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Comparison of the physical and thermodynamic stability of amorphous azelnidipine and its coamorphous phase with piperazine†

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A major challenge in drug development is that the majority of drugs are water insoluble, and a powerful method to conquer this obstacle is to transfer a crystalline drug into its amorphous phase (AP) or coamorphous phase (CAP) with a coformer. In the present study, the physical and chemical stabilities of an AP and a CAP based on the dihydropyridine calcium ion antagonist azelnidipine (AZE) were investigated using thermal analysis and a solution chemistry method. The identification of two APs (named α -AP and β -AP, from crystalline α -AZE and β -AZE, respectively) and one AZE-piperazine CAP was attempted using powder X-ray diffraction, temperature modulated differential scanning calorimetry and Fourier-transform infrared spectroscopy. The transition thermodynamics from the two APs and the CAP to stable crystalline β -AZE (β -Cry) were investigated using a solubility method. The solubility of the two APs, the CAP and β-Cry in 0.01 M HCl medium at 298, 304, 310, 316 and 322 K was investigated; the values obtained were used to calculate the thermodynamic parameters of the transition reaction. The transition temperatures of α -AP, β -AP and the CAP to form β -Cry in 0.01 M HCl were 237.7, 400.3, and 231.4 K, respectively. The glass transition temperature (T_q) values of α -AP, β -AP and the CAP were 365.5, 358.9 and 347.6 K, respectively, indicating a high physical stability for α -AP. However, β -AP proved to be the most thermodynamically stable form at room temperature compared with α -AP and CAP in the 0.01 M HCl medium. As evidenced by those observations, no general relationship occurred between the solid physical stability and the solution chemical stability for AP and CAP. The kinetics of the solidstate decomposition, studied using DSC analysis, showed that the activation energies for decomposition of α -AP, β -AP and CAP at high temperatures were 133.0, 114.2 and 131.6 kJ mol⁻¹, respectively.

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Introduction

In recent years, the magnitude of research emphasizing the importance of different solid-state forms of a drug, such as crystalline, amorphous, or hydrate forms, in pharmaceutical development and manufacturing has increased significantly.1-4 These forms have different physical, chemical and mechanical properties, which not only affect the processability and stability of the drug product but can also lead to differences in bioavailability. For example, using the amorphous phase (AP) of an active pharmaceutical ingredient (API) can improve the physicochemical properties of an insoluble API. However, the main drawback of using an amorphous API in dosage forms is the limited and largely unpredictable physical stability of the AP in the solid state during manufacturing or storage.5 Another

have been reported for indomethacin, dipyridamole, and carbamazepine.8 Amorphous indomethacin prepared using spray drying had some remaining residual crystallinity, and AP samples produced using a grinding procedure had small degrees of undetected crystallinity remaining or differences in the hydrogen bonding in the different APs.8 The results of an

investigation by Rades et al. indicate the existence of an AP for cryo-milled simvastatin that is spectrally distinguishable from the quench-cooled AP. The thermodynamic parameters suggest that this form is less disordered compared to the quench-cooled form.11 Furthermore, differences regarding the physical stability and recrystallization behavior for amorphous forms produced from different crystalline starting counterparts by ball-milling

have been reported for anhydrate piroxicam form I and form

problem of some AP systems is solvent-induced crystallization

upon oral application, that is, the thermodynamic stability in

the preparation method,8 thermal history,9 or starting crystal-

line form when a cryo-milling preparation method is used.^{3,10} Differences in the stability and physicochemical properties,

regarding APs produced using various preparation techniques,

The physicochemical properties of the AP can be affected by

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II.3 The results indicate that amorphous piroxicam prepared from crystalline form I had a lower physical stability than that of amorphous piroxicam prepared from form II, which was linked to the higher residual order in the amorphous samples prepared from crystalline form I; the recrystallization behavior of the amorphous samples prepared from anhydrate piroxicam form I was closely related to the starting material, whereas the recrystallization behavior of the amorphous samples prepared from form II resembled that of form III. Because these APs are formed via different intermediate states (melt, solution, solid) or different polymorphic forms, the question arises whether there are indeed differences in the amorphous solids formed. Hancock et al. used the term "pseudopolyamorphism" for those different amorphous forms.12 For true polyamorphs, a firstorder transition between them would have to be observed, which has never been seen in any organic substance.11 Although, using melting techniques, the amorphous state is obtained through destruction of the existing molecular order, it is not clear if the differences exist in APs produced from different crystalline starting counterparts by melting or if local molecular order may still be present in the APs.

However, an AP with a high internal energy and specific volume relative to the crystalline state has a possibility that it may undergo partial crystallization during processing (mechanical stress) and storage (temperature and humidity stress) or in a solvent medium. Hence, exploitation of the full potential of amorphous solids requires their stabilization in the solid state and in solvent media as well as during application. A coamorphous phase (CAP), an amorphous combination consisting of the active pharmaceutical ingredient (API) and a small molecule coformer, possesses special advantages in improving the solubility and dissolution rate of the API and enhancing the stability of single amorphous drugs. 13-15 For this reason, the CAP is considered as an efficient replacement for drug-polymer molecular association (solid dispersions) and cocrystals. This approach creates solubility advantages and stabilizes the amorphous system through intermolecular interactions, such as hydrogen bonds, and might conquer shortcomings associated with solid dispersions. However, there are very few reports on the thermodynamic mechanism for improving the stability of the CAP compared to the single AP.

Azelnidipine (AZE) is a new long-acting dihydropyridine calcium channel blocker for the treatment of patients with hypertension. It belongs to the group of polymorph drugs that have two, α and β , polymorphs, and the α form is a metastable phase that easily transforms to the β form under suitable conditions. In our previous studies, the thermodynamic stability of crystalline AZE polymorphs was investigated. 16 The present paper addresses the question of whether the different polymorphs result in differing forms of the AP using a melting technique. Furthermore, AZE is a good model API to screen APIcoformer amorphous combinations and many CAPs were obtained in our previous study.14,15

Based on the above considerations, the purpose of this study is to investigate the physical and thermodynamic stabilities of the AP and CAP and to find out if there is a general rule for the relationship between the physical stability in the solid state and

the thermodynamic stability in solution. In this study, we present our results on the measurement of solubility in a 0.01 M HCl solution for different APs prepared from two polymorphs and a new CAP produced from AZE and piperazine (PEZ); the solubility data at different temperatures were used to investigate and evaluate the thermodynamic stability of the CAP compared to the single AP. Furthermore, the physical stability in the solid state and the thermal decomposition of the two APs and the CAP were also investigated.

Experimental

Chemicals

The AZE polymorphs ($M = 582.64 \text{ g mol}^{-1}$), α (α -Cry, $T_{\rm m} = 123.0$ °C) and β (β -Cry, $T_{\rm m}=199.0$ °C), were purchased from the Wuhan Yuancheng Gongchuang Technology Co., Ltd. Hexahydrate PEZ was supplied by Damao Chemical Reagents Factory (Tianjin, China). All other chemicals in this work were used as received.

Preparation of the two APs from the AZE polymorphs

Chemically pure amorphous AZE, confirmed by high performance liquid chromatography (HPLC), was obtained by melting and quench cooling of the α and β crystalline forms. Crystalline form α or β was heated in a stainless steel beaker to 150 °C for α or 240 °C for β in an oven and held at this temperature for 3 min. Then, the resulting melt was taken out of the oven and quenched using liquid nitrogen at once. Subsequently, the resulting melt that had been cooled to room temperature was milled under the conditions of pouring liquid nitrogen; the obtained powder, α-AP or β-AP, was passed through BSS sieve no. 60. The as-prepared APs were retained in a vacuum desiccator until required for characterization.

Preparation of the CAPs of AZE and PEZ

The CAPs of α-Cry and PEZ were prepared by grinding a mixture of α -Cry and PEZ for 30 min in a laboratory-scale oscillatory disc mill (Mixer Mill RS200, Retsch GmbH, Dusseldorf, Germany) using a molar ratio of α-Cry/PEZ of 1:2 without any other solvents. The mixer mill was stopped every 10 min during the grinding process for 3 min, to scrape the wall of the steel jar and homogenize the mixture. Unnecessary heat generated during grinding was also dispersed during the stopping intervals.

Characterization

Powder X-ray diffraction (PXRD)

The solid-state of the crystalline forms, and the APs and CAPs produced from AZE were analyzed on a RINT 2100 X-ray diffractometer (Ultima Rigaku Co., Japan) using a Cu-Ka target with a 0.3 mm receiving slit. Approximately 50 mg of the sample powder was gently consolidated in a glass holder and scanned at 20 kV and 20 mA from 3 to 40° (2 θ) at a step size of 0.0002° (2 θ). The sample surface was flattened softly to avoid particle orientation before determination.

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Fourier-transform infrared (FTIR) spectroscopy

FT-IR spectra were recorded on a Bruker Equinox 55 FT-IR spectrometer in the 4000-400 cm⁻¹ range. The spectrum was an average of 40 scans with a spectral resolution of 2 cm⁻¹. The samples were prepared by grinding 4 mg of the sample with 160 mg of spectral grade KBr in an agate mortar.

Differential scanning calorimetry (DSC) and temperature modulated differential scanning calorimetry (TMDSC) measurements

DSC and TMDSC were performed using a DSC 214 (NETZSCH Company, Germany) equipped with a refrigerated cooling accessory (IC40) and a data analyzer (Proteus Analysis7.0) under a nitrogen atmosphere. For the TMDSC, the temperature was modulated by 0.5 K every 60 s with a heating rate of 2 K min⁻¹ from 273 to 473 K, to determine the glass transition temperature (T_{o}) of the APs and CAP. Accurately weighed specimens of about 5 mg were placed in hermetically sealed aluminum pans with empty aluminum pans used as a reference. Thermogravimetric (TG) analysis of the samples was performed using a TG 209 thermogravimetric analysis instrument (NETZSCH company, Germany) at a heating rate of 10 K min⁻¹ from 298 to 723 K with nitrogen purging at a flow rate of 20 mL min⁻¹. To explore the decomposition process of the two APs and the CAP, DSC measurements were performed at heating rates of 10, 20, 30, and 40°C min⁻¹ within the temperature range of 298–723 K. The relationship between the heating rate β and the temperature of solid-state decomposition is given by the Kissinger equation:¹⁷

$$\frac{d\left(\frac{\ln \beta}{T_{\rm m}}\right)}{d\ln\left(\frac{1}{T_{\rm m}}\right)} = -\frac{E}{R}$$

 β represents the heating rate (K min⁻¹), $T_{\rm m}$ is the temperature of the first endothermic decomposition peak in the DSC curve, and E is the activation energy of decomposition. Then, a curve was drawn with ln $\beta/{T_{\rm m}}^2$ as the Y-coordinate versus $1/T_{\rm m}$ as the abscissa. E, the activation energy of solid-state decomposition, can be calculated from the slope of this curve.

Physical stability

The two APs and the CAP samples were stored in a desiccator under 75% RH conditions (saturated NaCl solution) at 40 °C for three weeks. Samples were repeatedly analysed, once a week, using XRD to investigate possible recrystallization.

Analysis of the solubility, transition temperature, and thermodynamics

The saturated solubility values of the crystalline forms α and β , the α - and β -AP, and the CAP of AZE-PEZ in 0.01 M HCl medium were determined at 298, 304, 310, 316 and 322 K, respectively. All the APs and CAP were passed through a BSS sieve no. 60 after milling to exclude the impact of particle size on the solubility and dissolution rate. An excess amount of sample was placed in a 50 mL screw-capped vial containing 30 mL of 0.01 M HCl

medium under the conditions of oscillation at 150 rpm in a SPH-200B air-bath shaker (Shiping Tech. Co., Ltd., Shanghai, China) for 72 h. The concentration of AZE in the solvent was determined at 259 nm using an UV spectrophotometer (UV-160 A, Shimadzu Co., Japan) after filtering through a 0.45 μm membrane filter. All the samples were tested thrice.

As mentioned above, the solubility of the different forms of AZE was investigated; the values obtained were used to calculate the transition temperature and thermodynamic parameters of the transition reaction using the thermodynamics formulas in ref. 16.

Briefly, at standard conditions, when different forms of AZE coexist in solvent, the relationship between ΔG^{θ} , ΔH^{θ} , and ΔS^{θ} of the transition reaction of two forms A and B is given by eqn (1) and (2):

$$-RT \ln \frac{\alpha_{\rm A}}{\alpha_{\rm B}} = \Delta G_{\rm A,B}^{\theta} = G_{\rm A}^{\theta} - G_{\rm B}^{\theta} \tag{1}$$

$$-RT \ln \frac{\alpha_{\rm A}}{\alpha_{\rm B}} = \Delta G_{\rm A,B}^{\theta} = \Delta H_{\rm A,B}^{\theta} - T\Delta S_{\rm A,B}^{\theta} \tag{2}$$

 α_A and α_B are the activities of the two different forms. Approximately, c (solubility) is equal to α (activity) in an infinitely diluted solution. In this study, three different transition pairs, including α-AP/β-Cry, β-AP/β-Cry, and CAP/β-Cry pairs, are involved.

According to Henry's law, we get:

$$\frac{\partial \ln \frac{c}{c^{\theta}}}{\partial \frac{1}{T}} = -\frac{\Delta H^{\theta}}{R} \tag{3}$$

$$\frac{\partial \ln \frac{c_{\rm A}}{c_{\rm B}}}{\partial T} = \frac{\Delta H_{\rm A,B}^{\theta}}{RT^2} \tag{4}$$

T indicates the transition temperature when $c_A = c_B$. Then:

$$\Delta H_{A,B}^{\theta} = T \Delta S_{A,B} \theta \ T = T_{t}$$

Using eqn (3), a line was drawn with $\ln c$ as the Y-axis versus $\frac{1}{T} \times 10^3$ as the *X*-axis. The $T_{\rm t}$ can be obtained from the point of intersection of the two curves of the two different forms. Afterwards, using eqn (4), a curve was drawn with $\ln c_A/c_B$ as the Y-coordinate versus $\frac{1}{T} \times 10^3$ as the abscissa. The values of $\Delta H_{A,B}^{\theta}$ and $\Delta S_{A,B}\theta$ can be calculated according to the slope and intercept of the curves, and the value of $\Delta G_{A,B}\theta$ was obtained from:

$$\Delta G_{AB}\theta = \Delta H_{AB}^{\theta} - T\Delta S_{AB}\theta.$$

Results and discussion

Identification and characterization of the AP and CAP

PXRD. The PXRD patterns of the crystalline and amorphous forms of AZE are shown in Fig. 1. The two APs, which were prepared using a melting and quench cooling method from αPaper



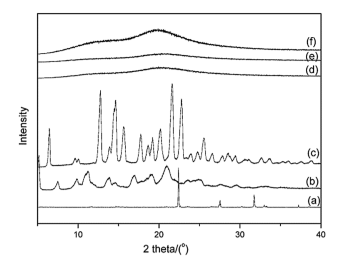


Fig. 1 PXRD patterns of the crystalline and amorphous phases of AZE. (a) PEZ; (b) α -Cry; (c) β -Cry; (d) α -AP; (e) β -AP; (f) α -AZE-PEZ CAP.

Cry and β-Cry, showed typical halo patterns, indicating that an AP was present (Fig. 1d and e). Furthermore, when α -Cry was coground with PEZ at a molar ratio of 1:2 for 30 min, the XRD peaks attributed to α-Cry disappeared and a clear halo pattern was observed (Fig. 1f), indicating the formation of a CAP of AZE and PEZ.

TMDSC. Generally, an amorphous phase can lead to the disappearance of the melting point and the emergence of a glass transition temperature (T_g) . The T_g is the single most defining characteristic of amorphous systems. We determined the T_g point of α -AZE-PEZ CAP using TMDSC and the TMDSC curves of the two APs were obtained from the data presented by Pan et al. 15 (Fig. 2). Tg values of 92.5 and 85.9 °C were observed for the α -AP and β -AP, respectively. In the present study, a melting and quench cooling method was used to prepare the AP, which can destroy the order of the crystalline phase.

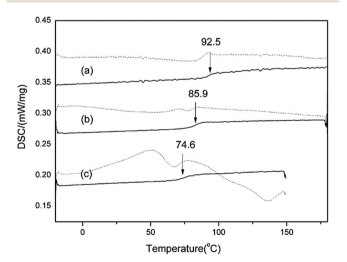


Fig. 2 TMDSC total heat flow signal (dash) and reversing heat flow signal (solid) for AZE amorphous systems: (a) α -AP; (b) β -AP; (c) α -AZE-PEZ CAP. TMDSC curves of the two APs were obtained from the data presented by Pan et al.15

However, different crystalline starting counterparts and different melting temperatures may lead to some different local and short-range order existing in the AP, which results in a minor difference in $T_{\rm g}$. Moreover, different APs with a different arrangement of the solid may result in a distinct difference in the weight loss rate, and we performed DTG analysis on the two APs to detect the difference (Fig. S2†). Unfortunately, the difference in the DTG results of the two APs was not enough to differentiate them. It was difficult to detect the difference in local and short order in the two APs by XRD, IR, DSC or DTG. In the case of the CAP with PEZ, the melting points disappear and a unique T_g appeared at 74.6 °C, indicating the formation of a homogeneous phase. The native AP of AZE showed a high T_g , but the CAP with PEZ at a ratio of AZE/PEZ of 1:2 showed a lower T_g , which seemingly followed the observation reported by Dengale *et al.*, 18 stating that the $T_{
m g}$ of a binary amorphous system tends to be closer to the T_g of the component present in excess within the mixture. So, although the T_g value of the native PEZ amorphous phase was not obtained in our study, the low T_{σ} value of the AZE-PEZ CAP depends largely on the PEZ. More evidence was deduced from the result of the T_g of a curcumin-PEZ CAP. Although pure amorphous curcumin has a higher $T_{\rm g}$ of 82 °C, the $T_{\rm g}$ of the curcumin-PEZ CAP was lowered to 33 °C owing to the introduction of PEZ.19 Fortunately, the T_g of the AZE-PEZ CAP is much higher than room temperature, which would make it possible for it to be developed as an actual pharmaceutical composition.

FT-IR. FT-IR spectroscopy can show the closely related structural compositions of different forms. The bands characteristic of AZE were in the range 3314-3455 cm⁻¹ where the N-H valence vibration occurred, in the range 1680–1700 cm⁻¹ where the -C=O-R stretching vibration is pronounced, then in the range 1000-1300 cm⁻¹ where C-O was observed, and in the range 800–860 cm⁻¹ where absorption bands were present due to C-H vibrations of benzene rings.16

Here, we focus on the differences between crystalline and amorphous forms. IR spectroscopy is used extensively to investigate hydrogen bonding because the peak position of the X-H stretch is very sensitive to the extent of association. Peak positions and assignments for selected interesting peaks which are related to hydrogen bonding for the crystalline and amorphous phases (AP and CAP) are shown in Fig. 3. The spectra of polymorphs α and β were compared with their amorphous counterparts to clarify the nature of the hydrogen bond interactions in the amorphous phase. Apparently, in addition to a broadening of the peaks in the amorphous sample, changes in peak shape, intensity, and position between the crystalline phases and their amorphous counterparts have occurred in the IR spectra. As can be seen in the spectrum of α -AP (Fig. 3d), the N-H peak has shifted slightly by 2 cm⁻¹ to a higher wavenumber; simultaneously, the peaks for the C=O groups merged into one peak. Generally, an unbonded X-H stretch gives rise to a relatively sharp peak, whereas on formation of a hydrogen bond X-H···Y (where Y is the acceptor atom), the peak shifts to a lower wavenumber and becomes much broader.20 The downward shift is caused by the lengthening of the X-H bond, which occurs on hydrogen bond formation. Based on this

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3440 1675 (f)
3444 1675 (e)
3444 1680, 1656 (b)
3205 (b)
4000 3800 3600 3400 3200 3000 2800 1500 1000 500 Wavenumbers/cm⁻¹

Fig. 3 FT-IR spectra of crystalline and amorphous phases of AZE. (a) PEZ; (b) α -Cry; (c) β -Cry; (d) α -AP; (e) β -AP; (f) CAP of AZE-PEZ.

consideration, a stronger hydrogen bond will lengthen X–H more and produce a shift to a lower wavenumber. In contrast to the crystalline state, a small upward shift of the N–H vibration occurred for the amorphous α , indicating that the hydrogen bond strength slightly increased or remained almost unchanged with loss of order in the crystalline structure.

A similar type of analysis of the IR spectral data was carried out for β -Cry and β -AP. On comparing with β -Cry, the N–H peak is broader and has shifted dramatically by 30 cm⁻¹ to a higher wavenumber for β -AP. Moreover, the peaks for the C=O groups also merged into one peak. A large upward shift of the N-H vibration in β-AP indicates that the hydrogen bond strength dramatically decreases with loss of the crystalline structure. It seems amazing that hydrogen bond strength slightly increases for form α but decreases for form β when the crystalline structure is destroyed. This difference may be derived from the differences in the crystalline structure between α -Cry and β -Cry. Single crystal XRD analysis revealed that β-Cry crystallized in an acentric space group of Pca21, in which hydrogen bonds played a crucial role in the 1D, 2D network in the crystal structure.14 When the crystal structure is destroyed, hydrogen bond interactions in β-Cry, which are responsible for stabilization the crystal structure, disappear. The reformed hydrogen bonds in the amorphous form are weaker than those in the crystalline form. Unfortunately, a single crystal of α-Cry has not been obtained and no crystal structure is referenced. Due to an almost unchanging hydrogen bond strength before and after transferring from the crystalline to amorphous phase, we can deduce reasonably that $\pi \cdots \pi$ intermolecular interactions but not hydrogen bonds play an important role in the crystal structure of α -Cry.

Using IR spectral data, the hydrogen bonding strength and pattern can be elucidated for the polymorphs and APs by monitoring the N–H (hydrogen bonding donor) and C=O (hydrogen bonding acceptor) stretching frequency. In α -Cry and β -Cry, the extent of variation in the N–H and C=O peak

positions suggests a difference in the hydrogen bonding patterns and thus in the strengths, which is consistent with the crystallographic data. However, the widths of the N-H and C=O peaks of the AP were greater than for the crystalline counterparts, which is consistent with a broader distribution of hydrogen bond lengths in the disordered phase. From Fig. 3, it was apparent, perhaps somewhat surprisingly, that form α appeared to have stronger hydrogen bonding on average in the amorphous state than in the crystalline state, while for form β, the opposite situation was observed, and the hydrogen bonding weakens on changing from the ordered to the disordered state. Obviously, there is no difference in the hydrogen bonding pattern and strength between α -AP and β -AP, but the opposite situation was observed for α -Cry and β -Cry. Different energetic barriers exist for rearrangement within the amorphous phase prior to crystallization depending on the differences in molecular association between α-Cry and β-Cry. There are few studies comparing the two polymorphs and amorphous forms. Based on the above observations, we can speculate that a weakening of the hydrogen bonding on progressing from an ordered to a disordered phase is probably more typical for form β than for form α because crystalline β is a stable form, in which hydrogen bonding interactions play a crucial role in the molecular packing.

When the small molecule PEZ was introduced to destroy the order of the crystal structure of AZE, the same peak broadening in the IR spectrum occurred for AZE-PEZ CAP. Furthermore, compared with the crystalline AZE, a downward shift of the C=O vibration and disappearance of the N-H vibration from PEZ for the AZE-PEZ CAP indicated that hydrogen bonding occurred in the AZE-PEZ heterodimer and that the hydrogen bond strength increased in the heterodimer compared to the homogenous dimer. Furthermore, as can be seen in the spectra of the AP and CAP, the vibrations derived from the hydrogen bonding donors and acceptors in the AP and CAP appear at almost the same wavenumbers, suggesting the random weak interactions in the disordered phases are somewhat identical regardless hetero (AP) or homogenous (CAP) combination.

Thermodynamic stability of the AZE-based AP and CAP

Following characterization of the AP and CAP of AZE, we tried to understand the thermodynamic relationships between the different forms of AZE, especially between the AP (or CAP) and crystalline phase. In the present study, solubility measurements were used to determine the transition temperature ($T_{\rm t}$) and relative stability of the AP and CAP. Temperature-dependent solubility data in a 0.01 M HCl medium at 298, 304, 310, 316 and 322 K for the AZE systems were determined and are shown in Table 1. Obviously, α -AP and CAP showed a higher solubility than that of crystalline AZE at the investigated temperatures. Unexpectedly, the solubility of β -AP was lower than that of crystalline AZE in the 0.01 M HCl medium.

Based on the solubility data, curves were drawn with $\ln c$ as the *Y*-coordinate *versus* $1/T \times 10^3$ as the abscissa (Fig. 4). $T_{\rm t}$ can be obtained from the point of intersection of the two curves. The values of $\Delta H^{\theta}_{\alpha\text{-AP}}$, $\Delta H^{\theta}_{\beta\text{-AP}}$ and $\Delta H^{\theta}_{\text{CAP}}$ obtained from the

Table 1 The solubility of the crystalline and amorphous phases of AZE at different temperatures in 0.01 M HCl medium ($\mu g \ mL^{-1}$, n=3, $x\pm s$)

Sample	Temperature (K)						
	298	304	310	316	322		
α-Cry	77.45 ± 2.85	132.38 ± 8.47	170.59 ± 13.30	240.18 ± 11.32	333.21 ± 5.47		
α-AP	122.32 ± 4.01	188.34 ± 9.65	255.86 ± 13.72	310.59 ± 16.11	469.47 ± 9.33		
CAP	117.97 ± 6.54	153.31 ± 3.82	217.76 ± 2.07	263.93 ± 14.35	420.81 ± 30.00		
β-Cry	57.09 ± 0.75	82.49 ± 3.79	100.87 ± 8.62	117.69 ± 5.98	166.47 ± 6.42		
β-АР	16.10 ± 1.22	24.37 ± 2.24	31.06 ± 0.70	38.19 ± 1.51	79.00 ± 5.59		

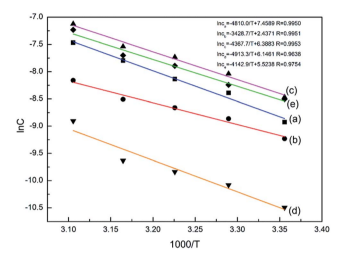


Fig. 4 Curves of $\ln c$ vs. 1000/T for the AZE systems in 0.01 M HCl medium: (a) α -Cry; (b) β -Cry; (c) α -AP; (d) β -AP; (e) α -AZE-PEZ CAP.

slopes of the curves were 36.31, 40.85, and 34.44 kJ mol^{-1} , respectively. Since β-Cry was thermodynamically stable at room temperature,12 we especially discuss the transition reactions of three different transition pairs α -AP/ β -Cry, β -AP/ β -Cry, and CAP/ β -Cry. Based on the solubility data of α-AP, β -AP, CAP and β -Cry, curves were drawn with $\ln c_A/c_B$ as the Y-coordinate versus $1/T \times$ 10³ as the abscissa (Fig. S1†); the corresponding thermodynamic parameters and Tt for the transition reactions are listed in Table 2. The T_t values of the transition pairs α -AP/ β -Cry and CAP/β-Cry were 237.5 K and 231.4 K, respectively, suggesting that α -AP and CAP are stable below those T_t values and β -Cry is stable above those T_t values, for example, at room temperature. While, the pair of β -AP/ β -Cry showed a T_t of 400.3 K, indicating that the stable temperature range of β -AP is below this T_t value. β-AP manifested a broader stable temperature range compared to α -AP and CAP, including room temperature ($\Delta G^{\theta} > 0$ meaning a nonspontaneous reaction of transition from β -AP to β -Cry at room temperature). However, α -AP and CAP were not stable at room temperature, they transformed to β-Cry spontaneously with a value of $\Delta G^{\theta} < 0$ (Table 2).

To verify the stability of the two APs and CAP in 0.01 M HCl solution, the remaining solids were collected and subjected to PXRD analysis after determination of the solubility of the crystalline and amorphous samples at room temperature; the patterns are shown in Fig. 5. For α - and β -Cry, the PXRD patterns showed subtle changes, such as the appearance of

small new peaks, before and after the dissolving-recrystallization process in HCl medium (Fig. 5c and d). This indicated that AZE is ionized partially in a strong acid medium owing to the existence of -N-H in the molecular structure. In the PXRD patterns of the solids from the suspensions after solubility determination of the α -AP and CAP, peaks assigned to β -Cry emerged, indicating that α -AP and CAP are unstable in the 0.01 M HCl medium and transformed to β -Cry at room temperature (Fig. 5e and g). β -AP had maintained its amorphous nature after the solution procedure (Fig. 5f), which is consistent with the results deduced from the $T_{\rm t}$ values that β -AP is stable at room temperature in the 0.01 M HCl medium.

Physical stability

All the APs and the CAP remained physically stable for at least one year at room temperature under dry conditions, which was evidenced by typical amorphous halo XRD patterns and a single $T_{\rm g}$ peak in the TMDSC scans (data not shown). Furthermore, the two APs and the CAP remained amorphous at 40 °C/75% RH during a three-week accelerated experiment (Fig. 6). As mentioned above, the physical stability of the amorphous solid state is relevant to the $T_{\rm g}$. The two APs and the CAP showed a high $T_{\rm g}$ of more than 70 °C, which is far from room temperature, suggesting a high physical stability under storage conditions. So, the CAP, with a high solubility and physical

Table 2 The transition temperatures and thermodynamic parameters of the AZE systems in 0.01 M HCl $\,$

	T(K)	ΔG^{θ} (kJ mol ⁻¹)	ΔH^{θ} (kJ mol ⁻¹)	ΔS^{θ} (J mol ⁻¹ K ⁻¹)	$T_{\rm t}$ (K)
α-AP, β-Cry	298	-1.89	9.24	37.36	237.5
,, ,	304	-2.11			
	310	-2.34			
	316	-2.56			
	322	-2.79			
β-AP, β-Cry	298	3.31	14.91	38.93	400.3
. ,, ,	304	3.07			
	310	2.84			
	316	2.61			
	322	2.37			
CAP, β-Cry	298	-1.61	7.77	31.47	231.4
,, ,	304	-1.80			
	310	-1.99			
	316	-2.18			
	322	-2.37			

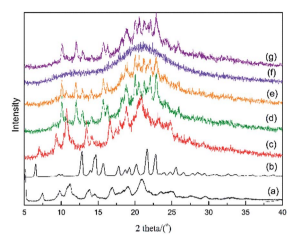


Fig. 5 PXRD patterns of solids from the suspensions after solubility determination in 0.01 M HCl medium at room temperature. (a) Initial α-Cry and (b) initial β-Cry; after solubility determination: (c) α-Cry; (d) β-Cry; (e) α -AP; (f) β -AP and (g) CAP.

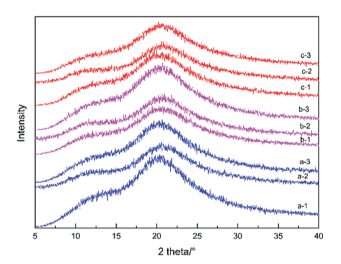


Fig. 6 PXRD patterns of solids from the stability study of the APs and CAP involving 3 weeks of storage at 40 °C/75% RH. (a-1-3) α -AP after 1-3 weeks storage; (b-1-3) β -AP after 1-3 weeks storage; (c-1-3) CAP after 1-3 weeks storage.

storage stability, was proven to be a potential solid form for application.

Kinetics of solid-state decomposition

DSC curves of α -AP, β -AP and CAP at various temperatures are presented in Fig. 7 and 8. Based on the exothermic peak temperature corresponding to decomposition, curves were drawn with 1000/T as the abscissa *versus* $\ln \beta/T_{\rm m}^2$ as the *Y*-coordinate. The thermal decomposition activation energies are listed in Table 3. The difference in the activation energy between the α -AP and β -AP can be obtained from Table 3. As mentioned above for the TMDSC, the α - and β -APs showed minor differences at $T_{\rm g}$ owing to different crystalline starting counterparts and different melting temperatures. So, we

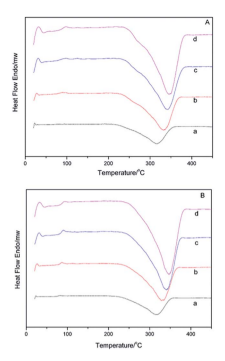


Fig. 7 DSC curves for the α -AP (A) and β -AP (B) systems at different heating rates: (a) 10 °C min⁻¹; (b) 20 °C min⁻¹; (c) 30 °C min⁻¹; (d) 40 °C min⁻¹.

speculated that some different local and short-range order exists in the α - and β -APs, which results in variation in the activation energy of solid-state decomposition. For the CAP, two endothermic peaks with less than a 10% weight loss were

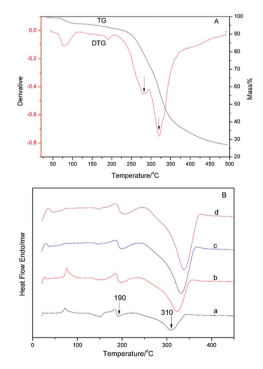


Fig. 8 (A) DTG and TG curves for the AZE-PEZ CAP ($20 \, ^{\circ}\text{C min}^{-1}$) and (B) DSC curves for the AZE-PEZ CAP at different heating rates: (a) $10 \, ^{\circ}\text{C min}^{-1}$; (b) $20 \, ^{\circ}\text{C min}^{-1}$; (c) $30 \, ^{\circ}\text{C min}^{-1}$; (d) $40 \, ^{\circ}\text{C min}^{-1}$.

Table 3 The decomposition temperature at different heating rates and the activation energy of solid-state decomposition for the APs and CAP

Sample	β (°C min ⁻¹)	$T_{ m m}/{ m K}$	$\frac{1/T_{\rm m}}{1000} \times$	$\ln \left(\beta/{T_{ m m}}^2 \right)$	r	E (kJ mol ⁻¹)
α-AP	10	590	1.695	-10.458	0.9874	133.0
	20	605	1.652	-9.816		
	30	615	1.625	-9.443		
	40	617	1.618	-9.164		
β-АР	10	587	1.704	-10.447	0.9944	114.2
	20	601	1.663	-9.802		
	30	614	1.629	-9.439		
	40	620	1.614	-9.170		
CAP	10	583	1.716	-10.43	0.9866	131.6
	20	595	1.682	-9.780		
	30	605	1.653	-9.409		
	40	611	1.636	-9.143		

observed before 200 °C in the DSC (Fig. 8B) and TGA-TG (Fig. 8A) curves, which were due to the loss of absorbed solvent/water and crystal water. The exothermic peaks about 310 °C were due to decomposition of the CAP. No obvious exothermic peak due to decomposition of the PEZ in the CAP appeared in the DSC scan (Fig. 8B), but appearance of a minor peak at 280 °C in the DTG curve indicated decomposition of the PEZ. Here, we focus on the decomposition of the AZE in the CAP, that is, the event corresponding to the peak at 320 °C in the DTG curve. Based on the decomposition temperatures of the AZE in the CAP at different scan rates, the activation energy of solid-state decomposition for the AZE in the CAP was calculated. Interestingly, the activation energies for solid-state decomposition of the α -AP and the AZE in the CAP were almost identical (Table 3). As mentioned above, the AZE-PEZ CAP was prepared using α -Cry as the precursor using a grinding method, while α-AP was obtained through a melting and quench cooling method. It seemed that the remaining AP in the CAP after decomposition of the PEZ had the same local and short-range order as the intact α-AP, because they came from same precursor, α-Cry.

Conclusion

Generally, APs prepared by grinding, unlike those prepared by melting, usually possess characteristics of the original polymorph, that is, "polymorphic memory", which would direct recrystallization toward the original polymorph.²¹ However, in the present study, two different APs were obtained and verified from two different polymorphs of AZE. Although, the two APs were obtained using a melting technique, in which the existing molecular long-range order was destroyed, the differences in the T_g , solubility, stability and thermal decomposition activation energy values of the two APs indicated differences in the local and short-range molecular order of the two APs. This phenomenon probably stems from the major difference in the melting points of the two polymorphs of AZE (129 vs. 199 °C). α -AP, with the highest T_g , was found to be thermodynamically

unstable in the HCl medium, suggesting that no general relationship occurred between the solid physical stability and solution chemical stability for the APs and CAP. The thermodynamic parameters and precipitation tests have demonstrated that the CAP and α -AP of AZE, with higher physical stabilities and solubilities, are interesting options to explore when formulating poorly soluble drugs.

Conflicts of interest

There are no conflicts to declare

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References

- K. Lehmkemper, S. O. Kyeremateng, O. Heinzerling,
 M. Degenhardt and G. Sadowski, *Mol. Pharmaceutics*, 2017,
 14, 157–171.
- 2 H. Meng-Lund, G. Kasten, K. T. Jensen, A. Poso, T. Pantsar, T. Rades, J. Rantanen and H. Grohganz, *Eur. J. Pharm. Sci.*, 2018, **119**, 31–38.
- 3 K. Naelapää, J. P. Boetker, P. Veski and J. Rantanen, *Int. J. Pharm.*, 2012, **429**(1–2), 69–77.
- 4 R. Paus, Y. Ji, F. Braak and G. Sadowski, *Ind. Eng. Chem. Res.*, 2015, 54(2), 731–742.
- 5 B. C. Hancock and G. Zografi, J. Pharm. Sci., 1997, 86(1), 1-12.
- 6 J. Aaltonen and T. Rades, *Dissolution Technol.*, 2009, **16**(2), 47–54.
- 7 B. Hancock and M. Parks, Pharm. Res., 2000, 17(4), 397-404.
- 8 M. Savolainen, A. Heinz, C. Strachan, K. Gordon, J. Yliruusi, T. Rades and N. Sandler, *Eur. J. Pharm. Sci.*, 2007, **30**(2), 113–123.
- 9 R. Surana, A. Pyne and R. Suryanarayanan, *Pharm. Res.*, 2004, 21(7), 1167–1176.
- 10 K. J. Crowley and G. Zografi, J. Pharm. Sci., 2010, 91(2), 492–507.
- 11 K. A. Graeser, C. J. Strachan, J. E. Patterson, K. C. Gordon and T. Rades, *Cryst. Growth Des.*, 2008, **8**(1), 128–135.
- 12 B. C. Hancock and M. Parks, *Pharm. Res.*, 2000, **17**(4), 397-
- 13 K. Lobmann, R. Laitinen, H. Grohganz, K. C. Gordon, C. Strachan and T. Rades, Mol. Pharm., 2011, 8(5), 1919– 1928
- 14 Y. Han, Y. Pan, J. Lv, W. Guo and J. Wang, *Powder Technol.*, 2016, **291**, 110–120.
- 15 Y. Pan, W. Pang, J. Lv, J. Wang, C. Yang and W. Guo, *J. Pharm. Biomed. Anal.*, 2017, **138**, 302–315.

- 16 D. Li, M. Wang, C. Yang, J. Wang and G. Ren, *Chem. Pharm. Bull.*, 2012, **60**(8), 995–1002.
- 17 P. Budrugeac, E. Segal and J. Therm, *Anal. Calorim.*, 2007, **88**(3), 703–707.
- 18 S. J. Dengale, S. S. Hussen, B. S. Krishna, P. B. Musmade, G. G. Shenoy and K. Bhat, *Eur. J. Pharm. Biopharm.*, 2015, **89**, 329–338.
- 19 W. Pang, J. Lv, S. Du, J. Wang, J. Wang and Y. Zeng, *Mol. Pharmaceutics*, 2017, **14**(9), 3013-3024.
- 20 X. C. Tang, M. J. Pikal and L. S. Taylor, *Pharm. Res.*, 2002, **19**(4), 477–483.
- 21 A. R. Sheth, S. Bates, F. X. Muller and D. J. W. Grant, *Cryst. Growth Des.*, 2005, 5(2), 571–578.