Surface Mediated Structures: Stabilization of Metastable Polymorphs on the Example of Paracetamol

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Supporting Information

ABSTRACT: The preparation of typically thermodynamically unstable polymorphic structures is a challenge. However, solid surfaces are well established aids for the formation and stabilization of polymorphic structures within, for instance, organic electronics. In this study, we report the stabilization of a pharmaceutically relevant substance via a solid surface at ambient conditions. Form III of paracetamol, which is typically unstable in the bulk at standard conditions, can be stabilized with a model silica surface by a standard spin coating procedure followed by rapid heat treatment. Such a preparation technique allows the use of atomic force microscopy and grazing incidence X-ray diffraction measurements revealing detailed information on the morphology and structure of the polymorph. Furthermore, the results exhibit that this polymorph is stable over a long period of time revealing surface mediated stabilization. These findings demonstrate a novel approach to provide thermodynamic stability when applied to similar molecules with specific applications.
rapid temperature increase was necessary to obtain form III. This allows the assumption that the crystallization in the oven occurs at the hot solid substrate further allowing the thermodynamic unstable polymorph to be entrapped and stabilized at the silica substrate even without the exclusion of air. It is well-known in the literature that form III is unstable at ambient conditions under air and interconverts into form II and I.\textsuperscript{16}

GIXD measurements of the three samples are shown in Figure 1. The GIXD measurement in general allows netplanes which are close to the surface-normal to be detected within thick films\textsuperscript{19} (up to hundreds of nanometers for organic layers) as well as within thin films consisting of a monolayer.\textsuperscript{20} High intensity spots correspond to Bragg reflections which can be used to index the pattern and thus to identify their crystal structures. The measurements of the sample containing form I reveal ring-like Bragg reflections showing that the crystallites arrange like a random oriented powder; i.e., no preferred orientation is observed. The indexation shows that all rings are a result of paracetamol being in the thermodynamic stable polymorph form I with a monoclinic unit cell (Figure 2). The GIXD pattern of form II shows defined spots at $q_z = 0.0, 0.8, 1.7$ nm\textsuperscript{-1} and various $q_{xy}$. The indexation reveals that these spots are a result of paracetamol being in form II conformation with an orthorhombic unit cell. In addition, the crystallites of form II show a preferred orientation with respect to the surface, whereby the 001 plane is in contact with the surface. However, rings are also present within the pattern indicating random oriented form I domains. This shows that the sample contains two polymorphs simultaneously, even if the amount of form I is very low compared to form II. The form III sample reveals a GIXD pattern with spots being distinct from the previously observed ones. The indexation of the spots can be achieved by introducing a 021 contact plane with respect to the surface with an orthorhombic unit cell (compare Figure 2). The smearing of the Bragg spots shows that the mosaicity of the crystalline needles is relatively high. This means that the molecules have a certain degree of freedom to assemble at the silica surface which is different compared to form II with a nearly perfect alignment.

In Figure 2, the visualization of the three different crystal structures of paracetamol form I,\textsuperscript{21} II,\textsuperscript{21} and III\textsuperscript{11} together with the contact plane with respect of the SiO$_2$ surface. The table below summarizes the used crystal lattice parameters with their corresponding CSD code, space group, and corresponding temperature at which the experiments were performed.
those. Form I is visualized with its monoclinic unit cell with no preferential alignment with respect to the surface due to its random orientation. The herringbone structure also is responsible for the mechanical stability and the disability for compaction required in tablet preparation.

In Figure 3 the morphologies of the three samples containing each polymorph of paracetamol at room temperature are shown with their corresponding height profiles. The AFM image of form I reveals the typical shape of monoclinic crystals with prismatic to plate-like morphology, whereby the crystallites are randomly rotated with respect to each other. Form II shows a distinct growth morphology, and plate-like structures are present. While the crystal morphologies are distinct, the height and consequently the coverage are very similar to form I and II, showing that diffusion in the upward direction from the surface is negligible. These two observed morphologies fit very well to previously observed experiments on paracetamol, where morphological differences are explained by different growth faces being dominant.22 The morphology of form III is distinct from the previous two forms, and needle-like structures are present. The coverage of the surface is strongly reduced, and consequently the heights of the needle-islands are higher compared to those of form I and II even though the nominal film thickness of the spin coated sample was very similar. The height of the needles is around 450 nm, which is about three times larger compared to the previous observed heights of form I and II.

Differences in the crystal structure and alignment are also reflected within their morphologies. The structure of form II is shown in a 001 orientation (Figure 2, green plane) which is parallel to the silica surface. This visualization reveals that the paracetamol molecules arrange in a way that maximizes their contact areas with the silicon surface. Further, the delocalized π-orbitals are slightly tilted toward the surface, but due to the large contact area, also the polar groups are in the highest possible contact to interact with the semipolar silica. Again intersheet H-bonds are visible, while intrasheet bonds of the hereby flatly arranged paracetamol sheets are not present (slipping plane of form II). According to the literature, form II exhibits a plate-like growth whereby the slowest growing crystal face is the 001, which is in excellent agreement with our observations.22 These findings indicate that growing into the a- and b-axis direction of the orthorhombic unit cell is equally likely and more favorable compared to the stacking into the c-axis direction. The arrangement of paracetamol molecules within the orthorhombic form III unit cell with respect to the 021 direction indicates that the molecules form hydrogen bonds toward the silicon surface and that the molecules stand nearly perpendicular to the substrate (Figure 2). Again the intersheet connection exhibits H-bonds, while the intrasheet stacking lacks any of those. The upright alignment of the form III sheets is in very good agreement with the observed needle-like morphology which was measured with AFM (Figure 3), whereby the paracetamol sheets seem to grow into the c-axis of the orthorhombic unit cell, or the molecules are perpendicular to the long needle axis. Further, the stability and reproducibility of the surface mediated stabilization of all three polymorphs of paracetamol was investigated. During the course of the synchrotron experiment, fresh prepared samples as well as 4-week-old samples were characterized revealing the same diffraction patterns. The samples were stored at ambient conditions (25 °C, relative humidity ~30%, 1 atm and under air environment), which reflects the potential of surface mediated stabilization. In addition, AFM investigations of the older samples did not reveal any significant change in the morphology, indicating that there is no rearrangement upon storage.

The surface mediated polymorph stabilization which was studied within this work using paracetamol is in excellent agreement with Ostwald’s step rule which states that the least thermodynamic stable polymorph crystallizes first.16 In particular, this means that a system moves to a thermodynamic equilibrium from an initial high energy state, whereby the least stable polymorph crystallizes first and rearranges stepwise into the different polymorphs (form III > form II > form I) due to changes within the free energy. Further, the different preparation conditions and the preferential alignment of the paracetamol molecules with respect to the surface indicate that the intensive (e.g., temperature, viscosity, chemical potential, density, etc.) and extensive (e.g., Gibbs free energy, entropy, mass, number of molecules, etc.) parameters of the solvent, analyte, and substrate in use are of crucial importance during surface mediated stabilization. EtOH exhibits a relatively low vapor pressure compared to THF. This suggests that EtOH solvent residues remain within the amorphous film, whereby molecule diffusion promotes the rearrangement in a thermodynamic stable configuration even if the viscosity of EtOH is about 2 times higher. Contrarily, the vapor pressure of THF is about 3 times larger compared to EtOH at standard conditions, meaning that THF molecules evaporate fast allowing the thermodynamic metastable polymorph II to be stabilized at the silica surface at 298 K. A shorter time frame typically means that formation of a thermodynamically less stable form is favored in accordance with other literature reports.23,24 The unstable forms II and III most likely develop due to the fast processing condition within the THF solutions. A rapid temperature increase to 383 K of an amorphous paracetamol film spin coated from THF results in polymorph III being stabilized at the silica surface. An increase in temperature typically reduces the H-bonding interaction strength which reduces the affinity for the molecules to interact with the substrate. Furthermore, higher temperatures mean that the
molecules require more space on account of molecular vibration. Both effects favor the formation of upright standing molecules on top of the substrate surface which in the case of paracetamol means additionally assembling into form II.

By comparing the observed alignment of paracetamol form III with different literature statements, it can be seen that the orientation of form III crystallites is highly influenced by the surface in use even if different preparation methods indicate similar growth kinetics of form III. In a recent publication,25 the authors state that form III, when confined in self-ordered anodic aluminum oxide nano tubes (AAO), preferentially crystallizes with its 001 plane being parallel to the AAO interface. This alignment indicates that no H-bonds are formed between the AAO host and the paracetamol molecules. Contrarily, our investigations strongly indicate the formation of H-bonds between the semipolar silica substrate and the paracetamol molecules on top of the substrate surface which in the case of paracetamol means additionally assembling into form III.

The authors declare no competing financial interest.

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