**Understanding the formation of apremilast cocrystals**

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**Abstract**

Apremilast (APR), an anti-psoriatic agent, easily forms isostructural cocrystals and solvates with aromatic entities, often disobeying at the same time Kitaigorodsky’s rule as to the saturation of possible hydrogen bonding sites. In this paper the reasons standing behind this peculiar behavior are investigated, employing a joint experimental and theoretical approach. This includes the design of cocrystals with coformers having high propensity towards the formation of both aromatic – aromatic and hydrogen bonding interactions, determination of their structure, using solid state NMR spectroscopy and X-Ray crystallography, as well as calculations of stabilization energies of formation of the obtained cocrystals, followed by crystal structure prediction calculations and solubility measurements. Our findings indicate that the stabilization energies of cocrystal formation are positive in all cases, which results from strain in the APR conformation in these crystal forms. On the other hand, solubility measurements show that the Gibbs free energy of formation of the apremilast : picolinamide cocrystal is negative, suggesting that the formation of the studied cocrystals is entropy-driven. This entropic stabilization is associated with the disorder observed in almost all known cocrystals and solvates of APR.

**Introduction**

Pharmaceutical cocrystals are usually defined as entities with at least two different solid components confined in a single crystal lattice, with at least one of these components being an active pharmaceutical ingredient (API), whereas the other one being either another API, or, more frequently, a substance from the generally-regarded-as-safe, GRAS, list (Bolla & Nangia, 2016). The components of a cocrystal are held together by non-covalent interactions, most frequently by hydrogen bonds, but also by halogen bonds or aromatic – aromatic interactions (Mukherjee & Desiraju, 2014; Wicker *et al.*, 2017). A significant increase of scientific interest in their formation and stability can be explained by the fact that cocrystals, just like any new solid crystal form, can offer different, sometimes improved physicochemical properties, including solubility, thermal stability and bioavailability (Karki *et al.*, 2009; Douroumis *et al.*, 2017; Grifasi *et al.*, 2015; Kaur *et al.*, 2017). For example, the formation of cocrystals of glibenclamide (Goyal *et al.*, 2018), epalrestat (Putra *et al.*, 2017), quercetin (Vasisht *et al.*, 2016), and cilostazol (Yoshimura *et al.*, 2017) has been recently proven to be profitable in enhancing water solubility of an API, when compared with the respective neat solid form.

One API with poor water solubility and short shelf-life is apremilast (APR), a phosphodiesterase-4 inhibitor used in psoriasis and psoriatic arthritis (Schafer, 2012). Its commercially available form B, characterized only by its powder X-Ray diffractogram, is regarded as the most thermodynamically stable one (US Patent 9850205B2, 2017), but suffers from the above mentioned unfavorable physicochemical properties (Shakeel *et al.*, 2017). It is not surprising, therefore, that new forms are sought after, and so far a number of better soluble isostructural solvates and cocrystals has been reported (Wu *et al.*, 2017a; Wu *et al.*, 2017b; Dudek *et al.*, 2018; Wang *et al.*, 2018; European Patent EP3339292A1, 2018). One of the most characteristic and unique features of these forms is the fact, that they are held together almost entirely by aromatic – aromatic interactions, even though the solvent and/or coformer molecules used for their formation, as well as APR itself, are capable of forming strong hydrogen bonds (Dudek *et al.*, 2018; Wang *et al.*, 2018).

A detailed analysis of functional groups present in APR indicates that, apart from the two sites engaged in an intramolecular hydrogen bond, *i.e.* CO and NH amide groups, there are at least 6 sites with strong propensity of acting as hydrogen bond acceptors: two oxygen atoms from the sulfonyl group, two methoxy oxygen atoms, and two carbonyl groups (Figure 1). A survey of the sulfonyl group propensity towards the formation of hydrogen bonds has shown that in all of 12 analyzed cases of newly synthesized sulfoxides, as well as those retrieved from the CSD database oxygen atoms from sulfonyl group acted as hydrogen bond acceptors, while in 10 cases methylene protons acted as hydrogen bond donors (Brondel *et al.*, 2010). In the cited reference the selection of structures was based on obtaining a wide variety of sulfur-containing organic compounds with different substituents, so that it enabled investigating the influence of the functional group on the intermolecular interactions of sulfoxides found in the crystals. Similarly, it has been demonstrated that oxygen from the methoxy group in methoxybenzenes is a strong acceptor of hydrogen bonds from various donor groups (Palusiak & Grabowski, 2002), not to mention the carbonyl oxygen from isoindoline-1,3-dione ring, which often acts as a hydrogen bond acceptor for C-H aromatic protons. Meanwhile, in APR cocrystals and solvates none of these hydrogen bond donors and acceptors are involved in the formation of such a bond, neither with another APR molecule nor a coformer (Wu *et al.*, 2017a; Dudek *et al.*, 2018). Clearly, certain factors capable of compensating the observed deviation from the maximum saturation of hydrogen bonds trend upon formation of cocrystals have to exist. This trend was observed as early as in 1973 by Kitaigorodsky and formulated by him into a rule, claiming that hydrogen bonds achievable for a given structure must be saturated as much as possible (Kitaigorodsky, 1973). We have recently shown that there is a strong preference for π – π interactions over hydrogen bond formation in the cocrystals and solvates of APR, and that it can be exploited to form a variety of cocrystals with pharmaceutically acceptable coformers (Dudek *et al.*, 2018), but the reason standing behind this preference remains unclear. This paper aims at clarifying this preference and explaining the factors that influence such a marked violation of Kitaigorodsky’s rule. To that purpose we account for three possibilities: (i) exceptionally favorable energy of aromatic – aromatic interactions in APR cocrystals and/or saturation of possible hydrogen bonding sites by CH∙∙∙O short contacts, (ii) an existence of a yet unknown stable crystal structure of APR cocrystals held together by hydrogen bonds, and (iii) a stabilizing role of disorder present in almost every known isostructural solvate and cocrystal of APR. As coformers, four different aromatic compounds were selected: resorcinol, picolinamide, imidazole, and hydroquinone, all being capable of acting as hydrogen bond donors, and all exhibiting a preference towards hydrogen bonds formation.

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**Figure 1**. Structure of APR with numbering and hydrogen bond (HB) sites marked with blue rectangles (HB acceptors), and with a red rectangle (HB donor).

**Experimental Section**

***Materials***

APR ((S)-N-(2-[1-(3-ethoxy-4-methoxyphenyl)-2-methane sulfonylethyl]-1,3-dioxo-2,3-dihy-dro-1H-isoindol-4-yl) acetamide) was purchased from Accel Pharmtech company and was recrystallized from acetone (POCh) *via* very slow evaporation of solvent to obtain suitable crystals of form B, which was identified by the comparison of the PXRD pattern (see Supporting Information, Figure S2) with patent literature data. Resorcinol (RES, Sigma Aldrich), hydroquinone (HQU, Sigma Aldrich), picolinamide (PIC, TCI Chemicals) and imidazole (IMI, Merck) were used as is.

***Cocrystal formation***

Cocrystals of APR were synthesized using a mechanochemical approach, which for the purpose of obtaining suitable crystals for single crystal X-Ray diffraction studies, was followed by solution recrystallization. In both cases the identity of the resulting crystalline forms was evaluated with 13C CPMAS NMR experiments. In the mechanochemical approach 100 mg of APR was ground for 1h in a ball mill set to 25 Hz frequency with an appropriate amount of a coformer, so that the resulting molar ratio was equal to 1:0.5. In all cases 20 μl of water was added to the reaction mixture to give liquid assisted grinding (LAG) conditions. All grinding experiments resulted in appropriate cocrystals formation, which was examined *via* solid-state NMR measurements. In order to obtain suitable crystals for single crystal X-Ray measurements ethyl acetate (STANLAB) was used to recrystallize cocrystals obtained *via* mechanochemistry. Note that direct crystallization of pure APR and appropriate coformers from ethyl acetate ends up in the formation of ethyl acetate APR solvate. In the case of APR:IMI cocrystals, we noticed that during grinding and crystallization experiments, a new, unknown polymorphic form appeared in small quantities. The crystal structure determination of this new form is beyond the scope of this manuscript and will be evaluated in our future work.

***Single crystal X-Ray measurements***

Single crystal diffraction experiments for form B of apremilast, as well as for the four cocrystals were carried out on Oxford SuperNova single-crystal diffractometer with micro source Cu Kα radiation (λ = 1.5418 Å) with a Titan detector. Diffraction data collection, cell refinement and data reduction were performed using the *CrysAlis PRO* program (Oxford Diffraction). The structure was solved by direct methods using SHELXS implemented in *OLEX2* package (Dolomanov *et al.*, 2009) and refined with SHELXL using full-matrix least-squares methods (Sheldrick, 2015). The crystal structures were deposited at the CSD under the deposition numbers 1916487, 1916491, 1916488, 1916489 and 1916490 for apremilast form B, APR:RES, APR:PIC, APR:IMI and APR:HQU, respectively. The hydrogen atoms attached to carbon and nitrogen atoms were set geometrically and refined as riding with thermal parameters equal 1.2 or 1.5 of thermal parameter of parent atom, depending on the type of geometrical restrains. Positions of hydrogen atoms connected to oxygen were localized on difference Fourier map and refined as riding.

***Crystal structure analysis***

Whenever hydrogen bonds are mentioned throughout the manuscript, they refer to O/N-H∙∙∙O/N interactions. All N/O∙∙∙N/O distances between intermolecular hydrogen bonded sites are given in Table S7. Other CH∙∙∙O/N interactions, in which the distance between interacting nuclei is smaller than the respective sum of their van der Waals radii are referred to as short-contacts. The distance between interacting aromatic rings in each aromatic – aromatic interaction in the studied crystals are presented in Table S7 as RCLOSEST value, which is the lowest C∙∙∙C distance found between interacting aromatic rings (see Figure S6). In addition in the case of displaced parallel aromatic – aromatic interactions, each is characterized by a value of γ angle, corresponding to the observed displacement between the interacting rings (see Figure S6 for graphic representation of γ angle).

***Solid-state NMR experiments***

13C and 15N CPMAS experiments have been performed with a Bruker Avance II 400 spectrometer equipped with a 4 mm probe-head, operating at 400.13, 100.90 and 40.56 MHz frequencies for 1H, 13C and 15N, respectively. Samples were spun at 8 kHz and a standard Bruker CP pulse programs with 2ms contact time and spinal decoupling during acquisition were used.

1H-13C inverse detected HETCOR and 1H-1H Back-to-Back correlation experiments were carried out on a Bruker Avance III 600 spectrometer equipped with 1.3 mm 1H/13C/15N triple resonance CP-MAS probe-heads with the 1H, 13C resonance frequencies of 600.13 and 150.90 MHz respectively. In all proton detected experiments samples were spun at 60 kHz.

Inverse detected HETCOR experiment have been carried out employing sequence described by Mao *et al.* (2009) and Althaus *et al.* (2014), using a RF equal to 100 kHz on 13C channel during contact time. For 1H linear ramp from 90 to 100% with maximum RF value near to 160 kHz have been used. Actual RF value of 1H RF during first and second contact have been precisely optimized on individual samples. Unwanted 1H magnetization from 12C isotopologues have been canceled out by applying two long (20 ms) pulses with phase X and Y and RF equal to 30kHz (HORROR condition) (Nieslen *et al.*, 1994). Low power 1H decoupling during t1 evolution and low power 13C decoupling during acquisition have been applied with an RF equal in both cases to 10 kHz applying swftpp decoupling sequence. For initial setup U-13C,15N-*l*-histidine hydrochloride monohydrate have been used. First contact time had always duration equal to 2 ms. Two experiments have been carried for individual samples with second contact time equal to 100 μs and 1ms. In the 1H-1H Back-to-Back experiments DQ excitation as well as reconversion time was 33.3 μs, while t1 increment was equal to 16.67 μs. 13C chemical shifts have been referenced according to secondary standard adamantane. 1H chemical shifts have been referenced according to solid sample of d4-TSP (methyl signal at 0 ppm).

***Powder X-Ray diffraction experiments***

Powder diffraction experiments were performed using a borosilicate glass capillary (0.5 mm diameter) and a Bruker D8 Discover powder diffractometer equipped with a sealed copper tube (Cu Kα radiation, λ = 1.5406 Å, formed by Goebel mirror monochromator). The registered data range was 5–50° 2θ, with a step of 0.05°.

***Quantum chemical calculations***

**CASTEP calculations.** For the assignment of the NMR parameters of form B of APR electronic structure calculations under periodic boundary conditions were performed for the obtained crystal structure with the CASTEP code (Clark *et al.*, 2005), using the PBE functional with D2 dispersion correction scheme (Grimme, 2006), ultrasoft pseudopotentials, plane wave energy cut-off of 600eV, and 0.07Å-1 separation to sample k-points in a Brillouin zone. All these parameters were primarily tested for convergence. In geometry optimization all atomic positions were allowed to relax, while keeping cell parameters rigid. After successful optimization NMR parameters were calculated, with a GIPAW approach (Yates *et al.*, 2007) at the same level of theory as stated above. The theoretical shieldings were subsequently used together with the experimental NMR data (2D correlations) to assign all the 1H, 13C and 15N resonances. For an assignment of 2D NMR spectra aided by quantum chemical calculations see Dudenko *et al.*, 2013, Abraham *et al.*, 2016, Czernek & Brus, 2013, Czernek *et al.*, 2018 and Webber *et al.*, 2010.

To evaluate the energy of cocrystal formation, geometry optimization at the same level of theory was performed for the crystal structures of the studied coformers and for the obtained cocrystals. In the case of the cocrystals due to the observed disorder there are two equally probable positions of coformers (designated as ‘a’ and ‘b’). Therefore, for each cocrystal two structures (‘a’ and ‘b’) were geometry optimized, each with one of the possible positions of a coformer, and mean energy was used in all the following calculations. Subsequently, single point energy calculations were carried out for each symmetry unrelated molecule of APR from form B, as well as from cocrystals, and for each of the coformers placed in a 20Å box, in order to calculate inter- and intramolecular energy contributions. To calculate the stabilization energies for the studied cocrystals (Estab), the following equations were used:

(1)

(2)

(3)

where denote total energies for a cocrystal, pure form B of APR and pure coformer, respectively, while and are the intra- and intermolecular energy contribution terms to the stabilization energy, calculated as differences between these terms obtained for cocrystals and pure forms, and *n*, *m* and *k* values denote number of molecules accounted for in a crystal, reflecting the ratio of particular components.

**Gaussian calculations.** Molecular electrostatic potential (MEP) surfaces were evaluated at the B3LYP-GD3BJ/6-311G(d,p) level of theory with Gaussian09 code (Frisch *et al.*, 2009), using molecular geometries from the optimized crystal structures. To evaluate the energetics of aromatic – aromatic and hydrogen bonds interactions, single point energy calculations for the optimized crystal geometries of appropriate molecular clusters, and corresponding isolated components were performed at the B3LYP-GD3BJ/6-311G(d,p) level of theory with the Gaussian09 code. The clusters were built in a way so that the molecules were interacting only *via* one of the regarded interactions. For aromatic – aromatic interactions two molecules of APR, forming a π–philic space with and without the presence of a coformer were used. Similarly, hydrogen-bonding interaction clusters were built. All the clustered arrangements are shown in Supporting Information (Figures S4 and S5). In that way an energy contribution of aromatic – aromatic and hydrogen bonds interactions to the total energy could be estimated separately, as most of the other intermolecular interactions are excluded from the system with such arrangements. Finally, the energy differences between the systems with and without the appropriate interactions were calculated in a way so that they would correspond to an asymmetric part of a unit cell in the studied cocrystals.

**Crystal Structure Prediction (CSP).** For CSP calculations, the optimized geometry of APR from the crystal structure of a cocrystal was used to generate trial crystal structures in three chiral space groups: *P*21, *P*212121 and *P*41, using the Global Lattice Energy Explorer code (Case *et al.*, 2016). The intramolecular energy contribution to the total energy of the trial structures were optimized using DMACRYS 2.2.1.0 (Price *et al.*, 2010), keeping molecular geometries rigid, and using atom-centred distributed multipoles up to hexadecapoles, calculated from the charge density obtained from Gaussian calculations using GDMA 2.2.11 (Stone, 2005). For repulsion-dispersion interactions, the FIT potential (Coombes *et al.*, 1996) with a 25Å cut-off was used. In each of the space groups the calculations were continued until 50000 successfully geometry optimized structures have been found. Subsequently, the structures were clustered on the basis of their PXRD pattern, energy and density similarities to remove duplicates.

***Solubility measurements***

The concentration of APR for solubility measurements of APR:PIC cocrystals and physical mixture was determined by ACQUITY UPLC I-Class chromatography system coupled with SYNAPT G2-Si mass spectrometer equipped with an electrospray source and quadrupole-Time-of-Flight mass analyser (Waters Corp., Milford, MA, USA). Acquity BEHTM C18 column (100 × 2.1 mm, 1.7 μm) maintained at 40°C temperature was used for the chromatographic separation of analyte. A gradient program was employed with the mobile phase, combining solvent A (0.1% formic acid in water) and solvent B (acetonitrile) as follows: 20% B (0–0.1 min), 20–95% B (0.1–2.3 min), 95–95% B (2.3–3.0 min), 95–20% B (3.0–3.1 min) and 20–20% B (3.1–4. min). The flow rate was 0.40 mL/min, and the injection volume was 1 μL.

For mass spectrometric detection the electrospray source was operated in a negative resolution mode. The optimized source parameters were: capillary voltage 2.5 kV, cone voltage 20 V, desolvation gas flow 600 L/h with temperature 350 °C, nebulizer gas pressure 6.5 bar, and source temperature 110 °C. Mass spectra were recorded over an *m/z* range of 100 to 1200. The system was controlled using MassLynx software (Version 4.1), while data processing (peak area integration, calibration curve) was performed by TargetLynxTM program.

The initial stock calibration solution of APR was created with a concentration of approximately 10 mg/mL of APR in acetonitrile. The stock solution was serial diluted (with acetonitrile) to obtain working solutions at several concentration levels. The calibration curves were prepared at six different concentrations of APR solutions and were linear over a concentration range from 10 ng/mL to 2000 ng/mL with a correlation coefficient of >0.987.

The samples for solubility measurement were prepared by dissolving an excess amount of each solid (APR:PIC cocrystals and a physical mixture in 1:0.5 ratio) in chloroform. The obtained suspensions were stirred at 25 °C for 24 hours. After filtrations through a 0.22 μm PTFE syringe filter the filtrates were diluted with acetonitrile to obtain appropriate concentration of solution (in the range of UPLC-MS calibration). The determined concentrations of APR were reported as the average of two replicated experiments for each sample of APR:PIC cocrystals and physical mixture and three calibration curves.

**Results and Discussion.**

*1) Structural characterization of pure APR and pure coformers.*

A first step to the understanding of cocrystals formation is the analysis of pure crystalline forms of both components. However, in the case of APR no pure crystal structure is available in the CSD database, but the patent literature indicates that APR can form at least 5 pure crystalline structures (patents no. US9850205B2, 2017;EP3339292A1, 2018; WO2009/120167A1, 2009; WO2017/196192A1, 2017). Due to the fact, that in this work commercially available form B of APR has been used in the co-crystallization experiments, and that it is regarded as the most thermodynamically stable form, we decided to put an extra effort to crystallize and solve its crystal structure.

Apremilast was crystallized from acetone by slow solvent evaporation to form needle-like (rod-like) crystals of monoclinic system with space group *P*21. The asymmetric unit of this crystal form contains four apremilast molecules which are assembled in two dimers. The main interactions responsible for this dimer creation are π stacking interactions of the substituted phenyl ring, with both dimers having similar architecture. Interestingly, in the solid-state NMR spectra of this form, only two to three sets of resonances are clearly distinguishable for most sites, with many 1H, 13C and 15N nuclei from corresponding sites in not symmetry-related molecules resonating at exactly the same frequencies (Figure 2). This is due to the presence of the previously mentioned dimers, each having very similar conformers of APR, with the molecular overlay RMSD for each pair being equal to 0.353 and 0.315 Å, respectively. These two dimers interact in a crystal mainly within themselves, forming layers held together *via* aromatic – aromatic interactions, while the interlayer interactions are mainly those between CH3-C=O and CH3-CH3 groups (Figure 2). Such close resemblance of the four symmetry-unrelated molecules results in poor resolution of the 1H MAS NMR spectrum with broad signals due to their mutual overlap, as well as in serious difficulties in the 1H and 13C resonances assignments to respective molecules, despite registering the spectra under very fast MAS conditions (that is with the spinning speed of 60 kHz). Here, to do the exact assignment, NMR calculations under periodic boundary conditions were necessary. The assigned resonances, together with 1H and 2D spectra are given in Supporting Information, while all crystallographic data are shown in Tables 1 and S1.

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**Figure 2.** Structural data obtained for form B of APR: crystal packing (a) and molecular overlays of two pairs of symmetry unrelated molecules (b) observed in the crystal, 13C and 15N solid-state NMR spectra (c and d, respectively). Colors of APR molecules denote symmetry-unrelated molecules.

**Table 1.** Crystallographic data for form B of APR, as well as for APR:RES, APR:PIC, APR:IMI and APR:HQU cocrystals.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| Compound | **APR:RES** | **APR:PIC** | **APR:IMI** | **APR:HQU** | **APR form B** |
| Empirical formula | C22H24N2O7S, 0.5(C6H6O2), 0.5(H2O) | C22H24N2O7S, 0.5(C6H6N2O) | C22H24N2O7S, 0.5(C3H4N2), 0.5(H2O) | C22H24N2O7S, 0.5(C6H6O2), 0.5(H2O) | C22H24N2O7S |
| Formula weight | 524.55 | 521.55 | 501.52 | 524.55 | 460.49 |
| Temperature | 100 K | 100 K | 100 K | 100 K | 293 K |
| Crystal system | tetragonal | tetragonal | tetragonal | tetragonal | monoclinic |
| Space group | *P*41212 | *P*41212 | *P*41212 | *P*41212 | *P*21 |
| a (Å) | 12.89179(7) | 12.81222(8) | 12.85808(14) | 12.88747(7) | 8.71383(13) |
| b (Å) | 12.89179(7) | 12.81222(8) | 12.85808(14) | 12.88747(7) | 29.4960(6) |
| c (Å) | 29.5087(3) | 29.7274(3) | 29.5715(4) | 29.5246(3) | 17.6717(2) |
|  (°) | 90 | 90 | 90 | 90 | 90 |
|  (°) | 90 | 90 | 90 | 90 | 97.9370(14) |
| (°) | 90 | 90 | 90 | 90 | 90 |
| Volume (Å)3 | 4904.29 | 4879.84 | 4889.07 | 4903.65 | 4498.36 |
| Density (g/cm3) | 1.421 | 1.420 | 1.363 | 1.421 | 1.360 |
| Z | 8 | 8 | 8 | 8 | 8 |
| R value\* | 0.047 | 0.058 | 0.063 | 0.060 | 0.045 |

\*listed R-factors are for data with F2 >2σ(F2)

In the case of coformers used in this study, all of them already have known crystal structures. Resorcinol and hydroquinone are closely related positional isomers, but they exhibit somewhat different crystallization preferences. For resorcinol there are three known crystal forms, *α*, *β*, and *ε* (with RESORA CSD refcode) (Bacon & Jude, 1973; Bacon & Lisher, 1980; Zhu *et al.*, 2016), two having *P*na21 and one *P*212121 space group symmetry. Similarly, hydroquinone crystallizes in one of the three known crystal forms: *α*, *β*, and *γ* (all having refcodes starting with HYQUIN) (Maartmann-Moe, 1966; Lindeman *et al*., 1981), with *P*21/c, *P*3 and *R*-3 space group symmetry, respectively. In each of these crystal forms both hydroxyl groups are engaged each in two hydrogen bonds: one as a donor and one as an acceptor of the bond. Also in picolinamide (refcode PICAMD) (Evora *et al*., 2012), as well as in imidazole crystals (refcode IMAZOL) (Craven *et al*., 1977), hydrogen bonds govern the crystal packing of these molecules (Figure 3). Our choice of coformers for the preparation of APR cocrystals was based on their tendency to form both hydrogen bonds and aromatic – aromatic interactions. However, as can be seen from the data presented above and in Figure 3, in all crystals of the selected coformers aromatic – aromatic interactions are either not observed, or arranged in such a way as to not distort the hydrogen bonding pattern. Contrarily, in APR molecules mainly aromatic – aromatic interactions govern the observed packing.

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**Figure 3.** Hydrogen bonding motifs in neat crystals of coformers used in this study: α-resorcinol (a), picolinamide (b), α-hydroquinone (c), and imidazole (d).

*2) Structural characterization of APR cocrystals.*

Despite strong propensity of coformers to form hydrogen bonds, definitely outweighing their tendency to interact *via* aromatic – aromatic interactions, as indicated by the analysis of their pure crystal structures, all four cocrystals formed by them with APR are isostructural and held together mostly by the aromatic – aromatic interactions. This can be concluded from a very similar outlook of their 13C CPMAS NMR spectra, very characteristic for systems with specific sandwich-like arrangement of molecules of APR and coformers (Figures S3 and S5, Supporting Information). In all the spectra the 13C signals assignable to coformers carbon atoms can be distinguished, with ca. 50% smaller signals intensities, as compared to those originating from APR, which reflects 2:1 APR:coformer ratio found in all APR solvates and cocrystals published so far. On the other hand, their 1H MAS spectra, as well as 1H-1H Back-to-Back (BaBa) correlation spectra are different to a certain extent (Figure 4), reflecting possible differences in the intermolecular interactions and close contacts of the 1H atoms.

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**Figure 4.** 1H VF MAS NMR spectra recorded with spinning rate 60 kHz for the cocrystals of APR with different coformers: 1H MAS NMR spectra (a), and SQ-DQ BaBa 2D NMR spectra (b) registered at 60 kHz spinning speed. Colors used for spectra plotting denote particular coformers.

A closer look at the differences in 1H chemical shifts between the studied cocrystals (Table 2) reveals that the most prominent changes concern H2 and H3 atoms (adjacent to C2 and C3 atoms, respectively, see Figure 1 for atom labeling scheme), with noticeably lower values found for a cocrystal with PIC, as well as NH hydrogen atoms from amide group of APR, which are similar for cocrystals with RES and HQU, but not so for the other cocrystals. As it is not likely for the intramolecular hydrogen bond of this NH proton with CO group to be broken, these latter differences may reflect a proximity of different functional groups originating from different coformers. On the other hand, the observed differences for H2 and H3 chemical shifts indicate somewhat similar interactions (or proximities) of APR molecule with particular coformers in IMI, RES and HQU cocrystals, distinct from those present in APR:PIC cocrystal. Not so prominent, but possibly important difference in the 1H chemical shifts concerns hydrogen atoms from methylsulfonyl group, namely methyl H13. For APR:HQU cocrystal its chemical shift is lower than for all the other cocrystals. The difference is small, but in all isostructural solvates and cocrystals studied so far this particular chemical shift value is well preserved and almost always equal to ca. 3.20 ppm. Its lower value in APR:HQU cocrystal may suggest that one of the sulfonyl group oxygen atoms is engaged in a hydrogen bond with a coformer.

**Table 2.** 1H chemical shifts of four isostructural cocrystals of APR and the studied coformers.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| atom | APR:RES | APR:HQU | APR:IMI | APR:PIC |
| 2  3  4  10  11  12  13  15  18  19  20  21  22  NH | 8.51  7.83  7.76  1.62  4.92  3.76 + 6.00  3.21  8.14  7.32  6.63  3.45 (2H)  0.91  4.03  7.43 | 8.43  7.63  7.73  1.62  4.78  3.73 + 5.83  3.13  8.15  7.18  6.49  3.43 (2H)  0.79  3.91  7.54 | 8.79  7.84  7.90  1.72  4.90  3.92 + 5.86  3.20  8.05  7.22  6.46  3.34 (2H)  0.56  3.97  8.15 | 8.08  7.32  7.73  1.82  4.82  3.85 + 5.88  3.22  8.11  7.31  6.63  3.47 (2H)  0.84  4.05  7.79 |

To follow up the conclusions drawn from solid-state NMR measurements, single crystal X-Ray diffraction experiments for the four cocrystals were performed (for crystallographic data see Table 1). Indeed, all the obtained crystals were found to be in the shape of tetragonal bipyramids, crystallizing isostructurally in the *P*41212 tetragonal space group, with aromatic – aromatic interactions playing the main role in holding together components in the crystal lattice. However, as was suggested by solid-state NMR results, some differences between these cocrystals in terms of intermolecular interactions have been found.

A first characteristic and distinctive feature is the presence of water molecule in three out of four cocrystals: APR:IMI, APR:RES and APR:HQU. In all cases water is located at the two fold axis and is strongly bonded with a coformer *via* a hydrogen bond, and as a consequence is located in the π-philic space formed by APR molecules, in a close proximity to H2 and H3 atoms of APR. Its presence can readily explain the higher values of 1H chemical shifts observed for H2 and H3 atoms of these cocrystals in comparison to that of APR:PIC, with no water molecule in its structure.

As to the formation of the hydrogen bonds engaging APR molecule itself, there are only two types of them observed in the obtained structures. The first one has been found in the APR:HQU structure, and is located between one of the oxygen atoms from APR methylsulfonyl moiety and a HQU hydroxyl proton, not engaged in the hydrogen bond with water (Figure 5a). Again, this is in agreement with the suggestion from the solid-state NMR results, according to which a small, but unusual change in the 1H chemical shift of the CH3 group from methylsulfonyl moiety indicated some differences in the local environment of this group. The second hydrogen bond motif is present in the three cocrystals containing water molecule, *i.e.* APR:RES, APR:HQU and APR:IMI. It is a bifurcated bond between water and an oxygen atom from either ethoxy or methoxy group (Figure 5b), although in APR:IMI cocrystal the O∙∙∙O distance between the interacting nuclei is rather long and equal to 3.96 Å, suggesting this interaction is a weak one (for all hydrogen bonds lengths see Table S7, Supporting Information). There are no intermolecular hydrogen bonds observed in the APR:PIC structure. It is clear, that even coformers with significant hydrogen bond formation propensity did not break the tendency of APR to form an arrangement in which aromatic – aromatic interactions are promoted, even though it left some of the coformer sites capable of acting as hydrogen bond donors unbounded. On the other hand, it should be noted that many possible hydrogen bonding sites are engaged in CH∙∙∙O short contacts, which additionally stabilize the studied structures. As a result only OMe and OEt groups of APR as well as NH group of PIC in APR:PIC cocrystal remain unbounded and without any significant short contacts.

Apart from saturating one of the hydrogen bonding sites in the three coformers, interaction with water also changes the molecular electrostatic potential (MEP) surfaces of the coformers, thus creating a possibility of stronger aromatic – aromatic interaction. Figure 6 and 7 show such surfaces mapped onto electron density isosurfaces for the APR (Figure 6), and coformers with and without the presence of water molecule (if it is present in the respective crystal structures) (Figure 7). From the analysis of the APR MEP surface, it can be said that the isoindoline-1,3-dione ring of APR, which is engaged in the aromatic – aromatic interactions in APR cocrystals with aromatic species, can be regarded as an electron deficient aromatic ring. Indeed, it has only one site with a small negative electrostatic potential value (meaning a slightly repulsive reaction towards the negatively charged species), while all the other sites of this ring exhibit positive electrostatic potential of up to +126 kJ/mol, indicating a strong attraction propensity towards the negative charge.

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**Figure 5.** Two hydrogen bond motifs observed in the cocrystals of APR and the studied coformers present in: (a) APR:HQU cocrystal (Ar-OH∙∙∙O2S-CH3); (b) APR:RES, APR:HQU and APR:IMI cocrystals (HO-H∙∙∙OMe/OEt)

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**Figure 6.** APR molecule viewed from the site engaged in the aromatic – aromatic interactions in the obtained cocrystals, and its molecular electrostatic potential mapped on the total electron density isosurface (0.005 a.u.). All electrostatic potentials are given in kJ/mol, MEP surfaces were calculated at the B3LYP-GD3BJ/6-311G(d,p) level of theory.

In the case of coformer molecules, RES, IMI and HQU are all electron-rich aromatic systems, which, upon hydrogen bonding to a water molecule, become even more enriched with a negative charge, leading consequently to a stronger attraction to the electron-deficient sites of isoindoline-1,3-dione ring of APR. Such cooperativity of hydrogen bonding and aromatic – aromatic interactions has been observed and quantified before (Ebrahimi *et al.*, 2009). The only coformer molecule for which there is no water present in the cocrystal of APR is PIC. However, even without the presence of water the electrostatic potential is already sufficiently negative to be inclined to interact with the electron-deficient aromatic systems.

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**Figure 7.** Molecular electrostatic potential surfaces mapped on the total electron density isosurfaces (0.01 a.u.) for coformers used in this study to form cocrystals with APR without (column B) and with the presence of water (column C). Column A indicates conformations of the coformers, which in all cases are derived from the respective cocrystals. All electrostatic potentials are given in kJ/mol, MEP surfaces were calculated at the B3LYP-GD3BJ/6-311G(d,p) level of theory.

*3) Evaluation of crystal energy landscape in the search for hydrogen bonded cocrystals of APR:PIC*

Among the studied cocrystals, APR:PIC is the only system for which no intermolecular hydrogen bonds have been found, despite the ability of both components to form such bonds. Therefore, crystal structure prediction (CSP) calculations were used to gain an insight into whether alternative crystal packing motifs may be energetically plausible, apart for the one found experimentally. Due to the large size of the system, the search was carried out for only three space groups *P*21, *P*212121 (which are the two most common chiral space groups describing 75% of all enantiomeric organic molecules) and *P*41. The *P*212121 and *P*41 space groups are subgroups of *P*41212 space group, in which the experimental crystal structure of APR:PIC was found. Figure 8 shows the resulting CSP landscape, and features one particularly stable crystal structure, which was found to be in an excellent agreement with the experimental structure; an overlay of a 15-molecule cluster from predicted and observed crystal structures leads to a root mean squared deviation in atomic positions of only 0.118 Å. The CSP search that has been performed cannot be regarded as a having sampled all possible co-crystal arrangements, due to limiting the search to one conformation of APR and the most common space groups, and therefore the results have to be treated cautiously. Nevertheless, the magnitude of the energy gap between the best and the next-best structure (76 kJ/mol) suggests that a possibility of the formation of other competitive structures is rather low.

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**Figure 8.** Crystal energy landscape for APR:PIC cocrystal tested in *P*21, *P*212121 and *P*41 space groups. An arrow indicates the lowest energy structure with an intermolecular hydrogen bond between APR and PIC.

An interesting feature of the predicted structures is that in *P*212121 and *P*41 space groups aromatic – aromatic interactions are preferred over hydrogen bonds. The lowest energy predicted structure in which APR and PIC interact *via* hydrogen bonds is in the *P*21 space group with an energy of -445 kJ/mol, 80 kJ/mol above the lowest energy structure, followed by a structure in space group *P*212121, with an energy of -440 kJ/mol. Here the hydrogen bond is formed between the NH2 group from PIC and oxygen atom of the OMe and OEt groups from APR (Figure 8). Such an interaction however, results in losing the possibility of exploiting not only aromatic – aromatic interactions, but also many Ar-H…O and CH3…O close contacts, maintaining at the same time favorable packing efficiency.

*4)* *Evaluation of the energetic contributions to the total energy of APR cocrystals*

The X-Ray, solid-state NMR and CSP results presented above have shown that aromatic – aromatic interactions are indeed a driving force in the formation of APR cocrystals, even in cases in which a coformer displays high propensity towards acting as a hydrogen bond donor. The next step is therefore to evaluate the energetic factors behind these interactions, starting with the calculations of stabilization energies of the cocrystals, *i.e.* the differences between the total energy of a cocrystal and the sum of total energies of its components in their pure forms. In a recent study on the energetics of cocrystals, it has been shown that for the vast majority (95%) of observed cocrystals, the stabilization energies have negative values, which means that cocrystal formation is thermodynamically driven (Taylor & Day, 2018). Surprisingly, this seems to be not the case for the cocrystals of APR. Table 3 presents the energy difference between the energy of a cocrystal and the sum of the energies of its pure components calculated at the PBE-D2 level of theory. In all cases the stabilization energies (calculated according to eq. 1) have positive values, and this is primarily the result of noticeably higher intramolecular energies of APR molecules in the cocrystals, as compared to these energies in the pure form of APR. On the other hand, the differences in the intermolecular contributions to the total energy are in all cases negative, meaning that it is not the lack of energetically favorable interactions between APR and the coformers that makes the stabilization energies positive, but the strains in the conformation of APR associated with cocrystal formation. Regardless of the cause, it seems at this stage, that the formation of cocrystals of APR with the studied coformers is not energetically favorable. It should be noted, that the particular components of the stabilization energy presented in Table 3, *i.e.* and were calculated separately from . As a result the sum of the mentioned terms agree with stabilization energy value only within the limits of computational error, which in the worst case is equal to 0.35 kJ/mol. This error may also originate from neglecting the intramolecular contribution from coformes, but still it does not influence the overall conclusions.

**Table 3.** Overall stabilization energies () of the formation of cocrystals of APR with RES, HQU, IMI and PIC, and the intra- and intermolecular energy contribution to .\*

|  |  |  |  |
| --- | --- | --- | --- |
| cocrystal | [kJ/mol] | [kJ/mol] | [kJ/mol] |
| APR:RES  APR:HQU  APR:IMI  APR:PIC | +3.14  +17.00  +16.79  +6.15 | -13.37  -1.72  -0.66  -9.54 | +16.51  +18.37  +17.44  +15.73 |

\*Note, that and values were calculated separately, according to equations (1-3), and as a result the two latter values add up to only within the limits of computational errors. Note also, that intramolecular contributions from coformers were neglected.

To gain a deeper understanding of the intermolecular energy contribution of the APR cocrystals, the energetics of aromatic – aromatic and HB interactions were evaluated (Table 4). As can be seen, the energetic contributions of aromatic – aromatic interactions are equal to 50 – 60 kJ/mol (per crystallographic asymmetric part of the unit cell of a cocrystal), which is comparable to the stabilization energy typically found for strong hydrogen bonds. It is noteworthy that these interaction energies are slightly stronger for the APR:RES and APR:HQU systems, which reflects their lower electrostatic potentials found on their MEP surfaces, in comparison to the values found for APR:IMI and APR:PIC systems. The energy contributions from the hydrogen bonds formed between APR and coformers, excluding the interactions ‘inside’ the coformer (*i.e.* between water molecule and an aromatic entity of a coformer), are half as large (as calculated per crystallographic asymmetric part of the unit cell), but become comparable to the Ar – Ar contributions after including HB interactions within the coformers. It should be noted, however, that these two calculated contributions to the overall intermolecular energy constitute in all cases only one third of the total lattice energy. Therefore, apart from these obvious intermolecular interactions, the sum of the energy contributions from weaker interactions, such as CH3- or Ar-H with SO2, C=O or OMe/OEt oxygen atoms, has to play an important role in the stabilization of the cocrystals. Obviously, such a sum would be very difficult to calculate, but it can be estimated by subtracting the sum of the energy contribution of the aromatic – aromatic and hydrogen bonds interactions from the overall intermolecular energies of the cocrystals. The obtained values, denoted as Eother, are included in Table 4. The obtain values indicate that together CH∙∙∙O short contacts give the highest energy contribution to the lattice energy of the studied cocrystals.

**Table 4.** Energy contributions of intermolecular interactions found in APR cocrystals: aromatic – aromatic interactions (Ar-Ar), hydrogen bonds between APR and a coformer (HB\_APR), hydrogen bonds “inside” a coformer, *i.e.* between water and an aromatic entity (HB\_coformer). Einter denotes total intermolecular energy for each of the cocrystals, whereas Eother is the difference between Einter and the sum of EAr-Ar, EHB\_APR and EHB\_coformer. All energy values are given in kJ/mol.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| Cocrystal | EAr-Ar | EHB\_APR | EHB\_coformer | Einter | Eother |
| APR:RES  APR:HQU  APR:IMI  APR:PIC | -62  -58  -48  -49 | -28  -39  -27  - | -23  -21  -19  - | -324  -314  -300  -278 | -210  -197  -206  -229 |

*5) Stabilizing role of disorder?*

Though the analysis of structure and energies of the studied cocrystals performed so far indicates preference for aromatic – aromatic interactions over hydrogen bonds, it does not answer the most important question: why the cocrystals are formed at all, despite positive stabilization energies? An answer to this may be provided by an evaluation of the entropy contribution to the Gibbs free energy of the cocrystals. Usually, when comparing energetic stability of crystals, their free energies are approximated by their enthalpies, and the entropic contribution to the free energy, *ΔG = ΔH – TΔS*, is neglected. However, in some cases this entropy contribution may play an important role in the stabilization of a crystal structure, especially in the crystals in which disorder is observed.

As mentioned before, in almost all known APR cocrystals and solvates (except for dichloromethane solvate) (Wu *et al.*, 2017b), a coformer or a solvent molecule is disordered, even at 100 K. The nature of this disorder is twofold: first is the result of equal probability of occupying two different sites (sometimes referred to as positional or static disorder), and the second one originates from vibrational motion of coformer/solvent molecules, which can be visualized by the respective thermal ellipsoids (Figure 9). Both types of disorder may have an impact on the entropy contribution to the total energy of the studied systems.

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**Figure 9.** Thermal ellipsoids in the studied cocrystals, drawn at 50% probability level.

In order to estimate this entropy contribution one can directly determine Gibbs free energy of formation of a cocrystal by measuring its solubility and comparing it with the solubility of physical mixture, according to the equation:

where R denotes the gas constant, T temperature of the measurement, while and are the concentration of apremilast from dissolution of a physical mixture and a cocrystal, respectively (Zhang *et al*., 2017; Ahuja *et al*., 2019). Since all enthalpies of formation of the studied cocrystals have positive values, negative Gibbs free energies indicate that the formation of cocrystals is indeed entropy driven. Such an experiment, however, can be easily performed only for the APR:PIC cocrystal, as in all other cases there is a water molecule present in the respective crystal structures, which is difficult to include in the solubility determination of a physical mixture, while possibly having significant influence on the experimental results. Therefore we decided to measure solubility only for the cocrystal and physical mixture of the APR:PIC system to gain at least an indication whether this above mentioned entropy term can indeed be responsible for the formation of APR cocrystals.

The obtained mean concentrations of APR dissolved from the physical mixture and the cocrystal were equal to 180.3 ± 6.3 and 171.2 ± 5.8 mg/ml for and , respectively, with the concentration for the physical mixture being always higher than for the cocrystal. This means that the free energy of cocrystal formation is negative and equal to –0.132 ± 0.002 kJ/mol (–0.29 J/g). As a result it may be concluded that the formation of APR:PIC cocrystals (and possibly also other APR cocrystals studied in this paper) is an entropy-driven process, with all probability associated with the observed disorder of the guest (coformer) molecules. Such an entropy-controlled formation of cocrystals has been already reported *e.g.* for celecoxib-nicotinamid cocrystals (Zhang *et al*., 2017), and the entropy gain upon formation of cocrystals has been associated with an increase in molar volume of a crystal (higher molar volume means more space, which in turn means more freedom of movement, leading to higher entropy) (Perlovich, 2018). Here, such an increase in volume is not observed. Contrarily, molar volume upon formation of the studied cocrystals decreases, so it seems that it is rather a lack of strong directional hydrogen bonds between host and guest molecules that enables guest molecules to be disordered.

**Conclusions**

Four new cocrystals of APR, formed with aromatic coformers prone to act as hydrogen bonds donors, were found to be isostructural to the known cocrystals and solvates of APR, and held together mainly by aromatic – aromatic interactions, despite strong propensity of the guest molecules to form hydrogen bonds. In addition, a significant contribution of CH∙∙∙O short contacts to the stabilization of the studied cocrystals have been observed. The calculated energy of these aromatic – aromatic interactions is comparable to the energy of hydrogen bonds. Interestingly, calculated stabilization energies of these cocrystals were found to be positive in all cases, which means that their formation is associated with a loss of enthalpy. This loss has been linked to the strains in the conformation of APR after cocrystal formation. At the same time, solubility measurements indicated that the observed loss in enthalpy is compensated by the gain in entropy, so that the overall Gibbs free energy of the cocrystal formation was found to be negative. Therefore, the stabilizing role of disorder, observed in almost all isostructural cocrystals and solvates of APR discovered so far, can be postulated.

**Acknowledgements**

This study was financially supported by the National Science Center, Poland (Grant No. UMO-2017/25/B/ST4/02684). The computational part of this work was carried out with a financial support of the Polish Ministry of Science and Higher Education, as a part of the fellowship ‘Mobility Plus’ (grant No 1664/MOB/V/2017/0). The Polish Infrastructure for Supporting Computational Science in the European Research Space (PL-GRID) and the IRIDIS High Performance Computing Facility, with associated support services at the University of Southampton are gratefully acknowledged for providing computational resources. We are also grateful to the UK Materials and Molecular Modelling Hub for computational resources, which is partially funded by EPSRC (EP/P020194/1).

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**Supporting Information**

to

**Understanding the formation of apremilast cocrystals**

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**Experimental data**

**Table S1.** Experimental details for crystallographic measurements of APR form B and APR:RES, APR:PIC, APR:IMI and APR:HQU cocrystals.

|  |  |
| --- | --- |
| APR FORM B cocrystal | |
| Crystal data | |
| Chemical formula | C22H24N2O7S |
| *M*r | 460.49 |
| Crystal system, space group | Monoclinic, *P*21 |
| Temperature (K) | 293 |
| *a*, *b*, *c* (Å) | 8.71353 (13), 29.4960 (6), 17.6717 (2) |
| b (°) | 97.9370 (14) |
| *V* (Å3) | 4498.36 (12) |
| *Z* | 8 |
| Radiation type | Cu *K*a |
| m (mm-1) | 1.68 |
| Crystal size (mm) | 0.31 × 0.06 × 0.02 |
|  | |
| Data collection | |
| Diffractometer | SuperNova, Dual, Cu at zero, TitanS2 |
| Absorption correction | Multi-scan  *CrysAlis PRO*, Agilent Technologies, Version 1.171.37.35h (release 09-02-2015 CrysAlis171 .NET) (compiled Feb 9 2015,16:26:32) Empirical absorption correction using spherical harmonics, implemented in SCALE3 ABSPACK scaling algorithm. |
| *T*min, *T*max | 0.817, 1.000 |
| No. of measured, independent and  observed [*I* > 2s(*I*)] reflections | 43359, 17606, 14660 |
| *R*int | 0.038 |
| (sin q/l)max (Å-1) | 0.628 |
|  | |
| Refinement | |
| *R*[*F*2 > 2s(*F*2)], *wR*(*F*2), *S* | 0.045, 0.119, 1.03 |
| No. of reflections | 17606 |
| No. of parameters | 1169 |
| No. of restraints | 1 |
| H-atom treatment | H-atom parameters constrained |
| Dρmax, Dρmin (e Å-3) | 0.17, -0.30 |
| Absolute structure | Flack x determined using 5723 quotients [(I+)-(I-)]/[(I+)+(I-)] (Parsons, Flack and Wagner, Acta Cryst. B69 (2013) 249-259). |
| Absolute structure parameter | 0.004 (8) |
| APR:RES cocrystal | |
| Crystal data | |
| Chemical formula | C22H24N2O7S·0.5(C6H6O2)·0.5(H2O) |
| *M*r | 524.55 |
| Crystal system, space group | Tetragonal, *P*41212 |
| Temperature (K) | 100 |
| *a*, *c* (Å) | 12.89179 (7), 29.5087 (3) |
| *V* (Å3) | 4904.29 (7) |
| *Z* | 8 |
| Radiation type | Cu *K*a |
| m (mm-1) | 1.66 |
| Crystal size (mm) | 0.48 × 0.30 × 0.16 |
|  | |
| Data collection | |
| Diffractometer | SuperNova, Dual, Cu at zero, TitanS2 |
| Absorption correction | Multi-scan  *CrysAlis PRO*, Agilent Technologies, Version 1.171.37.35h (release 09-02-2015 CrysAlis171 .NET) (compiled Feb 9 2015,16:26:32) Empirical absorption correction using spherical harmonics, implemented in SCALE3 ABSPACK scaling algorithm. |
| *T*min, *T*max | 0.250, 1.000 |
| No. of measured, independent and  observed [*I* > 2s(*I*)] reflections | 38451, 5001, 4965 |
| *R*int | 0.048 |
| (sin q/l)max (Å-1) | 0.626 |
|  | |
| Refinement | |
| *R*[*F*2 > 2s(*F*2)], *wR*(*F*2), *S* | 0.047, 0.132, 1.08 |
| No. of reflections | 5001 |
| No. of parameters | 360 |
| No. of restraints | 96 |
| H-atom treatment | H-atom parameters constrained |
| Dρmax, Dρmin (e Å-3) | 0.90, -0.63 |
| Absolute structure | Flack x determined using 1998 quotients [(I+)-(I-)]/[(I+)+(I-)] (Parsons, Flack and Wagner, Acta Cryst. B69 (2013) 249-259). |
| Absolute structure parameter | -0.001 (6) |
| APR:PIC cocrystal | |
| Crystal data | |
| Chemical formula | C22H24N2O7S·0.5(C6H6N2O) |
| *M*r | 521.55 |
| Crystal system, space group | Tetragonal, *P*41212 |
| Temperature (K) | 100 |
| *a*, *c* (Å) | 12.81222 (8), 29.7274 (3) |
| *V* (Å3) | 4879.84 (7) |
| *Z* | 8 |
| Radiation type | Cu *K*a |
| m (mm-1) | 1.65 |
| Crystal size (mm) | 0.41 × 0.28 × 0.25 |
|  | |
| Data collection | |
| Diffractometer | SuperNova, Dual, Cu at zero, TitanS2 |
| Absorption correction | Multi-scan  *CrysAlis PRO*, Agilent Technologies, Version 1.171.37.35h (release 09-02-2015 CrysAlis171 .NET) (compiled Feb 9 2015,16:26:32) Empirical absorption correction using spherical harmonics, implemented in SCALE3 ABSPACK scaling algorithm. |
| *T*min, *T*max | 0.746, 1.000 |
| No. of measured, independent and  observed [*I* > 2s(*I*)] reflections | 38165, 4980, 4953 |
| *R*int | 0.021 |
| (sin q/l)max (Å-1) | 0.627 |
|  | |
| Refinement | |
| *R*[*F*2 > 2s(*F*2)], *wR*(*F*2), *S* | 0.058, 0.160, 1.07 |
| No. of reflections | 4980 |
| No. of parameters | 363 |
| No. of restraints | 115 |
| H-atom treatment | H-atom parameters constrained |
| Dρmax, Dρmin (e Å-3) | 1.06, -0.63 |
| Absolute structure | Flack x determined using 2008 quotients [(I+)-(I-)]/[(I+)+(I-)] (Parsons, Flack and Wagner, Acta Cryst. B69 (2013) 249-259). |
| Absolute structure parameter | -0.002 (4) |
| APR:IMI cocrystal | |
| Crystal data | |
| Chemical formula | C22H24N2O7S·0.5(C3H6N2O) |
| *M*r | 501.52 |
| Crystal system, space group | Tetragonal, *P*41212 |
| Temperature (K) | 100 |
| *a*, *c* (Å) | 12.85808 (14), 29.5715 (4) |
| *V* (Å3) | 4889.07 (13) |
| *Z* | 8 |
| Radiation type | Cu *K*a |
| m (mm-1) | 1.62 |
| Crystal size (mm) | 0.47 × 0.24 × 0.12 |
|  | |
| Data collection | |
| Diffractometer | SuperNova, Dual, Cu at zero, TitanS2 |
| Absorption correction | Multi-scan  *CrysAlis PRO*, Agilent Technologies, Version 1.171.37.35h (release 09-02-2015 CrysAlis171 .NET) (compiled Feb 9 2015,16:26:32) Empirical absorption correction using spherical harmonics, implemented in SCALE3 ABSPACK scaling algorithm. |
| *T*min, *T*max | 0.576, 1.000 |
| No. of measured, independent and  observed [*I* > 2s(*I*)] reflections | 38007, 5005, 4888 |
| *R*int | 0.040 |
| (sin q/l)max (Å-1) | 0.627 |
|  | |
| Refinement | |
| *R*[*F*2 > 2s(*F*2)], *wR*(*F*2), *S* | 0.063, 0.179, 1.05 |
| No. of reflections | 5005 |
| No. of parameters | 335 |
| No. of restraints | 66 |
| H-atom treatment | H-atom parameters constrained |
| Dρmax, Dρmin (e Å-3) | 1.98, -0.96 |
| Absolute structure | Flack x determined using 1984 quotients [(I+)-(I-)]/[(I+)+(I-)] (Parsons, Flack and Wagner, Acta Cryst. B69 (2013) 249-259). |
| Absolute structure parameter | 0.006 (6) |
| APR:HQU cocrystal | |
| Crystal data | |
| Chemical formula | C22H24N2O7S·0.5(C6H8O3) |
| *M*r | 524.55 |
| Crystal system, space group | Tetragonal, *P*41212 |
| Temperature (K) | 100 |
| *a*, *c* (Å) | 12.88747 (7), 29.5246 (3) |
| *V* (Å3) | 4903.65 (7) |
| *Z* | 8 |
| Radiation type | Cu *K*a |
| m (mm-1) | 1.66 |
| Crystal size (mm) | 0.26 × 0.16 × 0.12 |
|  | |
| Data collection | |
| Diffractometer | SuperNova, Dual, Cu at zero, TitanS2 |
| Absorption correction | Multi-scan  *CrysAlis PRO*, Agilent Technologies, Version 1.171.37.35h (release 09-02-2015 CrysAlis171 .NET) (compiled Feb 9 2015,16:26:32) Empirical absorption correction using spherical harmonics, implemented in SCALE3 ABSPACK scaling algorithm. |
| *T*min, *T*max | 0.603, 1.000 |
| No. of measured, independent and  observed [*I* > 2s(*I*)] reflections | 38399, 5046, 5005 |
| *R*int | 0.027 |
| (sin q/l)max (Å-1) | 0.628 |
|  | |
| Refinement | |
| *R*[*F*2 > 2s(*F*2)], *wR*(*F*2), *S* | 0.060, 0.172, 1.06 |
| No. of reflections | 5046 |
| No. of parameters | 359 |
| No. of restraints | 114 |
| H-atom treatment | H-atom parameters constrained |
| Dρmax, Dρmin (e Å-3) | 1.23, -0.97 |
| Absolute structure | Flack x determined using 2039 quotients [(I+)-(I-)]/[(I+)+(I-)] (Parsons, Flack and Wagner, Acta Cryst. B69 (2013) 249-259). |
| Absolute structure parameter | -0.003 (4) |

In APR:IMI coccrystal relatively high residual density has been observed. This is the result of two effects: (i) a very strong hydrogen bond of imidazole nitrogen with water, which in consequence leads to a weak hydrogen contact of water with ethoxyl oxygen of APR, not listed in the checkcif contacts; (ii) relatively small imidazole ring could be modeled in more than two orientation but this interpretation did not result in better refinement statistics, so we decided to use similar disorder model as for all investigated cocrystals, which in turn left the observed residual density.

**Table S2.** 1H and 13C Signal assignment for APR form B.

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| atom | molecule A | | molecule B | | molecule C | | molecule D | |
| 1 | 135.7 |  | 135.7 |  | 136.3 |  | 134.6 |  |
| 2 | 120.1 | 6.5 | 120.7 | 6.9 | 123.1 | 6.7 | 120.1 | 6.5 |
| 3 | 137.4 | 7.1 | 137.4 | 7.1 | 137.4 | 7.1 | 135.7 | 6.9 |
| 4 | 119.0 | 6.8 | 119.4 | 6.6 | 119.4 | 6.6 | 119.4 | 6.6 |
| 5 | 130.0 |  | 129.2 |  | 130.0 |  | 130.0 |  |
| 6 | 169.2 |  | 167.2 |  | 167.2 |  | 167.2 |  |
| 7 | 167.2 |  | 169.2 |  | 169.2 |  | 167.2 |  |
| 8 | 113.7 |  | 114.8 |  | 113.7 |  | 115.2 |  |
| 9 | 167.2 |  | 166.0 |  | 166.0 |  | 166.0 |  |
| 10 | 21.6 | 0.3 | 24.1 | 0.5 | 21.6 | 0.3 | 25.1 | 0.8 |
| 11 | 47.9 | 5.0 | 48.3 | 5.3 | 46.8 | 5.4 | 47.9 | 5.0 |
| 12 | 52.8 | 4.8; 3.0 | 50.4 | 4.6; 3.6 | 49.4 | 4.6; 3.0 | 51.2 | 4.7; 3.0 |
| 13 | 39.8 | 2.4 | 40.1 | 2.8 | 37.8 | 2.6 | 41.6 | 2.8 |
| 14 | 131.9 |  | 130.0 |  | 130.0 |  | 131.9 |  |
| 15 | 109.0 | 5.7 | 107.9 | 5.9 | 106.8 | 5.6 | 110.6 | 5.6 |
| 16 | 147.7 |  | 146.8 |  | 146.8 |  | 147.7 |  |
| 17 | 146.8 |  | 147.7 |  | 146.8 |  | 146.8 |  |
| 18 | 112.1 | 7.4 | 110.7 | 6.9 | 110.7 | 6.9 | 111.9 | 7.4 |
| 19 | 117.7 | 6.3 | 123.1 | 6.7 | 122.4 | 6.7 | 119.0 | 6.8 |
| 20 | 64.5 | 3.7 | 63.6 | 2.9 | 63.2 | 3.7 | 63.6 | 3.7 |
| 21 | 15.5 | 1.0 | 14.0 | 1.0 | 14.3 | 1.0 | 14.3 | 1.0 |
| 22 | 54.5 | 3.5 | 54.7 | 3.8 | 55.4 | 3.5 | 55.4 | 3.8 |
| NH |  | 7.2 |  | 7.2 |  | 7.2 |  | 7.3 |

**Table S3.** Calculated 1H and 13C chemical shieldings for APR form B.

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| atom | molecule A | | molecule B | | molecule C | | molecule D | |
| 1 | 33,6 |  | 33,5 |  | 32,8 |  | 34,9 |  |
| 2 | 48,2 | 23,4 | 47,3 | 22,5 | 45,1 | 22,8 | 48,1 | 22,9 |
| 3 | 29,8 | 22,8 | 29,6 | 22,6 | 29,4 | 22,5 | 32,0 | 23,1 |
| 4 | 50,6 | 23,3 | 49,9 | 23,0 | 49,9 | 22,7 | 49,1 | 23,2 |
| 5 | 39,8 |  | 40,9 |  | 40,1 |  | 40,3 |  |
| 6 | -2,0 |  | 0,8 |  | 0,1 |  | -0,7 |  |
| 7 | -1,0 |  | -2,9 |  | -2,7 |  | -1,2 |  |
| 8 | 58,0 |  | 56,7 |  | 57,6 |  | 55,9 |  |
| 9 | 1,0 |  | 1,6 |  | 1,1 |  | 1,2 |  |
| 10 | 152,7 | 29,6 | 149,0 | 29,5 | 152,5 | 29,7 | 147,4 | 29,1 |
| 11 | 121,5 | 24,5 | 120,9 | 24,3 | 122,8 | 24,0 | 121,7 | 24,6 |
| 12 | 115,0 | 25,0; 26.8 | 118,4 | 25,4; 25.5 | 118,9 | 25,6; 26.3 | 116,7 | 25,0; 26.0 |
| 13 | 130,3 | 27,6 | 130,3 | 27,0 | 132,0 | 27,1 | 128,8 | 26,8 |
| 14 | 37,4 |  | 39,6 |  | 39,4 |  | 38,0 |  |
| 15 | 61,8 | 24,1 | 62,8 | 23,9 | 64,1 | 23,9 | 60,2 | 24,4 |
| 16 | 19,9 |  | 20,8 |  | 21,1 |  | 20,5 |  |
| 17 | 21,4 |  | 20,3 |  | 21,1 |  | 20,9 |  |
| 18 | 57,7 | 22,1 | 59,5 | 22,8 | 59,4 | 23,2 | 58,8 | 22,3 |
| 19 | 50,7 | 22,9 | 47,0 | 23,2 | 47,0 | 22,7 | 50,6 | 23,0 |
| 20 | 103,8 | 26,0 | 104,6 | 26,5 | 104,8 | 26,4 | 104,7 | 26,3 |
| 21 | 157,7 | 28,7 | 159,9 | 29,1 | 159,6 | 28,7 | 159,3 | 28,8 |
| 22 | 115,4 | 25,6 | 115,0 | 26,1 | 114,4 | 25,9 | 114,5 | 26,0 |
| NH |  | 21,9 |  | 21,7 |  | 21,8 |  | 21,6 |

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**Figure S1.** 1H and *inv*-1H-13C HETCOR solid-state NMR spectra registered under 60 kHz rotation for APR form B: a) 1H spectrum, b) HETCOR with short contact time (100 μs), c) HETCOR with long contact time (2 ms).

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**Figure S2**. Powder X-Ray diffraction pattern for form B

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**Figure S3.** 13C NMR spectra for the studied cocrystals.

**Data for calculation of energy contributions to the cocrystals formation (Gaussian calculations)**

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**Figure S4.** Clusters for the calculations of contribution of hydrogen bonds between APR and a coformer to the intermolecular energy of cocrystals: a) APR:HQU, b) APR:IMI, c) APR:RES.

**Table S4.** Data for the calculations of contribution of hydrogen bonds between APR and a coformer to the intermolecular energy of the studied cocrystals.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | ECLUSTER [Hartree] | EAPR [Hartree] | ECOFORMER [Hartree] | ECLUSTER – (EAPR + nECOFORMER) [Hartree] | ECLUSTER – (EAPR + nECOFORMER) [kJ/mol] | Final EHB\_APR per asymmetric unit [kJ/mol] |
| APR:HQU | -2806.15 | -1887.57 | -459.275 | -0.02951 | -77.4674 | -39 |
| APR:IMI | -2190.34 | -1887.57 | -302.748 | -0.02089 | -54.8486 | -27 |
| APR:RES | -2346.87 | -1887.57 | -459.28 | -0.02151 | -56.4792 | -28 |

**Table S5.** Data for the calculations of contribution of hydrogen bonds ‘inside’ a coformer to the intermolecular energy of the studied cocrystals.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | ECOF+H2O [Hartree] | ECOFORMER  [Hartree] | EH2O [Hartree] | ECOF+H2O – (EH2O + ECOFORMER) [Hartree] | ECOF+H2O – (EH2O + ECOFORMER) [kJ/mol] | Final EHB\_coformer per asymmetric unit [kJ/mol] |
| APR:HQU | -459.275 | -382.8123 | -76.4468 | -0.0158164 | -41.526 | -21 |
| APR:IMI | -302.7482 | -226.2863 | -76.4472 | -0.014673 | -38.524 | -19 |
| APR:RES | -459.2805 | -382.8156 | -76.447 | -0.017832 | -46.818 | -23 |

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**Figure S5.** Clusters for the calculations of contribution of aromatic – aromatic interactions between APR and a coformer to the intermolecular energy of cocrystals and corresponding ‘empty’ pockets formed by APR molecules: a) APR:RES, b) APR:PIC, c) APR:IMI, d) APR:HQU.

**Table S6.** Data for the calculations of contribution of aromatic – aromatic interactions between APR and a coformer to the intermolecular energy of cocrystals and corresponding ‘empty’ pockets formed by APR molecules. In each case two possible positions of a coformer was regarded (referred to as a and b).

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | E in [Hartree] | | | | E in [kJ/mol] | |
|  | E2APR+COF+H2O | EAPR\_EMPTY | ECOF+H2O | E2APR+COF+H2O – (EAPR\_EMPTY + ECOF+H2O) | E2APR+COF+H2O – (EAPR\_EMPTY + ECOF+H2O) | Final EAr-Ar per asymmetric unit |
| APR:RESa | -4234.475 | -3775.15 | -459.280 | -0.04733 | -124.3 | -62 |
| APR:RESb | -4234.475 | -3775.15 | -459.281 | -0.04739 | -124.4 | -62 |
| APR:PICa | -4192.317 | -3775.15 | -417.133 | -0.03748 | -98.4 | -49 |
| APR:PICb | -4192.316 | -3775.14 | -417.133 | -0.03727 | -97.8 | -49 |
| APR:IMIa | -4077.931 | -3775.15 | -302.748 | -0.03649 | -95.8 | -48 |
| APR:IMIb | -4077.93 | -3775.15 | -302.748 | -0.03642 | -95.6 | -48 |
| APR:HQUa | -4234.464 | -3775.14 | -459.275 | -0.04435 | -116.5 | -58 |
| APR:HQUb | -4234.464 | -3775.14 | -459.275 | -0.04438 | -116.5 | -58 |

**Data for calculation of energy contributions to the cocrystals formation (CASTEP calculations)**

**Table S6.** Data for calculations of stabilization energies of the cocrystals (in eV). In each case two possible positions of a coformer was regarded (referred to as a and b).

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | ECOCR | EAPR\_B | ECOF | EH2O | EAPR\_B + 0.5ECOF + 0.5 EH2O | ECOCR – (EAPR\_B + 0.5ECOF + 0.5 EH2O) |
| APR:RESa | -8988.0134 | -7790.53 | -1922.38 | -472.65 | -8988.0462 | 0.032762 |
| APR:RESb | -8988.0139 | -7790.53 | -1922.38 | -472.65 | -8988.0462 | 0.032324 |
| APR:PICa | -8808.5782 | -7790.53 | -2036.23 |  | -8808.6432 | 0.065036 |
| APR:PICb | -8808.5808 | -7790.53 | -2036.23 |  | -8808.6432 | 0.062349 |
| APR:IMIa | -8572.1329 | -7790.53 | -1090.90 | -472.65 | -8572.306 | 0.173079 |
| APR:IMIb | -8572.1312 | -7790.53 | -1090.90 | -472.65 | -8572.306 | 0.174804 |
| APR:HQUa | -8987.8373 | -7790.53 | -1922.32 | -472.65 | -8988.0142 | 0.17687 |
| APR:HQUb | -8987.8388 | -7790.53 | -1922.32 | -472.65 | -8988.0142 | 0.175408 |

**Intermolecular interactions found in form B of APR and in APR cocrystals**

**Table S7.** Intermolecular hydrogen bond lengths (in Å, expressed as O∙∙∙O/N distances) and geometry parameters for aromatic – aromatic interactions (RCLOSEST distance in Å and γ angle in °, calculated as shown in Figure S6). For form B of APR letters A, B, C and D denote 4 unique molecules in an asymmetric part of a unit cell.

|  |  |  |  |
| --- | --- | --- | --- |
| Hydrogen bonds | | | |
| crystal | interacting atoms | distance | |
| APR:RES | H2**O**∙∙∙**O**H-Ar (RES)  H2**O**∙∙∙**O**Me-Ar (APR) | 2.65  3.01 | |
| APR:IMI | H2**O**∙∙∙**N** (IMI)  H2**O**∙∙∙**O**Me (APR) | 2.05  3.96 | |
| APR:HQU | H2**O**∙∙∙**O**H-Ar (HQU)  H2**O**∙∙∙**O**Et-Ar (APR)  (HQU)Ar-H**O**∙∙∙**O**2S-CH3(APR) | 1.93  3.19  2.11 | |
| Aromatic – aromatic interactions | | | |
| crystal | interacting rings | RCLOSEST | γ angle |
| APR form B | Ar(OMe)(OEt) ∙∙∙ Ar(OMe)(OEt) (A:B)  Ar(OMe)(OEt) ∙∙∙ Ar(OMe)(OEt) (C:D)  Isoindol ∙∙∙ Isoindol (A:B)  Isoindol ∙∙∙ Isoindol (C:D) | 3.79  3.83  3.46  3.47 | 63.3  69.6  60.5  59.9 |
| APR:RES | Isoindol (APR) ∙∙∙ Phenyl (RES)  Isoindol (APR) ∙∙∙ Ar(OMe)(OEt) (APR) | 3.35  3.67 | 54.4  33.5 |
| APR:PIC | Isoindol (APR) ∙∙∙ Pyridine (PIC)  Isoindol (APR) ∙∙∙ Ar(OMe)(OEt) (APR) | 3.37  3.64 | 57.3  35.3 |
| APR:IMI | Isoindol (APR) ∙∙∙ Imidazole (IMI)  Isoindol (APR) ∙∙∙ Ar(OMe)(OEt) (APR) | 3.37  3.65 | 53.6  34.3 |
| APR:HQU | Isoindol (APR) ∙∙∙ Phenyl (HQU)  Isoindol (APR) ∙∙∙ Ar(OMe)(OEt) (APR) | 3.42  3.68 | 61.3  33.7 |

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**Figure S6.** Geometric parameters for aromatic – aromatic interact.ions given in Table S7. Purple balls mark centroids of the interacting aromatic ring.