NiAl intermetallics is considered to be a potential high temperature structural material based on a high melting point, low density, high thermal conductivity, excellent oxidation and hot-corrosion resistance at elevated temperatures [1–4], but its high temperature strength is insufficient for the majority of actual applications. The tensile ductility and fracture toughness at ambient temperatures are often also unsatisfactory [5, 6]. The solid solution strengthening, precipitation strengthening, directional solidification and single crystal technique have been explored to improve its high temperature strength. Although significant progress has been achieved, the ambient tensile ductility and fracture toughness often deteriorate as a result. On the other hand, the ductility and toughness can also be improved by microalloying, macroalloying and plastic secondary phase additions, but this at the expense of the strength [4–7]. In other words, it is very difficult to compromise its high temperature strength and ambient temperature ductility. This severely restricts its operating temperature and deteriorates its machinability. It is significant to explore novel fabricating techniques for NiAl intermetallic alloy to improve its comprehensive mechanical properties.

Preparing intermetallic matrix composite is an effective method to produce engineering NiAl intermetallic alloy and to improve its high temperature strength and retain ambient temperature ductility simultaneously [8, 9]. Good ductility can be obtained through adequately alloying the intermetallic matrix while the strength is improved with the reinforcements. The casting, combustion synthesis, powder metallurgy and mechanical alloying are major methods for preparing NiAl intermetallic matrix composite. Nevertheless, in many cases, the mechanical bonding due to interfacial reaction only result in weak interfacial bonding so that whole properties of the composite might even be worse than the matrix itself.

Based on clear and compatible interface, in situ reinforced composite generally shows better comprehensive mechanical properties than external additive ones. Laser cladding has been introduced to prepare particulate reinforced intermetallic composite coatings, for example, laser clad TiC/NiAl-Ni3(Al, Ti, C) wear-resistant coating [10]. However, this research was limited to the pure nickel substrate, elemental powder additives, pre-placed powder bed, definite matrix composition and a few reinforcement contents. Moreover, the matrix of the coating was mainly Ni3Al phase, no NiAl phase, and the research aimed at the coating preparation, no bulk materials.

In this investigation, laser powder deposition fabricated in situ TiC particulate reinforced Ni30Al20Fe intermetallic matrix composite was carried out with Ni/Al, Fe and TiC powders by coaxial powder delivery. A3k W continuous wave CO2 laser, attached with a powder feeder and a coaxial powder-feeding nozzle, was used for the laser deposition experiment. Ar was applied as the shield gas to avoid oxidation.
Figure 1 Morphologies of the original powders: (a) Ni/Al powders, (b) Fe powders and (c) TiC powders.

and contamination of the melted powders and pool. Through preliminary processing experiments, the optimized processing parameters were identified to ensure good metallurgical quality: laser power \( P = 1.5 \) kW, scanning velocity \( V = 4 \) mm/s, powder feeding rate \( Q = 4.5 \) g/min and laser beam diameter \( D = 5 \) mm. The bulk material was fabricated by track-by-track and layer-by-layer overlapping in the horizontal and vertical directions.

The microstructure of the bulk material was examined by a CSM950 scanning electron microscopy (SEM) and a JSM-6301F high-resolution scanning electron microscopy (HSEM). The material compositions were determined by a Link ISIS energy-dispersive spectrometry (EDS). The phase constituents were identified with a Rigida X-ray diffractometer (XRD). The microhardness was measured by HX-200 Vickers microhardness tester and a load of 100 g.

The surface and cross section morphologies of the laser deposited bulk material were examined with the naked eye and SEM. The results show that the material was fully dense and free of cracks and pores. This should mainly attribute to the optimization of processing parameters. Since slightly lower laser power, slower scanning velocity and larger beam diameter were used, the heating and cooling rates were slowed down and the temperature distribution was more uniform in the deposits. As a result, the thermal stress in the deposited material could be reduced considerably so that cracking was eliminated. Because the melting pool could be retained for a longer time, the gases dissolved could escape fully so that pores could be eliminated and the fully dense bulk material could be obtained.

The XRD profile of the laser deposited bulk material is given in Fig. 2. It is found that the actual position of every diffraction peak is exactly identical to JCPDS standard data of \( \beta \)-NiAl (B2 structure) and TiC (Cubic structure). Therefore it can preliminarily be considered that the laser deposited bulk material mainly consists of the \( \beta \)-NiAl intermetallics and TiC phase. Furthermore, it can be found later that the titanium carbides were precipitated \textit{in situ}.

Fig. 3 is the SEM images of the microstructure of the laser deposited bulk material. Fig. 3a is a full view of the morphology of the microstructure. A quantity of refined and dispersed particles, whose sizes are far smaller than the original TiC additives, homogeneously distributed on the matrix. The chemical compositions of the matrix and precipitated particles are given in Table I. The compositions of the matrix are in the range of \( \beta \) phase area in Ni-Al-Fe ternary phase diagram [11]. It can be considered as a Ni30Al20Fe alloy. It must be pointed out that Al concentration in the bulk material was reduced, compared to the original powders, due to oxidation and evaporation in the process of laser deposition. The dispersed particles mainly include Ti and C elements with a near iso-atomic proportion. A few Ni, Al and Fe

<table>
<thead>
<tr>
<th>Positions</th>
<th>Ni</th>
<th>Al</th>
<th>Fe</th>
<th>Ti</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>The matrix</td>
<td>49.32</td>
<td>29.86</td>
<td>19.13</td>
<td>1.69</td>
<td>–</td>
</tr>
<tr>
<td>The particle</td>
<td>2.52</td>
<td>2.81</td>
<td>5.63</td>
<td>47.79</td>
<td>41.25</td>
</tr>
<tr>
<td>The ( \gamma ) phase</td>
<td>45.28</td>
<td>7.94</td>
<td>46.26</td>
<td>0.52</td>
<td>–</td>
</tr>
</tbody>
</table>
particles are also found in the dispersed particles because they are so tiny that compositional information of the adjacent matrix was inevitably collected. According to Table I, in combination with the XRD results in Fig. 2, it is affirmable that the matrix is a NiAl intermetallic alloy dissolved with Fe and Ti atoms and the precipitated particles are titanium carbides. For comparison, the microstructure obtained under inappropriate processing conditions, where relatively low laser power density was used, is shown in Fig. 3b. In this case the TiC particles are apparently coarser than the ones in Fig. 3a and similar to original particles shown in Fig. 1c. This means that the majority of the TiC particles are unmelted. This also proves that the bulk material obtained under the optimum processing conditions, where relatively low laser power density was used, is shown in Fig. 3a. This means that the in situ TiC particle is much smaller than the additive TiC particle. The fine and dispersed TiC particles are favorable to the improvement of the whole properties of the NiAl intermetallic alloy.

Fig. 3c is the microstructure of the matrix of the laser deposited bulk material. It is composed of homogeneous equiaxed grains of 5 µm or so. A few super-superfine particles, whose compositions are given in Table I, can be recognized. According to other researches, some Ni-Al-Fe series intermetallic alloys included a few fcc γ-Fe-Ni phases that usually precipitated from β-NiAl-Fe phases and were distributed along crystal boundaries in appearance of lamellar or particle and could serve as the ductile improvement [11]. So they should be Fe-rich γ-Fe-Ni phase precipitated from β phase, although they were not enough to be identified by XRD analysis. Fig. 3d is a highly-enlarged SEM image giving a clear insight on the in situ TiC reinforcements and the adjacent matrix. The in situ TiC particles are characterized by their fine size, quasi-quadrangle shape, and cube spatial configuration.

The average microhardness of the laser deposited bulk material reached HV0.1595, which was much higher than HV250-350, the microhardness of Ni-Al-Fe intermetallic alloys. Based on the above research results, the following conclusions can be drawn. With the mechanical mixtures of Ni/Al cladding powders, Fe fused and crushed powders and TiC sintered and crushed powders, as well as coaxial powder feeding mode, in situ TiC particulate reinforced Ni30Al20Fe intermetallic matrix composite could be successfully synthesized and prepared by laser powder deposition. Under optimized processing parameters, laser deposited bulk material was fully dense and free of cracks and pores. The microstructure of in situ TiC particulate reinforced Ni30Al20Fe intermetallic matrix composite consisted of the refined β-NiAl-Fe intermetallic matrix, homogeneous and dispersive in situ TiC reinforcements and discontinuous γ-Fe-Ni phases. The microhardness of the bulk material, HV0.1595, was much higher than Ni-Al-Fe intermetallic alloys without reinforcements. This could mainly be attributed to in situ particulate reinforcement.
Figure 3 Microstructure of laser deposited particulate reinforced intermetallic matrix composite material: (a) In situ TiC particles dispersed on the matrix, (b) unmelted TiC particles and the matrix, (c) dense and homogenous intermetallic matrix and (d) fine in situ TiC particulates and the matrix.

Figure 4 A comparison of in situ and unmelted TiC particles and their interfaces with the matrix: (a) unmelted TiC particle, (b) in situ TiC particles, (c) interface between unmelted TiC particle and the matrix and (d) interface between in situ TiC particle and the matrix.
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