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Stability hierarchy between Piracetam forms I, II, and III from experimental pressure–temperature diagrams and topological inferences

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Graphical abstract

Stability domains of the piracetam polymorphs
Under ordinary conditions and with increasing temperature the stable phases are III, II, I, and the liquid
Abstract

The trimorphism of the active pharmaceutical ingredient piracetam is a famous case of polymorphism that has been frequently revisited by many researchers. The phase relationships between forms I, II, and III were ambiguous because they seemed to depend on the heating rate of the DSC and on the history of the samples or they have not been observed at all (equilibrium II-III). In the present paper, piezo-thermal analysis and high-pressure differential thermal analysis have been used to elucidate the positions of the different solid-solid and solid-liquid equilibria. The phase diagram, involving the three solid phases, the liquid phase and the vapor phase, has been constructed. It has been shown that form III is the high-pressure, low-temperature form and the stable form at room temperature. Form II is stable under intermediary conditions and form I is the low pressure, high temperature form, which possesses a stable melting point. The present paper demonstrates the strength of the topological approach based on the Clapeyron equation and the alternation rule when combined with high-pressure measurements.

Keywords: Crystal polymorphism, Solid state, Phase diagram, Phase transition, Physical stability, Preformulation
1 Introduction

1.1 The stability hierarchy of the three solid forms of piracetam

The stability hierarchy of polymorphs is often based on transition temperature measurements by DSC and illustrated by schematic diagrams representing the Gibbs energy as a function of temperature (Buerger, 1951, 1972). However, it may happen that this approach is incomplete and the trimorphism of piracetam is such an example for which the stability hierarchy remains unsolved. It is known for over a decade that forms II and III both transform endothermically into form I on heating; however, the onset temperatures of the two solid-solid transitions at equilibrium are not known, as they appear to depend on the investigator and on the sample used.

Kuhnert-Brandstätter et al. report an energy-temperature diagram in which the transition temperature of form II into form I ($T_{III-II}$) is about 348 K when measured by thermomicroscopy or about 377 K when measured by DSC (Kuhnert-Brandstätter et al., 1994). As the transition temperature of form III into form I ($T_{III-I}$) is about 388 K when measured by DSC, which is higher than $T_{III-II}$, the authors conclude that form III is the more stable phase at room temperature (Kuhnert-Brandstätter et al., 1994). They also mention that “the relative high stability of mod. II can be explained by the small difference of energetic data between mod. II and mod. III.”

Picciochi et al. concluded that “the results of solution and DSC experiments indicate that the stability hierarchy of forms I, II, and III at 298.15 K ... is III > II > I” (Picciochi et al., 2011).

Maher et al. reported a $\Delta G - T$ diagram exhibiting endothermic transitions $II \rightarrow I$ and $III \rightarrow I$ below 363 K and below 368 K respectively, thus leading to the conclusion that form III is the more stable form below 368 K (Maher et al., 2012b).
Céolin et al. reported that the endothermic transitions III→I and II→I occur at 392 K and 399 K respectively, thus inferring that form II is the more stable phase below 392 K (Céolin et al., 1996).

Although a tendency can be observed that form III is the more stable phase at room temperature, the large variation of solid-solid transition temperatures in the different papers warrants an alternative approach to determine the stability hierarchy of the polymorphs without ambiguity. In this paper, it will be demonstrated how the stability hierarchy can be inferred from experimental pressure-temperature diagrams.

1.2 Literature data

1.2.1 Crystallographic data

Three solid forms of piracetam (C₆H₁₀N₂O₂, M = 142.16 g mol⁻¹) have been structurally characterized under ambient conditions (Table 1 and Table S1 in the Supplementary Material), and two more forms as well as two hydrates have been obtained under pressure: form IV (monoclinic, P2₁/c) (Fabbiani et al., 2005), form V (triclinic, P-1) (Fabbiani et al., 2007), a monohydrate (Fabbiani et al., 2005), and a dihydrate (Fabbiani et al., 2007). From the data in Table 1, it can be inferred that form III is the denser form, while form I is the less dense one.

1.2.2 Calorimetric data

The thermal behavior of piracetam polymorphs has been described by several authors (see Table 2). From the data in Table 2, it can be inferred that ΔHII→I is consistently smaller than ΔIII→I by 3 to 6 J g⁻¹. The melting point of form II, TII→I, was reported to be 415.5 K (Kuhnert-Brandstätter et al., 1994) or 415 K (Céolin et al., 1996), but this information should be considered cautiously because it is based on the optical observation of a change in morphology (Kuhnert-Brandstätter et al., 1994) or the
presence of a very small endothermic peak in a few DSC curves (Céolin et al., 1996).

2 Material and Methods

2.1 Samples

Piracetam forms II and III of medicinal grade were obtained from Isochem, France. They were used as such after verification by differential scanning calorimetry and high-resolution X-ray powder diffraction. Form I, which is not persistent at RT, was obtained from either of these polymorphs by heating above the transition temperature and quenching (Céolin et al., 1996).

2.2 Differential scanning calorimetry

Three DSC analyzers were used to determine the temperatures and heats of transition: a TA Instruments (USA) DSC10 Differential Scanning Calorimeter, a Q100 thermal analyzer from TA-Instruments (USA), and a Mettler-Toledo (Switzerland) 822e thermal analyzer. Indium \( T_{\text{fus}} = 429.75 \, \text{K}, \Delta_{\text{fus}}H = 3.267 \, \text{kJ}\cdot\text{mol}^{-1} \) and zinc \( T_{\text{fus}} = 692.68 \, \text{K}, \Delta_{\text{fus}}H = 7.320 \, \text{kJ}\cdot\text{mol}^{-1} \) were used as standards for temperature and enthalpy calibration. Specimens were weighed with microbalances sensitive to 0.01 mg and sealed in aluminum pans.

2.3 High resolution X-ray powder diffraction

Two series of non-ambient X-ray powder diffraction data were obtained for the thermal expansion study.

A Bragg-Brentano optics was used with a Bruker D5005 powder diffractometer, equipped with an Anton Paar high-temperature oven camera HTK1200. Cu \( K\alpha_{1,2} \) X-rays were obtained with a diffracted-beam graphite monochromator. Powders were pressed
on a Ni grid sample-holder in order to reduce transparency effects. Samples of forms II and III were heated above 400 K and upon cooling form I was observed. It must be noted that volume changes during phase transformations may affect the quality of the sample surface (the volumes per formula units are 4.6% and 3.8% greater for form I than for forms III and II, respectively). The powder data were collected at selected temperatures under air. Previous diffraction studies showed a short life time of samples of form I obtained from samples of forms II and III quenched at room temperature (Céolin et al., 1996). To reduce the time required for data collection, the diffraction patterns were scanned over the angular range 10-35° (2θ), with a step size of 0.025° (2θ) and 3 s step⁻¹. Due to the low counting statistics, the accuracy of the results is moderate. The peak positions were evaluated with the software WinPLOTR (Roisnel and Rodriguez-Carvajal, 2001). For each pattern the better defined diffraction lines were selected and twenty peak positions were input in the indexing program DICVOL14 (Louër and Boultif, 2014). The cell parameters are least-squares refined, as well as the zero-point position of the patterns.

A Debye-Scherrer optics was used with an INEL powder diffractometer equipped with a cylindrical position-sensitive detector (CPS120) containing 4096 channels (0.029° 2θ angular step) and monochromatic Cu Kα₁ (λ = 1.54061 Å) radiation. For the measurements as a function of temperature, a liquid nitrogen 700 series Cryostream Cooler from Oxford Cryosystems (UK) was used. Slightly ground specimens were introduced in a Lindemann capillary (0.5 mm diameter) rotating perpendicularly to the X-ray beam during the experiments to improve the average over the crystallite orientations. For the temperature-dependent measurements in the range from 100 K up to the melting point, the sample temperature was equilibrated for about 10 min followed by an acquisition time of ca. 1 hour. The heating rate in between data collection was 1.33
K min⁻¹. The powder data were analyzed with the software FULLPROF using pattern matching (Rodriguez-Carvajal, 1993).

### 2.4 Specific volume of molten piracetam

10.1223 g of piracetam (weighed with a microbalance sensitive to 0.1 mg) was introduced in a cylindrical silica tube with a flat bottom, whose mean inner diameter (8.152 mm) was obtained with a DIGI-MET® 3-point alesometer with a precision of 0.003 mm (see below Table S7, Supplementary Material). After being sealed under vacuum, the tube was suspended in a XU 75/300 oven from Climats (France) whose inner temperature, controlled at ±1 K, was allowed to vary slowly between successive measurements in the 390 - 470 K range. The height of the melt was measured through the window of the door of the oven with a cathetometer sensitive to 2 μm.

### 2.5 Sublimation-condensation experiment

A specimen of form III was placed at one end of a cylindrical silica tube (29 cm length), which was sealed under vacuum. It was introduced in a horizontal furnace so that the specimen-containing end was maintained at about 414 K and the other end was outside the furnace at room temperature. Crystals grew by condensation of the vapor phase within 6 days (Figure 1).

### 2.6 Piezo-thermal analysis (PTA)

An in-house constructed piezo-thermal analyzer, whose operating principle and design have been described elsewhere (Petit and Ter Minassian, 1974; Ter Minassian and Milliou, 1992; Ter Minassian and Pruzan, 1977; Ter Minassian and Toscani, 1999), was used to record the thermal effects associated with phase transitions investigated isothermally as a function of pressure. Pressure was gradually decreased under
isothermal conditions at a rate of 20 or 50 MPa h\(^{-1}\) from a maximum pressure of 600 MPa down to atmospheric pressure. The measurement range for the temperature of the equipment is 298 to 490 K. The weight of the specimens (about 15 mg) was determined with a microbalance sensitive to 0.01 mg. It is important to point out that PTA allows precise entropy change measurements as a function of pressure because the fluxmeter of the device records enthalpy changes at constant temperature.

2.7 **High-pressure differential thermal analysis.**

The transitions in piracetam have been studied with high-pressure differential thermal analysis (HP-DTA). An in-house constructed HP-DTA, similar to the apparatus previously built by Würflinger (Würflinger, 1975) with temperature and pressure ranges from 203 to 473 K and 0 to 300 MPa, respectively, was used. Unweighed samples (about 10 to 50 mg) were sealed in cylindrical tin pans and to ensure that in-pan volumes were free from residual air, specimens were mixed with an inert perfluorinated liquid (Galden® from Bioblock Scientifics, Illkirch, France) before sealing. HP-DTA scans were carried out with a heating rate of 2 K min\(^{-1}\). In addition, DSC experiments at ordinary pressure (i.e., in standard aluminum pans) with mixtures of piracetam and perfluorinated liquid were carried out to verify that the liquid was inert.

3 **Results**

3.1 **Calorimetric data**

Two series of experiments were carried out to investigate whether the transition temperatures II→I and III→I depend on the history of the samples (see details in the Supplementary Material Tables S2 and S3). It can be seen in Figure 2a that the temperature intervals in which the two solid-solid transitions occur fully overlap at the
heating rates of 5 and 10 K min\(^{-1}\).

Hence, aliquots of forms II and III were ground to obtain fine powders with about the same grain size (verified by optical microscopy). Each has been heated at various rates to investigate their dependence on the heating rate and to compare the two samples. From the data, shown in Figure 2b (see also the Supplementary Material Table S2-S5), the following has been concluded: (1) the melting temperature of form I is virtually independent of the heating rate, (2) transitions III→I and II→I are heating-rate dependent, and (3) it is inconclusive which of the two forms transforms at the lowest temperature into form I. It is not uncommon to observe solid-solid transitions that are heating rate dependent (Barrio et al., 2012; Gana et al., 2015; Rietveld et al., 2011b); however a complete overlap in temperature range of two distinct transitions is rare.

A statistical analysis of the transition temperatures for the two solid-solid transitions obtained with heating rates of 5 and 10 K min\(^{-1}\) (see Supplementary Material Tables S2 to S5) is entirely inconclusive as to which of the two forms, II or III, transforms into form I first: \(T_{\text{II→I}}\) 390.62 K and \(T_{\text{III→I}}\) 389.04 K (\(T_{\text{II→I}}\): 12 values, min 372.35 K, max 402.82 K, \(\sigma = 9.4\) K and \(T_{\text{III→I}}\): 18 values min 363.33 K, max 406.86 K, \(\sigma = 12.5\) K). Average values of the transition quantities (compiled in the Tables S2 to S5 in the Supplementary Material) have been reported in Table 3.

Some unground specimens of form III crystals obtained from methanol did not transform to form I but melted. Tentatively, this could be explained by the presence of methanol in the crystal at a level of about 1 % of the initial mass, as determined by thermogravimetric analysis. Thus, methanol as an impurity may be able to stabilize form III up to its melting point. However, this also means that the specific melting enthalpy of form III obtained from these measurements needs to be corrected for the solvent weighed in before the measurement run. Thus, the apparent melting enthalpy for form
III from the measurements is 1 % smaller than the actual value for pure form III. The melting enthalpy has been corrected for this error as reported in Table 3. It is clear that the experimental $\Delta_{III \rightarrow \text{liq}}h$ virtually matches the value $214.62 \text{ J g}^{-1}$ calculated by adding $\Delta_{III \rightarrow \text{liq}}h$ to $\Delta_{I \rightarrow \text{liq}}h$.

### 3.2 Sublimation-condensation experiment

As can be seen in Figure 1, big crystals deposited in the temperature range from 390 down to 360 K, while small crystals had formed in the colder part of the tube below 360 K. X-ray diffraction of ground big crystals (390-360 K range) led to the diffraction pattern of form II. The small crystallites formed below 360 K exhibited a mixed diffraction pattern of forms II and III. Taking the calorimetric measurements into account, it would be expected that above 360 K form I deposits. After the tube is extracted from the furnace and cooled to room temperature, form I will have transformed relatively rapidly to form II. This has also been observed previously by X-ray diffraction (Céolin et al., 1996). The simultaneous observation of form II and form III crystals just above and at ambient temperature demonstrates that any stability difference between them is probably rather small.

### 3.3 Specific volume and thermal expansion of forms I, II, and III

Two series of experiments as a function of temperature have been carried out. The data are compiled in the Supplementary Material Tables S6a and S6b. Using all available data for the specific volumes ($v_i$, $v_{II}$, and $v_{III}$ in cm$^3$g$^{-1}$) as a function of temperature (K), the following three equations were obtained:

$$v_I = 0.7219 + 0.00016026 \ T$$

$$r^2 = 0.960$$

(1)
\[ v_{II} = 0.69661 + 0.00013037 \ T \quad r^2 = 0.992 \]  

\[ v_{III} = 0.69341 + 0.00011618 \ T \quad r^2 = 0.993 \]  

Expressed in expansivities, the expressions lead to \( \alpha_{v,II} = 2.22 \times 10^{-4} \ \text{K}^{-1} \), \( \alpha_{v,III} = 1.87 \times 10^{-4} \ \text{K}^{-1} \), and \( \alpha_{v,III} = 1.68 \times 10^{-4} \ \text{K}^{-1} \). These expansivities are all near the value of \( 2 \times 10^{-4} \ \text{K}^{-1} \) reported by Gavezzotti for organic molecular compounds (Gavezzotti, 2006) and for organic molecular APIs a similar trend has been observed (Céolin and Rietveld, 2015; Rietveld and Céolin, 2015). In addition, it can be seen in Figure 3 that the eqs. 1-3 are in close agreement with volumes obtained from single crystals at 100 and 150 K for all three forms.

### 3.4 Specific volume of molten piracetam as a function of temperature

The specific volume obtained by measurement of liquid piracetam as a function of temperature is presented in Figure 3 (compiled in the Supplementary Material in Table S7). The data is fit to the expression (units as in eqs. 1-3):

\[ v_{\text{liq}} = 0.6291 + 0.0005952 \ T \quad r^2 = 0.994 \]  

(4a)

Molecular organic compounds (including pharmaceuticals) often degrade in the molten state and therefore eq. 4a should be considered as approximate. However, this expression can be compared to the liquid expansion obtained via a different approach. The thermal expansion of the liquid can be estimated assuming that: (1) the specific volume of the melt at the temperature of fusion is in the order of 1.11 times the specific volume of its solid and (2) the mean expansivity \( \alpha_{v,\text{liq}} \) of the melt is \( 1.2 \times 10^{-3} \ \text{K}^{-1} \) (Céolin
and Rietveld, 2010; Céolin and Rietveld, 2015; Mahé et al., 2011; Rietveld and Céolin, 2015). With these two approximations, one obtains the dependence of the specific volume of the melt on the temperature (units as eqs. 1-4a):

\[ \nu_{\text{liq}} = 0.5810 + 0.0006971 T \]  \hspace{1cm} (4b)

This equation leads to values close to the experimental ones in the temperature range from 420 to 500 K (Figure 3).

3.5 Piezo-thermal analysis

The melting pressures \( (P / \text{MPa}) \) of forms I and III have been measured isothermically in decreasing pressure runs for a set temperature in the temperature range of 412 to 443 K together with the transition enthalpy, leading also to the entropy and specific volume changes (see the Supplementary Material Table S8 for numerical data). The \( P,T \) melting curves were fitted to straight lines:

\[ P_{\text{I-\text{liq}}} = -2529 + 5.956 T \quad r^2 = 0.998 \]  \hspace{1cm} (5a)

\[ P_{\text{III-\text{liq}}} = -1878 + 4.555 T \quad r^2 = 0.997 \]  \hspace{1cm} (6a)

When decreasing the pressure in the apparatus from a pressure higher than 200 MPa, the transition III→I was always observed, also for samples that were initially loaded as form II. This indicates that form II transforms into form III under a sufficiently high pressure i.e. \( >200 \) MPa. Measurement runs of form II starting below 200 MPa, in contrast, gave rise to the II→I transition on decreasing the pressure.
Transitions III→I and II→I could be easily identified by their respective enthalpies (see Table 3). The resulting $P$ (MPa), $T$ (K) data are reported in the Supplementary Material Table S9. They were fitted to the following straight lines:

$$P_{\text{III-I}} = -681.2 + 1.886 \, T$$
$$r^2 = 0.987$$

(7a)

$$P_{\text{II-I}} = -899.5 + 2.409 \, T$$
$$r^2 = 0.992$$

(8a)

The enthalpy changes obtained at high pressure (see the Supplementary Material Tables S8 and S9) are similar to the values obtained in a standard DSC apparatus and virtually no change is observed (within the experimental limits).

### 3.6 High-Pressure Thermal Analysis

The melting temperatures ($T/K$) of forms I and III measured at different pressures ($P/\text{MPa}$) are compiled in the Supplementary Material Table S10. Linear fits to the data lead to the following equations:

$$P_{\text{I-Iq}} = -3714 + 8.728 \, T$$
$$r^2 = 0.968$$

(5b)

$$P_{\text{III-Iq}} = -1978 + 4.814 \, T$$
$$r^2 = 0.993$$

(6b)

$$P_{\text{III-I}} = -785.7 + 2.118 \, T$$
$$r^2 = 0.964$$

(7b)

$$P_{\text{II-I}} = -1531 + 3.933 \, T$$
$$r^2 = 0.958$$

(8b)
4 Discussion

4.1 Preliminary phase diagram using only experimental data points

The aim of the discussion is to construct a complete pressure-temperature phase diagram of the trimorphism of piracetam in combination with the stability hierarchy of the different equilibria and phases. In first instance, the experimental $P$-$T$ data will be taken into consideration and this will subsequently be completed through thermodynamic arguments and other data.

For certain high-pressure measurements, different slopes can be observed for the same transition depending on the method used (cf. eqs. 8a and b). Nevertheless, irrespective of the method, the inequalities between the $dP/dT$ slopes of the two-phase equilibria is consistently $dP/dT_{\text{I-liq}} > dP/dT_{\text{II-liq}} > dP/dT_{\text{III-liq}} > dP/dT_{\text{I-II}} > dP/dT_{\text{I-III}}$. This is illustrated in Figure 4 with expressions using the combined data from both high-pressure methods:

$$P_{\text{I-liq}} = -3109 + 7.317 \times T \quad r^2 = 0.914$$  \hspace{1cm} (5c)

$$P_{\text{II-liq}} = -2022 + 4.904 \times T \quad r^2 = 0.990$$  \hspace{1cm} (6c)

$$P_{\text{III-I}} = -676.8 + 1.871 \times T \quad r^2 = 0.977$$  \hspace{1cm} (7c)

$$P_{\text{II-PTA}} = -899.5 + 2.409 \times T \quad r^2 = 0.992$$  \hspace{1cm} (8a)

$$P_{\text{II-HP-DTA}} = -1531 + 3.933 \times T \quad r^2 = 0.958$$  \hspace{1cm} (8b)

The high-pressure results from the two different methods could be smoothly combined
into single expressions with the exception of transition II→I whose HP-DTA curve is steeper and clearly separated from the curve obtained by PTA. Nevertheless, it can be observed that both lines converge at the exact same point (point ‘a’ in Figure 4), where the I-III equilibrium line intersects. It is therefore likely that point ‘a’ is the I-II-III triple point. Because piezothermal analysis is an isothermal method, the temperatures at which transitions are observed are not heating-rate dependent, therefore, it could be suggested that the line ‘a-a₁’ in Figure 4 coincides more accurately with the I-II equilibrium line.

It should also be noted that, although the whole line ‘b-c’ has been identified as the equilibrium I-III through its enthalpy value, once this line intersects the triple point I-II-III (point ‘a’) it must change its stability ranking.

If no hydrostatic pressure is applied (i.e. in a standard DSC) transitions III→I and II→I should ideally be observed at 361.7 K (eq. (7c)) and 373.4 K (eq. (8a)), respectively.

Since curve ‘a-a₁’ in Figure 4 converges with curves ‘a-a₁’ and ‘b-c’ at point ‘a’, it may be tentatively inferred that the influence of the heating rate on the temperature of transition II→I decreases with increasing pressure.

It is also clear that the I-III-liq triple point (where the equilibria I-III, I-liq, and III-liq intersect, point ‘b’ in Figure 4) will be located at a lower temperature than that of triple point I-II-liq (Not shown in Figure 4) because equilibrium I-III crosses equilibrium I-liq at a lower temperature than equilibrium I-II does.

4.2 Triple point I-II-III and equilibrium curve II-III

Triple point I-II-III is the intersection of the I-II and I-III equilibrium curves (point ‘a’ in Figure 4). Thus, by setting eqs. (7c) and (8a), or (7c) and (8b) equal, the coordinates of triple point I-II-III are found to be $T_{I-II-III} \approx 414$ K and $P_{I-II-III} \approx 95.5$ MPa.
Equilibrium II-III must pass through triple point I-II-III too, even though it has not been observed experimentally. To determine the value of its \( \frac{dP}{dT} \) slope, one can make use of the fact that the equalities \( \Delta_{\text{III} \to \text{II} \to \text{III}S} = \Delta_{\text{II} \to \text{III}S} - \Delta_{\text{II} \to \text{I}S} \) and \( \Delta_{\text{III} \to \text{II} \to \text{III}V} = \Delta_{\text{III} \to \text{IV}} - \Delta_{\text{II} \to \text{IV}} \) hold at the triple point I-II-III. Using mean \( \Delta s \) and \( \Delta v \) values calculated with the values compiled in Table S9 in the Supplementary Material, one finds \( \Delta_{\text{III} \to \text{II} \to \text{III}S} = 0.078 - 0.068 = 0.010 \) J K\(^{-1}\)g\(^{-1}\), and \( \Delta_{\text{III} \to \text{II} \to \text{III}V} = 0.0385 - 0.0282 = 0.0103 \) cm\(^3\)g\(^{-1}\), which results in a \( \frac{dP}{dT} \) of 0.97 MPa K\(^{-1}\) through the Clapeyron equation:

\[
\frac{dP}{dT} = \frac{\Delta s}{\Delta v} = \frac{\Delta h}{(T\Delta v)} \tag{9}
\]

Using the triple point as a coordinate on the II-III equilibrium line one finds for its \( P \) (/MPa) - \( T \) (/K) relationship:

\[
P_{\text{II-III}} = -306 + 0.97 \, T \tag{10}
\]

At \( P_{\text{II-III}} = 0 \) MPa, eq. 10 becomes 316 K. This is the temperature for triple point II-III-vap, as the vapor pressure of molecular substances around room temperature generally ranges between a few pascal up to a few hundred pascal and \( P \) in megapascal is approximately zero.

The slope of eq. 10 has been calculated using the entropy obtained by PTA. If instead one uses the mean enthalpy changes obtained in a standard DSC (Table 3), as would be the case for the topological approach, the entropy difference between forms II and III will be 0.0077 J K\(^{-1}\) g\(^{-1}\) at the triple point temperature of 414 K. Using the volume change obtained by X-ray diffraction \( \nu_{\text{II} \to \text{III}} = 0.00907 \) cm\(^3\)g\(^{-1}\) at 414 K (eqs. 2 and 3), the slope \( \frac{dP}{dT_{\text{II-III}}} \) calculated with eq. 9 is 0.85 MPa K\(^{-1}\), i.e. near the value in eq. 10 and clearly
the shallowest slope of all equilibrium curves.

The use of specific volumes obtained at ordinary pressure is validated by a reported
density increase of approximately 5 % between 0 and 0.5 GPa for form II. (Fabbiani et al.,
2007) Triple point I-II-III is found at about 0.1 GPa, thus it can be safely assumed that
the decrease in the specific volumes of the three solid phases is in the order of 1%.

4.3 Triple points I-III-liq and II-III-liq

The coordinates of triple point I-III-liq in the \( P,T \) diagram of piracetam can be most
easily found by setting the equations for the equilibria I-III and III-liq (6c and 7c) equal.
These two equilibria are experimentally better defined than the I-liq equilibrium curve.
The coordinates of triple point I-III-liq are found to be \( T_{I-III-liq} = 443.5 \) K and \( P_{I-III-liq} = 153 \)
MPa.

For the II-III-liq triple point, the intersection of curves II-III and III-liq, the \( P,T \)
coordinates are found by setting equations 10 (II-III) and 7c (III-liq) equal: \( T_{I-II-liq} = 436 \)
K and \( P_{II-II-liq} = 117 \) MPa. This is lower in temperature as well as in pressure than the
coordinates of triple point I-III-liq.

4.4 II-liq melting curve and triple point I-II-liq

The melting point of form II (or triple point II-liq-vap if the melting takes place under
saturated vapor pressure) has not yet been determined convincingly, as a clear melting
peak in the DSC has never been observed. An estimate can be obtained by determining
the slope of the II-liq equilibrium at the triple point II-III-liq (which has just been
obtained) and by extrapolating the obtained expression of the II-liq equilibrium
downwards to its saturating vapor pressure. Approximating the equilibrium curve by a
straight line, \( dP/dT \) can be determined following the same procedure as used above for
the II-III equilibrium line. At triple point II-III-liq, equalities \( \Delta_{II \rightarrow liq}s = \Delta_{II \rightarrow liq}s - \Delta_{III \rightarrow II}s \) and
\[ \Delta_{\text{II} \rightarrow \text{liq}} V = \Delta_{\text{III} \rightarrow \text{II} \rightarrow \text{liq}} V - \Delta_{\text{II} \rightarrow \text{III} \rightarrow \text{liq}} V \] hold. Because both enthalpy and volume changes for the melting of form III appeared to depend slightly on the temperature and pressure (see data in Table S8), their values have been calculated with equations S8a and S8b (derived from Table S8). \[ \Delta_{\text{III} \rightarrow \text{II} \rightarrow \text{III} \rightarrow \text{liq}} S = 0.01 \text{ J K}^{-1} \text{g}^{-1} \] (section 4.2) and \[ \Delta_{\text{III} \rightarrow \text{II} \rightarrow \text{liq}} S = 0.48 \text{ J K}^{-1} \text{g}^{-1} \] (eq. S8a evaluated at triple point II-III-liq) give \[ \Delta_{\text{II} \rightarrow \text{III} \rightarrow \text{liq}} S = 0.47 \text{ J K}^{-1} \text{g}^{-1} \], while \[ \Delta_{\text{II} \rightarrow \text{III} \rightarrow \text{liq}} V = 0.01030 \text{ cm}^3 \text{ g}^{-1} \] (section 4.2) and \[ \Delta_{\text{III} \rightarrow \text{II} \rightarrow \text{liq}} V = 0.10535 \text{ cm}^3 \text{ g}^{-1} \] (eq. S8b evaluated at triple point II-III-liq) result in \[ \Delta_{\text{II} \rightarrow \text{III} \rightarrow \text{liq}} V = 0.09505 \text{ cm}^3 \text{ g}^{-1} \]. Thus, \[ dP/dT_{\text{II} \rightarrow \text{III} \rightarrow \text{liq}} = 0.47/0.09505 = 4.94 \text{ MPa K}^{-1} \], and with \[ P_{\text{II} \rightarrow \text{III} \rightarrow \text{liq}} = 117 \text{ MPa} \] and \[ T_{\text{II} \rightarrow \text{III} \rightarrow \text{liq}} = 436 \text{ K} \], the linear equation for curve II-liq becomes:

\[ P_{\text{II} \rightarrow \text{liq}} = -2040 + 4.94 \, T \] \hspace{1cm} (11a)

This equation gives \[ T_{\text{II} \rightarrow \text{liq-vap}} = 413 \text{ K} \] at \( P = 0 \text{ MPa} \). This value, slightly higher than \[ T_{\text{III} \rightarrow \text{liq-vap}} = 412.34 \text{ K} \] (see Table 3), is in close agreement with the observation of Kuhnert-Brandstätter et al. by optical microscopy that the melting of crystals assumed to be forms II and III occurred at 415.5 \text{ K} and 413 \text{ K}, respectively (Kuhnert-Brandstätter et al., 1994).

The triple point I-II-liq is the intersection of equilibrium curves I-II, I-liq, and II-liq. Setting eqs. 5c and 11a equal for curves I-liq and II-liq, respectively, the triple point coordinates \( T_{\text{I} \rightarrow \text{II} \rightarrow \text{liq}} = 450 \text{ K} \) and \( P_{\text{I} \rightarrow \text{II} \rightarrow \text{liq}} = 182 \text{ MPa} \) are obtained. A second combination of equations 5a (I-liq) and 8a (I-II) from PTA data only leads to the coordinates \( T_{\text{I} \rightarrow \text{II} \rightarrow \text{liq}} = 459 \text{ K} \) and \( P_{\text{I} \rightarrow \text{II} \rightarrow \text{liq}} = 207 \text{ MPa} \), i.e. higher but still comparable to the foregoing results. Finally, using eqs. 8a and 11a (II-liq), \( T_{\text{I} \rightarrow \text{II} \rightarrow \text{liq}} = 451 \text{ K} \) and \( P_{\text{I} \rightarrow \text{II} \rightarrow \text{liq}} = 186 \text{ MPa} \) are found, which is also near the previous values. Moreover in all cases, the triple point coordinates are higher than those of triple point I-III-liq.
4.5 Construction of the complete $P-T$ diagram

A phase diagram involving three crystalline forms possesses ten triple points irrespective of their relative stabilities (Riecke, 1890). Three triple points, I-liq-vap, II-liq-vap, and III-liq-vap, are located on the liq-vap equilibrium, the saturating vapor pressure of the liquid. This equilibrium separates the phase diagram into a high-pressure and a low-pressure part. In the low-pressure domain, three more triple points are located, I-II-vap, I-III-vap, and II-III-vap, all part of the condensed-phase – vapor equilibria. In the high-pressure domain, another four triple points, I-II-III, I-II-liq, I-III-liq, and II-III-liq are located.

4.5.1 Regions involving only condensed phases

Using the results obtained or inferred from PTA and HP-DTA experiments, the diagram shown in Figure 5a represents the high-pressure domain with the condensed phases. It is important to realize that an equilibrium curve passing through a triple point must change its stability level. In fact, around a triple point, the stability ranking must alternate for each equilibrium line, as can be seen for triple point I-II-III, where line 4 below is stable, line 5 is metastable (broken line), line 6 is stable, continuing clockwise, equilibrium 4 is encountered again, which is now metastable etc. This fact, which is due to the intersecting Gibbs energy surfaces of which the pressure-temperature phase diagram is a projection, is called the alternation rule (Oonk, 1981; Oonk and Calvet, 2008). The alternation of the two-phase equilibria around the different triple points can be clearly observed in Figure 5a. Once the stability rank of one of the triple points has been determined, the ranks of the others are defined and therefore also those of the different two-phase equilibria (Figure 5).

The results presented in Figure 5 and obtained by high-pressure measurements and
thermodynamic inferences can be compared to a phase diagram obtained by the
topological method based on the Clapeyron equation and the assumption that all two-
phase equilibria can be represented by straight lines in a $P$-$T$ diagram (Barrio et al.,
2012; Gana et al., 2013; Gana et al., 2012; Perrin et al., 2013; Rietveld et al., 2011a;
Rietveld et al., 2011b).
The $dP/dT$ slopes of the $P$-$T$ melting curves can be calculated using the enthalpy and
volume changes at ordinary pressure ($P = 0$ MPa) from DSC data (Table 3) and specific
volume measurements (equations 1 to 4a). This results in $dP/dT_{l\text{-liq}} = 4.80$ MPa K$^{-1}$ and
$dP/dT_{\text{III-liq}} = 3.89$ MPa K$^{-1}$ and leads to the following equations ($P$ in MPa, $T$ in K, the
subscript $T$ in the equation number indicates “topological”):

$$ P_{l\text{-liq}} = -2037 + 4.80 \, T $$

(5\text{t})

and

$$ P_{\text{III-liq}} = -1604 + 3.89 \, T $$

(6\text{t})

for lines $l$-liq and $\text{III}$-liq, which intersect at $T_{l-\text{III-liq}} = 477$ K and $P_{l-\text{III-liq}}= 252$ MPa. This is
higher than the coordinates obtained through the high-pressure data points (section
4.3); however, it leads to the same topological picture of two converging equilibrium
lines with increasing pressure.

Because the $\text{III}$-$l$ equilibrium curve also passes through triple point $l$-$\text{III}$-liq its equation
($P$ and $T$ in MPa and K respectively)

$$ P_{\text{III}-l} = -315 + 1.187 \, T $$

(7\text{t}),
has been determined using the mean value $\Delta_{III \rightarrow II} h = 27.5 \text{ J g}^{-1}$ (Table 3) and $\Delta_{III \rightarrow IV} = 0.0486 \text{ cm}^3 \text{ g}^{-1}$ at 477 K (eqs. 1 and 3). Eq. 7$\tau$ leads to the transition point I-III at ordinary pressure of $T_{I-III-vap} = 266 \text{ K at } \rho = 0 \text{ MPa}.$

The experimental melting point of the II-liq equilibrium reported in the literature has not convincingly been determined. Nevertheless, if $T_{II-liq} = 413 \text{ K}$ is used together with $\Delta_{II \rightarrow liq} h = 211.5 \text{ J g}^{-1}$ (the algebraic sum $\Delta_{II \rightarrow III} h + \Delta_{III \rightarrow liq} h = 24.4 + 187.1$) and the volume change on melting, $0.1184 \text{ cm}^3 \text{ g}^{-1}$ calculated with equations 2 and 4$b$ (at 413 K), $dP/dT_{II-liq}$ becomes $4.32 \text{ MPa K}^{-1}.$ This leads to the following relation for the melting curve of form II ($P$ in MPa and $T$ in K):

$$P_{II-liq} = -1786 + 4.32 T \quad (11\tau)$$

Although this value of $dP/dT_{II-liq}$ is different from that reported in section 4.4, the topological method leads to the same inequality in the slopes of the melting equilibria (III-liq < II-liq < I-liq).

The coordinates of the “topological” triple points I-II-liq, I-III-liq and II-III-liq calculated with equations 5$\tau$, 6$\tau$ and 11$\tau$ are compiled in Table 4 and compared with the values obtained or inferred from the high-pressure data.

If the literature value of 415 K is used for $T_{II-liq}$, equation 11$\tau$ becomes

$$P_{II-liq} = -1769 + 4.26 T \quad (11\tau')$$

which leads in combination with equations 5$\tau$ and 6$\tau$, to the same inequalities $T_{I-II-liq} > T_{I-III-liq}$ and $P_{I-II-liq} > P_{I-II-liq}$ (Table 4). Clearly the melting point of form II must be located in
between those of forms I and III and it can thus be concluded that topological tools to construct a phase diagram can be a powerful means to verify and complete a phase diagram partially obtained by experimental data.

4.5.2 Triple points involving the vapor phase

Lines 4, 5 and 6 cross respectively triple points I-II-vap, I-III-vap and II-III-vap where the vapor pressure is the saturating vapor pressure of the involved condensed phases at the respective triple points. Their pressure coordinates can be approximately determined with the experimental vapor pressure results for form II obtained by Picciochi et al (Picciochi et al., 2011). These vapor pressures in the temperature range of 356.66 K to 374.44 K have been fitted with the Clausius-Clapeyron equation, leading to the following expression (P in pascal, T in kelvin):

\[
\ln(P_{\text{I-vap}}) = 36.7 - \left(\frac{14143}{T}\right) \quad \quad r^2 = 0.996 \quad \quad (12)
\]

From eq. 12, an average sublimation enthalpy is obtained \(\Delta_{\text{II-vap}}H = 14143 \text{ \, R} \) (the gas constant \(R = 8.3145 \text{ \, J \, K}^{-1}\text{ \, mol}^{-1}\)), which equals 117.6 kJ mol\(^{-1}\) (827.2 J g\(^{-1}\)).

At the temperature of the triple point I-II-vap, 373 K, the vapor pressure of form I equals that of form II. Thus, with \(\Delta_{\text{I-vap}}H = \Delta_{\text{II-vap}}H - \Delta_{\text{II-I}}H = 117592 - 3462 = 114130 \text{ \, J \, mol}^{-1}\), the vapor pressure of form I is given by (P/Pa, T/K):

\[
\ln(P_{\text{I-vap}}) = 35.6 - \left(\frac{13727}{T}\right) \quad \quad (13)
\]

This leads to a vapor pressure of 26 Pa at the melting point of form I or triple point \(T_{\text{I-liq-vap}} = 424.5 \text{ \, K}\).
At the temperature of triple point II-III-vap, 316 K, the vapor pressure of form III equals
that of form II. Thus, with $\Delta_{\text{III-vap}}H = \Delta_{\text{II-vap}}H + \Delta_{\text{II-III}}H = 117592 + 441 = 118033$ J mol$^{-1}$, the vapor pressure of form III is given by $(P/\text{Pa}, T/\text{K})$:

$$\ln(P_{\text{II-vap}}) = 36.9 - (14196/T) \quad (14)$$

The vapor pressure of form I at the temperature $(T_{\text{I-liq}} = 424.46$ K) of triple point I-liq-vap is the same as the vapor pressure of the liquid phase. With $\Delta_{\text{I-liq-vap}}H = \Delta_{\text{I-vap}}H - \Delta_{\text{I-liquid}}H = 114130 - 26598 = 87532$ J mol$^{-1}$, this leads to $(P/\text{Pa}, T/\text{K})$:

$$\ln(P_{\text{I-liq-vap}}) = 28.0 - (10528/T) \quad (15)$$

Eq. 15 gives a vapor pressure of 12 Pa at $T_{\text{II-liq}} = 412.3$ K.

By setting eqs. (12) and (15) equal, the pressure and temperature of the melting triple point of form II can be found: $P = 17.3$ Pa and $T = 418$ K, i.e. greater than $T_{\text{III-liq}}$ and close to the values of 413 K (see above) and 415 K (Kuhnert-Brandstätter et al., 1994) (If $T_{\text{I-liq}} = 413$ K is used, the vapor pressure is found to be 11.7 Pa with equation 12).

The last vapor pressure to be determined is the vapor pressure at triple point I-III-vap, whose temperature was found to be 362 K from PTA experiments. By setting equal equations 13 and 14, the temperature of this triple point (at which the vapor pressure of the two forms should be equal) is found to be 360.8 K (i.e. the same value as 362 K, the difference is due to rounding). Using either of equations 13 and 14, the vapor pressure is
found to be \(\approx 0.1\) Pa, slightly greater than the vapor pressure of form II of 0.093 Pa found with equation 12 (at 362 K), thus indicating that, at 362 K, the I-III-vap triple point is metastable with respect to the II-vap equilibrium. The results can be summarized topologically by representing the exponential curves as straight lines; the relative positions of the triple points involving the vapor phase are depicted in Figure 6.

5 Concluding remarks

Piracetam polymorphism is a case that clearly demonstrates how delicate the determination of the stability hierarchy of polymorphs can be, in particular if only transition temperature inequalities are used. Moreover, when a melting temperature is not known, the case becomes even more difficult.

It has been demonstrated how the stability hierarchy of forms I, II, and III was inferred from experimental results that do not rely on the measured solid-solid transition temperatures by DSC; the latter are heating-rate dependent and randomly distributed in relation to each other. High-pressure measurements (isothermal pressure dependent, and isobaric temperature dependent) provided the I-II-III triple point and the two equilibria II-I and III-I with which the position of the third solid-solid equilibrium III-II could be determined.

The inequalities between the slopes of the solid-solid equilibrium curves involving forms I, II and III were experimentally obtained and obey the sequence \(dP/dT_{I-II} > dP/dT_{I-III} > dP/dT_{III-II}\). These inequalities can also be inferred from calorimetric and volumetric data obtained at “ordinary” pressure, thus without explicit high-pressure measurements. Independent of the temperature coordinate, the inequality in the slopes can be calculated using the Clapeyron equation with the enthalpies of transition in J g\(^{-1}\) from Table 3 (\(\Delta_{I-II}h = 24.4\), \(\Delta_{III-II}h = 27.5\) and inferred \(\Delta_{III-I}h = 3.1\)) and with the
differences in the specific volumes in cm$^3$ g$^{-1}$ at for instance 100 K, 300 K and 500 K using equations 1, 2 and 3. In each case, one finds $T(dP/dT_{I-II}) > T(dP/dT_{I-III}) > T$ ($dP/dT_{II-III}$) (see calculations in Table S11 in the supplementary material). These three slopes will intersect at a single temperature, the triple point I-II-III, which leads to the unambiguous conclusion that the inequality in the temperatures of triple points I-II-vap, I-III-vap and II-III-vap (i.e. at $P = 0$ MPa) must be $T_{I-II-vap} > T_{I-III-vap} > T_{II-III-vap}$, without resorting to the temperatures found by DSC for transitions II → I and III → I.

To conclude, the $P$-$T$ diagram for the phase relationships involving the three solid forms (I, II, and III) of piracetam can be topologically drawn as presented in figure 7a, in which all two-phase equilibria are present, or in figure 7b in which only stable equilibria and subsequent stable phase regions are present. On increasing the pressure, forms I and II both transform to form III, which is the stable phase at low temperature, including room temperature (<316 K), and high pressure. Thus, the conclusion from previous studies that form III is the stable form at room temperature (Kuhnert-Brandstätter et al., 1994; Maher et al., 2012b; Picciochi et al., 2011) has been correct, but for the wrong reasons. Moreover, a stable domain for form II was in most cases overlooked (Figure 7b).

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References


Louër, D., Boultif, A., 2014. Some further considerations in powder diffraction pattern indexing with the dichotomy method. Powder Diffr. 29 (S2), S7-S12.
Petit, J.C., Ter Minassian, L., 1974. Measurements of $\left( \frac{\partial V}{\partial T} \right)_p$, $\left( \frac{\partial V}{\partial p} \right)_T$, and $\left( \frac{\partial H}{\partial T} \right)_p$ by flux calorimetry. J. Chem. Thermodyn. 6, 1139-1152.


**Figure Captions**

**Figure 1.** Sublimation-condensation of piracetam crystals in a silica tube subjected to a temperature gradient for 6 days. The initial sample (form III) was placed in the small container on the left-hand side and maintained at 414 K.

**Figure 2.** Onset temperatures of transitions II→I (solid squares), III→I (open circles) and melting of form I (solid diamonds) (A) of samples obtained by different methods and heated at 5 or 10 K min⁻¹ (see for details Table S2 to S5) and (B) as a function of the heating rate.

**Figure 3.** Specific volumes of piracetam forms I (open circles, eq. 1), II (solid diamonds, eq. 2) and III (solid circles, eq. 3) as a function of temperature from high-resolution X-ray powder diffraction and comparison with volumes from single crystal structures (a: (Fabbiani et al., 2007), b and c: (Chambrier et al., 2011)). The experimental (eq. 4a, open diamonds) and calculated (eq. 4b, dashed line, see text) specific volumes of liquid piracetam are also provided.

**Figure 4.** Pressure-temperature phase diagram of the phase transitions of piracetam with experimental data points only. Melting of form I: squares, melting of form III: diamonds, III→I: plus sign, II→I\text{PTA}: circles, and II→I_{HP-\text{DTA}}: triangles.

**Figure 5a.** Location of the triple points in the $P$-$T$ region involving only condensed phases. Two-phase equilibrium curves: 1: I-liq, 2: II-liq, 3: III-liq, 4: I-II, 5: I-III, and 6: II-III. Filled circles: stable triple points, half-filled circles: metastable triple points. Solid lines: stable equilibria, dashed lines: metastable equilibria, dotted lines: super –

**Figure 5b.** Detail of the P-T diagram with the three triple points along the liq-vap curve (= x-axis): solid circle: I-liq-vap, stable, half-filled circle: II-liq-vap, metastable, open circle: III-liq-vap, super-metastable. The diagram shows the same two-phase equilibrium lines as in Figure 5a. Stable phase regions: see Figure 5a.

**Figure 6.** Topological representation of the two-phase equilibria involving the vapor phase in the $P - T$ diagram of piracetam: 7 = liquid-vapor equilibrium, 8 = I-vapor, 9 = II-vapor, 10 = III-vapor

**Figure 7a.** Complete topological $P - T$ diagram for the phase relationships involving forms I, II and III of piracetam. Two-phase equilibrium curves: same numbering as in Figures 5 and 6. Stability ranking: same as in figure 6.

**Figure 7b.** Topological representation of the stable $P-T$ diagram. The stable phase regions are shown and delimited by the equilibrium curves connecting the five stable triple points (from ten in total). Form III is stable under ambient conditions. Heating the stable form III in the presence of its vapor phase (for example in a DSC capsule) will cause form II to become stable at point ‘j’, 316 K and 0 Pa (= vapor pressure of piracetam, eqs 14 or 12); at point ‘h’, 373 K and 0.3 Pa (eqs 12 or 13), form I will become the more stable form, which melts at point ‘a’, 424.5 K and 26 Pa (eqs 13 or 15) as the liquid becomes the more stable phase. Thus, all three solid forms of piracetam have a stable domain under ordinary conditions (i.e. in the presence of its vapor phase).
Figure 1
Figure 2
Figure 5.a
Figure 5.b
Figure 6
Figure 7.a
Figure 7.b
Table 1. Specific volumes ($v$) from the literature of the three solid forms of piracetam obtained under ambient conditions

<table>
<thead>
<tr>
<th>Form</th>
<th>System, space group</th>
<th>$v$ /cm$^3$g$^{-1}$</th>
<th>$T$ /K</th>
<th>Ref.</th>
</tr>
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<tbody>
<tr>
<td>I</td>
<td>Monoclinic, P2$_1$/n</td>
<td>0.7661</td>
<td>R.T.</td>
<td>(Louër et al., 1995)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.7518</td>
<td>150</td>
<td>(Fabbiani et al., 2005)</td>
</tr>
<tr>
<td>II</td>
<td>Triclinic, P-1</td>
<td>0.7383</td>
<td>R.T.</td>
<td>(Admiraal et al., 1982)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.7094</td>
<td>100</td>
<td>(Chambrier et al., 2011)</td>
</tr>
<tr>
<td>III</td>
<td>Monoclinic, P2$_1$/n</td>
<td>0.7216</td>
<td>R.T.</td>
<td>(Bandoli et al., 1981)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.7321</td>
<td>R.T.</td>
<td>(Admiraal et al., 1982)</td>
</tr>
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<td></td>
<td></td>
<td>0.7245</td>
<td>R.T.</td>
<td>(Galdecki and Glowka, 1983)</td>
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<td></td>
<td></td>
<td>0.7251</td>
<td>290</td>
<td>(Tilborg et al., 2011)</td>
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<tr>
<td></td>
<td></td>
<td>0.7058</td>
<td>100</td>
<td>(Chambrier et al., 2011)</td>
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Table 2. Calorimetric data for piracetam forms I, II, and III from the literature

<table>
<thead>
<tr>
<th>$T_{\text{II-I}}$ /K</th>
<th>$\Delta_{\text{II→I}} h$ /J g$^{-1}$</th>
<th>$T_{\text{III-I}}$ /K</th>
<th>$\Delta_{\text{III→I}} h$ /J g$^{-1}$</th>
<th>$T_{\text{I→liq}}$ /K</th>
<th>$\Delta_{\text{I→liq}} h$ /J g$^{-1}$</th>
<th>$T_{\text{III→liq}}$ /K</th>
<th>$\Delta_{\text{III→liq}} h$ /J g$^{-1}$</th>
<th>Ref.</th>
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<td>382</td>
<td>23.71</td>
<td>393</td>
<td>26.45</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>(Maher et al., 2012a)</td>
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<tr>
<td></td>
<td>23.92</td>
<td>392</td>
<td>26.73</td>
<td>423</td>
<td>179.38</td>
<td>-</td>
<td>-</td>
<td>(Maher et al., 2012b)</td>
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<tr>
<td>348 or 377</td>
<td>21</td>
<td>390</td>
<td>26.7</td>
<td>426</td>
<td>180.7</td>
<td>413</td>
<td>206</td>
<td>(Kuhnert-Brandstätter et al., 1994)</td>
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<tr>
<td>399</td>
<td>24</td>
<td>392</td>
<td>28</td>
<td>426</td>
<td>180</td>
<td>412</td>
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<td>(Céolin et al., 1996)</td>
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<td>26.24</td>
<td>425.9</td>
<td>176.6 or 182.9</td>
<td>-</td>
<td>-</td>
<td>(Picciochi et al., 2011)</td>
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Table 3. Summary of the calorimetric data obtained for forms I, II, and III of piracetam from 5 series of DSC experiments (see Tables S2 to S5 in the Supporting Information)

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<th>Number of values</th>
<th>Lowest value</th>
<th>Highest value</th>
<th>Mean</th>
<th>Standard deviation</th>
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<td>372.9</td>
<td>402.8</td>
<td>390.6</td>
<td>9.4</td>
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<td>( T_{\text{III-I}} / \text{K} )</td>
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<td>363.3</td>
<td>406.9</td>
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<td>12.5</td>
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<td>( \Delta_{\text{II-I}} h / \text{J g}^{-1} )</td>
<td>23</td>
<td>22.9</td>
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<td>0.9</td>
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<td>( \Delta_{\text{III-I}} h / \text{J g}^{-1} )</td>
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<td>25.2</td>
<td>30.1</td>
<td>27.5</td>
<td>1.2</td>
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<tr>
<td>( \Delta_{\text{I-liq}} h / \text{J g}^{-1} )</td>
<td>49</td>
<td>174.9</td>
<td>195.4</td>
<td>187.1</td>
<td>4.3</td>
</tr>
<tr>
<td>( T_{\text{I-liq}} / \text{K} )</td>
<td>49</td>
<td>423.3</td>
<td>425.6</td>
<td>424.5</td>
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</tr>
<tr>
<td>( \Delta_{\text{III-liq}} h / \text{J g}^{-1} )</td>
<td>7</td>
<td>(205.0)</td>
<td>(217.1)</td>
<td>(211.5)</td>
<td>4.4</td>
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<tr>
<td>( \Delta_{\text{I-liq}} h / \text{J g}^{-1} )</td>
<td>7</td>
<td>207.1(^b)</td>
<td>219.3(^b)</td>
<td>213.6(^a)</td>
<td>4.4</td>
</tr>
<tr>
<td>( T_{\text{III-liq}} / \text{K} )</td>
<td>7</td>
<td>411.9</td>
<td>412.8</td>
<td>412.3</td>
<td>0.34</td>
</tr>
</tbody>
</table>

\(^a\)Average for 5 and 10 K min\(^{-1}\) only

\(^b\)Values adjusted for the presence of 1 mass% of methanol in form III, the measured apparent melting enthalpies are provided in parentheses.
Table 4. Triple point coordinates I-II-liq, I-III-liq and II-III-liq determined directly from high-pressure data and by the Clapeyron equation (topological approach)

<table>
<thead>
<tr>
<th></th>
<th>I-II-liq</th>
<th>I-III-liq</th>
<th>II-III-liq</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$P$/MPa</td>
<td>$T$/K</td>
<td>$P$/MPa</td>
</tr>
<tr>
<td>High-pressure data</td>
<td>182</td>
<td>450</td>
<td>153</td>
</tr>
<tr>
<td>Topological data with TII-liq = 413 K</td>
<td>501</td>
<td>529</td>
<td>250</td>
</tr>
<tr>
<td>Topological data with TII-liq = 415 K</td>
<td>361</td>
<td>500</td>
<td>250</td>
</tr>
</tbody>
</table>