10). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.
Figure S23. Simulation snapshot of the quartz surface with pentyl concentration 3.182 groups per square nm (surface 11). View along x axis: top left; view along y axis: top right; view along z axis: bottom left; isometric view: bottom right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface is illustrated with square in light yellow color. Water molecules and pentyl groups are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms, light blue balls - carbon atoms.
Figure S24. Side views of a water droplet on the quartz surface with removed pentyl groups for pentyl concentration 0.289 groups per square nm (surface 1). View along x axis: left; view along y axis: right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface position is indicated by the lowest laying water molecules. Water molecules are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms.

Figure S25. Side views of a water droplet on the quartz surface with removed pentyl groups for pentyl concentration 0.579 groups per square nm (surface 2). View along x axis: left; view along y axis: right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface position is indicated by the lowest laying water molecules. Water molecules are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms.
Figure S26. Side views of a water droplet on the quartz surface with removed pentyl groups for pentyl concentration 0.868 groups per square nm (surface 3). View along x axis: left; view along y axis: right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface position is indicated by the lowest laying water molecules. Water molecules are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms.

Figure S27. Side views of a water droplet on the quartz surface with removed pentyl groups for pentyl concentration 1.157 groups per square nm (surface 4). View along x axis: left; view along y axis: right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface position is indicated by the lowest laying water molecules. Water molecules are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms.
Figure S28. Side views of a water droplet on the quartz surface with removed pentyl groups for pentyl concentration 1.447 groups per square nm (surface 5). View along x axis: left; view along y axis: right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface position is indicated by the lowest laying water molecules. Water molecules are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms.

Figure S29. Side views of a water droplet on the quartz surface with removed pentyl groups for pentyl concentration 1.736 groups per square nm (surface 6). View along x axis: left; view along y axis: right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface position is indicated by the lowest laying water molecules. Water molecules are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms.
Figure S30. Side views of a water droplet on the quartz surface with removed pentyl groups for pentyl concentration 2.025 groups per square nm (surface 7). View along x axis: left; view along y axis: right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface position is indicated by the lowest laying water molecules. Water molecules are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms.

Figure S31. Side views of a water droplet on the quartz surface with removed pentyl groups for pentyl concentration 2.314 groups per square nm (surface 8). View along x axis: left; view along y axis: right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface position is indicated by the lowest laying water molecules. Water molecules are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms.

Figure S32. Side views of a water droplet on the quartz surface with removed pentyl groups for pentyl concentration 2.604 groups per square nm (surface 9). View along x axis: left; view along y axis: right. The
sphere fitted to the iso-density chart is illustrated in light purple color. The surface position is indicated by the lowest laying water molecules. Water molecules are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms.

Figure S33. Side views of a water droplet on the quartz surface with removed pentyl groups for pentyl concentration 2.893 groups per square nm (surface 10). View along x axis: left; view along y axis: right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface position is indicated by the lowest laying water molecules. Water molecules are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms.

Figure S34. Side views of a water droplet on the quartz surface with removed pentyl groups for pentyl concentration 3.182 groups per square nm (surface 11). View along x axis: left; view along y axis: right. The sphere fitted to the iso-density chart is illustrated in light purple color. The surface position is indicated by the lowest laying water molecules. Water molecules are shown in VDW representation, white balls - hydrogen atoms, red balls - oxygen atoms.