Amorphous alloys and differential scanning calorimetry (DSC)

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Received: 2 November 2020 / Accepted: 21 August 2021 / Published online: 18 September 2021 © The Author(s) 2021

Abstract
A remarkable number of scientific papers are available in the literature about the bulk amorphous alloys and metallic glasses. Today, DSC is an essential tool for amorphous alloys research and development, and of course for quality assurance. In many cases, users seek to examine the determination of only one or two properties, although much more information can be obtained from the measurements. The research involved structural relaxation, Curie temperature, glass temperature, crystallization, phase separation, nanocrystalline volume fraction, melting point and liquidus temperature determination subjects and kinetics of microstructural transformations induced by thermal treatment. We collected and present the information that can be obtained with this technique and draws the reader’s attention to some potential problems related to data interpretation.

Keywords Thermal analysis · Amorphous alloy · Glass temperature · Phase separation · Liquidus temperature

Introduction
In the winter of 1782-83, Antoine Lavoisier and Pierre-Simon Laplace made the first known calorimeter, an ice-calorimeter [1]. Ever since, thermal analysis techniques, and DSC in particular, have been playing a significant role in the research and development in both non-metallic and metallic materials. Calorimetry has many advantages. Many changes in properties or structures in a material are accompanied by the liberation or consumption of energy. This energy is usually of sufficient magnitude and rate to be easily detected. Thanks to its speed, simplicity, and availability, DSC has become a universally used standard tool for characterizing thermodynamic and kinetic properties, phase transitions, and property evolution. Today DSC is the most often used thermal analysis method. Furthermore, DSC is the most commonly used device for the characterization of polymers, drugs, foods, geological materials, and metallic materials as well.

There are many types of DSCs today:

- Heat flux DSC
- Power compensated DSC
- Modulated DSC
- Pressure DSC
- Differential fast scanning calorimetry (DFSC)

The most frequently used DSCs are the heat flux and power compensated ones. The principle of heat flux DSC is that the difference in heat flow into the sample and the reference is measured, while the temperature of the sample is changed at a constant rate. The heat flow difference between the sample and the reference is:

$$\Delta \frac{dH}{dt} = (\frac{dH}{dt})_{sample} - (\frac{dH}{dt})_{reference}$$

The heat flow difference between the sample and the reference can be either positive or negative. In an exothermic process, like a crystallization, heat is released, and, therefore, heat flow to the sample is lower than that to the reference.
In an endothermic process, such as melting, the opposite is true: the heat flow is negative.

In power compensated DSC, the sample and the reference are separated into two furnaces and the temperatures of the sample and the reference are kept equal to each other, while both temperatures are increased or decreased linearly. The electric power needed to maintain equal temperatures for both the sample and the reference temperature is measured throughout the entire analysis.

For both types of equipment, the reference is an inert material such as alumina, or just an empty aluminum pan.

The results of the measurement are affected by calibration, sample pan, sample, experiment and environmental conditions, and most importantly the interpretation of the results [2–4].

The first metallic glass was published 60 years ago [5], since then one of the most important testing techniques of amorphous alloys is DSC. Numerous papers and reviews deal with the questions of amorphous alloys and their thermal behaviours [6–8]. Many glass-forming ability (GFA) factors are published in the literature. There are no rules on how to measure the thermal properties of amorphous alloys is missing in the literature. There are no rules on how to measure and evaluate the measurement curves for each property of amorphous alloys.

The goal of this paper is to show, what can be deduced from the measured results on amorphous materials from a process engineering point of view.

Materials and methods

By means of the standard procedure of rapid quenching of the melt on a rotating wheel (melt-spinning method), amorphous ribbon samples were prepared. The nominal composition of the as-prepared alloy samples can be represented as follows in atomic %: Fe_{48}Ni_{40}B_{16}Si_{4} and (Cu_{48}Zr_{48}Al_{4})_{96}Co_{2}Fe_{2}. The thickness of the ribbons was approximately 50 μm. FINEMET type alloy ribbons with Fe_{73}Si_{16}B_{7}Nb_{3}Cu_{4} nominal composition are commercial products.

The master alloy ingots were prepared by arc-melting from a mixture of high-purity elements under argon atmosphere (min. 99.9 mass%) with a Ti-getter. The ingots were re-melted at least four times in order to ensure chemical homogeneity. Bulk amorphous samples of Cu–Zr–Ag–Al system in a wedge and rod form have been prepared by the centrifugal casting in a copper mould. The ingots were re-melted by induction melting in a quartz crucible under argon atmosphere in the centrifugal casting machine. Before melting the chamber was evacuated and purged with high-purity Ar. The dimensions of the resulting wedge are: 30 mm high, 3 mm thick, 20 mm wide. The dimensions of the rod are 25 mm length, 3 mm diameter. The thermal transformation of the samples was studied by a NETZSCH-STA449C TG/DSC, NETZSCH 204 and 404 differential scanning calorimeters (DSC) under purified Ar atmosphere. The using a continuous heating regime with heating rates β from 2 to 90 °C min⁻¹. The temperature axis as well as the enthalpy axis was calibrated using indium, tin, bismuth, zinc, alumina and gold standards.

Structural relaxation

The metallic glasses and bulk amorphous alloys are thermodynamically unstable materials; when heating them structural relaxation and crystallization processes take place. If one heats a quenched metallic glass below the crystallization temperature, its structure transforms to a more relaxed (close to equilibrium) state, i.e. structural relaxation takes place. This phenomenon is a thermally activated process, and it happens below the glass transition temperature. During this structural relaxation short-term atomic rearrangements occur. Many properties change irreversibly or reversibly. Almost all physical properties change continuously: free volume, density, magnetic properties, electrical properties, ductility. Several models have been developed to describe and explain the structural relaxation of metallic glasses. Egami identified three mechanisms at the atomic level during annealing: diffusion, topological and chemical rearrangements [10]. The two rearrangements are not independent of each other. Topological structural...
relaxation (TSRO) and chemical short-range ordering (CSRO) have been observed by X-ray and neutron scattering measurements [11, 12] and by magnetic measurements [13] during the relaxation of metallic glasses. The free volume model was proposed by Turnbull and Cohen [14] and developed by Cohen [15] and Spaepen [16], and this model is one of the most commonly used. Heating of the material during the DSC experiment leads to an increase in the atomic mobility, and the excess free volume is annihilated. The decrease in the free volume $v_f$ is accompanied by a decrease in the enthalpy of the material. As a result, at a low temperature, a broad exothermic peak appears before the $T_g$ temperature can be observed in the DSC signal at a constant heating rate as an endothermic hump prior to the exothermic event. The amorphous structure has an equilibrium amount of free volume at $T_g^{\text{inset}}$ temperature.

The exothermic signal corresponds to the enthalpy release ($\Delta H_{f_v}$) due to the structural relaxation of the amorphous structure [17]:

$$\Delta H_{f_v} = \beta' \cdot \Delta v_f$$  \hspace{1cm} (2)

where $\beta'$ is a constant and $\Delta v_f$ is the change in free volume per atomic volume. Preheating of the sample at a different temperature or annealing for different times below the crystallization temperature, the low-temperature exothermic effect becomes weaker and shifts toward higher temperatures with increasing preheating temperature or time [17, 18]. A diagram of this phenomenon can be seen in Fig. 1. The change in material density is due to the change in free volume. Thanks to this, it can be determined that the change in free volume per atomic volume $\Delta v_f$ for sample was calculated as

$$\Delta v_f = \frac{\rho_i - \rho_0}{\rho_0}$$  \hspace{1cm} (3)

where $\rho_0$ is the density of the as-cast sample and $\rho_i$ is the density of the relaxed one. $\beta'$ constant can be determined by knowing the linear relationship between $\Delta H$ and $\Delta v_f$.

According to several researchers in the literature the plasticity and free volume are closely related [19–21, 21]. A large amount of free volume will enhance the plasticity of metallic glasses. Several articles can be found in the literature addressing the issue of relaxation and free volume [22–24]. It should be noted that in condensed matter physics structural relaxation is one of the investigated metallurgical problems.

**Structural relaxation and Curie transition in Fe-based metallic glasses with Cu content**

The Fe-based MGs have been widely applied as soft magnetic materials and the Co-based MGs as magnetic sensors. Fe-based amorphous alloys are ferromagnetic, and their soft magnetic properties make them useful for many applications. They are the most widely studied type of metallic glass.

During the relaxation of FINEMET type metallic glasses, other processes and changes in properties take place, which does not occur in the case of non-magnetic materials. During structural relaxation, not only a ferromagnetic Curie transformation occurs, but also the formation of clusters of Cu atoms, which is extremely important in the subsequent crystallization process. During heating a FINEMET type amorphous ribbon with a constant heating rate, a very small endothermic peak is visible at about 310 °C, indicating magnetic transformation at Curie temperature ($T_C$) (Fig. 2).

The Curie temperature of ferromagnetic glasses depends on the heating rate; therefore, it is recommended to use a lower rates—maximum of 80 °C min$^{-1}$. Furthermore, it is highly sensitive to relaxation and stress [25]. The cooling rate influences the structure forming and in this way it affects the Curie temperature of amorphous structure. The free volume also depends on the cooling rate and thus on the subsequent annealing process which changes the free volume. Annealing causes the $T_C$, and the density of metallic glasses to increase [26–29], yet compression causes a decrease in $T_C$ temperature [29]. Using a constant heating rate, a DSC measurement also provides both fast and accurate determination of $T_C$.

Using atom-probe field ion microscopy (APFIM) and extended X-ray absorption fine structure (EXAFS) it was confirmed that Cu atoms form Cu enriched clusters prior to the onset of crystallization [30, 31]. Furthermore, the three-dimensional atom probe (3DAP) technique made it
visible, that these Cu clusters serve as the heterogeneous nucleation sites for the bcc Fe(Si) primary crystals [32].

The exothermal peak of structural relaxation is broad and overlaps with the Curie temperature. Ohnuma et al. used an arbitrary baseline in order to roughly estimate the enthalpy for Cu clustering [33]. The enthalpy for clustering is less than 10% of the first crystallization enthalpy. In the case shown in Fig. 2, the enthalpy for clustering is ~ 5% of the α-Fe(Si) crystallization. The peak temperature of clustering (T_{cluster}) is higher in the case of lower Cu content, according to Ohnuma [33]. Our experiments showed that in the case where the difference between the Cu contents is very small (0.1 at.%), the measurement error can be compared with the degree of change. T_{cluster} increases with increasing heating rate. If the difference between the Tx (onset temperature of crystallization) and T_{cluster} is large, then there is enough time for Cu clusters to coarsen which favors obtaining the largest number of heterogeneous nucleation sites for bcc Fe(Si) crystallization.

For baseline correction, it should be taken into account that there is Curie temperature of bcc Fe(Si) crystallized phase formed during the first heating and it is visible as an endothermic signal for the 2nd measurement at above 600 °C.

**Glass transition temperature**

Glass transition temperature is defined as a single temperature which indicates the onset of the glass transition range for a glass. Chen and Turnbull already reported evidence of glass-liquid transformation in Au–Ge–Si alloy in 1968 [35]. Glass transition takes place at a temperature range rather than at a specific temperature [36, 37]. To date, the T_g temperature has been determined in several ways based on DSC measurements. The two most commonly used definitions can be observed in the inset to the Fig. 3: T_g-onset is the intersection of the baseline and the tangent with the greatest slope on the step, that is the most frequently used to indicate the glass transition temperature; the temperature lying midway between the extrapolated high- and low-temperature tangents on either side of the endothermic region indicated as T_g-mid [39]. However, conventionally T_g-onset is used as T_g temperature. Nowadays, the T_g temperature is most often and most rapidly determined by DSC measurement with continuous heating of the sample. But unfortunately, some alloys show no endothermic change in the DSC curve before crystallization, and even at higher heating rates (more than 20 °C min^{-1}) because T_g and Tx are so close to each other. For these alloys, the T_g temperature must be determined by another method (the electrical resistivity method). The glass transition temperature, crystallization temperature, and the width of the supercooled liquid region (ΔT_x = T_x-T_g) depend on the heating rate, i.e., they increase with an increase in heating rate [39, 41]. The supercooled liquid region of Fe_{40}Ni_{40}B_{16}Si_{4} metallic glass increases from 23 to 32 °C when the heating rate increases from 10 °C min^{-1} to 40 °C min^{-1} [39, 41].

The characteristic thermal properties of amorphous powders produced by high energy milling generally differ from the DSC analysis of metallic glasses and bulk amorphous alloys having the same composition. The amorphous powders produced by high energy milling do not show an endothermic change in the DSC curve before crystallization as can be seen in Fig. 4. Other authors have also
observed this phenomenon [40, 41]. However, the amorphous powders produced by the mechanical alloying or the process of gas-atomization behave like metallic glasses or bulk amorphous alloys.

**Crystallization temperature, and phase separation in the solid and liquid state**

In case of non-isothermal heating, the onset of crystallization temperature ($T_x$) can be determined by intersecting the tangents (Fig. 3). However, there are alloy systems for which it is not so easy to determine the real temperature $T_x$ on the DSC curve.

Phase separation can take place in the liquid and solid-state as well depending on the mixing enthalpy between the constituent element pairs. The separation occurs in a liquid state when it is large positive mixing enthalpy between the element pair. Since the discovery of amorphous alloys, phase separation has been observed in many alloy systems, and in glasses characterized by negative heat of mixing between the constituent elements or with a minor positive heat of mixing [43, 45].

The phase separation in metallic glasses characterized by negative or close to zero heat of mixing takes place mostly during heating of the as-quenched glass before the first crystallization [43, 44]. During the decomposition, nanoscale microstructure forms prior to the crystallization and this process follows the nucleation of nanocrystals on a few nm lengths [45–47]. The growth of nanocrystals is also an exothermic process, as is the crystallization of the amorphous structure.

Figure 5 shows the DSC curves of Cu$_{34.3}$Zr$_{37.2}$Ag$_{6.4}$Al$_{11.7}$ (at.%) bulk amorphous alloy before and after relaxation. The annealing was carried out at 300 °C of about 213 °C below $T_x$. The heat of mixing between Cu and Ag is +2 kJ mol$^{-1}$ [48]. In Fig. 5, the square area indicates the region, where the coarseness of the nanoparticles formed by phase separation.

In the case, when a phase separation takes place before the primary crystallization in the solid state, the $T_x$ temperature must be determined by the tangent method, but the tangent must not be drawn to the sharp exothermic peak, but to the small exothermic part (Fig. 6b). The DSC heating curve of (Cu$_{48}$Zr$_{48}$Al$_4$)$_{96}$Co$_{2}$Fe$_2$ alloy is shown in Fig. 6. This alloy system consists of atom pairs with a small positive heat of mixing in the liquid state: Cu–Fe (+13 kJ mol$^{-1}$ [48]) and Cu–Co (+6 kJ mol$^{-1}$ [49]).

It has long been known that in alloys with high positive mixing enthalpy, phase separation takes place in the liquid state. Below the separation temperature ($T_{sep}$) the homogeneous liquid separates into two liquids phases. These alloy systems are monotectic alloys, such as Al–Pb (+10 kJ mol$^{-1}$ heat of mixing), and in such systems it is a stable phase separation. Nakagawa [50] first reported that liquid separation occurs in the undercooled Fe–Cu system, in which a metastable miscibility gap exists.

In metallic glasses characterized by positive heat of mixing two immiscible liquids of different composition coexist in equilibrium between separation temperature ($T_{sep}$) and monotectic temperature ($T_M$). Bulk metallic glasses have high strength and good corrosion resistance and generally very limited ductility. When a monotectic alloy is undercooled into the miscibility gap, the liquid–liquid phase separation takes place and two liquids form. Drops of the new liquid, having a regular shape, begin generally with the nucleation of the liquid minority
phase. In immiscible alloys, a composite microstructure in situ forms with the crystalline or amorphous spheres embedded in the matrix. Amorphous/amorphous or amorphous/crystalline composites can be obtained owing to the liquid separation depending on the GFA of the immiscible liquids. Liquid separation provides an opportunity to produce in situ a composite microstructure, where the separated drops are ductile crystalline phases and the matrix has an amorphous structure.

The onset temperatures of phase separation $T_{\text{sep}}$ and monotectic reaction $T_M$ can be determined by DSC. The phase separation is manifested in the cooling curve (Fig. 7). The first small exothermic deviation from the baseline is attributable to liquid phase separation ($T_{\text{sep}}$) [50–52]. Figure 7 shows the heating and cooling curves of Cu$_{40}$Zr$_{20}$Ag$_{40}$ master alloy. At 1000 °C (1273 K) the homogeneous liquid separates into an Ag-rich liquid and a Cu–Zr–rich liquid. The primary solidification of the m-phase (AgCu$_4$Zr) from Ag-rich liquid starts at 986 °C (1259 K), and this is the $T_M$ temperature.

**Volume fraction of nanocrystals in an amorphous matrix**

Nano or/and microcrystals are solidified during the casting process when the cooling rate is slower than the critical cooling rate. DSC curves of Cu$_{48}$Zr$_{48}$Al$_4$ samples can be seen in Fig. 8. The three samples were cast with different cooling rates and, as a result, the samples cast with a slower cooling rate contained primary crystallized grains. The enthalpy in the first crystallization peak changes owing to nanocrystalline content, the enthalpy of the crystalline peak decreases with increasing nanocrystalline volume (Fig. 8). The peak temperature of the first crystallization slightly increases with increasing nanocrystalline volume fraction, and, because of the residual amorphous structure, it is more stable.

During annealing of an amorphous alloy at above the $T_g$ temperature, nanocrystallites crystallize, and these are essential for the enhancement of the mechanical or magnetic properties of an amorphous composite. It is important to know both the quantity of crystals formed during the casting process of an amorphous alloy and the quantity of crystals formed during the partial crystallization of the amorphous matrix. Quantitative determination of the volume fraction of the primary nanocrystalline phase is generally used in the TEM, XRD, or DSC methods. However, a precise TEM
Amorphous alloys and differential scanning calorimetry (DSC) study requires an expensive and difficult preparation and does not provide an overall landscape of a whole sample. DSC and XRD methods collect information about a larger volume, so these two methods are preferable to evaluate the volume fraction of primary nanocrystals.

For polymers and pharmaceuticals, the determination of the crystalline fraction by DSC measurement is accepted across the whole world \[54, 55\]. In the case of a sample with nanocrystallites in the amorphous matrix, the volume fraction can be estimated by using the following relationship \[56, 57\]:

\[
V_f = \frac{\Delta H_{\text{amorphous} + \text{nanocryst}}}{\Delta H_{\text{full amorphous}}}
\]

where \(\Delta H_{\text{full amorphous}}\) is the enthalpy change of a fully amorphous sample and \(\Delta H_{\text{amorphous} + \text{nanocryst}}\) is the enthalpy change of a partially crystallized sample during crystallisation.

According to several authors \[58, 59\], the values of the volume fraction of crystalline determined by DSC and XRD methods generally differ. In some cases, a lower value of \(V_f\) is obtained by DSC than that by XRD \[58\] and otherwise, the reverse occurs \[59\]. In my opinion, the agreement between DSC and XRD is good for small volume percentages, but the difference is significant for larger volumes. There can be several reasons for this discrepancy. The composition of the amorphous phase changes owing to the crystals, and therefore the residual amorphous structure has different properties. Furthermore, it is also difficult to find an accurate baseline. In turn, the XRD diffraction detects very small crystallites (~5 nm) as an amorphous phase. Summarizing, the determination of the volume fractions of crystals in the amorphous matrix is sensitive to the nature of the amorphous structure and the change of chemical composition during the nanocrystallization process. If we have a sample with a completely amorphous structure that was found to be completely amorphous based on an additional XRD measurement, we can decide very quickly whether another sample is completely amorphous or not by comparing their DSC measurements.

**Control of composition, quality assurance**

DSCs are applied to determine the impurity for many types of substances, e.g. pharma materials \[60\], organic matter \[61\], metals \[47, 62\]. Users of amorphous products and researchers need to know whether amorphous pieces with a nominally uniform composition actually do have the same composition. DSC measurement gives the fastest answer to this question, and thus DSC is required for quality assurance.

Two DSC curves of FINEMET type ribbons with the nominally equal composition are shown in Fig. 9. It can be observed that there is a significant difference in all crystallization processes. During the first crystallization step, the bcc Fe(Si) phase crystallizes. As we mentioned, Cu clusters serve as the heterogeneous nucleation sites for the primary precipitation of bcc-Fe(Si) crystals by heat treatment from the amorphous state. The peak temperatures do not significantly differ, but the difference in enthalpy values indicates that there is a difference in the volume fraction of bcc Fe(Si) phase. Diffusion of Nb outside the crystalline phase is very slow and Nb seems to control the growth rate of bcc-Fe(Si) precipitates. In the second and third crystallization steps, the intermetallic phases containing Fe, B, Nb and Si crystallize. The difference in peak temperature...
values also shows that the Nb and Si content also differ in the investigated ribbons. The ICP-OES measurement confirmed this observation because there was a difference in the Nb and Si content.

**Melting temperature and liquidus temperature**

Melting temperature and liquidus temperature are included in several GFA characterization study. It should be mentioned that some authors suggest that the solidus temperature should be used instead of the liquidus temperature \[63\]. Nonetheless, \( \gamma = T / (T_g + T_l) \) parameter \[64\], and a reduced glass transition temperature \( T_{rg} = T_g / T_l \) are included in a lot of articles.

The solidus temperature is the temperature below which there are only solid phases in equilibrium. The compositions of the best glass formers in an alloy system are quite close to the eutectic composition. In the case of non-eutectic composition, the onset temperature corresponds to melting in the eutectic reaction. If we see only one endothermic peak during the melting process, of course, it is not necessarily a pure eutectic composition. In many cases, the processes overlap and so we may see only one peak. It is highly recommended to study the first derivative of the heat flow curve and the cooling curve, because these curves help in the correct interpretation. In the Cu–Zr–Ag system, three different eutectic compositions were calculated by Kündig et al. \[65\], He et al. \[66\], and Kang et al. \[67\]. The DSC cooling curves of the three compositions revealed that these compositions are not eutectic (Fig. 10).

The liquidus \( T_L \) is the ultimate temperature of thermodynamic equilibrium between the solid and liquid phases of any material, above which crystals are unstable. The melting process is endothermic. The melting process begins at the temperature where the DSC signal deviates from the base-line (Fig. 11). However, this is not easy to determine, so the melting temperature (which is the solidification temperature) \( T_{onset} \) is defined by the intersection of tangents to the curve, traced on the baseline and the low-temperature peak side. The melting process finishes at a lower temperature than the endset temperature. Conventionally, the liquidus temperature is defined as \( T_{endset} \) temperature (Fig. 11) and is also determined by the tangent method, but it is not a standard method and its uncertainty is difficult to estimate. A heating speed of 0.5–10 °C min\(^{-1}\) should normally be used for the measurement, and by no means at the same heating rate as for \( T_g \) measurement (20–40 °C min\(^{-1}\)).

Because the onset temperature corresponds to melting in a eutectic reaction, thus, it does not change with the heating rate, while the endpoint temperature varies with the heating rate and largely depends on the mass, layer thickness and thermal conductivity of the sample. Therefore, there are several methods in the literature to determine the liquidus temperature with more physical content. G.W.H. Höhne et al. \[69\] proposed a method that takes into account the dependence of the temperature on the heating rate. E. B. Ferreira et al. \[70\] found that the endset temperature is directly proportional to the heating rate and the best approximation to the liquidus temperature is the endpoint of the DSC melting peaks measured at different rates and extrapolated to 0 °C min\(^{-1}\). H. Nishikawa et al. \[71\] found that the extrapolated end temperature of the endothermic peak is proportional to the square root of the DSC heating rate.

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**Fig. 10** DSC curves of eutectic compositions based on calculations in Cu-Zr-Ag system (heating and cooling rate: 5 °C min\(^{-1}\)) \[68\]

**Fig. 11** DSC curve of a Cu\(_{40}\)Zr\(_{34}\)Ag\(_{26\text{at.\%}}\) alloy (heating rate 10 °C min\(^{-1}\))
Amorphous alloys and differential scanning calorimetry (DSC)

The two proposed onset and endset temperatures for melting process determinations of Cu$_{35}$Zr$_{41.25}$Ag$_{23.75}$ (at.%) alloy are summarized in Fig. 12. It can be observed that the $T_{\text{onset}}$ temperature is independent of both the heating rate (Fig. 12a) and the square root of the heating rate (Fig. 12b). The y-axis intercepts for the peak and endset lines are very similar. The difference between the two extrapolated liquidus temperatures to 0 °C min$^{-1}$ of the two models is 5.6 °C. It is worth noting that the heating rate should only be less than 20 °C min$^{-1}$ because the temperature extrapolated to 0 °C min$^{-1}$ measured with a heating rate of 20 °C min$^{-1}$ and above differs from the value extrapolated from measurements carried out with a lower heating rate (Fig. 13). This extrapolation is also used for crystallization peaks. In Fig. 13 the crystallization temperature of Fe$_{40}$Ni$_{40}$B$_{16}$Si$_4$ metallic glass can be seen as a function of the heating rate. The heating rate was between 2 and 90 °C min$^{-1}$. It can be observed, that the above-mentioned correlations are not particularly valid for E. B. Ferreira et al. method [70] if one uses higher heating rates as we can see in many articles.

\[
\begin{align*}
T &= 838.7 \ °C + 1.61 \beta \ (R^2 0.96) \\
T &= 838.3 \ °C + 0.35 \beta \ (R^2 0.99) \\
T &= 817.9 \ °C + 0.37 \beta \ (R^2 0.22)
\end{align*}
\]

\[
\begin{align*}
T_{\text{onset}} &= 816.0 + 1.65 \beta \ (R^2 0.25) \\
T_{\text{peak}} &= 837.0 + 1.47 \beta \ (R^2 0.98) \\
T_{\text{endset}} &= 833.1 + 6.55 \beta \ (R^2 0.89)
\end{align*}
\]

Fig. 12 Correlation between the heating rate and $T_{\text{onset}}$, $T_{\text{endset}}$ and $T_{\text{peak}}$ measured from the DSC profiles of Cu$_{35}$Zr$_{41.25}$Ag$_{23.75}$ (at.%) alloy: (a) method by E. B. Ferreira et al. [70], (b) method by H. Nishikawa et al. [71]

Fig. 13 The crystallization peak temperature of Fe$_{40}$Ni$_{40}$B$_{16}$Si$_4$ (at.%) metallic glass as a function of heating rate
Kinetic analysis of crystallization processes

The phase transformation processes in solids are generally thermally activated processes, during which the system sitting in the local minimum of free enthalpy passes through a potential barrier with thermal fluctuation, thus reaching a lower energy level than in the case of glass metal crystallization. The crystallization of amorphous alloys involves nucleation and growth. The knowledge of the rate of transformation is very important because heat treatment affects the properties of the product. There are many methods in the literature to determine the crystallized volume, the local activation energy and the Avrami index \( n \), which provides information on the nucleation and growth mechanism and dimensionality. It should be noted that a method that is perfect for any process is not yet known. The reason is that the transformation rate depends exclusively on temperature, the degree of transformation and also the thermal history. Unfortunately, existing methods are also misapplied in several cases.

For both isothermal and continuous heating processes, kinetic analysis is possible. There are several kinetic models in the literature, and the most common ones will be presented here.

The Johnson, Mehl, Avrami and Kolmogorov model (JMAK) \[2, 72, 73\] describes the crystallization kinetics during an isothermal process. The model assumes that:

- The crystallization is isothermal, i.e. nucleation and growth occur at a constant temperature;
- The sample is assumed to be infinitely large and the nucleation is random throughout the bulk of the sample;
- Growth is isotropic until crystals impinge upon one another;
- The activation energy for crystallization and other model parameters are independent of time and temperature.

The volume fraction transformed during crystallization is written as:

\[
x(t, T) = 1 - \exp \left\{ -(k(T)t)^n \right\}
\]

where \( k(T) \) is the rate constant, \( t \)-time and \( n \)-the Avrami index. \( n \) is a parameter that depends on the nucleation and growth mechanism and dimensionality. Table 1 includes the characteristic \( n \) parameters for different reaction mechanisms as taken from Ref. [74].

The rate constant in the thermal activated process:

\[
k(T) = A \exp \left( \frac{-E_a}{RT} \right)
\]

where the pre-exponential factor \( A \) and activation energy \( E_a \) are kinetic parameters that should not depend on the temperature \( T \) and the fractional conversion \( x \).

A major disadvantage of this model is that the transformed fraction is given in the case of isotherms, and since measurements with DSC are often not made in isotherms, the terms must be generalized for the situation of non-isotherms. To describe non-isothermal reactions, there are several thermo-kinetic methods to evaluate the DSC curves. Kissinger [75], Takhor [76] and Ozawa [77] developed a method in which both \( E_a \) is determined at different heating rates. The basic equations of both methods are shown as Eq. 7 [75], 8 [76] and 9 [77]:

\[
\ln \left[ \frac{\beta}{T_{peak}^2} \right] = -\frac{E_a}{RT_{peak}} + \ln \left( \frac{B_0 R}{E_a} \right)
\]

\[
\ln \left[ \frac{\beta}{T_{peak} - T_{start}} \right] = -\frac{E_a}{RT_{peak}} + \ln \left( B_0 \right)
\]

\[
\ln = -\frac{E_a}{RT_{peak}} + \text{const.}
\]

### Table 1 Values of the exponent \( n \) for the JMA equation for different kinds of reaction mechanisms

| Type of nucleation                                      | Geometry of nuclei and their subsequent growth | Phase-boundary controlled | Diffusion controlled |
|--------------------------------------------------------|-----------------------------------------------|----------------------------|---------------------|
| Instantaneous nucleation (saturation of sites capable of nucleation prior to growth) | Bulk mechanism                               | 1                          | 0.5                 |
|                                                        | One-dimensional growth                        | 2                          | 1                   |
|                                                        | Two-dimensional growth                        | 3                          | 1.5                 |
|                                                        | Three-dimensional growth                      | ~2                         | 1                   |
|                                                        | Surface mechanism                             |                            |                      |
|                                                        | Two-dimensional growth                        | ~2                         | ~2                  |
| Constant rate of homogeneous nucleation during the process | Bulk mechanism                               | 1                          | 0.5                 |
|                                                        | One-dimensional growth                        | 2                          | 1.5                 |
|                                                        | Two-dimensional growth                        | 3                          | 2                   |
|                                                        | Three-dimensional growth                      | 4                          | 2.5                 |
|                                                        | Surface mechanism                             | ~3                         | ~2                  |
where \( \beta \) is the heating rate, \( E_a \) is the activation energy, \( R \) is the gas constant, \( T_{\text{peak}} \) is the temperature of a peak, \( T_{\text{start}} \) is the starting temperature of the peak, \( B_0 \) is the rate constant.

Drawing the function \( \ln (\beta / T_{\text{peak}}^2) - 1 / T_{\text{peak}} \) for the individual crystallization peaks, a straight line is obtained, the slope of which is \( E_a / R \). In the literature, these lines are called Kissinger's straight lines. Naturally, the Kissinger method can be used only if the peak appearing on the DSC curve belongs only to one process. The kinetic analysis of the crystallization is presented in the case of \( \text{Fe}_{40}\text{Ni}_{40}\text{B}_{16}\text{Si}_{4} \) metallic glass.

Heating rate \( \beta \) was changed between 2 K min\(^{-1}\) and 99 K min\(^{-1}\) in 13 steps. Kissinger's, Takhor's and Ozawa's lines are presented in Fig. 14.

The analysis gives 296.6 ± 11.3 kJ mol\(^{-1}\) 294.1 ± 11.3 kJ mol\(^{-1}\) and 302.1 ± 8.7 kJ mol\(^{-1}\) to the activation energy of crystallization based on the Kissinger, Takhor and Ozawa method. Only a small difference exists between the three values of the activation energy.

The crystallized volume fraction for the non-isothermal crystallization can be deduced as a function of temperature using the following equation:

\[
x = \frac{\int_{T_0}^{T_{\text{end}}} \left( \frac{dH}{dT} \right) dT}{\int_{T_0}^{T_{\text{end}}} \left( \frac{dH}{dT} \right) dT} = \frac{A}{A_0} \tag{9}
\]

where \( T_0 \) and \( T_{\text{end}} \) are the temperatures corresponding to the onset and end of crystallization and \( dH/dT \) is the heat capacity at constant pressure. \( A \) is the area under the DSC curve between the onset temperature and a given temperature, and \( A_0 \) is the area between the onset and end of crystallization. The crystallized volume fraction is plotted as a function of temperature for \( \text{Fe}_{40}\text{Ni}_{40}\text{B}_{16}\text{Si}_{4} \) metallic glass in Fig. 15. The variation of the crystallized volume fraction versus temperature follows a sigmoidal behavior.

The activation energy determined by Eqs. 7, 8 and 9 is an average value of the activation energy of the whole crystallization process. Unfortunately, the activation energy depends on the volume fraction. The \( E(x) \) corresponds to the appropriate values of activation energy in a wide range of crystallized volume fraction can be
evaluated from the following Kissinger–Akahira–Sunose (KAS) model [78]:

$$\ln \left( \frac{\beta}{T^2(x)} \right) = - \frac{E(x)}{RT(x)} + \text{const.}$$  \hspace{1cm} (10)

and the Ozawa–Flynn–Wall formula (OFW) [79]:

$$\ln \beta = -1.052 \frac{E(x)}{RT(x)} + \text{const.}$$  \hspace{1cm} (11)

where \(T(x)\) is a temperature corresponding to the certain crystallized volume fraction \(\beta\) at different heating rates. Figure 16 shows the relation between the local activation energy \(E(x)\) and the crystallized volume fraction calculated by KAS and OFW. The average activation energy \(E(x)\) between \(x = 0.1\) and \(0.9\) is 264.9 kJ mol\(^{-1}\) calculated by KAS and 263.1 kJ mol\(^{-1}\) calculated by OFW. These values show a very good agreement, i.e., either method can be used for this alloy. The average activation energy \(E(x)\) between \(x = 0.1\) and \(0.9\) is lower than the average activation energy of the first crystallization based on Kissinger, Takhor and Ozawa method \(E_a\) (see Fig. 14). In the initial stage, i.e., \(x < 0.1\), \(E(x)\) increases slightly, the crystallization is slow at this stage (see Fig. 15), a barrier of crystallization must be overcome. In the second stage, i.e., \(0.1 < x < 0.9\), the crystallized volume fraction increases rapidly, the local activation energy decreases monotonically. In the final stage, i.e., \(x > 0.9\), although the local activation energy decreases rapidly and reaches the lowest value, and reaches the lowest value and the degree of crystallization slows down.

The Avrami exponent is usually used to define the mechanism of nucleation and growth behavior during the crystallization process. Based on Kissinger’s work, Benett and Augis [78] created a reworking of the Kissinger method. The Avrami index \((n)\) can be estimated from the relation:

$$n = \frac{2 \cdot 5 \cdot T^2_{\text{peak}}}{\Delta T_{\text{FWHM}} E_a}$$  \hspace{1cm} (12)

where \(\Delta T_{\text{FWHM}}\) is the full width at half maximum of the DSC peak, \(E_a\) is the activation energy, \(R\) is the gas constant. The Avrami exponent of Fe\(_{40}\)Ni\(_{40}\)B\(_{16}\)Si\(_4\) metallic glass based on this analysis is 3.04. This relationship assumes that the activation energy is constant throughout the process and does not depend on the crystallized fraction. However, the activation energy depends on the volume fraction and consequently, the Avrami index \((n)\) also varies depending on the crystalline fraction. The local Avrami exponent can be determined from the relation proposed by J.S. Blázquez et al. [81]:

$$n(x) \cdot \left\{ 1 + \frac{E_a}{RT(x)} \left( 1 - \frac{T_0}{T} \right) \right\} = \frac{d \ln (-\ln (1-x))}{d \ln \left[ \frac{T-T_0}{\beta} \right]}$$  \hspace{1cm} (13)

According to Eq. (12), plotting \(\ln[-\ln(1-x)]\) versus \(\ln[(T-T_0)/\beta]\) will provide the \(n(x)\) values from the slope of the curves (see Fig. 17). The slopes of the curves vary in the case of larger crystalline volume fraction \((x > 0.8)\), which indicates that the nucleation and growth do not remain the same way during the whole crystallization process. The \(n(x)\) value reflects the crystallization mechanism. The local Avrami exponent \(n(x)\) as a function of the crystallized fraction \((x)\) at a heating rate of 5 and 30 K min\(^{-1}\) is shown in Fig. 17b. The crystallization of Fe-based amorphous alloys can be controlled by diffusion [82–84] or phase-boundary [85].

$$n = b + mp$$  \hspace{1cm} (14)

where \(b\) is the nucleation index \((b = 0\) for zero nucleation rate, \(0 < b < 1\) for decreasing nucleation rate, \(b = 1\) for constant nucleation rate, \(b > 1\) for increasing nucleation rate), \(m\) is the dimensionality of the growth \((m = 1, 2, 3\) for one-dimensionality, two-dimensionality, three-dimensionality growth respectively), and \(p\) is the growth index \((p = 0.5\) for diffusion-controlled growth and \(p = 1\) for interface-controlled growth). For diffused-controlled crystallization, \(n > 2.5\) indicates that the crystallization process is the different kinds of growth of small particles with an increasing nucleation rate. \(1.5 < n < 2.5\) implies the growth of precipitates with decreasing nucleation rate, and \(n < 1.5\) means that the growth of pre-existing nuclei [86].

In the case of the investigated Fe\(_{40}\)Ni\(_{40}\)B\(_{16}\)Si\(_4\) ribbon, the value of the local Avrami exponent at the first crystallization process is higher than 3 during the whole crystallization,
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which means that the crystallization mechanism of \( \alpha \)-Fe(Si) is dominated by three-dimensional growth with an increasing nucleation rate \((b = 3; p = 0.5\), considering this values, \(1.5 < n < 2.5\)).

Comparing the value of the Avrami exponent determined by the two methods, we can conclude that the value of Avrami index determined by Benett and Augis [78] is lower than the local n(x) value determined by J.S. Blázquez et al. [81], but in both cases greater than 3.

Thermal analysis methods are widely used to study crystallization kinetics in amorphous solids. There are opinions in the literature that the JMAK model has limited validity both in isothermal and non-isothermal conditions [87]. But many improved versions have been published [88, 89]. The validity of the Kissinger method is also disputed [90] and therefore authors have further developed the Kissinger method [91, 92]. Controversial results related to the activation energy of FINEMET alloys have been also reported. For example, the kinetics of nanocrystallization of \( \text{Fe}_{73.5}\text{Si}_{13.5}\text{B}_{9}\text{Cu}_{1}\text{Nb}_{3} \) alloy has been investigated by several authors. According to H.A. Shivaee et al. [93] for volume fraction in the range of 0.1–0.75, activation energy increased approximately linearly with increasing crystalline fraction. In another study, W.Lu et al. [94] have found that the activation energy is approximately constant in the range of 0.3–0.7. Differences in results are probably due to several reasons: the nanocrystallization complexity of the amorphous alloy, the state of the starting material may not be the same even though the chemical composition is the same (the amorphous structure and the volume fraction that crystallizes during the devitrification depends on the cooling rate [95]) and there may be nanoscale crystals in the material that are not visible on X-ray measurement.

**New developments-fast-scan DSC (FSDSC) and ultra-fast (Flash) DSC**

From the second half of the twentieth century, there has been a great interest in the fundamental study of the amorphization and partial to full crystallization of materials. It is required using high cooling and heating rates. The introduction of chip-based calorimeters allowed the application of a wide range of heating and cooling rates: from 0.1 to \( \sim 5000 \) K s\(^{-1}\) in the case of fast-scan DSC and up to 40,000 K s\(^{-1}\) in the case of ultra-fast DSC (Flash DSC). Using these instruments enable in-depth study of metastability and reorganization of samples with a 1 to 1000 ng scale. In addition, FSDSC will become a crucial tool for the optimization of processing methods at high speeds like injection molding [96, 97]. The first instrument to be commercially available was Flash DSC 1 (Mettler Toledo) in 2010. The apparatus of MultiSTAR USF1 (XI-400) works with the mechanical refrigerator for the effective range of measurement from \(-100\) to 450 °C. Nowadays it is widely
used for the study of polymers, pharmaceutical materials and even less so for metallic materials. The maximum temperature in the current equipment is 800 °C, so it is understood that it is currently used even less in the testing of metallic materials. However, this technique can detect the glass temperature, small and fast phase transitions, and it is possible to analyse non-equilibrium transformations at high heating or cooling rates in the case of metallic materials.

The most obvious advantage of using fast-scan DSC is that it reduces the instrumental analytical time giving a speedier collection of data. The mass of the sample is generally less than 500 μg, so it provides information on a very small volume. The good thermal contact between a sample and the pans is also important for normal DSC, but this is especially important for fast DSC. Flat pans should always be used with helium as a purge gas, which has a higher thermal conductivity than nitrogen or air and allows better heat transfer and peak resolution. As the heating rate increases, of course, the peak temperatures shift towards higher temperatures. The onset temperature of melting greatly increases above a heating rate of 1000 °C s⁻¹ [98]. S. Pogatscher et al. [98] determined the critical cooling rate required to reach the amorphous state from melt in the case of Au₄₉Ag₆₅Pd₂₃Cu₂₆₉Si₁₆₃ alloy using the fast scan DSC. Due to the use of fast and ultra-fast DSC, it is possible to study several things that have not been done so far with the use of traditional DSC: J. Schawe et al. [99] found that monotropic polymorphism exists in Au-based amorphous alloys, M. Gao et al. [100] determined the delay time for primary crystallization of Al-based metallic glasses with poor glass formation ability.

Kinetical studies show that the plots of the ultrafast DSC data exhibit non-Arrhenius behavior more or less, and the value of activation energy decreases continuously when the heating rate increases [101].

**Conclusions**

The differential scanning calorimetry can provide a lot of information about amorphous alloys, not only the characteristic temperatures and thermal events. Of course, there are some disadvantages of the DSC test, but compared to other measurement methods, it certainly has the advantage that the experiment is not complicated, and, after a calibration that was precisely carried out, much information can be obtained from a few measurements in a short time using a small amount of material. DSC analysis supplies both thermodynamics and kinetic data on these materials analysed and also provides a better insight into the short-range order—long-range order transition. Unfortunately, the DSC analysis of amorphous alloys is not standardized. A lot of the properties determined by DSC measurement strongly depend on the measurement conditions, so a uniform test would be needed. The authors hope that the remarks in this article will be helpful for the readers in their work.

**Acknowledgements** This work was carried out as part of the GINOP-2.3.2-15-2016-00027 project implemented in the framework of the Széchenyi 2020 program. The realization of this project is supported by the European Union.

**Authors contribution** All authors contributed to the study conception and design. Material preparation, data collection and analysis were performed by [Piotr Bala], [Maria Sveda], [Anna Sycheva] and [Ferenc Kristal]. The first draft of the manuscript was written by [Dora Janovszky] and all authors commented on previous versions of the manuscript. All authors read and approved the final manuscript.

**Funding** Open access funding provided by University of Miskolc.

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