Effects of Sintering Temperature on Densification, Microstructure and Mechanical Properties of Al-Based Alloy by High-Velocity Compaction

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Abstract: This present work investigates the effects of sintering temperature on densification, mechanical properties and microstructure of Al-based alloy pressed by high-velocity compaction. The green samples were heated under the flow of high purity (99.99 wt%) N2. The heating rate was 4 °C/min before 315 °C. For reducing the residual stress, the samples were isothermally held for one h. Then, the specimens were respectively heated at the rate of 10 °C/min to the temperature between 540 °C and 700 °C, held for one h, and then furnace-cooled to the room temperature. Results indicate that when the sintered temperature was 640 °C, both the sintered density and mechanical properties was optimum. Differential Scanning Calorimetry, X-ray diffraction of sintered samples, Scanning Electron Microscopy, Energy Dispersive Spectroscopy, and Transmission Electron Microscope were used to analyse the microstructure and phases.

Keywords: high-velocity compaction; Al-based alloy; sintering process

1. Introduction

The powder metallurgy (PM) route provides lower energy consumption, higher material utilisation and excellent machinability in aluminium [1]. PM technology can present a homogeneous microstructure with little segregation compared with the casting method [2]. Therefore, Al-based PM alloy is currently an area of rapid development. The automotive industry is the main consumer of Al-based PM parts because of the lightweight and cost-effective nature of this process [3]. Al-based PM camshaft bearing caps were firstly introduced with the commercial inception of this technology in the mid-1990s. In recent decades, Al-based PM materials have been developed to expand the application of the automobile field [4–6]. Efforts have largely addressed the development of PM counterparts to wrought or cast alloys that span a broad range of properties [7–9].

There are two main problems in the current research: 1) the effect of porosity of existing green on sintering performance. The existence of a few residual porosities in the sintered parts remains the hurdle to get over for many applications, although the newer Al-based alloys offer main strength improvements [10]. High-velocity compaction (HVC) has greatly enhanced the density of the green body [11,12]. Z. H. Zhang et al. [13] reported that the iron-base PM alloy parts with high performance and high density were obtained by the combination of HVC and powder annealing. K. Q. Zhang et al. [14] explored a data-driven model for green density prediction by the multilayer perceptron algorithm and optimised the processing parameters of HVC-based metallic powders. D. D. You et al. [15]...
reported that a green compact with a 98% relative density is produced using pure iron powders with HVC. 2) It is difficult to sinter Al alloy, because there is a stable alumina film which coats the powder particles and hinders the material transfer between particles during sintering. A lot of mechanisms for alumina film reaction or disruption have been offered. Z. Wang et al. [16] reported that the surface oxide layer of the Al alloy powders was broken during hot pressing by a processing route imposing local deformation between the nanoscale phases rather than between consolidated powders, and the mechanical properties were improved. A. Ibrahim et al. [17] studied sintering temperature, pressing pressure, and the effect of heat treatment to Alumix 321 and the theoretical density of 98% was achieved. Sintering in PM is generally defined as a process or phenomenon occurring with heating green powder compact under appropriate temperature and atmosphere. The sintering plays a decisive role in the performance of the final products [18]. The sintering temperature has great effects on the microstructures and the properties of the material. Moreover, an appropriate sintering temperature can contribute to the attainment of the economic sintering process since they can help to reduce the energy consumption of the sintering process [19]. The authors studied the effects of compaction velocity on sintered density and mechanical properties of Al-based alloy, and the results were published [20]. The previous study showed that the higher velocity of HVC contributed to the increasing of sintered density and mechanical properties of Al-based alloy.

The aim of this work was to study the best-sintered temperature for the Al-based alloy during the powder metallurgy process. The research was therefore performed to find the highest tensile strength of the sample while, at the same time, density remains in an acceptable range. Al-based alloy powder was high velocity compacted and sintered with different temperatures (540 °C–700 °C) for one h. The effect of sintering temperature on densification, mechanical properties and microstructure of Al-based alloy were studied.

2. Experimental Details

In this research, the compaction methods and raw materials of green were the same as the previous study. The main chemical compositions of the Al-based alloy powder were Al, Fe, Cr, Ti and chemical compositions (in wt%) were shown in [20].

Al-based alloy powder was pressed by the HYP 35-2 HVC machine (Hydropulsor Company, Karlskoga, Sweden). The flowchart of this process is illustrated in our previous research [11]. The green with compaction energy 1325 J was sintered in a tube furnace with flowing high purity nitrogen (99.99 wt%). The heating rate was 4 °C/min before 315 °C. For reducing the residual stress, the samples were isothermally holding for one h at 315 °C. Then, it was respectively heated at the rate of 10 °C/min to the temperature between 540 °C and 700 °C, held for one h, and then furnace-cooled to room temperature. The low-temperature heating rate, high-temperature heating rate, low-temperature holding temperature and other parameters used in this experiment are the best parameters optimised by previous experiments. The optimal sintering process is the one after the best sintering temperature is determined. As a comparison, the green with compaction energy of 1885 J was sintered under the optimal sintering process 7% of the ethylenediamine solution, powder particles, and 93% of the epoxy resin were mixed and then the mixture carefully into the mould of the round pipe. After leaving it for 24 h, the sample can be ground. The upper surface of this powder sample, green and sintered samples were ground smooth to observe the microstructure. The scanning electron microscope (SEM) of samples were observed by JSM-6510 (JEOL, Tokyo, Japan). The elemental distributions were analysed by energy dispersive spectroscopy (EDS) with the SEM. The evolution of phases was performed by field emission scan electron microscope (FE-SEM, ZEISS ULTRA55, ZEISS, Oberkochen, Germany). The phase analysis was performed by a transmission electron microscope (TEM, FEI Tecnai G2 F20 S-Twin, FEI, Hillsborough, OR, USA). The temperature range of melting point of the alloy was measured by a differential scanning calorimetry (DSC) method (Netzsch STA 449C, Selb, Germany) performed in the range of 40–900 °C at a constant heating rate of 10 °C/min. The Archimedean method, according to the ASTM
B962-08, was applied to obtain the sintered density. Phase identifications of samples were examined on an X-ray diffractometer (XRD, Siemens D5000, Munich, Germany) with a Cu Ka radiation source operated at a continuous scanning rate of 5 °C/min, current of 40 mA and a voltage of 40 kV.

The tensile tests were performed at room temperature. An Instron universal testing machine was used to stretch the samples until failure at a constant cross-head speed of 0.5 mm/min. All of the samples were cut with the GB/T 228.1-2010. The ultimate tensile strength (UTS), 0.2% yield strength (YS) and elongation (δ) of each temperature or compaction velocity were obtained as the average of their test results.

3. Results and Discussion

3.1. Effect of Sintering Temperature on Densification

Sintering is a heating process that causes particles to bond together, resulting in significant strengthening and improved properties [21]. The sintering temperature is the most important factor affecting alloying due to the thermal activation of diffusion [18].

In this experiment, the theoretical density of Al-based alloy was 2.92 g cm⁻³ [20], and, with the increasing of sintering temperature, the sintered density gradually increased. The experimental results are consistent with the Arrhenius diffusion coefficient equation. Figure 1a reveals the effect of sintering temperature on sintered density. The plots show that the sintered density of the samples increases with an increase in sintering temperature, with obvious densification occurring when the temperature increases from 620 °C to 640 °C. As shown in Figure 1a, the sintered density rapidly increases from 2.73 g cm⁻³ (relative density is 93.49%) to 2.8 g cm⁻³ (relative density is 95.89%) when the sintering temperature increases from 560 °C to 640 °C. The sintered density increases slowly from 2.8 g cm⁻³ (relative density is 95.89%) to 2.81 (relative density is 96.23%) g cm⁻³ when the sintering temperature increases from 640 °C to 700 °C. During the sintering process of the green, with the increasing of sintering temperature, the powder particles boundary developed from poor mechanical bonding to strong metallurgical bonding, and internal pores of the green become smaller or disappear. The driving force is generally considered as the reduction of free energy at the expense of minimising the internal and external surfaces [22]. While density increases significantly when the sintering temperature increase from 650 to 700 °C, serious deformation of the sample is also observed as shown in Figure 1a. In fact, the higher sintering temperature does not appear to be the best. Figure 2 is the DSC of Al-based alloy powder. Figure 2 reveals that the Al-based alloy begins to melt at 658 °C and the melting point is 658 °C. The samples began to deform at 650 °C due to the partial melting, which may hurt the mechanical properties of the compact, which occurred when the specimen was sintered at 650 °C. These were considered to be unsuitable for an industrial production environment because they negated the near net shape property for which PM manufacturing relies on [23]. Therefore, considering the influence of sintering temperature on sintered density and near net shape, 640 °C is the optimum sintering temperature. The effect of sintering temperature on the properties of the alloy will be further verified, and then the sintering temperature will be finally determined.

Nevertheless, the samples sintered at 640 °C had a lower sintered density 2.8 g cm⁻³ (relative density is 95.89%), because the green density was lower when the compaction energy is 1325 J as shown in Figure 1b. Figure 1b indicates that sintered density and green density increase as the compaction energy increase. When the compaction energy increased from 1325 J to 1885 J, sintered density increased by 0.05 g cm⁻³ (relative density is 2.11%). During the HVC, the compaction energy was transferred to the powder particles in a short time to make them densification. The higher the compaction energy contributed to enhancing the green density, which increased the contact area between adjacent powder particles and facilitated sintering densification. Therefore, HVC was beneficial to sintering,
3.2. Microstructure and Phases Constitution

Figure 3 shows the SEM microstructure of the atomised Al-based alloy powder with different magnification and the green sample with the compaction energy of 1325 J. From Figure 3a, it is clear that the large particle mainly contains a white phase and the little particle contains a grey phase with the dispersive distribution. Figure 3b shows the white phase in the large particles distributes as snowflakes. Figure 3c exhibits a large number of small pores which are irregular and uneven distribution and obvious particle boundaries. This is because the compaction energy 1325 J is not high enough. Figure 4 shows the XRD patterns of Al-based alloy powder and green. It mainly consists of the Al, Al$_3$Ti, and quasicrystalline phase (Q-phase). This assumption is in agreement with data reported by M. Galano [24], describing that the Q-phase is an icosahedral quasicrystalline phase. On the other hand, the peak width of each phase of the green is significantly wider than that of the powder, as shown in Figure 4. This confirms that the plastic deformation of the powder particles during the HVC. Under the action of external force, plastic deformation in different regions of the green is different, which leads to the inconsistency of microstrain, so that, for the same crystal surface index, the crystal surface spacing in different regions is different.
Figure 3. (a) lower amplification (b) higher amplification SEM/BSE of the atomised powder particles; (c) SEM/BSE of the green sample with the compaction energy of 1325 J.

Figure 4. X-ray diffraction (XRD) pattern of Al-based alloy powder and green.

Microstructure analysis of the sintered specimens was carried out to research the development of the evolution of the porosity and the microstructure. The microstructure of sintered samples with different sintering temperatures, as shown in Figure 5. The evolution of pores can be comparatively seen in Figure 5 with the sintering temperature increasing. Figure 5a exhibits the samples sintered at 560 °C have many pores and distinct original particle boundaries. This shows that the number of atoms participating in diffusion is small due to the low diffusion coefficient at 560 °C and the particles are still in contact which demonstrates a lack of metallurgical bonding. Clear original particle boundaries and many pores can be observed in Figure 5b when the sintering temperature is 580 °C, but the contact area between particles increases. This phenomenon indicates that the sintered neck began to form at 580 °C and the contact between particles began to change from mechanical meshing to metallurgical bonding with the sintered temperature and according to increase in the diffusion coefficient. Figure 5c shows that the samples have fewer pores and original particle boundaries, and the adjacent particles gain the contact area when the sintering temperature is 620 °C. This result exhibited that the sintered neck began to grow at 620 °C and the metallurgical bonding between particles gained. Figure 5d presents the sample has less porosity and original particle boundaries, and the contact area between the adjacent particles continue to gain when the sintering temperature is 640 °C. This result exhibited that the sintered neck continued to grow up at 640 °C and the metallurgical bonding between particles enhanced. On the other hand, gas atomised Al powder-coated amorphous Al$_2$O$_3$ (a-Al2O3) film [25], and L. Cao et al. [26] found that the a-Al$_2$O$_3$ turn into γ-Al$_2$O$_3$ nanoparticles during SPS (Spark plasma sintering) causing a dramatic increase in the elongation to fracture. For HVC, the high impact force of the HVC offered a mechanical means to disrupt the Al$_2$O$_3$ film due to the larger amount of deformation of the ductile Al-based powder [27]. The powder particles in this experiment were spherical, as shown in Figure 6a. The surface of small particles is smooth, and the surface of large particles is uneven, as shown in Figure 6b. When the spherical particles
were impacted by high energy, the impact force on the particles was greater due to the smaller contact area between the spheres, and it was easier to break the oxide film locally. When the compaction energy is 1885 J, as shown in Figure 5c, the elimination of porosity and the dissolution of the particles for the sintered sample at 640 °C can be observed.

Figure 5. FESEM/BSE micrograph of the green and sintered samples with different sintering temperature: specimens were compacted with compaction energy of 1325 J and sintered at temperature, (a) 560 °C (b) 580 °C (c) 620 °C (d) 640 °C; and specimens were compacted with compaction energy of 1885 J and sintered at 640 °C (e,f).

After sintering, the phase transition is very obvious with the increase in sintering temperature we can see the formation of white acicular phase and flaky grey phase, as shown in Figure 5a,b. As shown in Figure 5c–e, the white phase and grey phase evident grow up and the white phase moves to the particle boundary or across particle boundaries. From Figure 5f, it is clearly shown that the phases of the sample are composed of white phase 2, grey phase 3 and dispersed nano phase 1. Further analysis of the EDS presents the composition of phases, as shown in Table 1. XRD patterns of the sintered Al-based alloy samples, fabricated at 560 °C, 600 °C, 640 °C, and 660 °C, are shown in Figure 7. As it can be seen in Figure 7, the spectrums of all the samples have characteristic peaks of Al and a small amount of Al₃Ti, Al₁₃Fe₄, Al₁₃Cr₂, AlN and AlCr₂. The absence of quasicrystalline phase after sintering is due to the decomposition of the quasicrystalline phase at the high temperature [24]. The exothermic peaks of these phases decomposition are not observed on the DSC (Figure 2), because these phases only exist when the particle size is less than 25 µm [20]. The average particle size of the experimental powder is 25 µm, and the content of these phases is very small, which is almost invisible in the DSC experiment. There is no obvious new phase formation with the sintering temperature increasing, as shown in Figure 7. However, as the sintering temperature increasing, the angle of the peak becomes larger. For example, when the sintering temperature is 660 °C, peaks become wider.

Figure 6. SEM image (a) and particle size distribution (b) of atomised Al-based alloy powders.
and move to the right. This is because diffusion enhanced as the sintering temperature increasing. In Figure 8, SEM images together with detailed element distribution maps of Al, Fe, Cr, and Ti elements are shown. The EDS maps confirm that the main component of the white acicular phase is Fe, flake phase is Ti, and the finely dispersed nano phase is Cr. The combination of XRD and EDS results revealed the formation of the higher Cr content of phase $\text{Al}_{13}\text{Cr}_2$, the higher Fe content of phase $\text{Al}_{13}\text{Fe}_4$, and the higher Ti content of phase $\text{Al}_3\text{Ti}$. As shown in Figure 9, it can be seen that there are granular intermetallic compounds less than 0.5 µm and columnar intermetallic compounds. The constituent of 1, 2, 3 phases in TEM, as shown in Figure 9. TEM further verified the existence of intermetallic compounds. Phase 1 is $\text{Al}_{13}\text{Cr}_2$, phase 2 is $\text{Al}_3\text{Ti}$, and phase 3 is $\text{Al}_{13}\text{Fe}_4$. The results are consistent with the previous analysis [18]. There is a small amount of AlN and AlCr$_2$, and they are not verified because of the small quantity. Because the protective atmosphere of sintering is N$_2$, AlN was formed by the reaction of N$_2$ and Al, such as the reaction equation $2\text{Al} + \text{N}_2 \rightarrow 2\text{AlN}$. This is a strongly exothermic reaction. The heat released may cause local melting, which will be conducive to sintering [28].

Table 1. The energy dispersive spectroscopy (EDS) of the three phases of the arrow in the microstructures of the sample.

| Element | Phase | Al   | Ti    | Cr    | Fe   |
|---------|-------|------|-------|-------|------|
| Atomic% | 1     | 87.94| 2.08  | 6.88  | 3.10 |
| Atomic% | 2     | 77.52| 0.37  | 0.75  | 21.36|
| Atomic% | 3     | 77.64| 21.41 | 0.74  | 0.21 |

Figure 7. XRD of the sintered samples with different sintering temperature: (a) full scan from 10 to 100° (b) enlargement of the 30 to 50° section.

Figure 8. Al, Fe, Cr, Ti element EDS mapping of (a) flake phase and (b) acicular phase of samples with sintering temperature of 640 °C.
3.3. Mechanical Property

Tensile properties of the sintered samples with different sintered temperature are presented in Figure 10a. It can be seen from Figure 10a that the UTS, YS and δ all increase at first and then decrease with increasing the sintered temperature. Figure 10 displays UTS and YS reach the maximum (UTS 179 MPa and YS 148 MPa), and the maximum δ is 1% at the sintered temperature of 640 °C. Over 640 °C, UTS, YS, and δ tend to drop. The worst UTS, YS, and δ belong to the sample fabrication at the lowest sintering temperature of 560 °C. The tensile strength is 16 MPa which is consistent with the previously measured value [29] for the bending strength of the green is 14 MPa when the compaction energy is 1325 J. It seems that the mass transport phenomenon has slightly transpired at 560 °C. The YS of the sintered sample is zero before 600 °C, and it was sharply increased to 143 MPa at 620 °C. The δ is very low before 620 °C is zero. Over 640 °C δ is also zero. The samples sintered at 640 °C still had a lower UTS, YS and δ when the compaction energy was 1325 J which is related to the lower compaction energy of green, as shown in Figure 10b. We can see from Figure 10b, as the compaction energy increased from 1325 J to 1885 J, UTS increased by 44, YS increased by 12 MPa, and the maximum δ increased by 1.6% respectively. Hence, it seems that the tensile properties strongly depend on the compaction energy, similar to the trend was observed for sintered density. With compaction energy increasing, the porosity of the green decreased, and the contact area between particles increased, which was helpful to the mass diffusion between the particle boundaries during sintering. Although the quasicrystal phase decomposed, the tensile properties of samples still increased with the increase in temperature. Therefore, it seems that the decomposition of the Q-phase during the sintering did not have an important detrimental effect on the improvement in the strength of the Al-based alloy [30]. Intermetallic compounds got from the decomposition of the Q-phase are the one reason for increasing the tensile strength of the alloy. Tensile properties can be related to the porosity content, phase growth, and the particle boundary, respectively.

Figure 10. Ultimate tensile strength (UTS), yield strength (YS) and elongation versus sintering temperature (a); UTS, YS and elongation versus compaction energy (b).
3.4. Fractography

There is a noticeable evolution on a microscopic level, which can be appreciated by comparing the fracture surface morphology with the increase in temperature, as shown in Figure 11. The original particle boundary is evident before 620 °C, which is shown in Figure 11a,b. The existence of the original particle boundary was due to the inadequate sintering, and the diffusion between particles was prevented. At 620 °C (Figure 11c), the bond of particles began to appear, but the boundary of the original particle can still be seen, because the sintering temperature is not high, and the sintering is not sufficient. The fracture morphology of 650 °C (Figure 11e) and 660 °C (Figure 11f) is seen as clear gullies, which is due to excessively high sintering temperature promoting the formation of the local liquid phase. This is consistent with the results of the DSC of Al-based alloy powder (Figure 2). Therefore, over 640 °C, there was evidence of excessive melting phenomenon, which shows groove structure in the fracture morphology (Figure 11e,f). At 640 °C (Figure 11d), there is a good bond between the particles, and the original particle boundary cannot be seen, but a small number of pores can still be seen. We attribute the phenomenon to the reason for the lower compaction energy (1325 J) of green. With increasing compaction energy, the bond of the original particle boundary increased obviously for the sintered sample. Figure 12a is the fracture surface morphology of the sample with compaction energy 1885 J and sintered at 640 °C, there is a good bond between the particles, and basically cannot be seen the original particle boundary and pores. Higher magnification of the fracture surface morphology is shown in Figure 12b,c. From Figure 12b,c, fine and deep dimples are observed, indicating that the fracture mechanism is ductility fracture when the sample sintered at 640 °C with compaction energy 1885 J. This is consistent with the analysis of tensile properties. Combined with the sintered density, the microstructure and phase analysis, the particle boundary strengthening is helpful in developing tensile properties. Hence, 640 °C was selected as the optimum sintering temperature.

Figure 11. SEM/SE micrograph of fracture surface with different sintering temperature (a) 560 °C (b) 580 °C (c) 620 °C (d) 640 °C [20] (e) 650 °C (f) 660 °C.

Figure 12. SEM/SE micrograph of fracture surface with compaction energy 1885 J, (a-c) with different magnification.
4. Conclusions

Thus, based on the results of the conducted study on HVC and sintered Al-based alloy powder with different sintering temperatures, it is possible to draw the main conclusion that 640 °C is rather effective for obtaining high-quality samples for Al-based alloy powder with the compaction energy of 1885 J. More concrete conclusions are as follows:

1. For the green of Al-based alloy by HVC, the sintered density increased with the temperature increasing. When the compaction energy was 1885 J, the samples sintered at 640 °C with a relative density of 98%.

2. XRD of the sintered sample displayed the phases Al$_{13}$Fe$_4$, Al$_{13}$Cr$_2$, Al$_3$Ti, AlN, AlCr$_2$. TEM further verified the existence of intermetallic compounds Al$_{13}$Cr$_2$, Al$_3$Ti, and Al$_{13}$Fe$_4$.

3. The tensile properties of the sintered samples at 640 °C reached the maximum, UTS 222 MPa, YS 160 MPa, and the maximum δ 2.6% when the sample was compacted by the energy 1885 J.

4. Fracture mechanism of the Al-based alloy was ductility fracture.

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