Hidden amorphous phase and reentrant supercooled liquid in Pd-Ni-P metallic glasses

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An anomaly in differential scanning calorimetry has been reported in a number of metallic glass materials in which a broad exothermal peak was observed between the glass and crystallization temperatures. The mystery surrounding this calorimetric anomaly is epitomized by four decades long studies of Pd-Ni-P metallic glasses, arguably the best glass-forming alloys. Here we show, using a suite of in situ experimental techniques, that Pd-Ni-P alloys have a hidden amorphous phase in the supercooled liquid region. The anomalous exothermal peak is the consequence of a polyamorphous phase transition between two supercooled liquids, involving a change in the packing of atomic clusters over medium-range length scales as large as 18 Å. With further temperature increase, the alloy reenters the supercooled liquid phase, which forms the room-temperature glass phase on quenching. The outcome of this study raises a possibility to manipulate the structure and hence the stability of metallic glasses through heat treatment.

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d-Ni-P is an excellent glass-forming system, for which centimetre-sized metallic glasses can be made at a cooling rate as slow as 0.17 K s⁻¹ (refs 1–3). In 1976, Chen observed a broad exothermic peak, which is far below the crystallization temperatures in the differential scanning calorimetry (DSC) curves of (Pd₉₀Ni₅₀)₁₀₀₋ₓPₓ (x = 17–19) amorphous alloys. He proposed that this anomalous exothermal temperature, Tₓ, Pd₄₁.₂₅Ni₄₁.₂₅P₁₇.₅ and Pd₄₀Ni₄₀P₂₀. The heating rate was 20 K min⁻¹. The DSC scan of Pd₄₁.₂₅Ni₄₁.₂₅P₁₇.₅, and that the new phase, hereafter referred to as AP, corresponds to crystallization catalysed by amorphous phase separation. However, in situ small-angle X-ray scattering experiments by Yavari et al. could not confirm the development of compositional heterogeneities in the temperature range of interest. Subsequent study using atom probe tomography and element-specific transmission electron microscopy (TEM) also failed to provide evidence of compositional or microstructure modulations before the onset of crystallization. Together, these findings appear to rule out the possibility of nanoscale phase separation. Instead, changes in short-range ordering were suggested as a possible cause for the observed AEP. Indeed, there were hints in early TEM work, which suggested that phase separation was taking place, but at the atomic scale instead of the nanoscale as thought previously. 

A similar DSC anomaly has also been reported in a number of other metallic glass alloys, including Zr-Be-X (X = Ti, Nb)³⁹–¹², Zr-Ti-Cu-Ni-Be¹³,¹⁴, Gd-Zr-Al-Ni¹⁵, Ni-Zr-Nb-Al-Ta¹⁶, Ca-Zr-Al-Y¹⁷, Mg-Cu-Ag-Gd¹⁸,¹⁹, Fe-M-Y-B (M = Mo, W, Nb)²⁰–²², (Fe₀.⁹Co₀.₁)₆₇.₅Nb₄Gd₃.₅B₂₅ (ref. 2³), (Fe₀.₇₅Moₓ₀.₂₅)₆₇.₅Nb₄Gd₃.₅B₂₅ (ref. 2³), (Fe₀.₇₅Moₓ₀.₂₅)₆₇.₅Nb₄Gd₃.₅B₂₅ (ref. 2³), and (Fe₀.₇₅Moₓ₀.₂₅)₆₇.₅Nb₄Gd₃.₅B₂₅ (ref. 2³). Many of these alloys show good glass-forming abilities. Microscopic studies were carried out to investigate a scenario of amorphous phase separation before crystallization, but the results are mostly inconclusive due to, for example, possible artefacts arising from sample preparation. In situ studies are therefore essential to determine the structural origin of the DSC anomaly.

Here, we employ in situ high-energy synchrotron X-ray diffraction, and small-angle neutron scattering (SANS) with simultaneous DSC. Both are bulk techniques; consequently, sample preparation is not an issue and the data interpretation is straightforward. The scattering results were further checked and reconciled with TEM analysis. Our data for Pd-Ni-P glass alloys showed a change that is in perfect correlation with the DSC data, revealing a hidden amorphous phase, the associated polymorphic or liquid-to-liquid phase transition (LLPT) and a reentrant behaviour in the supercooled liquid region at higher temperatures. Although polymorphous phase transitions have been extensively studied, the experimental evidence has been rare and sometimes controversial. Evidence for LLPT has been reported in triphenyl phosphate, confined water and in metallic glasses close to the melting temperature. Polymorphic transition has been previously observed in metallic glasses as well, but under pressure. Nevertheless, the origin of polymorphous phase transition is far from clear, and has been a subject of intense debate. Our studies of Pd-Ni-P alloys show that at the transition temperature, the underlying atomic structure undergoes significant changes over the medium-range length scale, while the short-range order is little changed.

**Results**

The anomalous exothermal peak. Figure 1 shows the specific heat capacity, C_p, for two bulk metallic glass (BMG) samples, Pd₄₁.₂₅Ni₄₁.₂₅P₁₇.₅ and Pd₄₀Ni₄₀P₂₀. The heating rate was 20 K min⁻¹. The DSC scan of Pd₄₁.₂₅Ni₄₁.₂₅P₁₇.₅ exhibits an AEP at Tₓ = 612 K, which is ~41 K below the crystallization temperature, Tₓ = 653 K. The integral area for the AEP is 10.6 ± 0.1 J g⁻¹. This value can be compared to the total integral area of the crystallization peak which is 101.₄ ± 0.1 J g⁻¹. 

![Figure 1 | Specific heat (C_p) for as-cast BMGs on heating.](image)

The large exothermal peak at Tₓ is strongly indicative of a thermodynamic phase transition. Notice that the C_p for Pd₄₁.₂₅Ni₄₁.₂₅P₁₇.₅ approaches zero at Tₓ = 612 K, highlighted by the shadow, which is ~41 K below the crystallization peak at ~653 K. The C_p data for Pd₄₀Ni₄₀P₂₀, downward shifted by 80 J mol⁻¹ K⁻¹, are characteristic of a normal BMG, marked by a Tₓ and a crystallization peak; there is no DSC anomaly before crystallization. For both samples, the Tₓ ~ 570 K.

In situ synchrotron diffraction. Figure 2a,b show the evolution of X-ray structure factor, S(Q), for Pd₄₁.₂₅Ni₄₁.₂₅P₁₇.₅ and Pd₄₀Ni₄₀P₂₀ alloys, obtained by in situ high-energy synchrotron X-ray measurements, on heating the samples from room temperature to ~643 K. To detect the subtle structural changes during heating, the S(Q) at T = 303 K was subtracted, and the difference curves, ΔS(Q), are plotted in Fig. 2c,d, respectively. For Pd₄₀Ni₄₀P₂₀, the difference curves display a normal change with increasing temperature, in which the diffraction peaks shift to lower Q values accompanied by peak broadening and a decline in peak intensity. For Pd₄₁.₂₅Ni₄₁.₂₅P₁₇.₅, however, unusual changes can be readily identified in the vicinity of Tₓ. For example, on heating, the Q₁ peak initially broadens, but above Tₓ ~ 570 K, Q₁ begins to sharpen until the temperature reaches Tₓ ~ 612 K, and then resumes broadening at higher temperatures. This is evidenced by the development and disappearance of an extra peak at the position of Q₂ in the ΔS(Q) curves (Fig. 2c). These anomalous structure changes point to a phase transition in Pd₄₁.₂₅Ni₄₁.₂₅P₁₇.₅ and that the new phase, hereafter referred to SCL2, is metastable.

To correlate the observed anomalies in structure and thermodynamics, the evolution of the Q₂ peak intensity (integrated over a Q range of 4.98–5.00 Å⁻¹) and the C_p data are plotted together in Fig. 3. For Pd₄₁.₂₅Ni₄₁.₂₅P₁₇.₅, the intensity of Q₂₁ shows a rise and fall through TₓC, which correlates perfectly with the changes seen in C_p. For Pd₄₀Ni₄₀P₂₀, on the other hand, the intensity for Q₂₁ shows a continuous decline with temperature, except for a point of inflection at T_p. Subtle changes at other Q values can also be seen, showing similar temperature dependence. An example is shown for peak Q₁ in Supplementary Fig. 1. The position of Q₁ shows consistent changes, see Supplementary Fig. 2 and Supplementary Note 1.

The amorphous hidden phase and reentrant behaviour. Note that no crystalline peaks were detected in the X-ray diffraction...
patterns, even in the $\Delta S(Q)$ plots, which are sensitive to structure changes as low as $\approx 10^{-6}$ volume fraction transformed. High-resolution TEM was employed to confirm the amorphous nature of the new phase in the vicinity of $T_C$. Figure 3c shows a double Cs-corrected TEM image taken on a Pd$_{41.25}$Ni$_{41.25}$P$_{17.5}$ BMG, quenched from $T_C \approx 612$ K. The TEM image is typical of an amorphous alloy: uniform and maze-like. This result confirmed that the Pd$_{41.25}$Ni$_{41.25}$P$_{17.5}$ sample remained in an amorphous state at $T_C$.

At $T > T_C$, the extra intensities in the $\Delta S(Q)$ curves diminished with increasing temperature and disappeared entirely at $T = 643$ K. At this point the $S(Q)$ reverted back to the shape at $T = 575$ K. A high-resolution TEM was taken on a Pd$_{41.25}$Ni$_{41.25}$P$_{17.5}$ sample quenched from 643 K, after the transformation at $T_C$ ended; again, an amorphous structure is observed. The inset of Fig. 3d shows the selected area electron diffraction (SAED) pattern, further confirming the amorphous nature of the alloy at this temperature. These observations suggest that at $T > T_C$, after the SCL2 phase dissolves, the Pd$_{41.25}$Ni$_{41.25}$P$_{17.5}$ BMG reenters the supercooled liquid of the room-temperature glass phase.

Simultaneous SANS-DSC. SANS measurements, which are sensitive to composition or density fluctuations on the nanometer scale, provided further evidence of a reentrant supercooled liquid phase at $T > T_C$. The Pd$_{41.25}$Ni$_{41.25}$P$_{17.5}$ sample was heated, at a heating rate of 2.5 K min$^{-1}$, in a DSC furnace installed on the SANS instrument. SANS data were recorded in situ over a $Q$ range of 0.0065–0.1000 Å$^{-1}$. The integrated SANS detector count rate is plotted in Fig. 4a, along with the DSC data. The AEP is readily identified in the simultaneous DSC data, although the peak temperature $T_C$ is slightly lower due to the slower heating rate. The SANS intensity shows a rise and fall through $T_C$, once again demonstrating the development and disappearance of the SCL2 phase as the BMG sample proceeded through the phase transition. At $T = 623$ K, the SANS intensity dropped to the level at $T_p$, showing that the metastable SCL2 phase had dissolved and the phase transition was complete. The sample was then quenched to room temperature to retain its phase at $T = 623$ K (S1). Full $Q$ range (0.003–0.727 Å$^{-1}$) SANS data were collected and compared with that of an as-cast sample (C) (see Fig. 4b). Their SANS profiles almost overlapped completely with each other, indicating that samples C and S1 are identical, from both chemistry and density perspectives. This observation reinforces our conclusion that after the SCL2 phase dissolves, the Pd$_{41.25}$Ni$_{41.25}$P$_{17.5}$ sample indeed reenters the high-temperature supercooled liquid phase, which on quenching forms the room-temperature glass phase. In comparison, a crystallized sample was also measured and the results were superimposed in Fig. 4b. The crystallized sample shows significant SANS intensity over a $Q$ range of 0.002–0.02 Å$^{-1}$. At lower $Q$ values, the SANS intensity is lower, suggesting some growth of the nanoscale inhomogeneities in the as-cast samples at high temperatures when the sample crystallized.

Structure analysis in real space. Having established the hidden amorphous and the reentrant behaviour, we turn to real-space structure analysis to obtain atomistic insights of the structure evolution through the transition. The pair-distribution functions were obtained by Fourier transform of the structure factors. The reduced pair-distribution function, $G(r)$, is presented in Fig. 5a for selected temperatures through $T_C$. Clear changes can be seen in the differential $G(r)$ obtained by subtracting the reference data set at $T = 303$ K (see Fig. 5b). The first shell, R1 or the short-range order, shows a small change through the transition. Much more pronounced changes are seen in higher-order shells. For example, the intensity of the fifth shell, R5, clearly shows a rise and fall with

Figure 2 | In situ X-ray diffraction data. Evolution of synchrotron X-ray diffraction patterns for (a) Pd$_{41.25}$Ni$_{41.25}$P$_{17.5}$ and (b) Pd$_{40}$Ni$_{40}$P$_{20}$ BMGs during heating at a constant rate of 20 K min$^{-1}$. The structure factor is characterized by a first sharp diffraction peak $Q_1$, followed by the second diffraction maxima $Q_2$, and a shoulder $Q_{2.5}$. (c,d) are the respective difference plots obtained by subtracting the diffraction pattern at $T = 303$ K.
increasing temperature. In Fig. 5c, the integrated intensity of \( G(r) \) over the first and fifth shells are plotted for comparison (over comparable ranges, that is, \( \pm 20\% \) of the initial values). The intensity of R5 increases dramatically as \( T \) approaches \( T_C \), and starts to decrease at \( T < T_C \), and then returns to the level before AEP. The intensity of R1 shows similar, but much more subdued, temperature dependence. These results demonstrate that the phase transition through \( T_C \) is driven by the medium-range order; the short-range order is hardly changing. As a comparison, the temperature dependence of the \( G(r) \) intensity for Pd40Ni40P20 BMGs is shown in Fig. 5d. In metallic glasses, the first shell or short-range order is characterized by solute-centred clusters, while the higher-order shells describe the packing or connectivity of the clusters. Thus, the change of \( G(r) \) in the higher-order shells (for Pd41.25Ni41.25P17.5 alloys), which can be seen at a distance as large as 18 Å in Fig. 5b, indicates a rearrangement of the atomic clusters over the medium-range length scales. The characteristic temperature dependence of the intensity, that is, the rise and fall, confirms the reentrant behaviour.

**Figure 3 | Evidence for the hidden amorphous phase.** Temperature dependence of the \( Q_{21} \) peak intensity (integrated over a \( Q \) range of 4.98-5.00 Å\(^{-1}\) ) versus \( C_p \) for (a) Pd41.25Ni41.25P17.5 and (b) Pd40Ni40P20 BMGs. The \( T_g \) and \( T_C \) are marked by the vertical dashed lines. The solid lines are the results of a smooth spline fitting, which serve as a guide to the eyes. (c) Double Cs-corrected TEM image of Pd41.25Ni41.25P17.5 on a sample quenched at \( T_R \); scale bar, 5 nm. (d) High-resolution TEM of Pd41.25Ni41.25P17.5 on a sample quenched at 643 K, after the transition has ended; scale bar, 5 nm. The inset shows the SAED pattern; scale bar, 10 nm\(^{-1}\).

**Figure 4 | Simultaneous SANS-DSC data for Pd41.25Ni41.25P17.5.** (a) The integrated detector counting rates as a function of temperature. The simultaneous DSC scan is superimposed, which shows a \( T_C \sim 594 \) K at a heating rate of 2.5 K min\(^{-1}\). The coexistence of two phases can be clearly seen in the vicinity of \( T_C \). (b) Full-\( Q \)-range SANS data for three samples: as-cast condition (C), quenched from 623 K after the transformation had ended (S1), and quenched after crystallization at 673 K (X). The nearly complete overlap of the SANS profiles between the as-cast and S1 samples demonstrates the reentrant behaviour.
A systematic DSC study was completed for a series of Pd-Ni-P samples with varying degrees of P composition. Figure 6 shows a diagram illustrating the hidden phase and the sequence of phase transitions, constructed from characteristic temperatures determined from DSC scans (Supplementary Fig. 3). A sub-$T_g$ anomaly has been reported by Hu et al., with a strong cooling rate effect in that a hyperquenched glass exhibits a sub-$T_g$ exothermal peak, whereas a regular glass does not. Although this sub-$T_g$ exothermal peak is not directly related to our DSC anomaly (which is above $T_g$), it points out the clear contrast for samples prepared by different cooling rates. We have also studied the effect of cooling rate on AEP, and the results are shown and discussed in Supplementary Fig. 4 and Supplementary Note 2; the influence of cooling rate can be clearly seen. For clarity, we have specifically noted the applicable heating and cooling rates in Fig. 6, to indicate the dynamic nature of the diagram.

Two supercooled liquid phases are identified in Fig. 6. SCL1 is the high-temperature supercooled liquid phase. The room-temperature glass phase (G1) is the dynamically arrested or frozen state of SCL1. The region bounded by the dark turquoise lines is the low-temperature supercooled phase, SCL2, which is usually bypassed in as-cast samples due to rapid quenching. Heating the sample brought the hidden SCL2 phase to light.

**Discussion**

The energy landscape, a potential energy surface as a function of particle coordinates, offers a convenient way to consider and visualize the structural evolution in the supercooled liquid. Examination of Fig. 6 suggests a potential energy landscape depicted in Fig. 7. There are three deep minima or ‘megabasins’ in the potential energy landscape: two supercooled liquid phases (SCL1 and SCL2), and a crystalline phase (X). The crystalline phase has the deepest energy minima, but it is difficult to reach
enthalpy or the potential energy and ordering of a ferromagnet, where the paramagnetic phase at high configurations than SCL2. This scenario is analogous to the more disordered state SCL1 is illustrated with many more local entropy term. In Fig. 7, on the potential energy landscape, the transition (say important to consider the entropy term in the Gibbs free energy.

Experimental evidence has shown that when \( T > T_C \), the sample reenters into the SCL1 phase. This means that at high temperatures, SCL1, although of higher potential energy, is the thermodynamically stable state. To understand this paradox, it is important to consider the entropy term in the Gibbs free energy.

The Gibbs energy is given by \( G = H - TS \), where \( H \) is the enthalpy or the potential energy and \( S \) the entropy. Consider the Gibbs energy at a temperature \( T_1 \) just below the phase transition (say \( T = 570 \) K) and at \( T_C \), \( G_1 = H_1 - T_1 S_1 \) and \( G_C = H_C - T_C S_C \). Since SCL1 is the stable phase at \( T_1 \), \( G_1 < G_C \). However, \( H_1 > H_C \), because there is heat release on transforming to SCL2 as evidenced from the \( C_p \) data. Therefore, \( T_1 S_1 > T_C S_C \). It follows that \( S_1 > S_C \), because \( T_1 \) is lower. This analysis shows that SCL1 is more disordered, and that SCL1 is stabilized by the entropy term. In Fig. 7, on the potential energy landscape, the more disordered state SCL1 is illustrated with many more local configurations than SCL2. This scenario is analogous to the ordering of a ferromagnet, where the paramagnetic phase at high temperature is stabilized by the entropy of disordered spins.6

The presence of ‘megabasins’ had been previously discussed by Debenedetti and Stillinger. By comparing molten SiO2, a prototypical strong glass former, and o-terphenyl, a fragile glass former, they argued that the potential energy landscape of strong glass formers may consist of a single ‘megabasin’, whereas the fragile ones display a proliferation of well-separated ‘megabasins.’ Pd-Ni-P samples are known to form a fragile glass. In this regard, the potential energy landscape depicted in Fig. 7 is consistent with the general analysis by Debenedetti and Stillinger.

Structure wise, the phase transitions at \( T_C \) is accomplished through rearrangement of locally favoured structures, or atomic clusters. Figure 5 shows that the cluster rearrangement occurs most prominently at medium-range length scales, at the fifth shell \( (r = 11 \) Å) or longer. Zheng et al. observed three exothermal responses (peaks) in glassy ribbons of Cu42Zr42Al16Y5. Based on DSC analysis, they attributed the second peak to the formation of ordered clusters. This work is consistent with an earlier study by Kumar et al., who speculated that the anomalous DSC in Zr63Ti15Be40 could be due to development of short-range order towards the final crystallization product. The high-quality synchrotron data obtained in the present experiment allowed us to visualize the dramatic changes in medium-range order, at the fifth shell (R5) centred around \( r = 11 \) Å.

As stated earlier, the AEP is not unique to Pd-based BMGs; it has been found in a variety of metallic glass systems as well, although the effects are less pronounced and the interpretations vary widely. More interestingly, a systematic study has found a link between the AEP and glass-forming ability (GFA). This was demonstrated for Fe-based BMGs, which have broad applications in electricity delivery due to their exceptional soft magnetic properties. Fe-based BMGs with an AEP tend have better GFA than those without. Other BMGs with an AEP are reported to have improved plasticity. Here again, no evidence of nanoscale phase separation was found. Given the similarity, it is likely that the DSC anomalies observed in a variety of BMGs arise from the same structural origin, that is, an LLPT in the supercooled liquid region like the one found in Pd-Ni-P.

The GFA of a metallic glass is influenced by a number of factors. It is encouraging that in alloys exhibiting an AEP, there appears to be a correlation between GFA and the AEP. This finding can potentially provide a viable method to tune the GFA. However, additional studies (for example, \textit{ab initio} molecular dynamics simulations) will be required to fully establish the relationship and to clarify the underlying mechanisms.

In conclusion, we reveal a hidden amorphous phase and the associated LLPT in a Pd-Ni-P supercooled liquid. This finding explains four decades long puzzle about the thermodynamic anomaly in DSC scans. At the transition temperature, \( T_C \), the underlying atomic structure undergoes significant changes over the medium-range length scale, as evidenced by \textit{in situ} synchrotron diffraction data. In addition, the SANS data demonstrated the coexistence of two supercooled liquid states in the vicinity of \( T_C \). While the focus of the present study is on Pd-Ni-P alloys, the DSC anomaly has been reported in a number of other BMGs. That the appearance of this DSC anomaly has been linked to the excellent glass-forming ability or GFA in Fe-based BMGs raises a new possibility to control the GFA and properties of BMGs. Because \( T_C \) is rather low, close to \( T_g \) as opposed to the melting temperature, it is conceivable that conventional metallurgical methods, such as heat treatment, could be employed as a means to manipulate the phase and microstructure of the metallic glass of interest.

\section*{Methods}

\subsection*{Sample preparation}

A fluxing technique was employed to prepare Pd41.25Ni41.25P17.5 and Pd60Ni40P20 BMG ingots of diameter \( \approx 10 \) mm with air quenching. Pd-Ni-P BMG ingots were cut into small pieces of mass \( \approx 43 \) mg for heat capacity measurements. Heat treatment of Pd41.25Ni41.25P17.5 MGs around \( T_C \) is described in Supplementary Methods. The synchrotron X-ray diffraction results for the heat-treated Pd41.25Ni41.25P17.5 MGs are given in Supplementary Fig. 5.

\subsection*{Heat capacity measurements}

The specific heat capacity curves during heating were measured with reference to a sapphire standard using platinum crucibles with Netzsch DSC 404 F3 on \( C_p \) measurement mode in a high-purity Ar atmosphere.

\subsection*{In situ synchrotron diffraction measurements}

Time-resolved high-energy synchrotron X-ray measurements during constant heating were carried out on the beamline 11-ID-C at the Advanced Photon Source, Argonne National Laboratory. BMG plates with \( \approx 2 \) mm thickness were mounted in a Linkam THMS600 furnace for thermal scanning. High-energy X-rays with a beam size of \( 0.5 \) mm x \( 0.5 \) mm and wavelength of \( 0.18084 \) Å were used in transmission geometry for data collection. The measurement was conducted in a high-purity argon atmosphere. Two-dimensional diffraction patterns were obtained using a Perkin-Elmer amorphous silicon detector. The data acquisition time for each pattern is 1 s. However, including the time for saving data, the total acquisition time is \( \approx 6 \) s for each pattern. At a constant heating rate of \( 20 \) K min\(^{-1}\), the temperature precision for each pattern is estimated to be \( \approx 1-2 \) K. The static structure factor, \( S(Q) \) with \( Q_{\text{max}} \approx 30 \) Å\(^{-1}\), was derived from the scattering data by masking bad pixels, integrating images, subtracting the appropriate background and correcting for
oblique incidence, absorption, multiple scattering, fluorescence, Compton scattering and Laue correction using Fit2D and PDFgetX2. The details of condensation data reduction can be found in Supplementary Methods and Supplementary Figs 6 and 7.

Microscopy analysis. The double Cs-corrected TEM Titan3 G2 60–300 at Hokkaido University was employed to acquire high-resolution TEM images shown in Fig. 3c for a Pd$_{41.25}$Ni$_{41.25}$P$_{17.5}$ BMG air quenched from ~T$_c$ in SCL1. The high-voltage electron microscopy TEM-ARM 1300 with an accelerating voltage of 1.250 kV and Hokkaido University was used to acquire high-resolution TEM images and the SAED pattern for a Pd$_{65.21}$Ni$_{13.75}$P$_{21.04}$ BMG air quenched from ~643 K in SCL1.

**Time-resolved SANS-DSC measurements.** Simultaneous SANS-DSC studies during constant heating with a rate 2.5 K min$^{-1}$ were performed on the QUOKKA SANS instrument at the OPAL reactor, Australian Nuclear Science and Technology Organization (ANSTO)$^{48}$. A neutron beam with diameter of 5 mm and wavelength, $\lambda$, of 5 Å (10% resolution) was used. A DSC and Al crucibles designed by QUOKKA instrument group$^{50}$ were employed. The BMG plates with ~0.9 mm thick and with ~200 mg mass were used for simultaneous SANS-DSC studies. Three configurations were employed for room-temperature measurements covering a Q range 0.003–0.727 Å$^{-1}$, where Q is the magnitude of the scattering vector defined as $Q=\frac{4\pi}{\lambda}sin(\theta)/2$, and $\theta$ is the scattering angle. The configuration for in situ SANS-DSC study covers a Q range 0.0065–0.1 Å$^{-1}$. SANS data were reduced with data corrected for empty cell scattering, transmission and detector response and transformed onto an absolute scale by the use of an attenuated direct beam transmission measurement.

**Data availability.** The data that support the findings of this study are available from the corresponding author on request.

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Author contributions
S.L. and X.-L.W. designed the project; S.L. and Y.R. performed the in situ synchrotron experiments; S.L., X.Y.W., B.W. and E.P.G. performed the simultaneous SANS and DSC measurements; S.L., X.Y.W., T.S., S.W. and M.O. performed the TEM study; all authors analysed and reviewed the results, S.L. and X.-L.W. wrote the paper; X.-L.W. supervised the overall research activities.

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