Interfacial microstructure and mechanical properties of Ti–6Al–4V/Al7050 joints fabricated using the insert molding method

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Abstract: Ti–6Al–4V/Al7050 joints were fabricated by a method of insert molding and corresponding interfacial microstructure and mechanical properties were investigated. The interfacial thickness was sensitive to holding temperature during the first stage, and a good metallurgical bonding interface with a thickness of about 90 µm can be obtained at 750°C. X-ray diffraction, transmission electron microscopy, and thermodynamic analyses showed that the interface mainly contained intermetallic compound TiAl3 and Al matrix. The joints featured good mechanical properties, i.e., shear strength of 154 MPa, tensile strength of 215 MPa, and compressive strength of 283 MPa, which are superior to those of joints fabricated by other methods. Coherent boundaries between Al/TiAl3 and TiAl3/Ti were confirmed to contribute to outstanding interfacial mechanical properties and also explained constant fracture occurrence in the Al matrix. Follow-up studies should focus on improving mechanical properties of the Al matrix by deformation and heat treatment.

Keywords: interfacial microstructure; interfacial bonding mechanism; mechanical properties; insert molding method; coherent boundaries; Ti/Al joints

1. Introduction

Given the low density, high specific strength, excellent corrosion resistance, notable impact toughness, and significant thermal stability, titanium alloys are widely used in vehicle, aerospace, chemistry, and defense industries. However, unavoidable shortcomings of titanium alloys, such as high production cost, low thermal conductivity, and low shear strength, constrain their widespread applications [1]. Aluminum alloys, as primary structural materials for commercial and military applications, are known for their excellent mechanical properties, easy to design, and mature manufacturing and inspection techniques. However, these materials also face other challenges, such as low hardness and wear resistance, low tensile strength at high temperature, and high linear expansion coefficient [2]. To overcome disadvantages of Ti alloys and Al alloys and to utilize their respective advantages, Ti/Al bimetallic joints have attracted more attentions in recent years [3–12]. For these two alloys, significant differences in physical properties, such as melting point and heat conductivity, make conventional preparation techniques unsuitable for fabricating Ti/Al joints. Novel methods, such as accumulative roll-bonding [3–5], laser welding–brazeing [6–7], friction stir welding [8–9], powder metallurgy [10–11], and explosive cladding [12], were attempted. Despite these efforts, problems still exist with these fabrication methods: (i) poor interfacial bonding strength, (ii) complex processing procedures, and (iii) limited product sizes for industrial application.

Recently, a study of Ti/Al–7Si–0.3Mg joints prepared by a method of insert molding exhibited promising industrial application prospects due to low production cost, low energy consumption, simple production procedure, and high interfacial bonding strength [13–14]. The basic principle of this method is to immerse Ti insert into Al–7Si–0.3Mg melt at the optimized temperature. As a result, good metallurgical bonding...
2. Experimental

2.1. Specimen preparation

Joints were prepared by insert molding of commercial Al7050 (matrix) and Ti–6Al–4V (insert rods) alloys, whose compositions are illustrated in Table 1. The corresponding experimental set-up can be found in detail in Ref. [15]. During experiments, first, the aluminum alloy melt temperature was set to 790°C. Ti–6Al–4V rods were then inserted into the aluminum alloy melt. Such a high temperature was necessary to prevent the aluminum melt from solidifying around Ti-alloy rods upon their insertion into the Al7050 melt and to avoid foundry defects. Second, to ensure integrity of the Ti–6Al–4V/Al7050 interface, sufficient energy was provided for the interfacial diffusion reaction. Three temperatures (770, 750, and 730°C) with the same holding time (40 min) were selected to compare their influences on interfacial bonding behaviors. Third, to ensure secondary precipitation of the TiAl3 phase at the interface, the next process was set to 700°C for 10 min. Then furnace cooling was performed until 670°C to reduce residual stress induced by difference in thermal expansion coefficients between Ti–6Al–4V and Al7050. Finally, to minimize the presence of a compact layer in which microcracks can be easily initiated due to the formation of brittle intermetallic compounds (reported in Ref. [15]), air cooling was implemented until room temperature.

| Material           | Si  | Fe  | Cu  | Mg  | Zn  | Ca  | V   | Ti   | Al   |
|--------------------|-----|-----|-----|-----|-----|-----|-----|------|------|
| Al7050             | 0.04| 0.07| 2.49| 2.42| 6.3 | —   | 0.30 (max)| 0.03 | Balance |
| Ti–6Al–4V          | 0.09| —   | 2.49| 2.42| 6.3 | 0.30 (max)| 4.42 | Balance | 6.13 |

2.2. Interfacial characterization

After solidification, joints were sliced in a thickness of 5 mm by wire cut electrical discharge machining, and cross-sections of these slices were perpendicular to the rod axis. Surfaces of each specimen were then polished with 0.5 μm diamond paste and utilized for interface microstructure characterization, chemical composition analysis by scanning electron microscopy (SEM), and electron probe micro-analysis (EPMA). Transmission electron microscopy (TEM), high-resolution TEM (HRTEM), and energy-dispersive X-ray spectroscopy (EDS) were performed to characterize interfacial microstructures and analyze phase constituents using a Tecnai G2 F30 microscope operated at 300 kV. TEM and HRTEM specimens were mechanically ground, polished, dimpled to 60 μm, and then thinned by ion milling using a Gatan precision ion polishing system (Gatan model 691.CS).

2.3. Mechanical properties

Interface shear strength was measured using classical push-out tests, as described in Ref. [13]. A universal testing machine was used to conduct push-out tests. Interfacial shear strength (τ) can be calculated using the following equation [13,15]:

\[ \tau = \frac{F_{\text{max}}}{2\pi rt} \]  

where \( F_{\text{max}} \) is the maximum load, \( r \) refers to the radius of Ti–6Al–4V insert (3 mm), and \( t \) denotes the specimen thickness (5 mm). Vickers hardness was measured across the interface layer under an indentation load of 2.94 N for...
15 s. The micro hardness value was evaluated by averaging three mediate values of five indentation measurements. Following the ASTM standard, plate-type tensile specimens with the gauge length of 25 mm, gauge width of 6 mm, and thickness of 3 mm were machined with axes perpendicular to the interface. Tensile tests were carried out on a universal testing machine, and the strain rate was $5 \times 10^{-5} \text{s}^{-1}$. A cylindrical cast sample of 10 mm in diameter and 20 mm long was cut for the uniaxial compression experiment at room temperature.

3. Results

3.1. Interface characterization

Fig. 1 shows the interfacial morphologies of samples at different holding temperatures (770, 750, and 730°C). It also shows that at 730°C, the interface thickness was 3 µm owing to slow elemental diffusion. With increasing holding temperature, the interface thickness correspondingly increased. At 750°C, a uniform thin interfacial layer with a thickness of 90 µm was attained. However, at 770°C, a thick interface with many shrinkage voids was observed. These voids were reported to exert detrimental effects on the mechanical characteristics of bimetallic joints [13]. Thus, the interfacial morphologies of Ti–6Al–4V/Al7050 joints strongly depended on the holding temperature in the first stage, and good interfacial metallurgical bonding can be possibly attained by parameter optimization, as shown in Fig. 1(b). For deep analysis of interface microstructure, the interface layers of specimens at 750°C are enlarged in Fig. 1(d). A large number of irregular intermetallic particles were distributed in the black alloy matrix. Meanwhile, small amounts of white and gray phases were observed around intermetallic particles. To confirm the phase constituents at the interface shown in Fig. 1(d), EPMA analysis was performed. Corresponding results from spot scan and area scan are shown in Table 2 and Fig. 2, respectively. As shown in Fig. 2, Ti, Al, and V were more abundant in the newly formed intermetallic particles than those in the black alloy matrix. However, Mg, Zn, and Cu in the newly formed intermetallic particles were less than those in the black alloy matrix. Combined with the spot scan results in Table 2, irregular intermetallic particles and the black alloy matrix are the TiAl$_3$(V) phase and the Al-alloy matrix, respectively. Similarly, white and gray particle phases were confirmed to be S(CuMgAl$_2$)Zn and Al$_3$Fe phases, respectively. The interfacial

![Fig. 1. Interface SEM images of the samples fabricated at different holding temperatures: (a) 730°C, (b) 750°C, and (c) 770°C; (d) enlarged SEM images of the Ti–6Al–4V/Al bonding interface at 750°C for studying the precipitation behavior of phases.](image-url)
Table 2. Spot scan results of irregular intermetallic compound particles, black alloy matrix, and white and light gray particle phases contained at the interface of Ti–6Al–4V/Al7050 joints fabricated by the insert molding method.

| Phases                        | Al   | Cu   | Ti   | Mg   | Zn   | Fe  | V  |
|-------------------------------|------|------|------|------|------|-----|----|
| Intermetallic compound particles | 74.28| 0.09 | 22.19| 0.15 | 0.17 | 0   | 3.12|
| Black alloy matrix             | 93.31| 0.78 | 2.11 | 1.32 | 2.33 | 0   | 0.15|
| White particle phase           | 46.82| 14.17| 0.0068| 24.92| 4.24 | 0   | 0  |
| Gray particle phase            | 78.03| 3.74 | 0.68 | 0    | 0    | 17.55| 0  |

Fig. 2. Area scan spectrum maps of the Ti–6Al–4V/Al7050 reaction interface analyzed by EPMA.

energy for nucleation in the Al matrix is larger than that for nucleation in the interface of Ti/Al intermetallic compounds. This observation explains why S(CuMgAl₂)Zn and Al₃Fe phases were easily precipitated around TiAl₃(V) particle phases rather than in the Al-alloy matrix. Besides the above four phases, a small number of tiny titanium particles were also observed at the interface around Ti–6Al–4V alloy rods, as shown in Fig. 2.

Fig. 3 shows the XRD patterns at the interface. Peaks of Al and TiAl₃(V) phases can be found in the same figure. Besides the two phases, smaller amounts of Ti and S(CuMgAl₂)Zn phases were also confirmed in the XRD patterns, in accordance with the aforementioned EDS analysis results shown in Table 2. However, the Al₃Fe phase cannot be observed from the XRD patterns, and this result is speculated to be related to its low concentration in the sample.

Fig. 3. XRD patterns of the Ti–6Al–4V/Al bonding interface at a holding temperature of 750°C.
We investigated the interfacial bonding behavior of pure Al and pure Ti joints fabricated by the insert molding method [15]. The interfacial bonding layer comprised TiAl3 and TiAl2 phases, with the TiAl2 phase existing around pure Ti rods. To investigate the interface phase constituents of Ti–6Al–4V/Al7050 joints, TEM analysis was carried out (Fig. 4) around Ti–6Al–4V insert rods. Results from selected area diffraction patterns (SADP) and EDS (shown in Fig. 4(d))

Fig. 4. TEM, SADP, and EDS analyses around Ti–6Al–4V insert rods: (a) TEM interface image approaching Ti insert rods; (b) enlarged TEM image of the TiAl3 phase from region A shown in (a); (c) TEM image of Ti particulates around Ti insert rods; (d) SADP for the TiAl3 phase shown in (b); (e) EDS analysis for the Ti region shown in (a); (f) EDS analysis for the TiAl3 region shown in (a); (g) EDS analysis for the Ti particulate shown in (c).
and Fig. 4(f), respectively) confirmed that only the TiAl3 phase existed around Ti-alloy inserts, and this condition differs from the interfacial microstructures of previous pure Ti/pure Al joints. The cause of sole precipitation of TiAl3 phases (instead of other Ti–Al intermetallic compounds) around Ti-alloy rods will be discussed later. Tiny Ti particles (Fig. 4(g)) were confirmed around the Ti-alloy rod, indicating accordance with aforementioned EPMA and XRD analyses.

### 3.2. Mechanical properties

Ti−6Al−4V/Al7050 joints fabricated by the insert molding method also exhibited outstanding mechanical performance. Fig. 5(a) shows the hardness measured across the interface. Hardness of the Ti−6Al−4V rod was measured to be HV 320, which is nearly three times that of the Al7050 matrix (HV 106). As the interface was mainly composed of TiAl3 (V) and the Al7050 alloy matrix, interface hardness reached a mediate value between those of titanium insert and aluminum matrix. As depicted in Fig. 5(a), intermetallic compounds formed on the insert side occupied a major portion of the interface, increasing the hardness value to approximately HV 210, which is higher than HV 180 on the Al matrix side.

Fig. 5(b) shows the load-displacement curves of Ti−6Al−4V/Al7050 specimens. Shear strength can be calculated from the push-out load and surface circumference of the insert according to Eq. (1) and as illustrated in Table 3. Table 3 also illustrates the shear strength values for some typical Ti−6Al−4V/Al-alloy joints fabricated by different methods [17–25]. The shear strength value of Ti−6Al−4V/Al7050 joints reached 154 MPa, which is superior to the values attained from other fabrication methods, such as transient liquid phase (TLP) bonding, indicating remarkable advantage in improving interfacial bonding strength by the current insert molding method. A push-out test of the Ti−6Al−4V/Al7050 joint was also interrupted upon application of 80% of the maximum load value. Slices of this specimen were later cut with a diamond wire and polished to a finish better than 0.5 µm to characterize the damage along the vertical section. Damage characterization performed by SEM (Fig. 5(c)) indicated that cracks can initiate at the bottom face of the specimen in the aluminum matrix rather than at the bonding interface, further confirming high interfacial bonding performance when insert molding was applied.

To further determine the interfacial mechanical performances of Ti−6Al−4V/Al7050 samples, traditional tensile and compressive tests were performed. Fig. 6 shows the macroscopic and microscopic scanning images of tensile and compressive fracture including stress–strain curves. Fig. 6(a) indicates that no significant differences (within experimental error) were observed in tensile strength between as-cast Al7050 (219 MPa) and Ti−6Al−4V/Al7050 samples fabricated by insert molding (215 MPa), confirming
occurrence of fracture in the Al7050 alloy matrix. Fig. 6(c) displays a typical SEM fracture morphology of the Ti–6Al–4V/Al7050 bimetallic composite obtained after tensile tests. Some shallow dimples were detected on fracture surfaces, and some coarse iron-rich phases appeared in these shallow dimples, further confirming fracture of the as-cast Al7050 alloy matrix. Based on the compression curve shown in Fig. 6(d), non-proportional compressive stress ($\sigma_{pc}$) of Ti–6Al–4V/Al7050 reached 283 MPa. Fig. 6(e) shows lateral views of the macroscopic compressive fracture

Table 3. Comparisons of the mechanical performance of Ti–6Al–4V/Al alloy joints fabricated in the present study and those by other fabrication methods

| Material       | Fabrication method     | Maximum shear strength / MPa | Interface hardness, HV | Ref.    |
|----------------|------------------------|------------------------------|------------------------|---------|
| Ti–6Al–4V/Al7050 | Insert molding          | 154                          | 180-210                | This study |
| Ti–6Al–4V/Al7075 | TLP bonding             | 30                           | 50-360                 | [17]    |
| Ti–6Al–4V/Al7075 | TLP bonding             | 36                           | 104-330                | [18]    |
| Ti–6Al–4V/Al7075 | TLP bonding             | 42.3                         | 57-413                 | [19]    |
| Ti–6Al–4V/Al7075 | TLP bonding             | 19                           | 170-400                | [20]    |
| Ti–6Al–4V/Al7075 | TLP bonding             | 36                           | 124                    | [21]    |
| Ti–6Al–4V/Al1060 | Ultrasonic-assisted brazing in air | 68 ± 2.3                   | —                      | [22]    |
| Ti–6Al–4V/Al2024 | TLP bonding             | 37                           | 118-153                | [23]    |
| Ti–6Al–4V/Al2024 | Liquid state diffusion bonding | 36                           | 75-344                 | [24]    |
| Ti–6Al–4V/Al2024 | TLP bonding             | —                            | 139                    | [25]    |

Fig. 6. Tensile and compressive test results of Ti–6Al–4V/Al7050 joints prepared by the insert molding method: (a) tensile stress–strain curves; (b) macroscopic tensile fracture morphology; (c) microscopic tensile fracture morphology; (d) compressive stress–strain curve; (e) lateral view of the macroscopic compressive fracture morphology; (f) microscopic compressive fracture surface morphology.
morphism caused by the uniaxial compression experiment at room temperature. The 45° compressive fracture plane can be observed throughout the aluminum matrix. Therefore, compressive fracture was mainly caused by shear stress, and the common failure mode was shear fracture. As a result, fibrous fracture surfaces can be observed in the SEM images of compressive samples (Fig. 6(f)). Different from tensile tests, the compression curve directly shows the yielding process. Based on the above results, interfacial bonding performance was enhanced by insert molding to prevent fracture at the interface bonding layer. Instead, fracture was initiated in the Al-alloy matrix. Tensile or compressive fracture characteristics are those of the Al7050 matrix alloy. Chen et al. [7] reported that tensile strength of Al5A06/Ti6Al4V prepared by the laser welding–brazing method can reach 292 MPa, which is higher than the current value observed. This result can be attributed to different treatment states of Al-alloy samples. Instead of the as-cast state of the Al7050 alloy used in this study, Chen et al. used Al-alloy matrices with different heat treatment states and as-rolled conditions. Instead of the tensile fracture observed in the Al alloy matrix in this study, tensile fracture from Chen’s study was mainly located at the joint interface. Thus, mechanical performances of Ti–6Al–4V/Al7050 joints prepared in this study exhibit higher potential than those prepared by Chen et al. after further rolling and heat treatment of the Al-alloy matrix.

4. Discussion

4.1. Interfacial bonding mechanism

When temperature at the interface is above the melting point of aluminum alloy, both liquid and solid surfaces can experience separation of the surface oxidation layer and spreading of liquid aluminum due to capillary pressure [26–27]. Compared with the solubility of aluminum in titanium within the experimental temperature range, titanium in aluminum presents lower solubility [28], indicating that formation of intermetallic compounds at the interface is mainly related to the degree of diffusion of aluminum before complete solidification. As the lattice misfit between TiAl3 and Ti or Al is low, TiAl3 will preferentially nucleate and induce minimal increase in interfacial energy in Ti–Al compounds [29–30]. Within the experimental temperature range, sufficient Al atoms were dissolved into titanium inserts, guaranteeing the formation of TiAl3 phase until chemical equilibrium. As a result, the interface layer was mainly composed of TiAl3. To confirm this speculation, the thermodynamic behavior of Ti–Al intermetallic compounds were compared. Fig. 7 exhibits the free energy-temperature line of Ti–Al intermetallic compounds and diffusion between Ti and TiAl3 from 630 to 900°C [30–31]. As shown in Fig. 7, free energies of TiAl3, TiAl2, and Ti2Al5 phases are lowest among Ti–Al intermetallic compounds at the experimental temperature. Though free energies of TiAl2 and Ti2Al5 are lower, intermetallic compound at the interface layer still mainly comprised TiAl3 phase. This result can be explained by the following reasons. First, phase-transition temperature of Ti2Al5 is beyond the experimental temperature, as observed from the Ti–Al binary phase diagram [32]. Second, during the formation of Ti2Al5 and TiAl2 phases, TiAl often acts as the intermediate phase [29,32]. TiAl and Ti2Al phases can only be formed in extreme absence of aluminum atoms near the titanium insert (Figs. 7(b)–7(c)). However, in this study, adequate aluminum atoms were supplied to participate in interface reactions during casting. Therefore, the TiAl phase was not easily formed, implying that Ti2Al5 and TiAl2 phases can also be ruled out in this study. Finally, given the slower diffusivity of titanium atoms than aluminum atoms, thicknesses and phases of interface layers depend on the degree and distance of diffusion of aluminum atoms. In other words, compounds

![Fig. 7. Free energy–temperature spectrum line of Ti–Al intermetallic compounds (a) and diffusion process between Ti and TiAl3 from 630 to 900°C (b–c).](image-url)
present in reaction layers are determined by interdiffusion of aluminum atoms along grain boundaries, and this process is directly affected by interfacial heat content and element distribution [29]. From Refs. [30−31], TiAl2 and Ti2Al5 are mesophases of interfacial reactions and thus, they can also be ruled out considering thermodynamic effects.

4.2. Strengthening mechanism on interfacial mechanical properties

As mentioned earlier, cracks were only initiated in the Al matrix for Ti−6Al−4V/Al7050 joints fabricated by insert molding. To further elucidate the strengthening mechanism on interfacial mechanical properties, HRTEM was used to examine interfacial microstructures. The HRTEM images of Al/TiAl2 and Ti/TiAl2 interfaces are illustrated in Figs. 8 and 9, respectively. EDS was used to characterize the elemental distribution. Based on HRTEM images, intermetallic compounds at the interface layer are evidently the TiAl3 phase and Al-alloy matrix, in accordance with Figs. 1 and 3, respectively. Indexed electron diffraction patterns indicate...
the coherent boundary between \( [0\overline{1}1]_{\text{Al}} \) and \( [\overline{1}12]_{\text{TiAl}_3} \) and between \( [011]_{\text{TiAl}_3} \) and \( [0\overline{1}1]_{\text{Ti}} \). As a result of coherent Al/TiAl\(_3\) and TiAl\(_3\)/Ti boundaries, stress will concentrate at smaller casting flaws in the aluminum alloy matrix during deformation. This condition will explain why fracture constantly occurred in the Al matrix during tensile or compressive tests (Fig. 6). Therefore, to improve the mechanical performances of Ti–6Al–4V/Al7050 joints, mechanical performance of the aluminum alloy matrix should be enhanced through various strengthening methods, such as rolling and aging. Aside from TiAl\(_3\) intermetallic compounds, Ti particles (Fig. 4(c)), S(CuMgAl\(_2\)), and Al\(_3\)Fe phases (Table 2) were also present in the interface layer. However, their levels were very low at the interface, and they cannot influence the interfacial mechanical performance observed in this study.

5. Conclusions

In this study, Ti–6Al–4V/Al7050 joints with outstanding interfacial mechanical properties were successfully fabri-
cated by the insert molding method. Conclusions can be summarized as follows:

1. Good interfacial metallurgical bonding can be formed between two dissimilar alloys of Ti−6Al−4V and Al7050, ensuring high mechanical properties, i.e., shear strength of 154 MPa, tensile strength of 215 MPa, and compressive strength of 283 MPa.

2. The corresponding interfacial bonding mechanisms were revealed. From thermodynamic analysis, phases formed at the interface of Ti−6Al−4V/Al7050 joints were mainly composed of TiAl3 and the Al matrix.

3. Coherent Al/TiAl3 and TiAl3/Ti boundaries were confirmed, explaining constant mechanical fracture in the Al matrix.

4. Follow-up studies will focus on improving mechanical properties of the Al matrix by heat treatment and deformation.

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