Bonding SiCp/Al Composites via Laser-Induced Exothermic Reactions

Yifeng Wang, Guangjie Feng *, Yan Wei, Bingxu Hu and Dean Deng

College of Materials Science and Engineering, Chongqing University, No. 174, Shazhengjie, Chongqing 400044, China; wangyf0902@cqu.edu.cn (Y.W.); 201909131163@cqu.edu.cn (Y.W.); 202009021009@cqu.edu.cn (B.H.); deandeng@cqu.edu.cn (D.D.)
* Correspondence: fenggj@cqu.edu.cn

Abstract: In this paper, the SiCp/Al composites were bonded via laser-induced exothermic reactions of a Ni–Al–Zr interlayer. The Ni–Al–Zr interlayer was designed based on its exothermic property and chemical compatibility with the SiCp/Al composites. The influences of the interlayer composition and bonding pressure on the joint microstructure and shear strength were investigated. Results indicated that high exothermic reactions occurred in the Ni–Al–Zr interlayer and realized the reliable bonding with the SiCp/Al composites. The interlayer products were the eutectic structure of NiAl + Ni$_2$Al$_3$Zr + Ni$_3$Al$_5$Zr$_2$. NiAl$_3$ and Ni$_2$Al$_3$ reaction layers were formed at the bonding interfaces. The interlayer composition and the bonding pressure determined the morphology and distribution of the voids and the reaction layers, thus controlling the joint shear strength. When the SiCp/Al composites were bonded using the interlayer with the Zr content of 15 wt.% under the bonding pressure of 3 MPa, the joint shear strength reached the maximum of 24 MPa.

Keywords: SiCp/Al composites; bonding; laser-induced exothermic reactions; microstructure; mechanical property

1. Introduction

Particle-reinforced aluminum matrix composites, such as SiCp/Al composites [1], Si$_3$N$_4$/Al composites [2], SiO$_2$/Al composites [3], Al$_2$O$_3$/Al composites [4], B$_4$C/Al composites [5], and TiC/Al composites [6], process excellent electrical and chemical properties, as well as the comprehensive mechanical performances, and have been important structural materials in aerospace, electronics, and automobile industries. Among them, the SiCp/Al composites are particularly attracting attention and are usually used to manufacture components with complex structures in practical applications [7,8]. However, the poor processibility of the SiCp/Al composites have greatly limited their potential applications. The deformation processing will damage the interfacial bonding between the reinforced SiC particles and the aluminum matrix, thus causing the degeneration of mechanical properties [9]. Due to the SiC particles with high hardness, machining is also not suitable for the SiCp/Al composites [10]. One solution for this issue is that the component with complex structure can be manufactured through bonding modular SiCp/Al composite pieces together. From the above viewpoint, developing a technique for reliable bonding SiCp/Al composites can further broaden their applications and is of great importance.

Since there are great differences in the chemical and physical properties between the reinforcements andmatrix, it is extremely difficult to realize the reliable bonding of SiCp/Al composites [11]. It mainly has three difficulties. First, at high temperature, harmful interfacial reactions will occur between the SiC particles and aluminum matrix, forming a large quantity of brittle Al$_4$C$_3$ compounds and resulting in the degeneration of mechanical properties. Long et al. [12] investigated the laser welding of SiCp/Al composites. They found that the high temperature during the laser welding led to the evaporation of the aluminum matrix and the excess interfacial reactions, forming large numbers of pores...
and brittle Al$_4$C$_3$. Adding appropriate Zr could partly inhibit the generation of large-size brittle phase Al$_4$C$_3$ in the joint. Second, at low temperature, the SiC particles can hardly react with surrounding materials and the bonding at the interface is very weak. Third, the alumina film on the surface of aluminum matrix also blocks the atomic diffusion at the bonding interface. To achieve a sound joint, during the welding, the interfacial reactions between the reinforcements and matrix should be inhibited to prevent the degradation of materials properties. At the same time, the metallurgical reactions between the SiC particles and surrounding materials must be enhanced to form a reliable bonding at the interface. To meet the above requirements, the chemical reactions in the bonding couple must be controlled precisely. Unfortunately, although great efforts have been done on the bonding of SiCp/Al composites, these problems have not been solved fundamentally.

Compared with conventional welding techniques, such as fusion welding [13], brazing [14,15], diffusion bonding [16], etc., the exothermic bonding has some inherent advantages on the bonding of SiCp/Al composites. It utilizes the exothermic reactions in the interlayer as the heat source to enhance the atomic activities and promote the metallurgical reactions at the bonding interface [17]. It can quickly ($10^5$–$10^6$ K/s) heat the narrow area at the bonding interface to a high temperature (about 2000 K), and the adjacent part of substrates remains a low temperature [18,19]. The transient high temperature could not only significantly increase the chemical activity of SiC particles at the bonding interface and enhance the metallurgical reactions with the filler metals, but also avoid the harmful thermal damage on the substrates. Swiston et al. [20] bonded the bulk metallic glass with the assistance of the exothermic reactions in the reactive multilayer foils. They found that during the bonding, the rapid heating and cooling made crystallization of metallic glasses unlikely. The thermal damage on the metallic glasses was voided successfully. Lin et al. [21] bonded carbon–carbon composites using the Ti/Ni–Al/Ti interlayer. They found that the exothermic reactions in the Ni–Al interlayer enhanced the metallurgical reactions and formed the NiAl$_x$ and TiC$_y$(O$_z$) phases at the bonding interface. In our previous study [22], the C$_f$/Al composites were bonded to the TiAl alloys using the exothermic bonding technique. A defect-free joint was achieved successfully. Although many efforts have been done on this bonding method, the exothermic bonding of SiCp/Al composites has been seldom reported.

As a fundamental research, this work is to explore the feasibility of bonding SiCp/Al composites via laser-induced exothermic reactions. A Ni–Al–Zr interlayer was designed and used as the bonding interlayer. The exothermic characteristics of the interlayer were analyzed. The typical joint microstructure was characterized. The influences of the bonding parameters on the microstructural evolution and mechanical property of the joint were investigated systematically.

2. Experimental Procedures

The SiCp/Al composites applied in this study were fabricated using the squeeze casting method. The volume fraction of SiC particles was 50% and the aluminum matrix was the 6063 aluminum alloy. The microstructure of the SiCp/Al composites was given in Figure 1. Prior to the bonding, the SiCp/Al composites were cut into small pieces with the sizes of 12 mm × 6 mm × 2 mm and 5 mm × 5 mm × 4 mm, respectively. Then, all the bonding surfaces were polished by the diamond abrasive discs to 600#, and cleared in acetone for 10 min with the assistance of ultrasonic to remove the impurities.

The raw materials to prepare the powder interlayer were the commercial Ni (99.5%, 28 µm), Al (99.5%, 28µm), and Zr (99.5%, 28 µm) powders purchased from Beijing Xingrongyuan Technology Co., Let. The weighted powders were put in an agate jar and milled with alumina balls for 60 min. Ten (10) mL Hexane was added in the agate jar as the process controlling agent and to prevent the oxidation of the powders. The rotational speed was set as 300 rpm and the mass ratio of the grinding media to the material was 10:1. Then, 0.5 g milled powders were cold-pressed to a cylindrical compact (Φ 10 mm). The bonding process was conducted in the argon atmosphere. The joining schematic diagram
is represented in Figure 2. During the bonding, the exothermic interlayer was applied between the SiCp/Al substrates. A laser beam (YAG-W100E, P = 200 W, Han’s Laser Technology Industry Group Co., Ltd, Shenzhen, China) heated the interlayer and ignited the interlayer. Then, the combustion flame propagated to the other side and finished the bonding process.

![Figure 1. Microstructure of the SiCp/Al composites.](image1)

![Figure 2. Schematic diagram of the bonding process.](image2)

The exothermic performance of the interlayer, the joint interfacial microstructure, and interlayer products were characterized by the differential scanning calorimetry (DSC, NETZSCH STA449C, Erich NETZSCH GmbH & Co. Holding KG, Selb, Germany), scanning electron microscopy (SEM, FEI Nova400, FEI Company, Hillsboro, OR, USA) equipped with the energy-dispersive spectrometer (EDS, FEI Company, Hillsboro, OR, USA), and X-ray diffraction (XRD, D8 ADVANCE, Bruker Company, Karlsruhe, Germany). The joint shear strength was tested at room temperature using the universal test machine (Instron-1186, Instron Corporation, Canton, OH, USA) with a shear rate of 0.5 mm·min⁻¹. The schematic diagram of the shear test is shown in Figure 3.
characteristic. It requires that the interlayer should have a high exothermicity and a rapid heat releasing characteristic. Meanwhile, the high temperature residence time should also be very short to avoid the excess interfacial reactions in the SiC particles and the surrounding materials. On the other hand, the high-temperature region should be localized around the bonding interface and should be as smaller as possible. Meanwhile, the high temperature residence time should also be very short to avoid the excess interfacial reactions in the SiC particles and the surrounding materials. On the other hand, the high-temperature region should be localized around the bonding interface and should be as smaller as possible. Meanwhile, the high temperature residence time should also be very short to avoid the excess interfacial reactions in the SiC particles and the surrounding materials. On the other hand, the high-temperature region should be localized around the bonding interface and should be as smaller as possible.
Figure 4. (a) Ni–Al phase diagram; (b) Ni–Al–Zr ternary phase diagram.

The calculation results suggest that the equimolar Ni–Al system has the highest \( T_{\text{ad}} \) (1912 K). During the exothermic bonding, the combustion stability is also very important. Merzhanov et al. put forward an empirical formula to evaluate the self-sustained reaction [32]. It indicates that when the exothermic reaction has a \( T_{\text{ad}} \) higher than 1800 K, the exothermic reaction can self-sustained. According to the empirical formula, the equimolar Ni–Al system can self-sustained once being ignited.

During the welding of the SiC\(_p\)/Al composites, one problem is the weak bonding between the SiC particles and filler metals owing to the high chemical inertness of the SiC reinforcements. To increase the bonding quality, the chemical affinity of the SiC reinforcements and the interlayer products should be promoted. In the brazing of SiC ceramics, the active filler metals (containing Ti or Zr) were usually used. It indicated that the Zr element could react with the SiC ceramics and form the ZrC and Zr\(_2\)Si reaction layers. By means of the reaction layers between the active elements and the SiC ceramics, a reliable bonding could be ensured. Thus, in this study, the Zr element was added into the Ni–Al system, forming the Ni–Al–Zr interlayer. There are two reasons to the addition of the Zr element. First, the Zr element is the active element and the carbide-forming element. It can easily react with the SiC particles and form a reliable metallurgical bonding. Second, according to the Ni–Al–Zr ternary phase diagram (Figure 4b), the Ni–Al–Zr system can form a eutectic product of NiAl + Ni–Al–Zr. The formation of the eutectic products can lower the solidification temperature and prolong the existence time of the liquid phase. Thus, under the action of the bonding pressure, the interlayer products can be densified much more easily, which can largely decrease the formation of the voids in the joint.

To evaluate the exothermic property of the interlayer, the DSC test was conducted on the Ni–Al and Ni–Al–Zr interlayer (Figure 5). For the Ni–Al interlayer, there were two sharp exothermic peaks, which were located at the heating temperature of 840 K and 916 K, respectively. The main exothermic peak was at the heating temperature of 840 K. After adding a small amount of Zr element, the position of exothermic peaks changed slightly and shifted to 762 K and 904 K, respectively. The height of the exothermic peaks decreased slightly. This is mainly caused by the addition of Zr acting as a diluent and absorbing some reaction heat. Meanwhile, after the addition of Zr in the Ni–Al interlayer, the position of the main exothermic peak decreased by 78 K, which meant that the exothermic reactions in the interlayer could be ignited more easily. The additive Zr is necessary since it can largely enhance the metallurgical bonding and increase the bonding quality. If the Zr content is in an appropriate range, the Ni–Al–Zr interlayer still can maintain a high exothermicity and can provide sufficient heat for the bonding. According to above calculation method and the previous study [17], when the Zr content ranges in 0 wt.% ~36.08 wt.%, the \( T_{\text{ad}} \) of the Ni–Al–Zr system was still maintained at a high temperature (1912 K) and the high exothermicity of the Ni–Al–Zr interlayer could be ensured.
Figure 4. (a) Ni–Al phase diagram; (b) Ni–Al–Zr ternary phase diagram and previous studies [17,22], they were supposed to be the NiAl and NiAl2 phases, respectively. The white needle phase in Fig. 2a was Al3C6. The black plate-like phase was the dark gray layer (point C) and the gray layer (point D), were formed at the bonding interface, there were some pores and some residual Al. This phenomenon was mainly owing to the particularity of the exothermic bonding. During the bonding, the exothermic reactions occurred in the interlayer and released a great amount of heat. At the center of the interlayer, the interlayer products were maintained at a high temperature and were in a solid–liquid state. Under the bonding pressure, they could be easily densified and the pores were largely eliminated. However, in the regions near the bonding interfaces, the SiCp/Al substrates were maintained at the room temperature and absorbed some reaction heat from the interlayer. So, the interlayer products cooled rapidly and could hardly be fully densified, forming the voids in the region near the bonding interface.

3.2. Joint Microstructure

Figure 6 shows the microstructure of the SiCp/Al joint bonded using a Ni–Al–Zr interlayer under a pressure of 3 MPa. It could be seen from the overall morphology in Figure 6a that the interlayer reacted completely and formed a good bonding with the substrates. In the center of the joint, the interlayer products were quite dense. Near the bonding interface, there were some pores and some residual Al. This phenomenon was mainly owing to the particularity of the exothermic bonding. During the bonding, the exothermic reactions occurred in the interlayer and released a great amount of heat. At the center of the interlayer, the interlayer products were maintained at a high temperature and were in a solid–liquid state. Under the bonding pressure, they could be easily densified and the pores were largely eliminated. However, in the regions near the bonding interfaces, the SiCp/Al substrates were maintained at the room temperature and absorbed some reaction heat from the interlayer. So, the interlayer products cooled rapidly and could hardly be fully densified, forming the voids in the region near the bonding interface.

Figure 6. Typical microstructure of the SiCp/Al joint bonded with a Ni–Al–Zr interlayer (Zr 15 wt.%, 3 MPa). (a) The overall joint; (b) the bonding interface; (c) the interlayer products.
To analyze the characteristics of the joint in detail, the morphologies at the bonding interface and the interlayer products were enlarged in Figure 6b,c. It could be seen from Figure 6b that at the bonding interface, the interlayer products bonded well with the SiCp/Al composites. No cracks and voids can be observed at the bonding interface. In the SiCp/Al composites near the bonding interface, some dark gray block phase (point A) and white needle-like phase (point B) were formed. Two continuous reaction layers, which were the dark gray layer (point C) and the gray layer (point D), were formed at the bonding interface. Near the reaction layers, some off-white phases (point E) were also observed. In Figure 6c, the interlayer products presented an eutectic feature. It consisted of large dark block phase (point F), gray reticular phase (point G), and white small block phase (point H).

To determine the phase composition of the joint, the EDS analysis was conducted on the above zones and the results were given in Table 1. The dark gray phases (point A and C), gray reaction layer (point D), and off-white phase (point E) consisted of Ni and Al elements. The ratios of Ni:Al were about 1:3, 2:3 and 1:1, respectively. According to the Ni–Al phase diagram and previous studies [33,34], they were supposed to be the NiAl, Ni2Al3, and NiAl, respectively. The white needle-like phase (point B) consisted of Ni, Al, and Zr elements and was confirmed as the (Ni,Zr)Al3 phase. The interlayer products were a mixture of three phases. According to the phase diagram and previous studies [33,34], the dark phase (point F) was confirmed to be the NiAl phase. The gray reticular phase (point G) and white small block phase (point H) were supposed to be the Ni3Al5Zr2 and Ni2AlZr, respectively. The formation of Ni3Al5Zr2 and Ni2AlZr was mainly caused by the Zr atoms replacing the places of Ni or Al atoms in the NiAl space lattice. To further confirm the phase composition of the interlayer products, the XRD test was carried out. The peak refinement was conducted with the assistance of the MDI Jade software. According to the study by Chérif et al. [35] and the peak refinement results in Figure 7, the interlayer products were confirmed as a mixture of NiAl + Ni3Al5Zr2 + Ni2AlZr, which proved the above analysis.

Table 1. EDS results on zones in Figure 6 (at.%).

| Zone | Ni     | Al     | Zr     | Possible Phase     |
|------|--------|--------|--------|--------------------|
| A    | 26.63  | 73.37  | —      | NiAl3              |
| B    | 6.03   | 84.95  | 9.02   | (Ni,Zr)Al3        |
| C    | 28.07  | 71.67  | 0.26   | NiAl3             |
| D    | 42.73  | 57.08  | 0.19   | Ni2Al3           |
| E    | 60.33  | 39.67  | —      | NiAl              |
| F    | 50.45  | 47.51  | 2.04   | Ni3Al5Zr2        |
| G    | 37.93  | 42.17  | 19.9   | Ni2AlZr         |
| H    | 53.03  | 25.39  | 21.59  | Ni2AlZr         |

Figure 7. XRD results of the interlayer products.
3.3. Influence of the Interlayer Chemical Composition on the Joint Microstructure

With the increase of Zr content in the interlayer, the joint morphology underwent a dramatic change, as shown in Figure 8. These changes were mainly displayed in two aspects. First, with the increase of the Zr content, the density of the joint had a significant improvement. The amount and the size of the voids in the joint decreased obviously. Second, with the increase of the Zr content, the residual white Ni particles gradually increased. When the Zr content was 20 wt.%, there were large numbers of the residual Ni particles around the bonding interfaces and some large voids were also observed at the bonding interfaces.

Figure 8. Microstructures of the joints bonded with different interlayers. (a) Zr 5 wt.%; (b) 8 wt.%; (c) 15 wt.%; (d) 20 wt.%.

To analyze the microstructure of the joints in detail, the regions of the interlayer products and the bonding interfaces were enlarged in Figures 9 and 10, respectively. It could be seen that the interlayer products with different Zr contents all presented a eutectic morphology of mixed dark NiAl phase and white Ni–Al–Zr IMC, as shown in Figure 9. When the Zr content was 5 wt.%, the Ni–Al–Zr IMC was less and separated by the dark block NiAl phase. With the increase of the Zr content, its volume fraction increased gradually. The Ni–Al–Zr IMC became continuous gradually and the grain of the NiAl phase decreased obviously. Compared with the NiAl phase, the ternary Ni–Al–Zr compounds had a much higher hardness and brittleness [36]. Thus, with the increase of the Zr content, the high-volume fraction of the Ni–Al–Zr IMC increased the brittleness of the interlayer products. On the other hand, the higher volume fraction of the Ni–Al–Zr IMC also meant the increase of the eutectic liquid in the interlayer products. During the exothermic bonding process, the higher volume fraction of the eutectic liquid promoted the densification process and was conducive to eliminating the voids in the joints. Thus, the amount and the sizes of the voids in the joint gradually decreased.
The interlayer had a slight decrease. The decrease of the interlayer and no defects were observed at the bonding interface. However, with the increase of the Zr content, the voids and the unconnected areas gradually disappeared. The reaction products were the eutectic structure of NiAl+NiAl. The density and bonding quality of the joint increased. Under high bonding pressure, the reaction products were in the solid state, but the atomic activity at the bonding interface was reduced due to the exothermicity of the Zr element. It provided the required energy for the bonding process. During the bonding process, the interlayer products were a mixture of brittle NiAl and Ni IMC. As the Zr content increased, the voids and the unconnected areas gradually disappeared. The joint density was quite high. The residual Ni particles became more and more disordered. The density and bonding quality of the joint increased.

![Figure 9. Microstructures of the different interlayer products.](image)

(a) Zr 5 wt.%; (b) 8 wt.%; (c) 15 wt.%; (d) 20 wt.%

![Figure 10. Interfacial bonding characteristics of the joints bonded with different interlayers.](image)

(a) Zr 5 wt.%; (b) 8 wt.%; (c) 15 wt.%; (d) 20 wt.%
The interfacial microstructures of the different bonding interfaces with different Zr contents were demonstrated in Figure 10. With the increase of the Zr content, the interfacial microstructures at the bonding interfaces mainly changed in two ways. On one hand, the NiAl$_3$ and Ni$_2$Al$_3$ reaction layers at the interfaces became discontinuous gradually. On the other hand, the amount and the size of the residual Ni particles increased gradually at the bonding interfaces. These changes were mainly due to the different exothermocities of the interlayers with different Zr contents. The analysis in Section 3.1 indicated that the exothermic reactions between the Ni and Al could release a large quantity of heat and provided the required energy for the bonding process. During the bonding, the additive Zr absorbed some reaction heat and acted as the coolant. Thus, with the increase of the Zr content, the exothermicity of the interlayer had a slightly decrease. The decrease of the exothermicity shorted the high-temperature residence time of the bonding interface and reduced the atomic activity at the bonding interface, thus affecting the interfacial reaction. The reaction layers became thinner and continuous gradually. Meanwhile, since the interlayer exothermicity decreased with the increase of the Zr content and the SiCp/Al composites absorbed some heat, the cooling rate at the bonding interface increased significantly. It affected the exothermic reactions in this region. The reactions in this region were incomplete and some residual Ni particles were observed. With the increase of the Zr content, the residual Ni particles became more and more plentiful. Zr element is a carbide-forming element. Qi et al. [37] and Zhou et al. [38] brazed SiC ceramics with filler metals containing Zr element. They suggested the Zr element promoted the metallurgical reactions between the SiC ceramic and the filler metals. In this study, the higher Zr content in the interlayer also enhanced the bonding quality between the SiCp/Al composites and the interlayer. As the Zr content increased, the voids and the unconnected areas gradually disappeared. When the Zr content was 15 wt.%, the SiCp/Al composites had a reliable bonding with the interlayer and no defects were observed at the bonding interface. However, with the further increase of the Zr content, the voids appeared again due to the low exothermicity of the interlayer.

3.4. Influence of the Bonding Pressure on the Joint Microstructure

The microstructures of the SiCp/Al joints bonded at different bonding pressures were shown in Figure 11. Under different bonding pressures, the density of the joints changed significantly. The reaction products were the eutectic structure of NiAl + Ni–Al–Zr IMC, as shown in Figure 6. During the bonding process, the interlayer products were in the solid–liquid state. Under the action of the bonding pressure, the interlayer products were squeezed by the adjacent substrates, and the original pores in the interlayer could be largely eliminated. The higher the bonding pressure was, the denser the interlayer products were. When the bonding pressure was low (1.5 MPa), a large number of the voids were observed in the joint. With the increase of the bonding pressure, the amount and size of the voids decreased significantly. When the bonding pressure reached 3 MPa, large voids disappeared in the joint. There were only some small voids distributed in the region adjacent to the bonding interface. The density and bonding quality of the joint had a dramatic improvement. When the bonding pressure further increased to 4 MPa, the small voids around the bonding interfaces almost disappeared. The joint density was quite high. However, since the interlayer products were a mixture of brittle NiAl and Ni–Al–Zr IMC, under high bonding pressure, a fracture occurred in the interlayer products and propagated to the SiC/Al composites, as shown in Figure 11d.

3.5. Joint Shear Strength

The shear strength was used to evaluate the joint mechanical property. With the increase of the Zr content in the interlayer, the joint shear strength first increased and reached the highest value (24 MPa) when the Zr content was 15 wt.%, then it decreased to 15 MPa, as shown in Figure 12. The joint shear strength depended on the joint microstructure [39]. As analyzed in Section 3.3, the Zr content influenced the joint microstructure in three ways,
which were (i) the joint density, (ii) the residual Ni particles, and (iii) the reaction layers at the bonding interfaces, respectively. When the Zr content was 5 wt.%, the interlayer products were mainly the NiAl phase. The low-volume fraction of the Ni–Al–Zr IMC led to less eutectic liquid during the bonding process. Thus, the original pores in the interlayer were unable to be eliminated effectively. The large number of the voids in the joint greatly affected the mechanical property of the joint. With the increase of the Zr content, the amount and size of the voids in the joint decreased significantly. When the Zr content was 8 wt.%, the voids were small and mainly distributed around the bonding interfaces. The joint shear strength slightly increased to 16 MPa. When the Zr content reached 15 wt.%, the voids in the joint further decreased. However, the high Zr content slightly reduced the exothermicity of the interlayer. Since the adjacent SiCp/Al substrates absorbed some reaction heat from the interlayer, the reactions in the region near the bonding interfaces were incomplete. Some residual Ni particles were observed around the bonding interfaces. The decrease of the interlayer exothermicity also affected the interfacial reactions at the bonding interfaces. The reaction layers became thinner and uncontinuous gradually. It should be noticed that there was a competitive relationship between the joint density and the reaction layers. With the increase of the Zr content, the joint density gradually increased, which was beneficial to the joint shear strength. Meanwhile, the reaction layers became thinner and uncontinuous gradually, which had unfavorable effects for the bonding quality. The joint mechanical performance was a combined result of these two aspects. When the Zr content was 15 wt.%, these two aspects were in a balance. Thus, the joint shear strength reached the highest value (24 MPa). With the further increase of the Zr content, the joint density did not continue to increase. The high fraction of Ni–Al–Zr IMC greatly increased the brittleness of the interlayer products. The large amount of the residual Ni particles and the thin reaction layers greatly affected the bonding quality. Thus, the joint shear strength decreased to 15 MPa when the Zr content was 20 wt.%.

Figure 11. Microstructures of the joints bonded under different pressures. (a) 1.5 MPa; (b) 2 MPa; (c) 3 MPa; (d) 4 MPa.
The influence of the bonding pressure on the joint shear strength was similar to that of Zr content, as shown in Figure 12. The bonding pressure mainly influenced the joint density. When the bonding pressure was 1.5 MPa, the interlayer products only had small extrusion deformation. The original pores in the interlayer were largely kept in the joint. The large amount of the voids decreased the joint’s ability to withstand loads. The joint shear strength was as low as 4 MPa. With the increase of the bonding pressure, the interlayer products suffered a larger extrusion deformation, which effectively reduced the amount and size of the voids. In consequence, the joint shear strength had an obvious increase. When the bonding pressure was 3 MPa, there were only some small voids distributed around the bonding interfaces. The joint shear strength reached the maximum of 24 MPa. When the bonding pressure further increased to 4 MPa, the high pressure led to the fracture of the brittle interlayer products. Continuous cracks were formed in the joint. It significantly affected the joint mechanical performance. The joint shear strength decreased to 21 MPa.

4. Conclusions

(1) The equimolar Ni–Al system was chosen as the fundamental exothermic system due to its high exothermic property. The active Zr element was added to promote the metallurgical reactions between the SiC particles and the interlayer. NiAl$_3$, Ni$_2$Al$_3$ reaction layers were formed at the bonding interfaces and ensured the reliable bonding of the SiCp/Al composites.

(2) The additive Zr transformed the interlayer products from the NiAl to the eutectic organization of NiAl + Ni$_2$Al$_3$Zr + Ni$_3$Al$_2$Zr. Higher Zr content increased the joint density, but slightly decreased the interlayer exothermicity and led to the thin reaction layers. Higher bonding pressure increased the extrusion deformation of the interlayer products and effectively eliminated the original pores in the interlayer products.

(3) With the increase of the Zr content and the bonding pressure, the joint shear strength first increased and then decreased. When the Zr content was 15 wt.% and the bonding pressure was 3 MPa, the joint shear strength reached the maximum of 24 MPa.

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