Nucleation behaviour and microstructure of single Al-Si12 powder particles rapidly solidified in a fast scanning calorimeter

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ABSTRACT

The understanding of rapid solidification behaviour, e.g. the undercooling versus growth velocity relationship, is crucial for tailoring microstructures and properties in metal alloys. In most rapid solidification processes, such as additive manufacturing (AM), in situ investigation of rapid solidification behaviour is missing because of the lack of accurate measurement of the cooling rate and nucleation undercooling. In the present study, rapid solidification of single micro-sized Al-Si12 (mass%) particles of various diameters has been investigated via differential fast scanning calorimetry employing controllable cooling rates from 100 to 90,000 K s\(^{-1}\) relevant for AM. Based on nucleation undercooling and on microstructure analysis of rapidly solidified single powder particles under controlled cooling rates, two different heterogeneous nucleation mechanisms of the primary \(\alpha\)-Al phase are proposed. Surface heterogeneous nucleation dominates for particles with diameter smaller than 23 \(\mu\)m. For particles with diameter larger than 23 \(\mu\)m, the nucleation of the primary \(\alpha\)-Al phase changes from surface to bulk heterogeneous nucleation with increasing cooling rate. The results indicate that at large undercoolings (\(> 95\) K) and high cooling rates (\(> 10,000\) K s\(^{-1}\)), rapid solidification of single particle can yield a microstructure similar to that formed in AM. The present work not only proposes new insight into rapid solidification processes, but also provides a
Introduction

Rapid solidification of metal particles is an important process in many production chains of metal components. Recently, rapid solidification has attracted further interest as it is intrinsically linked to the evolving field of additive manufacturing (AM). AM of metals holds great potential for improving materials’ efficiency, reducing life-cycle impacts, enabling greater engineering functionality and allowing fast reaction to customers’ demands [1, 2]. Al-based alloys such as Al–Si alloys are important engineering materials in general and are frequently used in AM. Attention has therefore been given to understand the rapid solidification processes of micro-sized metal particles in AM processes and their influence on the developing microstructures [3–7]. Undercooling and nucleation influence practically relevant microstructural features such as grain size and hot cracking susceptibility. The solidification cooling rates in AM processes range from about 100 K s\(^{-1}\) up to \(10^6\) K s\(^{-1}\) [7–11]. For example, the cooling rate of Al-Si12 (mass\%) in laser powder bed fusion (LPBF, sometimes also called laser beam melting or selective laser melting, LBM/SLM) was estimated (based on simulations) to be above \(10^3\) K s\(^{-1}\) for most parts of the melt pool [9]. Besides, by using other rapid solidification techniques, similar high cooling rates can be achieved. Numerous studies have been reported on the rapid solidification of Al–Si alloys by melt spinning, gas atomisation, laser melting, spray forming, etc. [12–15], especially for the nucleation undercooling in Al–Si alloy particles [13, 16–18]. However, the in situ investigation of the rapid solidification behaviour of powder particles under such conditions is not straightforward. Hence, so far ex situ investigations and numerical simulation are commonly applied to study the transient, non-equilibrium states during rapid solidification under such extreme conditions far away from thermodynamic equilibrium [3, 4, 7, 13]. Unfortunately, the quality of those estimates is hard to evaluate since particularly the occurrence and lifetime of transient non-equilibrium states are not easy to verify experimentally. Hence, the predictions of the solidification mechanisms have been supported by only a few quantitative studies [19–22]. Thus, in situ investigations of rapid solidification revealing the solidification temperature intervals, and its nucleation mechanisms (e.g. nucleation kinetics) are lacking. However, for a proper description of rapid solidification, these results are crucial and fundamental for proper parameter selection in additive manufacturing. A better understanding of rapid solidification at such high cooling rates requires accurate measurements of the cooling rate and solidification temperature intervals and an accurate description of its mechanisms.

In the present work, a non-adiabatic differential fast scanning calorimeter (DFSC) [23] with a wide range of controllable cooling rates ranging from 100 to \(90,000\) K s\(^{-1}\), covering the cooling rate in AM, was utilised for in situ analysis of rapid solidification. This thermal analysis device provides the opportunity to investigate the fundamental issues related to rapid solidification of single micro-sized, near-spherical particles, such as the solidification temperature intervals and nucleation kinetics. Particle size is a crucial factor in the nucleation kinetics, especially for determining nucleation undercooling [17, 24, 25]. Therefore, micro-sized Al-Si12 (mass\%) particles with diameters from 6 to 34 \(\mu\)m were used. By combining DFSC analysis and microstructure analysis, the rapid solidification behaviour of single Al-Si12 particles, specifically the nucleation mechanism of the primary \(\alpha\)-Al phase, is discussed. A modified classical heterogeneous nucleation theory, which assumes surface nucleation in a spherical cavity and/or bulk nucleation on a convex spherical interface, is developed.

Experimental procedure

Gas-atomised Al-Si12 (mass\%) powders supplied by Nanoval GmbH & Co. KG (Germany) were used in the present work. The composition of the as-received powders was analysed by X-ray fluorescence spectroscopy (XRF) (see Table 1). The composition is close to the nominal composition. The microstructure of
the cross section of as-received Al-Si12 powders was characterised using a scanning electron microscope (SEM) (Zeiss Merlin VP Compact, Germany), as shown in Fig. 1. The microstructure is composed of α-Al dendrites and (α-Al + Si) eutectics, where the grey and bright areas are Al and Si, respectively.

DFSC based on thin film chip sensors [23] was used for the in situ measurements of the melting and solidification temperatures. The sensor, UFH1 (Mettler Toledo, Switzerland), consists of an amorphous silicon–nitride membrane with film thermopiles and resistive film heaters placed at the centre of the membrane, as shown in Fig. 2. Particles of near-spherical shape with diameters ranging from 6 to 28 μm were chosen for DFSC measurements. The particle sizes were characterised by optical microscopy (Olympus BX41, Japan) before and after the measurements. A single micro-sized particle was deposited in the centre of the measurement area (see Fig. 2) by a thin copper wire under a stereomicroscope (Olympus SZX16, Japan). To improve the heat contact between the sample and membrane, a tiny amount of silicone oil was used. The boiling temperature of silicone oil is ca. 250 °C at low heating rates. However, due to the high heating rates investigated and thus short times at high temperatures, the silicone oil was expected to be stable up to high temperatures. After the first melting and solidification cycle, an improved thermal contact between the sample and membrane was established due to slight flattening of the particle.

To investigate the rapid solidification of Al-Si12 particles by DFSC, the single particle was heated at 5000 K s\(^{-1}\) from 37 to 707 °C and held for 0.01 s under argon atmosphere at a constant pressure of 100 kPa. As the liquidus temperature of Al-Si12 is near the eutectic temperature of the Al-Si-system 577 °C [27], this maximum heating temperature ensures a certain overheating even at high heating rates. Afterwards, the single particle was cooled to 37 °C at cooling rates from 100 to 90,000 K s\(^{-1}\). The temperature dependence of the accessible scanning rate of a particle of 7 μm in diameter (from 100 to 90,000 K s\(^{-1}\)) is demonstrated in Fig. 3a, indicating that cooling rates as high as 90,000 K s\(^{-1}\) are controllable in the aimed temperature range of 600–350 °C. For improved reproducibility, identical conditions, i.e. heating rate, overheating and time above liquidus temperature of the single particle before subsequent cooling, were applied. For each cooling rate, ca. 100 repeated cooling–heating cycles were performed on each single particle to appropriately account for the stochastic nature of the nucleation process. A representative temperature–time profile (with a heating and cooling rate of 5000 K s\(^{-1}\)) is shown in Fig. 3b. During this heating–cooling process, the heat flow was recorded as a function of time. During the heating–cooling cycle, the Al-Si12 particle

Table 1 Mass fractions of alloying elements in the as-received Al-Si12 powders

| Alloy | Al | Si   | Fe  | Cr  | V  |
|-------|----|------|-----|-----|----|
| Al-Si12 | ca. 88% | 11.80% | 0.16% | 0.02% | 0.01% |

Figure 1 SEM secondary electron images of the cross section of an as-atomised Al-Si12 particle, etched with Weck’s reagent [26].

Figure 2 Single Al-Si12 powder particle of 13 μm in diameter on a UFH1 DFSC sensor.
underwent melting and solidification. The melting onset temperature, $T_{m_{\text{onset}}}$, and the solidification onset temperature, $T_{s_{\text{onset}}}$, were determined [28]. The temperature calibration and correction (accounting for a static temperature offset and a dynamic temperature offset due to thermal lag) for each particle were performed by the eutectic temperature of Al-Si12 alloy upon heating (for details see “Appendix”). According to the Al–Si phase diagram [27], the liquidus temperature, $T_l$, is approximated to the eutectic temperature, $T_e$, for Al-Si12 particles. Therefore, the nucleation undercooling, $\Delta T$, can be calculated by

$$\Delta T = T_l - T_n = T_{m_{\text{onset}}} - T_{s_{\text{onset}}}$$

(1)

where $T_n$, $T_{m_{\text{onset}}}$, and $T_{s_{\text{onset}}}$ are the nucleation temperature, the corrected melting and solidification onset temperatures for each scanning rate, respectively. It should be noted that the melting onset temperature did not change in the course of the repeated experiments, as shown in Fig. 3c. This indicates not only the repeatability of the DFSC analysis but also the stable shape and composition of the single Al-Si12 particle during all the experiment cycles. Thus, the influence of the particle shape on the nucleation undercooling can be neglected. The scatter of the onset temperatures of the solidification peaks even with the same cooling rate in Fig. 3c comes from the stochastic nature of the nucleation process.

**Figure 3**  
(a) Scanning rates of a single Al-Si12 particle (7 μm in diameter) by DFSC indicating that cooling rates as high as 90,000 K s$^{-1}$ are controllable in the temperature range of 600 °C–350 °C.  
(b) Typical DFSC measurement curve of the melting and solidification processes of a single Al-Si12 particle. The dashed curve is the temperature–time profile, and the solid curve is the heat flow curve. The melting onset temperature, $T_{m_{\text{onset}}}$, the solidification onset temperature, $T_{s_{\text{onset}}}$, and the uncorrected undercooling, $\Delta T^*$, are indicated in the figure.  
(c) 100 measured curves of a single Al-Si12 particle (14 μm in diameter) at 5000 K s$^{-1}$ heating/cooling rate.
Finally, the microstructures of single rapidly solidified particles with controlled cooling rates by DFSC were investigated. According to the analysis of the DFSC results (for details see next section “Undercooling dependence on cooling rate and particle size”), two sets of particles with certain sizes were chosen, i.e. ca. 20 µm and ca. 32 µm in diameter. Four cooling rates were chosen, namely 100 K s⁻¹, 500 K s⁻¹, 5000 K s⁻¹ and 50,000 K s⁻¹, respectively. It should be mentioned that one specific particle was chosen for each selected cooling rate and size for microstructure characterisation. After solidification by DFSC at a controlled cooling rate, resulting in a known undercooling, the single particle was transferred from the DFSC sensor and placed into a cone-shaped bore in an epoxy block with a thin copper wire under a stereomicroscope. Then, the bore was carefully filled with epoxy resin, avoiding movement of the particle. After hardening the epoxy resin in the bore, the epoxy block was ground and polished carefully until obtaining the particle’s cross section. Then the specimen was etched with Weck’s reagent (100 mL water, 4 g KMnO₄ and 1 g NaOH) for 4 s. Details of the specimen preparation for microstructure characterisation of one single micro-sized particle can be found in [29]. The cross-sectional microstructure of the particle was characterised by SEM. A linear intercept method [30] was applied to measure the α-Al dendrite arm spacing (DAS). Therefore, about 18 lines in different directions with a total length of about 250 µm were placed on the cross section of each single particle.

**Results and discussion**

**Undercooling dependence on cooling rate and particle size**

Solidification of metals requires a certain degree of nucleation undercooling. By controlling undercooling, different scenarios of crystal nucleation, resulting in a variety of solidification structures with superior material properties, can be obtained [31, 32]. During rapid solidification by DFSC, the solidification onset temperature (e.g. shown in Fig. 3c) is lower than 500 °C. According to the calculated Al-rich boundary of the coupled zone in laser-treated Al-Si alloys [33], during rapid solidification of an Al-Si12 alloy, the primary α-Al phase solidifies first from the melt, followed by the formation of (α-Al + Si) eutectic due to the coupled zone [33, 34]. Therefore, the solidification onset upon cooling corresponds to the nucleation of the primary α-Al phase, and the nucleation undercooling of the primary α-Al phase can be calculated by Eq. (1). It should be noted that the undercooling of the second eutectic phase cannot be evaluated because of the overlapping solidification peaks, as shown in Fig. 3c. The dependence of the nucleation undercooling on the cooling rate of an Al-Si12 particle of 7 µm in diameter is shown in Fig. 4a. For a sound statistical basis, the heating–cooling process was repeated 100 times for each cooling rate. As expected, due to the stochastic nature of nucleation, nucleation occurs in a temperature range where the mean value of the undercooling.
corresponding to a specific cooling rate is indicated by the blue circle symbol. The mean nucleation undercooling increases with increasing cooling rate. The variance of undercooling because of the stochastic nature of nucleation is about 40 K for each cooling rate between 100 and 90,000 K s\(^{-1}\).

The dependence of the mean nucleation undercooling of single Al-Si12 particles with different diameters (6 to 28 \(\mu m\)) on the cooling rate from 100 to 90,000 K s\(^{-1}\) is illustrated in Fig. 4b. The undercooling is in a similar range as the estimated undercooling of atomised Al-Si eutectic alloy particle [13]. For particles larger than 23 \(\mu m\) in diameter, no sufficient data set can be obtained up to the cooling rate of 10,000 K s\(^{-1}\). This is because of the serious dynamic temperature offset due to the thermal lag at higher cooling rates, resulting from high heat capacities of large particles. This cooling rate of 10,000 K s\(^{-1}\) is still relevant for L-PBF processes with typical cooling rates above 1000 K s\(^{-1}\) [9]. Even though measured undercooling is not reliable for particles larger than 23 \(\mu m\), they can be cooled at rates up to 50,000 K s\(^{-1}\) and their microstructure becomes accessible. For smaller particles with diameters from 6 to 21 \(\mu m\), cooling rates of up to 90,000 K s\(^{-1}\) can be achieved. For these smaller particles (Group I) a linear increase in nucleation undercooling with increasing cooling rate (logarithmic scale) was observed. Further, undercooling decreases with increasing particle size. This is similar to the observation on pure Sn particles [24], pure Al particles [17], Al–Si alloy particles [18] and Al–Cu alloy particles [25]. However, for particles with diameters larger than 23 \(\mu m\) (Group II), a transition of the undercooling dependence was observed, marked as regions i, ii and iii in Fig. 4b. The change in slope indicates a transition of the nucleation mechanism at a certain cooling rate, which is also dependent on the particle size. The two groups of undercooling behaviour are termed as Group I and Group II.

For rationalising the nucleation behaviour dependent on the cooling rate and particle size, a theoretical interpretation based on the classical nucleation theory (CNT) is proposed. For continuous cooling, the nucleation rate becomes time, \(t\), respectively, temperature, \(T\), dependent. Taking into consideration both heterogeneous mechanisms of bulk and surface nucleation, the total number of nuclei, \(N\), of a single Al-Si12 particle at a certain nucleation undercooling \(\Delta T\) (Eq. 1), at a constant cooling rate, \(\beta_c\), is given by [19]

\[
N = \int_0^t (J_V V + J_S S) dt = \frac{1}{\beta_c} \int_{T_{onset}}^{T_{onset} + \Delta T} (J_V V + J_S S) dT
\]

\[
= \frac{1}{\beta_c} \int_0^{\Delta T} (J_V V + J_S S) d\Delta T
\]

where \(J_V\) and \(J_S\) are the bulk and surface nucleation rates, respectively; and \(V = \pi d^3/6\), where \(d\) is the particle diameter and \(S = \pi d^2\) are the volume and surface area of a particle, respectively. It is assumed that at first only one nucleus forms at the temperature \(T_{onset}\), i.e. \(N = 1\). Thus, the nucleation undercooling \(\Delta T\) (and the temperature \(T_{onset}\)), as well as the time when the first supercritical nucleus forms, can be determined by Eq. (2).

Based on the spherical cap model of CNT, the expressions for the heterogeneous surface and bulk nucleation rates of a single particle during a rapid cooling process are derived in [35, 36] as

\[
J_S = n_S \frac{D_l}{a_0} \sqrt{\frac{\sigma_k}{k(T_{onset} - \Delta T)}} \exp\left(-\frac{\Delta G^* f_S}{k(T_{onset} - \Delta T)}\right)
\]

(3)

and

\[
J_V = n_V \frac{D_l}{a_0} \sqrt{\frac{\sigma_k}{k(T_{onset} - \Delta T)}} \exp\left(-\frac{\Delta G^* f_V}{k(T_{onset} - \Delta T)}\right)
\]

(4)

with

\[
\Delta G^* = \frac{16\pi\sigma_k^2}{3\Delta H_V}
\]

(5)

where \(n_S\) and \(n_V\) are the numbers of potential surface and bulk heterogeneous nucleation sites per unit surface area and volume (densities of surface and bulk heterogeneous nucleation sites), respectively, \(D_l\) is the liquid diffusivity, \(a_0\) is the atomic spacing, \(\sigma_k\) is the solid–liquid interfacial energy, \(k\) is the Boltzmann constant, \(\Delta G_V\) is the free energy difference between the melt and the solidified phase, and \(f_S\) and \(f_V\) are the shape factors for surface and bulk heterogeneous nucleation, which describe the potency of the heterogeneous nuclei. In the absence of specific heat capacity data, especially in the undercooled region, \(\Delta H_V\) is estimated from the melting enthalpy \(\Delta H_V\) and
the melting temperature $T_{\text{m onset}}$. $\Delta G_V$ can be approximated by the expression of Thompson and Spaepen [37]

$$\Delta G_V = \frac{\Delta H_V}{2} (\frac{T_{\text{m onset}} - \Delta T}{2T_{\text{m onset}} - \Delta T})$$  \hspace{1cm} (6)

Assuming that a surface nucleus for $x$-Al forms in a spherical cavity at the surface of the particle (surface nucleation on the Al$_2$O$_3$ oxide layer) or on a convex spherical interface inside the particle (bulk nucleation on impurities), as illustrated in Fig. 5a, the shape factors, $f_S$ and $f_V$, according to [38], can be expressed by

$$f_S = \frac{1}{4\rho_S} \left[ -1 + 2\rho_S + \sqrt{1 + \rho_S^2 + 2\rho_S \cos \theta_S} \right]$$

$$\left( 1 + \rho_S - \sqrt{1 + \rho_S^2 + 2\rho_S \cos \theta_S} \right)^2$$  \hspace{1cm} (7)

and

$$f_V = \frac{1}{4\rho_V} \left[ 1 + 2\rho_V - \sqrt{1 + \rho_V^2 - 2\rho_V \cos \theta_V} \right]$$

$$\left( -1 + \rho_V + \sqrt{1 + \rho_V^2 - 2\rho_V \cos \theta_V} \right)^2$$  \hspace{1cm} (8)

with

$$\rho_{SV} = \frac{r^*}{R_{SV}}$$  \hspace{1cm} (9)

where $\theta_S$ and $\theta_V$ are the contact angles between the undercooled melt and the heterogeneous nucleus for the surface and bulk heterogeneous nucleation, respectively; $R_S$ and $R_V$ are the radii of the spherical cavity and the convex spherical interface, respectively; and $r^*$ is the critical radius of a nucleus based on CNT, which is expressed by

$$r^* = \frac{2\sigma_b}{\Delta G_V}$$  \hspace{1cm} (10)

To complete the description of this model, the density of bulk heterogeneous nucleation sites can be taken to follow the form [39]

$$n_V = n_V^0 \exp(b\Delta T)$$  \hspace{1cm} (11)

where $n_V^0$ is the base level of nucleation sites ($= 10^{13}$ m$^{-3}$), and $b$ is a positive constant ($= 0.059$) [39]. For all the particles in this study, $n_V$ is in the range of $10^{14}$-$10^{16}$ m$^{-3}$. During the DFSC measurements, an inherent Al$_2$O$_3$ layer always exists. Therefore, assuming all the Al$_2$O$_3$ on the surface of the Al-Si12 particle are possible nucleation sites of surface heterogeneous nucleation, the density of surface heterogeneous nucleation sites can be calculated by

$$n_S = \frac{N_A h}{V_m}$$  \hspace{1cm} (12)

where $N_A$ is Avogadro constant, $h$ is the thickness of Al$_2$O$_3$ layer (= 1 nm) [40], and $V_m$ is the molar volume of Al$_2$O$_3$ (= 25.7 $\times 10^{-6}$ m$^3$ mol$^{-1}$) [40]. Thus, $n_S$ equals $2.343 \times 10^{19}$ m$^{-2}$.

Based on this model, the change of the nucleation activity of a certain heterogeneity for different undercooling can be estimated. For Al-Si12 particles, $T_{\text{m onset}} = 577.5$ °C, $\gamma_{st} = 0.169$ J m$^{-2}$ (Al in Al–Si system) [41], $\Delta H_V = 12.4 \times 10^5$ J m$^{-3}$ (measured by DSC for as-atomised Al-Si12 powders), $D_l = 5 \times 10^{-9}$ m$^2$ s$^{-1}$ [42], and $a_0 = 2.6 \times 10^{-10}$ m (for pure Al) [43]. Employing these parameters, the data were fitted for cooling rates ranging from 100 to 90,000 K s$^{-1}$. Here $\theta_S$, $\theta_V$, $R_S$ and $R_V$ are fitting parameters. Boundary conditions were set for each parameter: $20^\circ \lesssim \theta_{SV} \lesssim 180^\circ$ (for the spherical cap model, the contact angle is larger than 20° [44]); 0.1 nm $< R_{SV} < 10$ nm. Nonlinear least squares curve fitting (Levenberg–Marquardt algorithm) was utilised. The fitted curves are indicated by dashed lines in Fig. 4b and the results as well as the coefficients of determination, $R^2$, are displayed in Table 2. No value given in Table 2 means that this part of the model, i.e. bulk nucleation, does not fit the data. For particles with larger size (> 23 µm in diameter, Group II), two nucleation mechanisms are suggested, while for smaller size particles (< 23 µm in diameter, Group I) only surface heterogeneous nucleation dominates. For each particle, the contact angles of surface and bulk nucleation are the same for different cooling rates: approximately 103°–114° and 34°, respectively. The contact angle of surface nucleation determined is consistent with the contact angle between Al and Al$_2$O$_3$ in [45]. This supports the assumption of surface nucleation occurring on the Al$_2$O$_3$ oxide layer. For bulk nucleation, the nuclei probably form on impurities inside the particle.

The dependence of the nucleation undercooling on the cooling rate for particles of 7 µm, 17 µm and 27 µm in diameter is shown in Fig. 5b. Undercooling for the particle of 27 µm in diameter is determined by two different nucleation mechanisms. Similar behaviour was observed for pure Sn particles solidified at
cooling rates from 20 to 40,000 K s\(^{-1}\) \cite{24, 46}. Based on the theoretical approach mentioned above, surface heterogeneous nucleation is dominant at low cooling rates, and is replaced by bulk heterogeneous nucleation: 

**Figure 5**  
(a) Schematic of the nucleation mechanisms of rapidly solidified Al–Si particles.  
(b) Dependence of undercooling on cooling rate for studied particles of 7 µm, 17 µm and 27 µm in diameter: solid curves—surface and bulk nucleation, dashed curves—surface nucleation, short dashed curves—bulk nucleation.  
(c) Dependence of the surface and bulk nucleation rates on the undercooling for the studied particle of 27 µm in diameter: solid curve—surface nucleation, dashed curve—bulk nucleation.
nucleation at high cooling rates. In between, there is a transition region, e.g. from 300 to 1000 K s\(^{-1}\) for the particle of 27 \(\mu\)m in diameter, as shown in Fig. 5b. In order to understand this behaviour, the dependence of the heterogeneous surface and bulk nucleation rates on the undercooling is calculated by Eqs. (3 and 4) with the fitted parameters listed in Table 2. The results are shown in Fig. 5c. At low undercoolings/cooling rates, the bulk heterogeneous nucleation rate is lower than the surface heterogeneous nucleation rate, and vice versa. Therefore, surface heterogeneous nucleation is preferred at low cooling rates, whereas bulk heterogeneous nucleation is preferred at high cooling rates. For the particles of 7 \(\mu\)m and 17 \(\mu\)m in diameter, surface nucleation dominates in the whole investigated cooling rate range. With decreasing particle size, fewer impurities are expected inside the particle [35, 47], i.e. fewer bulk heterogeneous nucleation sites, as illustrated in Fig. 5a. For example, the number of bulk nucleation sites is even less than 1 inside the particle of 6 \(\mu\)m in diameter, calculated by Eq. (11). Therefore, surface heterogeneous nucleation dominates for small particles.

**Table 2** Parameters obtained by fitting the model to different particle sizes

| \(d\) (\(\mu\)m) | \(\theta_s\) (°) | \(R_s\) (nm) | \(\theta_V\) (°) | \(R_V\) (nm) | \(R^2\) |
|-----------------|---------------|-------------|---------------|-------------|--------|
| 6               | 103.10        | 0.96        |               |             | 0.999  |
| 7               | 104.66        | 0.95        |               |             | 1.000  |
| 9               | 104.97        | 0.94        |               |             | 0.996  |
| 14              | 106.56        | 0.92        |               |             | 0.996  |
| 15              | 103.09        | 0.98        |               |             | 0.992  |
| 17              | 105.16        | 0.94        |               |             | 0.982  |
| 18              | 113.60        | 0.81        |               |             | 0.999  |
| 21              | 113.23        | 0.82        |               |             | 0.999  |
| 23              | 112.31        | 0.82        | 34.28         | 7.45        | 0.954  |
| 27              | 112.26        | 0.82        | 33.90         | 7.45        | 0.975  |
| 28              | 112.45        | 0.82        | 34.02         | 7.45        | 0.990  |

for the two undercooling groups which were solidified at 100 K s\(^{-1}\), 500 K s\(^{-1}\), 5000 K s\(^{-1}\) and 50,000 K s\(^{-1}\), respectively. Although DFSC data could not be evaluated properly for large particles and high cooling rates (e.g. 31 \(\mu\)m at 50,000 K s\(^{-1}\)) due to thermal lag, particle microstructures from rapid solidification experiments at high cooling rates still provide useful information understanding the rapid solidification behaviour of Al-Si\(_{12}\) powder particles. The particle diameters and associated undercooling from DFSC are shown in Fig. 6 for Group I (left part, from top to bottom): 20 \(\mu\)m and 89 K, 22 \(\mu\)m and 113 K, 19 \(\mu\)m and 115 K, 21 \(\mu\)m and 137 K, respectively. For Group II (Fig. 6, right part, from top to bottom), the particle diameters and associated undercooling from DFSC are 33 \(\mu\)m and 76 K, 34 \(\mu\)m and 91 K, 32 \(\mu\)m and 93 K, and 31 \(\mu\)m and 96 K, respectively. In all cases, the microstructure consists of \(\alpha\)-Al dendrites and (\(\alpha\)-Al + Si) eutectic due to the coupled zone [33, 34], where the grey and bright areas are Al and Si, respectively. As expected, the microstructure becomes finer with increasing cooling rate. It can be concluded in Fig. 6 (left, Group I) that the \(\alpha\)-Al dendrites have formed from the particle surface. This result confirms the surface heterogeneous nucleation, which is consistent with the suggested theoretical model based on the DFSC results. For example, for the particle of 19 \(\mu\)m in diameter and 115 K undercooling (left middle in Fig. 6), the microstructure is similar for the case in which nucleation events occur at the surface as illustrated in [13, 48] schematically. During cooling, the particle nucleated from the surface (left side of the particle), leading to finer microstructure of \(\alpha\)-Al in the left side of the particle than that in the right side. As the solidification proceeded further away from the nucleation region, the microstructure of \(\alpha\)-Al becomes coarser due to the influence of the recalescence effect inside the particle. Similar behaviour can be found for the particle of 22 \(\mu\)m and 113 K. For Group II (Fig. 6, right), different microstructural features were observed. With increasing cooling rate, \(\alpha\)-Al dendrites have formed simultaneously at the surface and on inclusions inside the particle. For example, for the particle of 31 \(\mu\)m and 96 K (right bottom in Fig. 6), \(\alpha\)-Al dendrites grow from the surface as well as from an inclusion in the particle volume (marked by a red circle). This is consistent with the transition of the nucleation mechanism from surface heterogeneous nucleation to bulk heterogeneous nucleation.

**Microstructure of single rapidly solidified particles at controlled cooling rates**

In order to verify the nucleation mechanisms of the primary \(\alpha\)-Al phase in Al-Si\(_{12}\), the microstructures of single particles treated by DFSC at different cooling rates were investigated by SEM. Figure 6 shows the cross-sectional SEM images of single Al-Si\(_{12}\) particles...
Moreover, the increasing cooling rate contributes to finer Si in the eutectic, while the Si morphology seems to change from more fibrous to more globular.

To examine the change of morphologies of the primary α-Al in more detail, the α-Al DAS is plotted in Fig. 7a depending on the cooling rate. Each point corresponds to one particle shown in Fig. 6. The solidification microstructures are dependent on the cooling rate and nucleation undercooling of the primary α-Al. When the particle solidifies at high cooling rates with large nucleation undercooling, the nucleation rate of primary α-Al increases, and the solidification time is shorter. Thus, the α-Al DAS is reduced, and the microstructure is finer, which can
promote the alloy properties. Moreover, the primary α-Al size in the smaller particles (ca. 20 μm in diameter, Group I) is smaller than that in the larger particles (ca. 32 μm in diameter, Group II) at a particular cooling rate.

In general, the relationship between the dendrite arm spacing and cooling rate can be expressed by

\[ \lambda = B \beta_c^{-n} \]  \hspace{1cm} (13)

where \( \lambda \) is the dendrite arm spacing in μm, and \( B \) and \( n \) are alloy-dependent fitting parameters. The fitted curves are also plotted in Fig. 7 for the two undercooling groups I and II. \( B \) and \( n \) of rapidly cooled aluminium alloys via splat quenching were reported to equal 43.2 and 0.324 [49], which is also plotted in Fig. 7a for comparison. It looks as if the influence of the cooling rate on the DAS is underestimated in the present work, compared to [49]. Such low values of the exponent \( n \) (0.08 and 0.09 for Groups I and II, respectively) indicate that the DAS is not only governed by coarsening, but also influenced by the high nucleation undercooling and heat distribution effects in the constrained volume of the particle. In splat quenching as well as AM, the solidified volume is significantly larger than that of individual particles. In this case, bulk nucleation is more relevant than surface nucleation. This is in general agreement with the results in Fig. 7, where the DAS decreases with increasing cooling rate. The results in [49] are close to the Group II results at a high cooling rate of 50,000 K s\(^{-1}\). Deviations are attributed to the differences between splat quenching and single particle solidification as described above.

To represent AM microstructures, in which bulk heterogeneous nucleation dominates, it is suggested that Group II particles, i.e. particles larger than about 30 μm, should be used in DFSC. Further, the dependence of the α-Al DAS on the nucleation undercooling is shown in Fig. 7b. The α-Al DAS decreases rapidly in the moderate undercooling range below ca. 95 K, and then decreases slowly at larger undercoolings. This result indicates that the α-Al DAS may be related not only to the cooling rate but also to the nucleation undercooling and volume constrictions. In comparison, similar α-Al DAS of about 1 μm surrounded by (α-Al + Si) eutectics was observed in AM components, e.g. LPBF of AlSi10Mg [50, 51], laser deposited Al4047 [52] and LPBF of Al-Si12 [53]. This proves the feasibility of DFSC studies at larger undercoolings (above ca. 95 K) and higher cooling rates (above ca. 10,000 K s\(^{-1}\)) to represent typical AM microstructures.

**Figure 7** α-Al dendrite arm spacing as functions of the cooling rate **a** and the nucleation undercooling **b** for single Al-Si12 particles as shown in Fig. 6. The results from rapidly cooled aluminium alloy by Matyja et al. [49] are also plotted in **a**. The dashed curves in **a** show the fitted curves by Eq. (13). The dashed line in **b** is a guide for the eye. The error bar shows the standard deviation.

**Conclusions**

The rapid solidification process of single micro-sized Al-Si12 (mass%) particles was successfully investigated in a wide range of cooling rates from 100 to 90,000 K s\(^{-1}\) by in situ DFSC. With increasing cooling rate, the nucleation undercooling increases for all particles irrespective of their diameter. According to a modified theoretical model based on CNT, the α-Al nuclei either form at the surface (Al₂O₃ layer) or on impurities inside the Al-Si12 particle. For smaller size particles (< 23 μm in diameter), surface heterogeneous nucleation dominates in the whole
investigated cooling rate range. For single Al-Si12 particles with larger size (> 23 μm in diameter), the
dominant nucleation mechanism of the primary α-Al phase changes from surface to bulk heterogeneous
nucleation with increasing cooling rate.

Undercooling of rapidly solidified single micro-sized particles treated by DFSC was correlated with a
microstructure investigation. The results reveal that the solidification microstructure consists of fine
primary α-Al phase and (α-Al + Si) eutectic. Moreover, the dendrite arm spacing (DAS) is found to notice-
ably decrease as the cooling rate/undercooling increases. The morphology change of the primary α-
Al phase gives some hints on the nucleation mechanism (surface/bulk) change with cooling rate, as
described by the theoretical model.

Combining in situ DFSC with microstructure characterisation, it is possible to observe rapidly
solidified structures of single metallic particles by controlled rapid cooling and known nucleation
undercooling. The results indicate that large nucleation undercoolings above 95 K, high DFSC cooling
rates above about 10,000 K s⁻¹ and particle diameters above about 30 μm should be used to represent AM
microstructures successfully. Therefore, this is an effective approach for the investigation of the influ-
ence of the cooling rate and undercooling on microstructure evolution under conditions of rapid
solidification, such as in AM processes.

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Declarations

Conflict of interest  The authors declare that they
have no conflict of interest.

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Appendix

Temperature calibration and correction
for DFSC measurements

Primary temperature calibration

Primary temperature calibration of the whole DFSC
instrument (including the thermopile of the sensor,
electronics, etc.) was performed prior to the experi-
ments by measuring the phase transition of standard
material, in other words to obtain the function of the
thermopile voltage reading from the instrument in
dependency on the certified temperatures of the
standard under given experimental conditions (e.g.
heating rate equals zero, certain purge gas, given
surrounding temperature, etc.). The temperature of
the central area of the sensor is influenced by the
surrounding temperature of the sensor housing,
which was at room temperature in this study, and the
measured thermopile voltage originating from the
temperature difference between hot and cold junc-
tions [54]. This voltage has to be calibrated in the
temperature range from room temperature up to the
required maximum temperature, which was 707 °C in this study. The temperature was therefore calibrated under the same experimental conditions as in the solidification measurements, applying the melting temperatures of indium (156.6 °C), tin (231.9 °C) and zinc (419.5 °C), and the solid-state transition temperature of K2SO4 (585 °C). It should be pointed out that the thermopile voltage at the phase transition temperature of a calibration material is correlated with a heating rate of 0 K s⁻¹. This value was obtained from the measured thermopile voltage of the phase transition at different heating rates, extrapolating to the zero heating rate (e.g. see Fig. 8). Then, the functional relationship between the thermopile voltage and temperature was approximated by a quadratic polynomial function [54, 55],

\[ E = c_0 + c_1 T + c_2 T^2 \]  

(14)

where \( E \) is the thermopile voltage, \( T \) is the temperature and \( c_0, c_1 \) and \( c_2 \) are fitting parameters determined as 0, 6.5 \times 10⁻⁴ and 4.2 \times 10⁻⁷, respectively. With the resulting function, extrapolated to 707 °C, the DFSC instrument was calibrated for the temperature range from room temperature to 707 °C. The sensors used in this study came from the same wafer and were therefore almost identical. Consequently, this calibration function was used for all DFSC measurements.

### Temperature correction for each individual Al-Si12 particle

Apart from primary temperature calibration of the DFSC instrument, the measured temperature should also be corrected for each individual particle. Therefore, before investigating the rapid solidification of each particle by DFSC, the individual particle was heated and cooled at different rates (from 1000 to 10,000 K s⁻¹) from 37 to 707 °C and back to 37 °C under an argon atmosphere at a constant pressure of 100 kPa. As an example, the measured curves at different heating–cooling rates from 1000 to 10,000 K s⁻¹ for one single Al-Si12 particle are shown in Fig. 8a. As expected, the leading edges of the melting peaks overlap due to the thermal resistance of the sensor/sample system, which limits the heat flow during melting [28, 56]. The measured temperature at different scanning rates has to be corrected regarding three sources of temperature errors:

(i) The temperature gradient inside the particle due to the limited heat conductivity of the particle. For metallic particles, this temperature gradient is relatively small (a few Kelvins) [46, 57]. Therefore, this error is neglected for the temperature correction in this study.

(ii) The dynamic temperature offset (\( \Delta T_d \), due to thermal lag), which can be expressed as in [28, 58] for heating

\[ \Delta T_d = \tau_{lag} \rho_n \]  

(15a)

and for cooling...
\[ \Delta T_d = -\tau_{\text{lag}} \beta_c \]  

(15b)

where \( \beta_h \) and \( \beta_c \) are the heating and cooling rates, and \( \tau_{\text{lag}} \) is the thermal lag. The thermal lag is determined by the thermal resistance, i.e. the thermal contact, between the particle and the sensor and the thermal resistance inside the sensor membrane. For Al-Si12 particles in the micron range, no size or rate dependence for the eutectic temperature, i.e. the melting onset temperature, is expected. According to Eq. (15a), the dynamic temperature offset \( \Delta T_d \) increases linearly with the rate. Consequently, the apparent melting onset temperature measured should also increase linearly with the increasing heating rate. Figure 8b shows this dependency for a 13 \( \mu \)m diameter particle. The apparent melting onset temperatures are on a straight line, confirming the validity of Eq. (15a) for the case under investigation.

(iii) The static temperature offset due to a balance between the heat provided by the heater and the heat losses from the sample to the surroundings, even under isothermal conditions [56, 59]. For example, for a spherical particle, the larger additional heat-exchanging surface compared to a flat sample causes significantly higher heat losses into the surrounding gas. Besides, the small contact area between the spherical particle and the membrane results in a relatively large thermal resistance between the sample and membrane. In addition, the particle position relative to the sensor heater and the thermopile also has an influence. If the particle is placed away from the heater, say close to the thermopile in the centre, the sample temperature will be lower compared to a situation where the particle is closer to the heater. This offset is independent of the heating or cooling rate and can reach some few tens of Kelvin. Therefore, it is important to place the sample properly and to correct this offset after the measurements. For the Al-Si12 alloy, particles start to melt at the eutectic temperature during heating, i.e. the melting onset temperature equals the eutectic temperature. Therefore, it is assumed that the static temperature offset at this temperature is the offset between the true eutectic temperature of the Al–Si alloy and the measured apparent melting onset temperature at 0 K s\(^{-1}\). The eutectic temperature can be found in Ref. [27] (577 ± 1 °C) or measured by differential scanning calorimetry (DSC) (577.5 °C for the as-atomised Al-Si12 powders investigated here). The measured melting onset temperature at 0 K s\(^{-1}\) can be obtained from the melting onset temperatures measured at different heating rates extrapolating to zero heating rate by Eq. (15a). For the particle shown in Fig. 2, this offset is \(-11 \)K (see Fig. 8b: axis intercept 566.5–577.5 °C). In other words, the extrapolated \((\beta_h = 0)\) measured melting onset temperature for the particle shown in Fig. 2 is lower than the real temperature. A possible reason is that the particle is “thermally” much closer to the heater than to the thermopile, as shown in Fig. 2. Hence, particle placement is essential for accurate absolute temperature measurements for such particles. For simplicity and because of the dominance of heat conduction [57], it is assumed that the static offset is proportional to the temperature difference between room temperature and the measured apparent temperature. Thus, the static offset, \( \Delta T_s \), is expressed by

\[ \Delta T_s = \left( T_{m_{\text{DSC}} - \text{onset}} - T_{m_{\text{DSC}} - \text{onset}} \right) \frac{T_{\text{measure}} - T_{RT}}{T_{m_{\text{DSC}} - \text{onset}} - T_{RT}} \]  

(16)

where \( T_{m_{\text{DSC}} - \text{onset}} \) and \( T_{m_{\text{DSC}} - \text{onset}} \) (577.5 °C for as-atomised Al-Si12 powders) are the melting onset temperatures at 0 K s\(^{-1}\) heating rate measured by DFSC and DSC, \( T_{\text{measure}} \) is the measured thermopile (apparent) temperature and \( T_{RT} \) is room temperature.

Each individual measured particle was different in size, position on the sensor and thermal contact. Therefore, the measured temperature of each particle must be corrected in a proper manner. Since solidification (nucleation) is a stochastic process, a comparison of the characteristic temperature of solidification obtained during cooling with the respective literature data is impossible. Instead, assuming good symmetry of the DFSC instrument (in terms of thermal behaviour during heating and
cooling [28, 58]), the correction function (Eq. 15a) on heating can be utilised on cooling as well (Eq. 15b). Thus, the measured apparent onset temperatures during heating and cooling can be evaluated with respect to a correction function:

for heating

\[
T_{\text{correct}} = T_{\text{measure}} - \tau_{\text{lag}} \beta_h - \left( \frac{T_{m_{\text{onset}}}}{T_{m_{\text{onset}}}} - \frac{T_{KF}}{T_{KF}} \right) \tag{17a}
\]

and for cooling

\[
T_{\text{correct}} = T_{\text{measure}} + \tau_{\text{lag}} \beta_c - \left( \frac{T_{m_{\text{onset}}}}{T_{m_{\text{onset}}}} - \frac{T_{KF}}{T_{KF}} \right) \tag{17b}
\]

where \(T_{\text{correct}}\) is the corrected temperature. The measured apparent melting and solidification onset temperatures were therefore corrected by Eqs. (17a and 17b).

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