Phase Transformations in Thermally Treated Gas-Atomized Al 7075 Powder

Caitlin Walde1 · Kyle Tsaknopoulos1 · Victor Champagne2 · Danielle Cote1

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Abstract
Al 7075 is a heat-treatable Al–Mg–Zn alloy widely used in the aerospace industry. Recently, it has found application as feedstock for metal additive manufacturing (MAM). It has been shown that wrought alloy compositions in powder form differ in microstructure and properties from their conventional form. Given this, it is important to understand the microstructure of the powders prior to use in MAM processes. This work studies as-atomized gas-atomized Al 7075 powders and the effect of thermal treatments on microstructure. Extensive electron microscopy revealed the presence of T-phase, Al7Cu2Fe, and Mg2Si in the as-atomized condition. Thermal treatments were performed at 465 °C and 480 °C to homogenize the microstructure; however, S-phase was unexpectedly present in the samples treated at 465 °C. In both 465 °C and 480 °C treatments, T-phase was not fully dissolved after the 60-min treatment. Guided by thermodynamic modeling, these results indicate a shift in local equilibria in these powders.

Keywords Powder metallurgy · Modeling · Microstructure · Aluminum alloys

Introduction
Al 7075 is a heat-treatable Al–Mg–Zn alloy that is widely used in the aerospace industry in compressively loaded parts for its combination of high specific strength and resistance to stress corrosion cracking [1]. Al 7075 is available in multiple forms; it is typically cast and then worked to a wrought condition, though it is also available in wires and powders. The latter forms are used mainly as feedstock in metal additive manufacturing (MAM) processes. Powder-bed processes are more common, with selective laser melting (SLM), powder bed fusion (PBF), and selective laser sintering (SLS) having perhaps the most widespread use [2]. In liquid-state processes, where the powder is melted during consolidation, features present in the feedstock powder—such as surface oxides and internal porosity—can be retained during consolidation, while the rest of the features are melted and reform. In solid-state processes, where the powder is not melted during consolidation, the retention of feedstock characteristics is more significant. Features such as secondary phases, in addition to surface oxides and internal porosity, can be retained during consolidation. For example, Rokni et al. [3] showed that the feedstock powder microstructure is retained in cold spray deposits. With this in mind, it is important to understand the microstructure of the powders prior to consolidation [4, 5].

The feedstock powders are often gas-atomized, experiencing cooling rates on the order of $10^6$ °C/s during solidification [6]. This categorizes the powders as rapidly solidified, which results in microstructures very different than conventional castings, where the cooling rate is on the order of less than $10^2$ °C/s [5]. Minimal segregation, nonequilibrium phases, and microstructure homogeneity are characteristics of rapidly solidified structures [6–8].

The three primary steps in a heat treatment are solutionization, quenching, and aging [9]. The purpose of the solutionizing step is to dissolve secondary phases and achieve a homogenous microstructure; this is typically performed at elevated temperatures in the 450–550 °C range, depending on the alloy. This step is followed by a rapid quench, to maintain the supersaturated metastable microstructure at room temperature. The aging step is designed to homogeneously nucleate and grow the primary strengthening phase...
in the alloy; this is usually performed at lower temperatures in the 150–250 °C range, depending on the alloy. For Al 7075, the strengthening sequence is GP zones (MgZn) → η′ → η → (MgZn2) → T′ → T → (Al2Mg3Zn3) [9]. Whether the sequence finishes at η′ or at T-phase depends on the Mg/Zn ratio. Natural aging typically continues for years, so artificial aging steps are important to achieve stable precipitates. Additionally, as Fe and Si are impurities in Al, Al7Cu2Fe, Al12(Fe,Mn)3Si, Al6(Fe,Mn), Mg2Si, SiO2, and Al23CuFe4 are also common phases in Al 7075, in addition to the η and T strengthening phases. Depending on the exact alloy compositions, some of these phases may be insoluble. Of note is that Mg2Si is virtually insoluble in Al 7075 [1].

Because features in the powder are retained during consolidation, the as-atomized microstructure can be controlled using thermal treatments. Recent research has shown the retention of feedstock powder microstructure during consolidation for other alloys, though that research included a limited analysis of the internal microstructure of the powders, not fully investigating the effect of the thermal treatments on the powders [10–14]. These researchers assumed the network of solute elements at the boundaries in the as-atomized structure, as seen in scanning electron microscopy (SEM) and energy-dispersive X-ray spectroscopy (EDS), was segregation. While the presence of secondary phases was noted in the thermally treated powders, no further in-depth analysis was performed to identify the phases. However, in-depth analysis here indicates the as-atomized microstructure of the powders is actually distinct phases, as revealed using transmission electron microscopy (TEM), consistent with what is expected in rapid solidification [4, 15]. Additionally, it was recently noted that these gas-atomized powders have both granular and subgranular features. The secondary phases exist at all boundaries, with groups of subgrains having the same orientation, comprising larger grains [4, 5, 15]. The size of the substructure is typically what is reported for these powders; when etched, it is impossible to distinguish between the high-angle and low-angle boundaries that differentiate the grains and subgrains.

Solutionization times for Al 7075 are typically on the order of 4–16 h for wrought components, depending on part size [9]. Given this timescale, other researchers chose to use long heat treatment times for their powders; however, other researches performed on Al 6061 indicated faster diffusion times in powders, resulting in the need for shorter thermal treatments [5, 10].

This research investigates the effect of thermal treatment on the transformations of the secondary phases present in gas-atomized Al 7075 powders through the use of TEM and EDS.

| Element | Wt.% | ASTM B209 |
|---------|------|-----------|
| Cr      | 0.210| 0.18–0.28 |
| Cu      | 1.700| 1.2–2.0   |
| Fe      | 0.180| 0.50      |
| Mg      | 2.300| 2.1–2.9   |
| Mn      | 0.031| 0.30      |
| Si      | 0.110| 0.40      |
| Ti      | 0.031| 0.20      |
| Zn      | 5.600| 5.1–6.1   |
| Other   | 0.15 |           |
| Al      | 89.8 | Remainder |

Fig. 1 Equilibrium diagram, as predicted by thermo-calc simulations. Amount of phases present as a function of temperature for the specific composition of the alloy investigated given in Table 1

**Method**

The studied powder was gas-atomized Al 7075 (Valimet, Inc., Stockton, CA) with a \( d_{10} \) of 26.1 μm, \( d_{50} \) of 37.1 μm, and \( d_{90} \) of 52.5 μm. The composition of the powders, as determined by direct-current plasma emission spectroscopy and shown in Table 1, is within the acceptable ranges for Al 7075 [16, 17].

Thermal treatment parameters were determined with the assistance of computational thermodynamic and kinetic software (Thermo-Calc, Stockholm, Sweden), shown in Fig. 1. Thermal treatments were performed with the goal of homogenizing the microstructure—dissolving T-phase while avoiding S-phase growth, avoiding Al7Cu2Fe coarsening, and avoiding melting. The temperature of 465 °C was selected to reduce the coarsening effect of Al7Cu2Fe by maintaining a lower temperature to slow diffusion while still avoiding the formation of S-phase, which is present below 450 °C. The temperature of 480 °C was chosen.
because it is the conventional solutionization temperature used for Al 7075. Treatment times ranged from 0 to 4 h. The “as-atomized” indicates an untreated condition, while the thermally treated conditions are noted by the time at treatment temperature; the thermally treated samples all experienced the ramp to elevated temperature. Samples were treated in a nitrogen environment in a differential scanning calorimeter (DSC) because of its high thermal accuracy. One sample was left untreated, in the as-atomized condition.

Samples were prepared for scanning electron microscopy (SEM) and energy-dispersive X-ray spectroscopy (EDS) analyses by mounting the powders in a hot-cure epoxy and grinding and polishing incrementally with a final 0.25 μm colloidal silica suspension.

SEM and EDS analyses were performed on all samples using a tungsten-filament SEM at 10-15 kV. Powders in both the as-atomized and thermally treated conditions of 465 °C for 60 min and 480 °C for 60 min were prepared for transmission electron microscopy (TEM) analysis using focused ion beam (FIB) milling (Scios DualBeam and Helios 660 Nanolab, Thermo Fisher Scientific, Waltham, MA) [15]. Lamellae were polished with a final beam current of 0.1 nA and had a final thickness of less than 200 nm. TEM analysis was performed using a probe-corrected TEM (Titan Themis with ChemiSTEM, Thermo Fisher Scientific, Waltham, MA) and energy-dispersive X-ray spectroscopy (EDS) (Super-X, Thermo Fisher Scientific, Waltham, MA) at an accelerating voltage of 300 kV. The amounts of secondary phases present in the conditions were quantified using image contrast thresholding. All EDS maps are in wt. %.

Additionally, the as-atomized condition was evaluated using electron backscatter diffraction (EBSD). FIB was used to prepare powder particles for EBSD analysis, and an accelerating voltage of 20 kV with a step size of 0.3 μm was used for analysis.

**Results and Discussion**

Figure 2 displays an overview SEM micrograph and EBSD micrograph of the as-atomized condition. The continuous network structure present at both the granular and substructure boundaries, as seen in the SEM micrograph, in the as-atomized condition is consistent with what is seen in other gas-atomized powders [15]. Note that when orientation is evaluated, as with EBSD in Fig. 2b, there are regions of the substructure that have the same orientation and comprise grains, again consistent with what is seen in similar powders [4, 5, 15].

Figure 3 shows overview SEM micrographs of the thermally treated conditions. As seen in Fig. 3a and b, the network structure present in the as-atomized condition begins to dissolve simply by bringing the powder up to an elevated temperature. With increased treatment times, note the presence of both larger phases and smaller phases in two different regions; the large phases are present at the grain boundaries, such as seen in Fig. 2b, while the smaller phases are present at only the substructure boundaries.

Figure 4 shows the area percent of secondary phases present as a function of treatment time for both treatment temperatures.

In the treatment performed at 465 °C, there is an initial decrease in secondary phase amount after the sample reaches the set temperature, consistent with Fig. 3. Then, there is a slight decrease in amount of light-contrasting secondary phases after 20 min of treatment, stabilizing after 60 min of treatment. The dark-contrasting phases, while not detected in the as-atomized condition, are stable in very low amounts across all treatment temperatures.

In the treatment performed at 480 °C, there is an initial decrease in secondary phases after increasing the sample to the set temperature, consistent with Fig. 3d, after which there is a slight increase in the light-contrasting phases.
until 60 min of treatment. After 60 min, the light-contrasting phases decrease. The dark-contrasting phase, while not detected in the as-atomized condition, is stable across all treatment temperatures.

When comparing the two treatment temperatures, it is of note that the dark-contrasting phase area percent is the same, whereas the light-contrasting phase trends vary. There is a similar reduction in the light-contrasting phase in both treatments when at the set temperature. However, after approximately 20 min of treatment, the trends diverge; at 480 °C, the light-contrasting phase increases, while at 465 °C the light-contrasting phase decreases. After approximately 120 min of treatment, the trends converge again, appearing to reach an equilibrium.

Due to the large interaction volume in the SEM, EDS was not successful in further differentiating the light- and dark-contrasting phases into the constituents predicted in Thermo-Calc. Hence, TEM evaluation was performed. Diffraction was attempted, but due to the small size of the phases and overlap in diffraction patterns of the phases, Al matrix was unsuccessful in confirming the phases. EDS was performed in a TEM to better understand which phases may be forming.

Figure 5 shows overview TEM micrographs of the as-atomized and thermally treated conditions for 60 min at each temperature. Figure 5a shows a similar structure as seen in the SEM in Fig. 2a; on the other hand, Fig. 5b and c provides much more detail than their SEM counterparts in Fig. 3. Figures 6, 7 and 8 show higher magnification TEM micrographs and corresponding EDS maps to give further insight into the specific constituents present in each condition.

Figure 6 shows TEM micrograph and EDS maps for the as-atomized condition. Note the presence of two phases—an Al-Cu-Mg-Zn network phase and discrete Fe-rich phases. EDS point quantification results are shown in Table 2, while the predicted compositions are shown in Table 3. Figure 1 is consulted to evaluate which phases may be present, and then, Tables 2 and 3 are compared, looking for overlaps in proportions of elements present. For example, the network phase at the boundaries in the as-atomized condition is rich in Al–Cu–Mg–Zn, and T-phase and V-phase are the only phases that fit the composition for that but T-phase is the only phase predicted at equilibrium (Fig. 1). Additionally, discrete Fe-rich phases present at the boundaries are also rich with Al and Cu. Consulting Table 3, it can be seen that Al2Cu3Fe is the only phase rich in just Al, Fe, and Cu. This indicates the Al–Cu–Mg–Zn network is likely T-phase and...
the discrete Fe-rich phases are likely Al$_7$Cu$_2$Fe, which is consistent with what is seen in literature [1]. Additionally, the dark-contrasting phase seen in the thermally treated SEM micrographs (Figs. 3 and 4) is also found in the as-atomized structure using TEM and has been identified as Mg$_2$Si through similar comparison of Tables 2 and 3.

Figure 7 shows TEM micrograph and EDS maps for the sample thermally treated at 465 °C for 60 min. Note the presence of four phases—an Al–Cu–Mg–Zn phase, an Al–Cu–Mg phase, an Fe-rich phase, and a Si-rich phase. EDS point quantification results are shown in Table 2. The compositions of the predicted secondary phases, shown in Table 3, compared to the results in Table 2, indicate the Al–Cu–Mg–Zn phase to likely be residual T-phase that had not fully dissolved, the Al–Cu–Mg phase to be S-phase, the Fe-rich phase to be the Al$_7$Cu$_2$Fe, and the Si-rich phase to be Mg$_2$Si. Based on the equilibrium diagram in Fig. 1, S-phase is not expected to be present at this temperature; this indicates a difference in the local equilibrium for the powder in comparison with the global powder composition as measured by direct-current plasma emission spectroscopy.

Figure 8 shows a TEM micrograph and EDS maps for the sample thermally treated at 480 °C for 60 min. Note the presence of three phases—an Al–Cu–Mg–Zn phase, an Fe-rich phase, and a Si-rich phase. EDS point quantification is shown in Table 2. The compositions of the predicted secondary phases, shown in Table 3, compared to the results in Table 2, indicate the Al–Cu–Mg–Zn phase to likely be residual T-phase that had not fully dissolved, the Fe-rich phase to be the Al$_7$Cu$_2$Fe, and the Si-rich phase to be Mg$_2$Si. Note the lack of the S-phase that was present in the 465 °C thermally treated condition.

An unexpected similarity between the treatment temperatures is the presence of T phase seen in Figs. 7 and 8. The equilibrium predictions indicate that T-phase should not be present at either temperature; this indicates that 60 min of treatment at either temperature is not the sufficient time to dissolve the T-phase. As Scheil simulations predict T-phase to be the most abundant phase, it is expected that it may take additional time to dissolve. Unfortunately, it is not possible to distinguish between the T-phase and Al$_7$Cu$_2$Fe in the SEM (Fig. 3), so further TEM work would be needed to determine whether 120 min or 240 min of treatment is sufficient to fully dissolve T-phase.

Additionally, large and small phases can be seen in both the SEM and TEM micrographs; they have been identified as both T-phase and Al$_7$Cu$_2$Fe, with sizes corresponding to different boundary types. The high-angle boundaries have more open space and hence the phases can be larger there, whereas the phases at the low-angle boundaries have less open space for growth.

Of significant note is the difference in precipitate locations between the 465 °C and 480 °C treatment temperatures;
Fig. 6 TEM micrograph and EDS maps for the as-atomized condition. (a) HAADF, (b) Al, (c) Cu, (d) Fe, (e) Mg, (f) Si, and (g) Zn

Fig. 7 TEM micrograph and EDS maps for the 60 min at 465 °C condition. (a) HAADF, (b) Al, (c) Cu, (d) Fe, (e) Mg, (f) Si, and (g) Zn
the 465 °C treatment has precipitates both at the boundaries and in the bulk of the grains, whereas the 480 °C only has precipitates at the boundaries. Diffusion and nucleation energy are both driving forces in determining both the type and location of precipitates that form. Because the grains are so small in these gas-atomized powders and there is such a small diffusion distance to the substructure boundaries from the center of the grains, the driving force for precipitation on the boundaries is easily achievable; thus, precipitation will occur first and most frequently on the boundaries. However, it is possible for the boundaries to become saturated, forcing precipitation to occur within the bulk of the matrix phase.

This is presumed to be the case in the 465 °C treatment. The boundaries are saturated with T-phase, which is in the process of dissolving, and with Al₇Cu₂Fe, so the S-phase can only precipitate in the bulk matrix as Al₇Cu₂Fe is more stable at this temperature. This can be seen in both the SEM and TEM micrographs. In order to prove the precipitation sequence, additional SEM or TEM studies should be performed of heat treatments leading up to 465 °C. Ideally, in situ hot-stage STEM studies could be performed on a grain boundary, similar to the work of Vijayan et al. [18], which would enable direct observation of precipitation and dissolution.
Given the increase in understanding of the phases present in each condition from study in TEM, it is beneficial to revisit Fig. 4 to further understand the trends found in the SEM analysis. Recall the increasing trend in the light-contrasting phases in the 480 °C treatments at 60 min; in TEM, two different light-contrasting phases were identified at the boundaries: Al₇Cu₂Fe and T-phase. As previously stated, the T phase is in the process of dissolving, so this increase is likely due to the coarsening of the Al₇Cu₂Fe at a faster rate than the T-phase dissolution. It is expected that after 120 min of treatment, the T phase would be fully dissolved and the Al₇Cu₂Fe reached a stable size and fraction, reflected in the plateau in Fig. 4. In comparison, at 60 min of treatment at 465 °C, there is a decreasing trend in the light-contrasting phases in the SEM. This trend is surprising because based on the TEM results, there are three different light-contrasting phases present—Al₇Cu₂Fe and T-phase at the boundaries and S-phase in the bulk of the grains. Because S-phase is forming, Al₇Cu₂Fe cannot coarsen. In SEM, S-phase was not detected in the bulk, but this is likely due to the limiting resolution of the SEM; this causes the area fraction to appear lower than expected based on TEM results. With this in mind, it is important to understand the limitations of individual characterization techniques and utilize multiple techniques.

### Conclusions

Extensive electron microscopy in combination with computational thermodynamic and kinetic modeling indicated the presence of T-phase, Al₇Cu₂Fe, and Mg₂Si in the as-atomized condition of this gas-atomized Al 7075 powder. Thermal treatments were performed with the goal of homogenizing the microstructure—dissolving T-phase while avoiding S-phase growth, avoiding Al₇Cu₂Fe coarsening, and avoiding melting. The temperature of 465 °C was chosen because the Al₇Cu₂Fe should coarsen slower at a lower temperature, and this is as low as possible without containing S-phase. The temperature of 480 °C was chosen because it is the conventional solutionization temperature used for Al 7075. In the samples treated at 465 °C, S-phase was present, indicating a shift in local equilibrium of the powders as compared to the global composition of the powder batch. In both treatments, 60 min was insufficient to fully dissolve the T-phase. Due to the large interaction volume of the SEM as compared to the size of the precipitates present in these powders, it is not possible to accurately differentiate these phases in the SEM; additional TEM would be necessary to evaluate the time required to fully dissolve the T-phase.
It is important to understand the benefits, as well as the limitations, of different microscopy techniques and utilize different techniques for different purposes. The SEM is useful for high-throughput measurements and total secondary phase area quantification. However, its lower resolution does not allow for tracking individual secondary phases. The TEM is useful for its high resolution, allowing for better quantification of individual phases. However, the time required for sample preparation limits its feasibility for high-throughput evaluations. Leveraging the strengths of each technique, and coupling both of these microscopy techniques with computational thermodynamic and kinetic modeling, allows for enhanced understanding of the microstructural evolution with decreased time spent on data collection.

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