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The role of fluids in high-pressure polymorphism of drugs: different behaviour of β -chlorpropamide in different inert gas and liquid media

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The hydrostatic compression of β -chlorpropamide gives different high-pressure phases, depending on the choice of pressure-transmitting fluid (paraffin, neon and helium). This is particularly surprising as none of these fluids interact obviously with the solid at ambient pressure. This phenomenon is not related to dissolution and recrystallization, in contrast to what has been previously observed for β -chlorpropamide in a 1:1 pentane-isopentane mixture.

Introduction

The last decade has seen an explosive increase in the number of publications on the effect of hydrostatic pressure on organic crystals ^{1–8}. One of the reasons for this is that new polymorphs can be formed. This is of fundamental interest for the design of crystal structures and understanding the mechanisms of crystallisation and solid-state transformations. High-pressure polymorphism is also of practical importance, with new phases being used as obtained or as seeds for subsequent masscrystallization. These applications are particularly relevant if the high-pressure phase can be preserved on decompression. Polymorph discovery is of special interest to the pharmaceutical industry ^{7,9–11}, since different polymorphs differ not only in their physical and biological properties, but are also separate legal entities, are patentable and are thus important for intellectual property ^{12,13}.

It is generally not possible to predict which phase will be formed at a selected (T, P) point based solely on a thermodynamic phase diagram. Instead, there exists a complex interplay of nucleation and growth kinetics, alongside thermodynamics, that leads to unpredictable results ¹⁴. As a result of the kinetic control of nucleation and nuclei growth ¹⁴, different phases can form depending on the choice of the

starting polymorph $^{15-19}$, the compression/decompression protocol $^{20-23}$, and the choice of hydrostatic medium $^{24-28}$. The latter is of particular interest to the present study.

Hydrostatic conditions cannot be achieved without a hydrostatic medium – a fluid phase, i.e. a gas or a liquid. In high-pressure mineralogy, the importance of selecting a pressure-transmitting fluid is well understood. This has become particularly important in relation to microporous materials, zeolites, which are characterized by an "open" system of interconnected channels and cavities²⁹. For example, when present in the hydrostatic medium (even in trace quantities), molecules of H₂O can enter zeolite channels upon compression; this process is reversible ³⁰. The so-called "pressure-induced hydration" 31 can cause either volume increase (abrupt as in 31,32 or gradual as in 33), or simply a decrease in compressibility as compared with the nonhydrated species, due to the site occupancy increase of the already existing water sites ^{34,35}. It is important to note, however, that this pressure-induced hydration effect is not always observed ^{36,37}. Gasses used as pressure-media are generally inert with respect to minerals, but gases such as hydrogen or helium can also penetrate into the solid. This can result in a decreased compressibility of the material, ³⁸⁻⁴¹ or also in the structural transformations, as for cristobalite ⁴², or vitreous silica ⁴³ compressed in helium. Despite these effects being known for minerals, the choice of pressure-transmitting medium (PTM) for high-pressure research of organic molecular crystals has been rather arbitrary. Most commonly, a PTM is chosen for practical reasons, often simply based on which are physically available in the laboratory. Gas loading is not widely available, and similarly, not all groups have the facilities to work with low boiling point fluids, such as pentane-isopentane ⁴⁴. In practice, a fluid is selected not only for availability, but i)



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Figure 1. Molecular structure of chlorpropamide. Carbon atoms - grey, hydrogens - light grey, oxygens - red, sulfur - yellow, chlorine – green

to ensure hydrostatic conditions are maintained up to the desired pressure point, and ii) to avoid obvious chemical interaction with the immersed solid, while simultaneously avoiding its dissolution. For the former, there exist large repertoires of data summarizing the hydrostaticity limits of various fluids. 45,46 Recent studies with precise pressure measurements across multiple sites in a pressure cell show that a notable loss of ideal hydrostaticity (thus possible shear stresses in the medium) occur at considerably lower pressures than was previously expected from either the broadening of Xray diffraction peaks or from the measurement of a single pressure point in the cell ⁴⁵. In interactions of the PTM with the sample, it is very difficult (if not impossible) to predict a priori whether the solubility of a substance will change with pressure. This is particularly problematic when a compound is insoluble in the PTM under ambient conditions, but dissolves under pressure. It is also difficult to predict surface interactions between a solvent and the crystal material, an interaction that may increase with pressure. One must therefore always consider the possibilities of high-pressure dissolution, re-crystallisation, and solvate formation at elevated pressures.

A number of studies dedicated specifically to the role of fluids in pressure-induced solid-state transformations have been ^{6,21,23-28,47-83}. Fluids have been documented to published reversibly penetrate into crystal structures to form inclusion compounds, especially if the solid structure has channels or large cavities, such as MOFs ^{47–55} or zeolites ^{30–35}. Additionally, solids have been reported to recrystallise into solvates at high pressure ^{6,21,23,28,56–70}. It was also shown that some solid-state pressure-induced phase transitions can be solvent-assisted, i.e. observed in some fluids, and non-observed in others ^{24–27}. This phenomenon is also known for ambient-pressure crystallisation ⁷¹⁻⁸². Transformations in liquids, in which solids can dissolve only under pressure - often recrystallising as a new high-pressure phase - are sometimes compared with those in inert fluids (i.e. those in which samples do not dissolve under pressure, so that real solid state transformations are possible). A systematic investigation into the variety of possible effects fluids may have on pressure-induced transformations of solids is just emerging. To the best of our knowledge, there are currently no documented examples that compare the effect of pressure on the same solid phase when immersed in different non-dissolving fluids.

The aim of the present work was to compare the effect of pressure on a selected organic crystal using several "inert"



Figure 2. Molecular packing in $\beta\mbox{-chlorpropamide}.$ Hydrogen bonds are shown by blue lines

hydrostatic fluids, none of which can visibly dissolve the solid under ambient conditions. The present work investigates (4-chloro-N-(propylaminochlorpropamide, carbonyl)benzenesulfonamide, $C_{10}H_{13}CIN_2O_3S$) (Fig. 1), an antidiabetic drug which is prone to forming new polymorphs 84,85 under ambient and high pressures $^{25,26,86-89}.$ The α polymorph, which is the stable form under ambient conditions ⁹⁰, undergoes at least one phase transition on increasing pressure to give a high-pressure phase, α' -polymorph ²⁶. This high pressure phase is the same as that obtained when the PTM used is either a saturated ethanol solution (recrystallisation possible) ²⁶ or a 1:1 pentane-isopentane mixture (no visible dissolution)⁸⁹. The same phase transition seems to also take place in dry powder samples without the addition of any pressure-transmitting fluids ²⁵, albeit at a considerably slower rate. This has led to the suggestion that the high-pressure transformation in chlorporopamide is kinetically hindered and thus solvent-assisted. ²⁵. When different polymorphs (α -, β -, γ -, δ - forms) are compressed in pentane-isopentane, different high-pressure phases are formed at different pressures ⁸⁹. At the same time, when compressing a sample of the β -polymorph in pentaneisopentane (1:1), different high-pressure phases are formed, depending on the presence of other chlorpropamide polymorphs as seeds ⁸⁹. This suggests that the pressureinduced structural transformations of the β -polymorph are related to recrystallisation from pentane-isopentane at high pressure, even though this solvent does not visibly dissolve any of the polymorphs at ambient pressure, or the α -, γ -, and δ forms at high pressure ⁸⁹. This system therefore offered an intriguing opportunity to compare the effect of an inert liquid (paraffin) with that of inert gases (Ne, He) on the structural transformations in the β -polymorph (Fig. 2). These results could then be compared with those observed on compression in a 1:1 pentane-isopentane mixture, in which the sample starts dissolving as pressure increases⁸⁹.

Medium	A high-pressure phase	SG and cell parameters	Pressure range when	Polycrystalline or single
			formed, GPa	crystal?
Ne	α-polymorph	P2 ₁ 2 ₁ 2 ₁ , 4.9582(4) Å,	Between ambient	Several domains in what
		26.502(13) Å, 8.9098(7)	pressure and 0.6 GPa	was originally a perfect
		Å) <i>,</i>		single crystal
	α'-polymorph?	Cell could not be found	Between 2.3 and 2.6 GPa	Strongly distorted crystal
		(reflections are strongly		split into several large
		distorted)		fragments

Table 1. A summary of the pressure-induced phase transformations observed on compressing a single crystal of β -chlorpropamide immersed in different media*

He	A new phase, β^{I}_{HP}	Monoclinic, 14.349(5) Å, 9.2192(17) Å, 18.84(4) Å, 90.35(8)°	Between 0.3 and 0.5 GPa;	Single crystal
	A new phase, β^{II}_{HP}	Monoclinic, 14.041(6) Å, 9.1411(19) Å, 18.71(4) Å, 94.63(11)°	Between 0.7 and 1.0 GPa	Several domains in what was originally a perfect single crystal
Liquid paraffin	A new phase; β^{I}_{HP} ?	Monoclinic, 14.399(3) Å, 9.2288(18) Å, 18.87(4) Å, 91.61(3)°	Between ambient pressure and 0.1 GPa	Several domains in what was originally a perfect single crystal
	A new phase; β^{III}_{HP} ;	Triclinic, 14.200(2) Å, 9.2010(17) Å, 18.858(16) Å, 89.45(3)°, 86.66(3)°, 89.980°	Between 0.1 and 0.3 GPa;	Several domains
	Not identified	Cell could not be found (reflections are strongly distorted)	Between 1.6 and 2.2 GPa	Strongly distorted crystalline domains
1:1 pentane-isopentane mixture ⁸⁹	γ-polymorph	Monoclinic, 6.1040(5) Å, 8.9243 Å, 12.0304(14) Å, 99.516(8)°	At 0.1-0.2 GPa	Multiple new crystals
1:1 pentane-isopentane mixture, δ - and α - phases also present	γ + δ	Not measured, crystal growth was followed visually	0.3-0.5 GPa	γ- polymorph as new crystals + δ- polymorph seed growing; no growth of the α-polymorph;
		Non-identified phases after transformation of both forms, characterized by Raman spectra	2.4-3.3 GPa	Single crystals visually preserved

*- Data for compression in pentane-isopentane are from $^{\mbox{\tiny 89}}$ and given for comparison



Figure 3. β -CPA loaded in neon immediately after loading at BGI (0.6 GPa) and after high-pressure experiment at ESRF (6.0 GPa); the sample at the left photo looked the same two days later when examined at ESRF after transport

Results and Discussion

The main results of the experiments are summarised in Table 1. Microphotographs are given for crystals of β -chlorpropamide loaded in a DAC in neon, immediately after compression to the first pressure point (0.6 GPa) and after the phase transformations in neon, Fig. 3.

Compression in Ne

A single crystal of the β -polymorph was loaded into a DAC in Ne, with a pressure of 0.6 GPa immediately after closing the cell. The loaded DAC was then transferred to ESRF to perform diffraction experiments. The time interval between DAC loading and the diffraction experiments was therefore two days. The initial diffraction data obtained at the synchrotron source corresponded to the α -polymorph; the high-quality single crystal originally loaded had transformed into several singe-crystalline domains. The pressure measured after the transfer of the DAC to ESRF and the diffraction experiment was 0.5 GPa. Unfortunately, it was not technically possible to repeat the loading of a DAC with a crystal of the β -polymorph in Ne and to track any phase changes at pressures below 0.5 GPa. Therefore, it was not possible to determine the phase transition point more precisely, or investigate any kinetics effects, i.e. if the high-pressure transformation takes place after some delay when kept at a certain pressure. On further compression in steps of 0.5 GPa, the α -polymorph (which formed from the original β -form) transformed into another phase. The transition pressure range (2.3-2.6 GPa) was similar to that at which the transformation of the α -form into the high-pressure $\alpha'\text{-polymorph}$ was previously observed on compression in saturated ethanol solution ²⁶ (2.6-2.9 GPa), or in pentane-isopentane ⁸⁹ (2.4-3.3 GPa). Obviously, the diffraction pattern no longer corresponded to that of $\alpha\text{-}$ polymorph. However, the quality of the diffraction data was not sufficient to unambiguously decide whether the new phase was the same α' -polymorph described in ²⁶ based on single-crystal diffraction, or any other phases which are claimed to exist. Unfortunately, many of these phases have not been characterized, even by cell parameters, ⁸⁷ and have formed apparently without PTM. No obvious transitions were



Figure 4. Dependencies of cell parameters on pressure for $\beta\mbox{-chlorpropamide loaded}$ in He

observed on further compression up to 6.0 GPa (the last pressure point achieved in the experiment). In an independent experiment, a crystal of the original α -polymorph in Ne, which was compressed immediately to 4.6 GPa, also transformed into a high-pressure phase. As before, it was not possible to unambiguously identify this phase as the α' -polymorph, or another phase, as the crystal was strongly distorted.

Compression in He

When compressing a crystal of the β -polymorph in He, pressure was increased in small steps. The diffraction data at pressures below 0.3 GPa corresponded to the orthorhombic βpolymorph. The first changes in cell parameters, which might be a manifestation of a phase transition into a monoclinic phase (designated β_{HP}^{I}), were observed between 0.3 and 0.5 GPa (Fig. 4). Although the crystal was not visibly destroyed, and the β -angle was still very close to 90° (90.35(8)° at 0.5 GPa), the diffraction data could no longer be described using the structure of β -chlorpropamide as a starting model. Attempts to solve the crystal structure or to refine in any of the other previously known models also failed. The cell parameters of the new monoclinic phase did not correspond to the α -polymorph, or to any of the other previously reported chlorpropamide polymorphs. This suggests that the structural transformation must have been significant and a new β'_{HP} phase has been formed. The next jump-wise change in the

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on compression has been documented

Pressure transmitting media (further PTM)	Не	Ne	Paraffin	Pentane- isopenta ne
Atomic	0.31	0.38	>15	~5
radius/Molecule size,	(1.40	(1.54		
Å	VdW)	VdW)		
Possible dissolution of PTM in β-CPA	+	±	-	-
interstitial voids?				
Interaction of β-CPA	-	-	+	+
alkyl tales with PTM				
Volume of voids in	764.26	667.15	-	-
unit cell of β-CPA	(28.9 %)	(25.2 %)		
available for sphere of				
a certain atomic				
radius, Å ³ (% of unit				
cell)				
Volume of voids in	29.11	25.06	-	-
unit cell of α-	(17.0 %)	(14.6 %)		
cristobalite available				
for sphere of a certain				
atomic radius, Å ³				
Volume of voids in	312.69	297.08	-	-
unit cell of arsenolite	(23.0 %)	(21.9 %)		
available for sphere of				
a certain atomic				

value of the β -angle was observed between 0.7 and 1.0 GPa (Fig. 4). The second new high-pressure phase was designated β^{II}_{HP} ; the cell parameters of this phase were very close to those of the β^{II} -chlorpropamide low-temperature phase, which forms on cooling of the β -polymorph to about 257 K ⁹⁴. Again, although the crystal was not visibly destroyed, attempts to solve the crystal structure of this new phase, or to refine it using the structure of β^{II} -chlorpropamide as a starting model failed, this indicating that structural distortion / rearrangement was very significant.

So, the response of the crystals of the β -polymorph to pressure was different, depending on the choice of the inert gas. One can suppose that this difference can be a consequence of the difference in the atomic radii of Ne and He. Helium was shown to penetrate into voids in various solids, such as, e.g. crystalline SiO₂ (cristobalite) ⁴², arsenolite^{92,95}, or ice ⁹⁶. Cristobalite gave a new phase at about 8 GPa which was supposed to have a molar volume of about 30 % larger, than cristobalite, suggesting the dissolution of helium atoms in its interstitial voids. As₄O₆-2He clathrate was shown to appear at the surface layer of arsenolite and penetrate in depth on increasing pressure. Experimental and computational results provide evidence for ordered helium trapping above 3 GPa between adamantane-type As₄O₆ cages ⁹⁵.

Analysis of voids sizes in β -chlorpropamide (Fig. 2) shows that they are larger than the voids in α -cristobalite $^{^{93}}$ and



Figure 5. Dependency of unit cell volume on pressure for β -chlorpropamide loaded in paraffin

arsenolite ⁹² (Table 2). It therefore follows that small He atoms can penetrate into the crystal structure of the β chlorpropamide, filling the cavities between chlorpropamide molecules (Fig. 2). The changes in cell parameters and volume of the β -chlorpropamide crystals on compression in helium can be caused by two phenomena. First, these changes may be caused by helium inclusions that occur as a result of continuous structural strain (itself being introduced by the hydrostatic compression), or secondly, they may occur as a result of a combination of strain from external hydrostatic compression and internal strain from helium guest atoms. In this respect it may be relevant to note that helium penetration into the interstitial free volume of the glass network of vitreous silica has been documented to induce deformation that accounts for the small apparent compressibility of silica ³⁹.

Compression in paraffin

As the third inert fluid, liquid paraffin was used. The values of volume vs. pressure measured are plotted in Fig. 5. Transformation of the β -polymorph into a monoclinic phase with cell parameters similar to those of β'_{HP} chlorpropamide was observed to have occurred by 0.1 GPa. As pressure was increased to 0.3 GPa, another high-pressure phase was formed, with a diffraction pattern that could be indexed by assuming a triclinic cell that does not correspond to any previously reported crystal structures of chlorpropamide (β^{III}_{HP}) . Further compression shows that the *V(P)* curve does not have discontinuities up to 1.6 GPa. On further pressure increase to 2.2 GPa the crystal was destroyed, yielding a polycrystalline phase. Attempts to index the diffraction pattern and find the cell parameters failed, as the reflections were strongly distorted. This is presumably due to structural distortion. This again shows that the pressure-induced transformation must be related to a rather significant structural reorganisation.

It has been shown that the behaviour of β -chlorpropamide on increasing pressure in several "inert" gas and liquid media is different. Helium is small enough to penetrate into the crystal structure and subsequently trigger a solid-state polymorphic

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Table 3. Summary of the details related to loading the samples of $\beta\mbox{-}polymorph$ into the DACs

Medium	DAC type	Gasket and crystal details	
Ne	BX-90 type ⁹⁷ , an	Crystal size 0.16×0.14×0.02 mm	
	opening cone of 80°.	Gasket:	
	Culet size 600 µm.	Thickness: 100 μm	
		Hole diameter: 250 μm	
Liquid	BX-90 type ⁹⁷ , an	Crystal size 0.10×0.05×0.05 mm	
paraffin	opening cone of 80°.	Gasket:	
(ROTH	Culet size 600 µm.	Thickness: 85 μm	
GmbH)		Hole diameter: 300 µm	
He	A membrane-driven	Crystal size 0.05×0.04×0.02 mm	
	DAC ⁹⁸ with Boehler-	Stainless steel Gasket:	
	Almax seats ⁹⁹ , an	Thickness: 80 μm	
	opening cone of 64°.	Hole diameter: 350 µm	
	Culet size 600 µm.		

transformation. The transformations observed in the presence of Ne can be expected to be of a solid-to-solid type and be defined solely by intracrystalline interactions. In the absence of He as a guest molecule, the solid-state transformation in Ne gives the α -polymorph below 0.6 GPa (the thermodynamically stable form at ambient conditions ⁹⁰), and not the highpressure phases that are formed in He. Pentane-isopentane starts dissolving β -chlorpropamide at high pressure such that a high-pressure recrystallisation, and not a polymorphic solidstate transformation, takes place ⁸⁹. The most interesting observation is that even the high-pressure transformations in paraffin and Ne are different, despite the fact that the two media neither dissolve nor penetrate the chlorporpamide crystal. The reason for this difference likely rests in different van der Vaals interactions between the surface of chlorpropamide crystals and the pressure transmitting media in the two cases. In particular, the alkyl tails of chlorpropamide molecules can be supposed to interact with the paraffin molecules^{*}.

Experimental

 β -Chlorpropamide was crystallised by slow evaporation of saturated ethanol solutions prepared by dissolving a powder

sample of α -chlorpropamide. The chlorpropamide polymorphs were loaded into DACs of different types, depending on the choice of the pressure medium (Table 3). Small ruby spheres were used as pressure calibrants ^{101,102}. The gas loading of Ne and He did not require any preliminary cooling of the samples in the DACs. He or Ne was loaded into the DAC as a PTM using a high pressure gas loading apparatus in BGI ¹⁰³. Pressure was further increased as long as the structural integrity of the crystal and the quality of diffraction data permitted determination of cell parameters. These upper limit pressure values were different for different media (see below). All diffraction data (excluding the experiment with β chlorpropamide compressed in helium) were collected at the Swiss-Norwegian Beamline BM01A at the European Synchrotron Radiation Facility (ESRF, Grenoble, France, experiment CH-4526). A parallel monochromatic X-ray beam (E = 17.8 keV, λ = 0.69783 Å) cropped to 200 × 200 μm^2 on the sample was used. Single-crystal data were collected by a horizontal-acting ω -axis rotation with an integrated step scan of 0.5° and a counting time of 2s per frame. A PILATUS 2M hybrid pixel detector was used with sample-to-detector distance 196 mm. The data were converted and integrated using the SNBL toolbox ¹⁰⁴ and CrysAlisPro ¹⁰⁵ software packages. X-ray diffraction data for β -chlorpropamide loaded with helium were collected at ID09A synchrotron beamline (ESRF, Grenoble, France) using a parallel monochromatic X-ray beam (E = 30 keV, λ = 0.413 Å) focused to 30 × 30 μ m² on the sample. Single-crystal data were collected by a vertical-acting ω -axis rotation with an integrated step scan of 0.5° or 1° and a counting time of 1s per frame. A MAR555 flat-panel detector was used for recording the diffraction intensities. The sampleto-detector distance was 250 mm. The data were then processed according to 106 and integrated using the CrysAlisPro 105 software package. Mercury 107 was used to visualise the crystal structures and calculate the size of voids. The contact surfaces were calculated using the following parameters: probe radius equal to 0.3 Å for helium and to 0.4 Å for neon, approximate grid spacing is 0.3 Å.

Conclusions

This study has shown that the choice of pressure-transmitting fluid can be critical for the outcome of a high-pressure experiment, and not only when the fluid visibly dissolves the solid. As shown in the present contribution, recrystallisation is not the only possible mechanism through which a fluid can influence a solid-state transformation in an organic molecular solid, to justify classifying the transformation as "fluidassisted". Other mechanisms include the penetration of the medium into the solid, generating additional inner stress and thus triggering a structural transformation, or surface interactions that can influence the mechanical properties (macroscopic level of consideration). Alternatively, fluid molecules may induce conformational changes in molecules, and thus their rearrangement into a new structure (microscopic level of consideration). The ability to change mechanical properties of solid materials (plasticity, brittleness)

[‡] It has been shown previously that ball-milling of α -chlorpropamide results in a polymorphic transformation into the ϵ -polymorph only at 77 K, but not at ambient temperature. It has been supposed, that a change in the orientation of the alkyl tails in chloropropamide molecules on cooling can trigger a polymorphic transformation on milling at low temperatures¹⁰⁰

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through addition of fluids at the surface has long been known for metals and ionic salts (loffe¹⁰⁸, Rebinder¹⁰⁹ or Rosko¹¹⁰ effects). The same effects can be expected also for organic molecular crystals. All the various types of solid-fluid interactions, not only the possibility of dissolution or penetration, must be taken into account in any polymorph screening experiments involving high pressure as a varied parameter. Further research of the solid-fluid interfaces in these systems, of the effect of fluids on the mechanical properties of the chlorpropamide crystals, as well structure solution of the new high-pressure phases will shed more light on the mechanisms of the solid-state transformations in the presence of a fluid in these and other organic crystals. This is an area of critical importance for many fields of chemical, materials and pharmaceutical technologies.

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