Effects of temperature on interfacial evolution and mechanical properties of pure titanium and carbon steel sheets bonded via new multi-pass continuous hot-roll diffusion with nickel interlayer

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Abstract
This study performed experiments and thermodynamic calculations to elucidate the effects of diffusion temperature on interfacial evolution and mechanical properties of pure titanium and carbon steel (i.e., steel) sheets bonded via a new type of multi-pass continuous hot-roll diffusion with nickel interlayer. The interfacial evolution results revealed that this new type of multi-pass continuous hot-roll diffusion treatment showed a very good adherence due to its metallurgy bonding, because it made a remarked improvement between compound and intermetallic compound relationship. Secondly, in mechanical properties, results revealed that the highest shear strength (∼470 MPa) was obtained at a processing temperature of 850°C. The highest peel strength (∼21 N/mm) was obtained in the sample processed at 900°C. Bonding temperatures above and below these levels reduced the bond strength respectively due to poor atom diffusion and excessive compound formation, resulting in joint failure at the Ti-Ni interface. Extensive cleavage planes with various alignments were observed on the fracture surfaces in these cases. Overall, a hot-rolling temperature of 850°C was found to provide the optimal tradeoff between interfacial bonding strength and ductility. This work provided an economical and convenient solution for broadening the engineering application of interface between sheets of pure titanium and steel.

Keywords Diffusion bonding · Phase reaction · Microstructure · Mechanical properties

1 Introduction

The light weight, high strength-to-weight ratio, high toughness, high creep resistance, and excellent corrosion resistance of titanium (Ti) and its alloys have led to their extensive use in the aerospace, chemical, marine, nuclear, and medical industries [1, 2]. In many applications, it is necessary to bond Ti with some other metal or alloy in order to achieve the desired functional performance [3]. However, Ti is extremely reactive at high temperatures, and hence the bonding process represents a significant challenge since the Ti readily reacts with oxygen and/or nitrogen in the atmosphere [4, 5]. Nonetheless, bi-metallic joints containing Ti are extremely attractive due to their superior mechanical properties and potential to reduce the amount of materials needed to realize the joint, thereby reducing the cost. Many methods are available for joining Ti to steel, including welding [6–8], brazing [9–11], and soldering [12, 13]. However, these methods have several drawbacks, including a large difference in the thermal expansion coefficients of Ti and steel, respectively, which may cause significant distortion, and the rapid formation of intermetallic layers.
in the interfacial region [11]. The intermetallic layers typically consist of FeTi and Fe$_2$Ti phases, both of which have high hardness and brittleness, and therefore reduce the strength of the joint [14–16].

To improve the practical applicability of Ti-steel sheet (SS) bonded joints, it is necessary to increase their plasticity by reducing the volume fraction of intermetallic compounds (IMCs) using some form of solid-state connection method or selecting a suitable interlayer material for the Ti alloy and stainless steel so as to form a solid solution between them during the bonding process. The use of hot-roll diffusion bonding to form dissimilar joints has attracted significant attention in the recent literature [17–19]. Hot-roll bonding provides an effective means of joining similar or dissimilar materials without gross microscopic distortion or large dimensional tolerances [20, 21]. Furthermore, virtually no phase transformation or microstructural change occurs during the bonding process, and the inherent mechanical properties and behaviors of both metals are preserved [22]. In addition, the diffusion-bonding process minimizes structural inhomogeneities due to the negligible effect of the temperature gradient and yields significant cost and weight savings compared to traditional joining techniques [16]. Finally, the bonding time is very short, and thus the volume fraction of IMCs formed at the bonding interface is greatly reduced. Many studies have shown that the inclusion of an aluminum (Al) [23–25], copper (Cu) [26, 27], nickel (Ni) [22], or silver (Ag) [16] interlayer increases the strength of hot-roll diffusion-bonded Fe-Ti joints [17–19, 28] and suppresses the formation of IMCs [27–32]. However, even though a small improvement in the bonding strength is achieved, the bonding strength at the joints is still limited by the formation of brittle IMCs at either the Ti-interlayer or the interlayer/SS interface. Thus, it is essential to eliminate, or at least suppress, the formation of these IMCs. In this regard, Ni is one of the most effective intermediate materials for the diffusion bonding of Ti-SS since, according to the Ni-Fe and Ni-Cr binary phase diagrams, Ni does not form IMCs with any of the main elements of SS [33]. In addition, even though TiNi-TiNi$_2$ and TiNi-TiNi$_3$ compounds may be formed at the Ti-Ni interface, previous studies [34, 35] have shown that these phases are not as brittle as many other Ti-based intermetallic phases. Moreover, they coexist within the matrix (microstructure or interface) and have good toughness, thereby improving the bonding strength.

Hot rolling is an effective means of fabricating metallic composite sheets consisting of pure titanium and carbon steel with a nickel interlayer. Many studies have shown that hot rolling has the advantages of high efficiency, a low cost, and an improved product quality due to its effects in improving the distribution of the intermediate material, increasing the relative density, and refining the grains and precipitates [4, 6, 7, 36]. However, if the volume fraction of the intermediate material is too high (i.e., hard brittle metals), the edge of the composite rolled sheet may be cracked [37, 38]. Therefore, hot rolling is more suitable for the processing of low-volume fraction intermediate materials (i.e., ductile metals). Nonetheless, the combination of hot rolling and diffusion bonding enables the fabrication of metallic composite sheets consisting of Ti-SS and a nickel interlayer with an excellent bonding strength.

As described above, the bonding strength of Ti-SS composite sheets can be improved by the introduction of two or more reinforcing phases. However, the interfacial strength of such tri-metal systems (i.e., the tri-metal composite sheet) is heavily dependent on their microstructural evolution during the bonding process. The evolution process is dependent in turn on the temperature at which rolling is performed. Therefore, it is essential to confirm the quantitative relationships among the diffusion-controlled reaction, the rolling temperature, and the hot deformation behavior of the constituent materials in the hot-rolling process. However, the literature provides relatively little information pertaining to the effects of processing temperature on the microstructure and mechanical properties of hot-roll diffusion-bonded joints fabricated using interlayers. This study performed experiments and thermodynamic calculations to elucidate the effects of diffusion temperature on microstructural evolution at the interfaces, including phase transformation, the growth of the diffusion layers, and the sequence of compound formation. Thermodynamic calculations focused on the Gibbs free energy and interface energy to predict the sequential phase formation at the interfaces. In the current study, we investigated the microstructural evolution and metallurgical reactions that occurred in joining Ti and carbon steel (i.e., steel) via hot-roll diffusion bonding with a nickel interlayer at temperatures of 800, 850, and 900°C, respectively. Our findings related to the influence of diffusion temperature on interface evolution provide important theoretical insights and practical guidance for the development of components in medical devices, actuators, and gas turbine engines.

2 Experimental procedure

2.1 Base metal

The hot-roll diffusion-bonding trials were performed using pure Ti and carbon steel plates (All Metal Sales Inc., Cleveland, OH) with dimensions of 150 mm × 100 mm × 2 mm. The chemical compositions of the two plates were as follows: 0.08C-0.12O-0.013H-0.05N-0.15Si-bal Ti (wt%) and 0.12C-0.50Mn-0.38Si-0.034P-0.035S-bal Fe (wt%), respectively. In fabricating the joints, the polished surfaces of the two plates were separated by an interlayer of pure Ni foil (purity &gt;99.95 at%) as follows: 0.01C-0.001Sn-
0.0018O-0.0005Mn-0.01Si-0.001Cr-0.001Fe-0.01Si-
0.015Ti-0.0005Co-0.0015V-bal Ni (at%) with the thickness
of 800–850 μm and the surface roughness Ra of 0.036 μm.

2.2 Hot-roll diffusion-bonding process

Prior to bonding, the Ti and carbon steel plates were cleaned
using emery paper and then respectively scratch-brushed
using circumferential brushes of pure Ti and carbon steel
(i.e., steel) to remove the oxidization layers. The edges of
the two plates were then mechanically cleaned using abrasive
paper. Finally, the plates were placed in an acetone-containing
vessel to undergo degreasing under ultrasonic agitation for 20
min. Test specimens were assembled by inserting Ni foil be-
tween the two plates. The specimens were then placed in a
furnace at 450°C for 600 s (preheating), before they
underwent hot rolling using a rolling mill to form sheets of
the tri-metal composite (i.e., Ti-Ni-steel). Specifically, the
sheets were fixed in place under 3–5 MPa uniaxial load (ap-
plied along the longitudinal direction of the specimens),
whereupon roll-bonding was multi-pass continuous hot
rolling process with the aim of reducing the thickness of
the sheets respectively by 5%, 15%, 25%, 30%, and 45% between
passes, each of which was performed at rolling quantity
should be gradually increased (i.e., this is a key point for the
experimental process) (from an initial thickness of 16.2 mm to
a final thickness of 11.34 mm). Note that the tri-metal com-
posite sheets underwent annealing at 400 °C for 5 min be-
tween passes, each of which was performed at rolling speed
of 0.2 m/s. Annealing was meant to remove work hardening
cased by the rolling process and improve deformability in
order to prevent severe edge cracking. Note also that three sets
of samples were fabricated at rolling temperatures of 800°C,
850°C, or 950°C under an inert atmosphere (high-purity argon
gas; 99.99%) to inhibit oxidation. Rolling was performed
using 100-mm rollers with a maximum capacity of 20 t (see
Fig. 1), and the roll mills were lubricated using industrial oil.
After the final rolling, the specimens were left in the furnace to
cool naturally to room temperature and then annealed at
550°C in a muffle furnace for 3 h in an inert atmosphere
(high-purity nitrogen gas; 99.99%).

2.3 Microstructural characterization

Following the bonding trials, the samples were sectioned
using an abrasive cutting tool to form specimens with a
size of 20 mm × 13.75 mm × 3.15 mm. The specimens
were etched using 3% Nital solution and Kroll’s solution
(92 ml H₂O + 2 ml HF + 6 ml HCl) for 45 s to reveal the
cross-sectional microstructure. The interfacial regions of
the joint were analyzed using an optical microscope (OM,
Olympus GX-51) and scanning electron microscope
(SEM, JEOL JSM-5410). In addition, the chemical com-
positions and element distributions were examined using
energy dispersive X-ray spectrometry (EDS). Finally, the
phase structures of the IMCs were identified by X-ray
diffraction (XRD, Rigaku, Japan) using Cu Kα radiation
with an operating voltage and current of 30 kV and 20
mA, respectively.

Fig. 1 Schematic illustration
showing structure of hot-roll dif-
fusion-bonded joints.
2.4 Mechanical properties

2.4.1 Shear strength tests

Shear tests were performed in accordance with the ASTM-D 1002-10 standard [39] (see Fig. 2a) using a universal testing machine (UTM; Santan STM-50) with a 5-kN load cell. The tests were performed at room temperature using a cross-head speed of 0.1 mm/min. The bond strength was calculated as the load at fracture (N) divided by the interfacial area of the joint (mm\(^2\)) and was recorded in megapascals (MPa). For each specimen, the shear strength was reported as the average value obtained over five separate tests. The fracture surfaces of the tested samples were examined by SEM and XRD.

2.4.2 Peel strength tests

In accordance with the ASTM-D 1876-01 standard [40] peeling test (see Fig. 2b), specimens with gauge dimensions of 25 mm (width) by 100 mm (length) were machined from the bonded specimens in a direction vertical to the Ni interlayer. Tensile tests were performed at room temperature using a UTM (Hounsfield H50KS) with a constant force of 50 kg and a crosshead speed of 20 mm/min. After each test, the break-off force was plotted against the peeling distance. The average peel strength was then evaluated as:

\[
\text{Average peel strength} = \text{average load/bond width (N/mm)}
\]

For each specimen, the peel strength was reported as the average value obtained over five separate tests.

3 Results and discussion

3.1 Interfacial microstructural evolution

For diffusion-bonded joints such as those prepared in the present study, the interfacial microstructure, diffusion layer thickness, and phase composition of the reaction zone are all critically dependent on the bonding temperature. Figure 3a–c present SEM cross-sectional images of the Ti-Ni-steel samples diffusion bonded at temperatures of 800, 850, and 900°C, respectively. For all three samples, the diffusion interface between the carbon steel plate and the Ni interlayer shows some defects, discontinuities, and voids. In the sample diffusion bonded at 800°C, the interfacial region between the Ti plate and the Ni interlayer also shows obvious defects (discontinuities and voids). However, these defects are less evident in the samples processed at higher temperatures of 850 and 900°C, respectively. In addition, Table 1 shows the thicknesses of the NiTi and remnant NiTi layers of the bonded joints formed at different temperatures. It is seen that the thickness of the NiTi layer increases with an increasing temperature. The SEM images (see Fig. 3) confirm that a higher holding diffusion temperature increases the diffusion layer thickness. This finding is reasonable, since the use of a Ni interlayer harmonizes the rolling amount and rolling temperature and consequently increases the strain value of the Ti-Ni diffusion interface. Note that the application of roll-bonding only once seldom results in a perfect bond without fractures or defects. This can be attributed primarily to the large amount of stress produced when the materials are subjected to the initial compression of the rollers. We therefore performed rolling multiple times at various bonding temperatures to reduce stress. This also...

![Fig. 2](https://example.com/figure2.png)

**Fig. 2** Schematic illustration showing a shear strength tests and b peel strength tests of hot-roll diffusion-bonded joints.
helped to increase the dislocation density and stored energy in the Ni interlayer [41] as well as increase the thickness of the diffusion layer at the Ti-Ni interface. Moreover, according to the dislocation hypothesis [41, 42], dislocations at free surfaces undergoing plastic deformation break the oxide layers and produce stairs at the atomic scale, which increase the involvement of the diffusion bonded joint parts. The phenomena are discussed later in relation to the microstructural observations.

As shown in Fig. 3, all three of the samples displayed compound layers (i.e., phases) at the interface between the carbon steel plate and the Ni interlayer. Microstructural analysis of these specimens revealed that the Ni/Ti/steel interfaces were well-differentiated. Table 2 lists the EDS analysis results showing the atomic percentages of Ti and Ni in the various diffusion specimens bonded at temperatures of 800, 850, and 900°C. In other words, only limited element diffusion occurs at the Ni-Ni-steel interface, even at the highest temperature of 900°C. However, the compound layers are apparent in the interfacial region between the Ti plate and the Ni interlayer in every case. At a lower bonding temperature (800°C, Fig. 3a), the compound layers have the form of isolated phases and are present in only relatively small amounts. However, as the bonding temperature increases, the number of compound layers also increases, and discrete reaction layers are formed (see Fig. 3b and c). In particular, on the basis of the difference in contrast, the pure Ti-Ni interlayer interface region contains a NiTi2 compound layer (i.e., dark gray region) next to the Ti plate and a Ni3Ti compound layer (i.e., gray-white region) adjacent to the Ni interlayer. The two compound layers are separated by a NiTi compound layer (i.e., gray region) (see Fig. 3c, for example). According to [43, 44], the formation process for the Ni3Ti, NiTi, and NiTi2 phases can be elaborated as follows. Under the effects of the elevated bonding temperature, a Ni3Ti compound layer is rapidly formed during the initial stage of the bonding process due to the relatively low diffusion speeds of Ti and Ni, respectively.

The strong affinity of Ti-Ni atom pairs (i.e., the high negative mixing enthalpy of the principal elements in the alloy system) caused the Ni atoms to diffuse quickly into the Ti (i.e., β-Ti) (D = 3 × 10⁻¹² m² s⁻¹ at 900°C) [45]. Isothermal transformation under the effects of Ni atom diffusion led to the formation of Ni3Ti and NiTi compounds, followed by the formation of NiTi2. These reactions resulted in the formation of a Ni3Ti compound layer, which is separated from the Ni interlayer by a NiTi compound layer and an NiTi2 compound layer. The formation of these compounds was confirmed by X-ray diffraction analysis, and the thicknesses of the NiTi and Ni3Ti layers were measured using a scanning electron microscope with an energy-dispersive X-ray analysis system. The results are listed in Table 1.

**Table 1** Thicknesses of NiTi and remnant NiTi layers in samples processed at temperatures of 800, 850, and 900°C.

| Unit: μm | 800°C | 850°C | 900°C |
|---------|-------|-------|-------|
| NiTi    | 5.8   | 12.2  | 29.1  |
| Remnant NiTi layers | Ni3Ti | 5.6   | 6.3   | Ni3Ti | 8.4   |
| NiTi2   | 3.5   | 5.8   | NiTi2 | 6.8   |
formation of β-Ti phase in accordance with the Ti-Ni binary phase diagram [46]. As the holding time increased, the inward diffusion of Ni atoms also increased. The α-Ti phase transformed into NiTi₂ through an isothermal reaction of β-Ti → NiTi₂, due to similarities in the crystal structure and composition of NiTi and NiTi₂. The β-Ti phase and disordered α-Ti phase dissolve a greater amount of Ni than the ordered NiTi₂ compound. As a result, excess dissolved Ni is expelled from the NiTi₂ phase. The segregated Ni atoms combined with the Ni atoms diffused from the Ti substrate to form Ni₃Ti. NiTi₂ with almost no Ni continuously formed adjacent to the Ni₃Ti compound, likely due to the diffusion of Ti atoms toward the NiTi at high temperatures. This balanced the diffusion of Ni in the opposite direction, which limited the formation of NiTi₂. The shorter diffusion distance of Ti should be limited to multi-layers with a short period, as shown in Table 1. The chemical affinity of Ni₃Ti phase is greater than that of NiTi₂ phase, manifesting as a difference in chemical potential. Consequently, NiTi₂ compound precipitates are formed in the α-Ti matrix in accordance with the eutectoid reaction β-Ti → α-Ti + NiTi₂ during the cooling process [46, 47]. A close inspection of Fig. 3b and c reveals the presence of α-Ti phase + β-Ti phase in the form of discrete needle-shaped particles in the matrix of the Ni₃Ti compound and β-Ti phases in the form of discrete islands in the matrix of the NiTi₂ phase in the compound reaction layers. This can be attributed to the high negative mixing enthalpy of Ni-Ti, which led to the formation and subsequent segregation of atom pairs within the matrix (NiTi: −40 kJ/mol, NiTi₂: −33 kJ/mol, and Ni₃Ti: −53 kJ/mol) [48–50]. Under these conditions, the probability of compound formation was as follows: Ni₃Ti → NiTi₂ → NiTi.

In NiTi, NiTi₂ and Ni₃Ti compounds form as a result of inter-diffusion of the Ti and Ni elements during the bonding process. The associated chemical reaction equation can be stated as follows:

\[
\text{Ni} + \text{Ti}(s) = \text{NiTi} \quad (R1)
\]
\[
\text{Ni}(s) + 2\text{Ti}(s) = \text{NiTi}_2 \quad (R2)
\]
\[
3\text{Ni}(s) + \text{Ti}(s) = \text{Ni}_3\text{Ti} \quad (R3)
\]
\[
\text{Ti} \text{Ni} + \text{Ti}(s) = \text{NiTi}_2 \quad (R4)
\]
\[
\text{TiNi} + 2\text{Ni}(s) = \text{Ni}_3\text{Ti} \quad (R5)
\]
\[
\text{NiTi}_2 + \text{Ni}(s) = \text{NiTi}_2 \quad (R6)
\]
\[
\text{Ni}_3\text{Ti} + 2\text{Ti}(s) = \text{Ni}_3\text{Ti} \quad (R7)
\]
\[
\text{NiTi}_2 + 3\text{Ni}(s) = \text{NiTi}_2 + \text{Ni}_3\text{Ti} \quad (R8)
\]
\[
\text{Ni}_3\text{Ti} + 3\text{Ti}(s) = \text{NiTi}_2 + \text{NiTi} \quad (R9)
\]

Based on the Table of Thermodynamic Properties in [51], we calculated the change in Gibbs free energy (ΔG) involved in the various reactions at the Ti-Ni diffusion interface under various temperatures. As shown in Table 3, we then estimated the likelihood of each reaction occurring based on the ΔG related to the formation of the compound phases described in reaction (R) Eqs. 1-9.

Figure 4 shows the ΔG as a function of temperature in the nine reaction processes described above. ΔG was less than zero in all three of the reactions over a temperature range of

| Temperature | ΔG 1 | ΔG 2 | ΔG 3 | ΔG 4 | ΔG 5 | ΔG 6 | ΔG 7 | ΔG 8 | ΔG 9 |
|-------------|------|------|------|------|------|------|------|------|------|
| 550°C       | -14.140 | -17.355 | -35.975 | -3.214 | -21.835 | -10.926 | -6.446 | -32.761 | -9.661 |
| 600°C       | -14.001 | -17.125 | -35.031 | -3.183 | -21.246 | -10.837 | -5.621 | -31.123 | -9.435 |
| 650°C       | -13.804 | -16.963 | -36.172 | -3.159 | -22.368 | -10.645 | -5.240 | -33.013 | -8.398 |
| 700°C       | -13.661 | -16.794 | -36.250 | -3.133 | -22.588 | -10.529 | -4.735 | -33.117 | -7.867 |
| 750°C       | -13.517 | -16.622 | -36.324 | -3.105 | -22.807 | -10.412 | -4.227 | -33.219 | -7.332 |
| 800°C       | -13.324 | -16.465 | -36.863 | -3.012 | -23.023 | -10.772 | -3.362 | -34.351 | -6.993 |
| 850°C       | -13.061 | -16.232 | -37.154 | -2.962 | -23.452 | -10.541 | -3.320 | -34.415 | -6.521 |
| 900°C       | -12.865 | -15.821 | -37.834 | -2.953 | -23.932 | -10.332 | -3.246 | -34.861 | -6.21 |
550–900 °C (i.e., 773–1173 K). The second law of thermodynamics stipulates that the probability of a compound forming via spontaneous reaction is closely related to its $\Delta G$ [52]. Thus, $\Delta G$ also determines the sequence of compound formation in any given reaction. Figure 4 and Table 3 illustrate the fluctuations in $\Delta G$ as a function of temperature in all of the reactions that occurred at the Ti/Ni interface. Note that all of the compounds associated with Ni$_3$Ti, NiTi$_2$, and NiTi formed within a temperature range of 550–770 °C (i.e., 773–1043K) [52, 53]. From this, we can infer that the Ni$_3$Ti compound (reaction 3 (i.e., R3)) has a lower $\Delta G$ (more negative) than those of NiTi (R1) and NiTi$_2$ (R2). Based on the principle of minimum energy, Ni$_3$Ti forms before NiTi$_2$ and NiTi. Thus, from the perspective of $\Delta G$ only (i.e., disregarding the impact of all other factors), we derive the following sequence for the formation of Ti-Ni compounds: Ni$_3$Ti &gt; NiTi$_2$ &gt; NiTi.

Table 3, the $\Delta G$ in R3 decreased (i.e., became more negative) with an increase in temperature, whereas the $\Delta G$ in R1 and R2 increased. We can see in Fig. 4 that in the second diffusion stage at 800–900 °C (i.e., 1073–1173 K), R5 and R8 took precedence over all other reactions except R1, R2, and R3, resulting in a rapid increase in the amount of Ni$_3$Ti compound. Subsequent long-range atomic diffusion initiated other reactions, such as the reaction of Ni$_3$Ti with Ti to form stable compounds of NiTi and NiTi$_2$. Eventually, all of the pure Ti, Ni, and Ni$_3$Ti were consumed. Note that the Ni$_3$Ti and NiTi$_2$ compounds formed before NiTi, due to their higher $\Delta G$ (i.e., more negative) and less pronounced increase in surface energy [53]. We were unable to determine the sequences of Ni$_3$Ti and NiTi$_2$ formation, due to similar surface energy values [54].

It is noted that this inference is consistent with that of previous studies [55–60], which also reported the preferential formation of Ni$_3$Ti compound. The formation of a high-volume fraction of NiTi$_2$ compound implies that a large quantity of Ni is depleted during the diffusion-bonding process. Therefore, the capabilities of the different diffusion temperatures in dissolving the NiTi$_2$ compound are limited, if the fundamental mechanism accounting for the severe Ni loss is not uncovered. However, the serious loss of Ti content in the diffusion process implies that it is impossible to attain a full NiTi compound. In addition, some irregular-shaped voids can be seen in Fig. 3a–c arising from the higher diffusion coefficient of Ni compared to that of Ti, which results in an imbalance in the mass flow across the interface. The volume fraction of these voids reduces with an increasing bonding temperature, which suggests that a higher temperature enhances the thermal excitation of the atoms and hence facilitates an improved inter-diffusion and reaction process across the interface [47, 61–65]. It is noted that the present observations are consistent with those of previous studies for diffusion couples of Ti-Ni and microduplex stainless steel-Ti alloy processed at temperatures in the range of 700–850°C, respectively [47, 66].

It is also important to consider the atomic ratio of Ti and Ni when seeking to elucidate the formation of Ni$_3$Ti, NiTi, and NiTi$_2$ compounds at the Ti-Ni diffusion interface at various temperatures. Including the Gibbs free energy of mixing ($\Delta G_{mix}$) in the binary (transition) metal-metal system helped to explain why homogeneous NiTi formed when the atomic ratio of Ni to Ti was 1:1. $\Delta G_{mix}$ refers to changes in the energy of the system under the effects of diffusion [67]. Diffusion for a sufficient duration permits the formation of mixed phases B$_1$...A$_x$ at the interface of the two original phases A (Ti) and B (Ni), where variable $x$ indicates the atomic percentage of the corresponding element. Note that $\Delta G_{mix}$ is a function of $x$ and temperature of $T$. Mixing enthalpy can be estimated using the method outlined in [68] and is taken as configurational entropy. Thus, $\Delta G_{mix}$ can be calculated as follows [67, 69]:

$$\Delta G_{mix}(x, T) = x F_A^A(x) \Delta H_{A/B}^{\text{interface}} - T \Delta S_{\text{conf}} (1)$$

where $\Delta H_{A/B}^{\text{interface}}$ indicates the change in enthalpy resulting from the dissolution of 1M of A in B and $\Delta S_{\text{conf}}$ indicates the change in configurational entropy. $F_A^A(x)$ is the degree to which A atoms are surrounded by B atoms, which can be accurately estimated as follows [67]:

$$F_A^A(x) = C_A^A(x) = \frac{(1-x) V_A^2}{V_A^2 + (1-x) V_B^2} \quad (2)$$

where $C_A^A(x)$ indicates the volume fraction of B. $V_A$ and $V_B$ refer to the molar volumes of pure crystalline solids (i.e., $V_{Ni} = 10.6 \text{ cm}^3/\text{mol}$, $V_{Ni} = 6.5 \text{ cm}^3/\text{mol}$) [70, 71].
The $\Delta H_{\text{A in B}}$ and $\Delta S_{\text{conf}}$ in the binary crystal system are defined as follows [70, 72]:

$$\Delta H_{\text{A in B}} = 2 p f A V_A \left[ -\left( \Phi_A^* - \Phi_B^* \right)^2 - \frac{R}{P} \right]$$ (3)

$$\Delta S_{\text{conf}} = -R[\ln(x) + (1-f_x)\ln(11x)]$$ (4)

where $n_A$ and $n_B$ are the electron density parameters (i.e., $n_A^2 = 1.47, n_B^2 = 1.75$) [70] used to describe phenomena occurring at an interface at the atomic scale. Parameters $\Phi_A$ and $\Phi_B$ are used to describe ionic bonds in metals (i.e., $\Phi_A = 3.65, \Phi_B = 5.20$) [70, 71]). $Q$, $R$, and $P$ are constants with the following values: $Q/R (9.4)$, $R/P (1.9)$, and $P (12.35)$.

The above equations were used to calculate $\Delta G_{\text{mix}}$, and the results are shown in Fig. 5 in terms of variations in $\Delta G_{\text{mix}}$ as a function of the atom percentage of Ti in Ni$_{1-x}$Ti$_x$. We can see that when the atom percentage of Ti was fixed, there was a decrease in the $\Delta G_{\text{mix}}$ of Ti-Ni diffusion with an increase in temperature. When $x$ was increased from 0.2 to 0.7, the value of $\Delta G_{\text{mix}}$ first decreased and then increased. The lowest $\Delta G_{\text{mix}}$ value was obtained when the value of $x$ was roughly 0.5, which indicates the point at which NiTi formation occurred (0.48–0.51). However, the formation of Ni$_3$Ti and NiTi is easy to understand from the perspective of metallurgical characteristics or thermodynamics. We speculate that the respective $\Delta G_{\text{mix}}$ values of Ni$_3$Ti and NiTi during Ti-Ni diffusion decreased with an increase in temperature; i.e., moving toward dynamic thermal equilibrium. Ni$_3$Ti generation (near the Ni side) was followed by NiTi$_2$ (near the pure Ti side). This was followed by the formation of NiTi at the interface between the two. During the initial formation of Ni$_3$Ti, NiTi$_2$, and NiTi, diffusion was still ongoing. This is particularly true of the Ti (rapid diffusion), which resulted in the steady growth of NiTi. The unsteady growth of NiTi$_2$ was the result of Ti diffusion toward the NiTi and pure Ti sides. Note that the NiTi$_2$ region in phase images was extremely narrow. Nonetheless, the formation of NiTi$_2$ cannot be used to analyze the mutual influence between NiTi$_2$ and Ni$_3$Ti or between NiTi$_2$ and NiTi. Further analysis is required.

### 3.2 Mechanical properties

#### 3.2.1 Interfacial hot-roll diffusion-bonded joint quality

**Shear strength of hot-roll diffusion-bonded joints** The shear strength is an important index for evaluating the interfacial bonding quality of hot-roll clad composites. Figure 6 shows the interfacial shear strengths of the present hot-rolled joints processed at temperatures of 800°C, 850°C, and 900°C, respectively. For a rolling temperature of 800°C, the shear strength is equal to approximately 343 MPa. However, the shear strength increases to a maximum value of 470 MPa in the sample processed at a rolling temperature of 850°C. This value is markedly higher than the values reported in previous studies [73]. This can be explained by the increase in the thickness of the NiTi layer and the fact that the $\beta$-Ti phase exists as discrete needle-shaped particles in the matrix of the Ni$_3$Ti phase, whereas $\alpha$-Ti phase exists as discrete islands. The atomic migration of chemical species across the bond line increased with an increase in bonding temperature. The presence of Ni in the interfacial zone adjacent to the Ti substrate stabilized the $\beta$-Ti phase, which led to the formation of a plastic zone capable of bridging cracks and thereby maintaining the integrity of the interfacial microstructure. Furthermore, a lower bonding temperature suppressed the migration of atoms from the pure Ti substrate toward the carbon steel substrate (and vice versa). As a result, an incomplete coalescence of the mating surfaces occurs [15, 74]. However, at higher
rolling temperatures, the shear strength is significantly improved as a result of both a greater inter-diffusion of elements from the Ti and carbon steel substrates and the temperature-induced plastic collapse of the mating surface asperities, which results in a more intimate contact of the two surfaces and an enhanced bonding outcome as a result. Interestingly, however, the shear strength reduces to approximately 386 MPa as the bonding temperature is further increased to 900°C. The reduction in the shear strength implies that, at a high bonding temperature, the Ni interlayer fails to completely block the diffusion of carbon atoms to the Ti side. Consequently, compound phase reaction layers with a relatively large thickness are produced in the interfacial region between the Ti substrate and the Ni interlayer (see Fig. 3). As the rolling process proceeded, compound phase particles tended to disintegrate under the effects of embrittlement. The resulting formation of irregular micro-voids led to a significant reduction in shear strength, thereby demonstrating that shear strength depends heavily on the microstructure at the interface.

Fracture morphology of hot-roll diffusion-bonded joints

The fracture surfaces of the hot-roll diffusion-bonded joints were observed via SEM, as shown in Fig. 7. (Note that for each sample, fracture occurred at the Ti side due to the formation of compound phases.) The dark region in Fig. 7a confirms the presence of micro-voids due to the minimal diffusion of intermetallic elements at 800°C. These micro-voids reduce the contact area between the mating surfaces and therefore degrade the shear strength of the joint. For the sample bonded at a higher temperature of 850°C, a larger number of intermetallic phases are formed in the interfacial region. The fracture surface shows evidence of a ductile behavior (see Fig. 7b), and hence it is inferred that these intermetallic phases enhance the bonding strength of the joint without any embrittlement effect. However, for the sample bonded at 900°C (see Fig. 7c), the fracture surface contains multiple cleavage planes with different alignments. In other words, the thick compound phase reaction layers impart a brittleness to the interfacial region and cause the joint to fail accordingly. In addition, the river pattern differs from grain to grain, and the size of the river patterns increases with an increasing bonding temperature.

Figure 8 shows the X-ray diffraction patterns of the three fracture surfaces in Fig. 7. For the specimen processed at 800°C, the diffusion zone contains NiTi₂, NiTi, Ni₅Ti, α-Ti, and β-Ti phases (see Fig. 8a). However, at 850°C, the diffusion zone consists mainly of NiTi₂, α-Ti, and β-Ti (see Fig. 8b). Finally, for the sample processed at 900°C, the diffusion zone consists predominantly of NiTi₂, α-Ti, and β-Ti phases. Notably, titanium carbide (TiC) is not observed in the diffusion zones of any of the samples. Hence, it is inferred that the Ni interlayer serves as an effective barrier in preventing the diffusion of Ti atoms from the Ti plate to the carbon steel side. The SEM images in Fig. 3 show the presence of α-Ti and β-Ti phases in the Ti-Ni interface region of all three samples. However, α-Ti and β-Ti phases are not evident in the corresponding X-ray diffraction patterns since the bonded samples fracture through the NiTi₂ and β-Ti regions, and the X-ray diffraction depth is insufficiently large to perform subsurface detection. However, NiTi₂ phase is evident in the XRD patterns of the joints processed at temperatures of 850°C and 900°C, which confirms the migration of Ti elements from the pure Ti plate toward the carbon steel substrate at higher temperatures.

3.2.2 Peel strength of hot-roll diffusion-bonded joints

The interfacial bonding strength is one of the most important mechanical characteristics of clad composites. In the present study, the interfacial bonding strength was evaluated by means of ASTM-D 1876-01 standard [40]. The corresponding results are shown in Fig. 9 for both the mean peel strength of each sample (see Fig. 9a) and the variation of the peel strength.
with the peeling displacement. The samples processed at temperatures of 800°C and 850°C have peel strengths of 10 N/mm and 18 N/mm, respectively. However, the peel strength increases to 21 N/mm in the sample processed at 900°C. In other words, the average peel strength increases by approximately 110% as the bonding temperature is increased from 800 to 900°C.

Several factors affect the interfacial bonding strength of hot-roll diffusion-bonded joints, including the formation of compound phases, cracking, and the extent of elemental diffusion. Intermetallics generally have a complex crystal structure and are brittle due to their ionic and covalent bonding [75]. When Intermetallics are present in the interfacial regions of joints such as those prepared in the present study, and they serve as inclusions and prompt crack. Crack is followed by a region of stable crack propagation and later fast fracture. Furthermore, defects in the interfacial region act as stress concentrators, which increase the local stress intensity and promote crack nucleation [47]. However, as the rolling temperature increases, a greater number of atoms gain the activation energy required to undergo diffusion. Consequently, the peel strength increases, as shown in Fig. 9a. However, a higher temperature also increases the rate of atomic diffusion of Ni atoms toward the pure Ti plate. This in turn increases the size and number of the compound phases formed in the interfacial region, which increases the local stress and prompts crack nucleation and growth. As a result, the ductility reduces, as shown in Fig. 9b.

Overall, the present results show that the interfacial bonding ability is related to the formation and distribution of compound phases between the Ti and Ni layers in the hot-roll diffusion-bonded joints. Moreover, for the present samples, a hot-rolling temperature of 850°C results in the optimal tradeoff between the interfacial bonding strength and the ductility.

4 Conclusions

We applied a hot-roll solid-state diffusion-bonding technique to join commercially pure Ti plates to carbon steel substrates with a Ni interlayer (i.e., the tri-metal composite) at processing
temperatures of 800, 850, and 900°C, respectively. The experimental results support the following conclusions:

1. Inter-diffusion between the Ni interlayer and pure Ti substrate resulted in the formation of layers of NiTi2, NiTi, and Ni3Ti phase at the Ti-Ni interface. Note that the formation sequence of the three intermetallic compounds was as follows: (1) Ni3Ti formed first; (2) NiTi2 formed when the Ni3Ti layer reached a certain thickness; and (3) Ti and Ni diffused into the NiTi2 and Ni3Ti layers to form NiTi. As the thickness of the NiTi layer increased, a portion of the Ni3Ti layer was consumed. This can be attributed to the fact that the nucleation of Ni3Ti and NiTi2 occurred prior to that of NiTi, due to their lower Gibbs free energy. β-Ti phases additionally exist as discrete needle-shaped particles in the matrix of the Ni3Ti phase, while α-Ti phase exists as discrete islands. The atomic migration of chemical species across the bond line increases with an increasing bonding temperature. The presence of Ni in the interfacial zone adjacent to the Ti substrate stabilizes the β-Ti phase, leads to the formation of α-Ti phase, and acts as a diffusion barrier which prevents the migration of Ti atoms to the carbon steel substrate. The Ni-carbon steel interface is thus free from any reaction products for the considered bonding temperatures of 800–900°C.

2. The maximum shear strength (~470 MPa) is obtained in the sample processed at 850°C due to a better coalescence of the mating surfaces and an improved diffusion of the atomic species. For a lower bonding temperature of 800°C, the shear strength is reduced due to a more limited plastic deformation of the mating surface asperities. By contrast, for a higher bonding temperature of 900°C, the shear strength is degraded due to the formation of brittle TiNi3-type micro-intermetallic layers.

3. The peel strength increases with an increasing bonding temperature and has a value of 21 N/mm in the sample processed at 900°C. The enhanced peel strength at higher temperatures can be attributed to an increase in the intermetallic compounds and a greater composition homogeneity in the diffusion zone between the pure Ti and carbon steel plates and Ni interlayer. Overall, the results suggest that the failure behavior of the joints is directly related to the formation and distribution of compound phases at the bond interface.

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Author contribution All authors contributed to the research, writing, and reviewing of the paper. Professor Chun-Ming Lin analyzed the experimental results and reviewed and edited the manuscript. Mr. Mohsen Saboktakin Rizi and Mr. Chia-Kai Chen conceived the experimental design, conducted the experiments, and prepared the draft manuscript.

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Data availability The authors confirm that the data supporting the findings of this study are available within the article. The raw data are available from the corresponding author upon a request.

Declarations

Ethical approval The work contains no libelous or unlawful statements, does not infringe on the rights of others or contains material or instructions that might cause harm or injury.

Consent to participate and publication The authors consent to participate. In addition, consent for publication informed consent was obtained from all individual participants involved in the study.

Conflict of interest The authors declare that they have no conflict of interest.

References

1.  Weng F, Chen C, Yu H (2014) Research status of laser cladding on titanium and its alloys: a review. Mater Des 58:412–425
2.  Karolczuk A, Kowalski M, Barinski R, Zok F (2013) Fatigue phenomena in explosively welded steel-titanium clad components subjected to push-pull loading. Int J Fatigue 48:101–108
3.  Lee WB, Lee CY, Chang WS, Yeon YM, Jung SB (2005) Microstructural investigation of friction stir welded pure titanium. Mater Lett 59(26):3315–3318
4.  Kahraman N, Gulenc B, Findik F (2007) Corrosion and mechanical-microstructural aspects of dissimilar joints of Ti6Al4V and Al plates. Int J Impact Eng 34:1423–1432
5.  Sun Z, Annergren I, Pan D, Mai TA (2003) Effect of laser surface remelting on the corrosion behavior of commercially pure titanium sheet. Mater Sci Eng A 345(1-2):293–300
6.  Auwal ST, Ramesh S, Yusof F, Manladan SM (2018) A review on laser beam welding of titanium alloys. Int J Adv Manuf Technol 97(4):1071–1098
7.  Gao M, Chen C, Wang L, Wang Z, Zeng X (2015) Laser-arc hybrid welding of dissimilar titanium alloy and stainless steel using copper wire. Mater Trans A 46(5):2007–2020
8.  Wang T, Zhang B, Feng J, Tang Q (2012) Effect of a copper filler metal on the microstructure and mechanical properties of electron beam welded titanium-stainless steel joint. Mater Charact 73:104–113
9.  Soltani Tashi R, Akbari Mousavi SAA, Mazar Atabaki M (2014) Diffusion brazing of Ti-6Al-4V and austenitic stainless steel using silver-based interlayer. Mater Des 54:161–167
10. Lee MK, Park JJ, Lee JG, Rhee CK (2013) Phase-dependent corrosion of titanium-to-stainless steel joints brazed by Ag-Cu eutectic alloy filler and Ag interlayer. J Nucl Mater 439(1-2):168–173
11. Elrefaey A, Tillmann W (2009) Correlation between microstructure, mechanical properties, and brazing temperature of steel to titanium joint. J Alloys Compd 487(1-2):639–645
12. Norouzi E, Atapour M, Shaminan M, Allafchian A (2016) Effect of bonding temperature on the microstructure and mechanical properties of Ti-6Al-4V to AISI 304 transient liquid phase bonded joint. Mater Des 99:543–551

13. Zakipour S, Halvaea A, Amadeh AA, Samavatian M, Khodabandeh A (2015) An investigation on microstructure evolution and mechanical properties during transient liquid phase bonding of stainless steel 316L to Ti-6Al-4V. J Alloys Compd 626:269–276

14. Srikanth V, Laik A, Dey GK (2017) Joining of stainless steel 304L with Zircaloy-4 by diffusion bonding technique using Ni and Ti interlayers. Mater Des 126:141–154

15. Kundu S, Sam S, Chatterjee S (2011) Interface microstructure and strength properties of Ti-6Al-4V and microduplex stainless steel diffusion bonded joints. Mater Des 32:2997–3003

16. Ghosh M, Chatterjee S (2003) Diffusion bonded transition joints of titanium to stainless steel with improved properties. Mater Sci Eng A 358(1-2):152–158

17. Yang DH, Luo Za, Xie GM, Wang MK, Misra RDK (2018) Effect of vacuum level on microstructure and mechanical properties of titanium-steel vacuum roll clad plates. J Iron Steel Res Int 25(1):72–80

18. Akramifard HR, Mirzadeh H, Parsa MH (2014) Cladding of aluminum on AISI 304L stainless steel by cold roll bonding: mechanism, microstructure, and mechanical properties. Mater Sci Eng A 613:232–239

19. Zhao DS, Yan J, Liu YJ (2013) Heat resistance of hot roll bonded titanium alloy-stainless steel transition joint. Trans Nonferrous Metals Soc China 23(7):1966–1970

20. Kundu S, Chatterjee S (2006) Interfacial microstructure and mechanical properties of diffusion-bonded titanium–stainless steel joints using a nickel interlayer. Mater Sci Eng A 425(1-2):107–113

21. Arik H, Aydin M, Kurt A, Turker M (2005) Weldability of Al4C3-Al composites via diffusion welding technique. Mater Des 26:555–560

22. Ghosh M, Bhanumurthy K, Kale GB, Krishnan J, Chatterjee S (2003) Diffusion bonding of titanium to 304 stainless steel. J Nucl Mater 322(2-3):235–241

23. Yazdani Z, Toroghinejad MR, Edris H, Negan AHW (2018) A novel method for the fabrication of Al matrix nanocomposites reinforced by mono-dispersed TiAl3 intermetallic via a three-step process of cold-roll bonding, heat-treatment and accumulative roll bonding. J Alloys Compd 747:217–226

24. Mahdavian MM, Khatami-Hamedani H, Abedi HR (2017) Microstructure evolution and mechanical properties of accumulative roll bonded Al/Cu/Sn multilayer composite. J Alloys Compd 703:605–613

25. Dong HG, Yang ZL, Yang GS, Dong C (2013) Vacuum brazing of TiAl alloy to 40Cr steel with Ti60Ni22Cu10Zr8 alloy foil as filler metal. Mater Sci Eng A 561:252–258

26. Song TF, Jiang XS, Shao ZY, Fang YJ, Zhu MH (2017) Microstructure and mechanical properties of vacuum diffusion bonded joints between Ti-6Al-4V titanium alloy and AISI316L stainless steel using Cu/Nb multi-interlayer. Vacuum 145:68–76

27. Hosseini M, Danesh Manesh H (2015) Bond strength optimization of Ti/Cu/Ti clad composites produced by roll-bonding. Mater Des 81:122–132

28. Balasubramanian M (2016) Characterization of diffusion-bonded titanium alloy and 304 stainless steel with Ag as an interlayer. Int J Adv Manuf Technol 82:153–162

29. Zhao DS, Yan J, Wang CW, Wang Y, Yang SQ (2008) Interfacial structure and mechanical properties of hot roll bonded joints between titanium alloy and stainless steel using copper interlayer. Sci Technol Weld Join 13:765–768

30. Yan JC, Zhao DS, Wang CW, Yang SQ (2009) Vacuum hot roll bonding of titanium alloy and stainless steel using nickel interlayer. J Mater Sci Technol 25:914–918

31. Zakipour S, Halvaea A, Amadeh AA, Samavatian M, Khodabandeh A (2015) An investigation on microstructure evolution and mechanical properties during transient liquid phase bonding of stainless steel 316L to Ti-6Al-4V. J Alloys Compd 626:269–276

32. Balasubramanian M (2015) Application of Box-Behnken design for fabrication of titanium alloy and 304 stainless steel joints with silver interlayer by diffusion bonding. Mater Des 77:161–169

33. Massalski TB (1990) Binary alloy phase diagrams, 2nd edn. ASM Internalional, Materials Park, OH

34. Verdian MM, Raeissi K, Salehi M, Sabooni S (2011) Characterization and corrosion behavior of TiNi-Ti3Ni-TiN3 multiphase intermetallics produced by vacuum sintering. Vacuum 86:91–95

35. Hu LF, Xue YZ, Shi FG (2017) Intermetallic formation and mechanical properties of Ni-Ti diffusion couples. Mater Des 130:175–182

36. Feng B, Xin Y, Hong R, Yu H, Wu Y, Liu Q (2015) The effect of architecture on the mechanical properties of Mg-3Al-1Zn rods containing hard Al alloy cores. Scr Mater 98:56–59

37. Wang XB, Humbeeck Jv, Verlinden B, Kustov S (2016) Thermal cycling induced room temperature aging effect in Ni-rich TiNi shape memory alloy. Scr Mater 113:206–208

38. Fan QC, Zhang YH, Wang YY, Sun MY, Wen YH (2017) Influences of transformation behavior and precipitates on the deformation behavior of Ni-rich TiNi alloys. Mater Sci Eng A 700:269–280

39. ASTM-D 1002-10 standards, ASTM, PA, 2010

40. ASTM-D 1876-01 standards, ASTM, PA, 2015

41. Verdier M, Groma I, Flandin L, Lendavi J, Brechet Y, Guyot P (1997) Dislocation densities and stored energy after cold rolling of Al-Mg alloys: investigations by resistivity and differential scanning calorimetry. Scr Mater 37:449–454

42. A.A. Shirzadi (1998) Ghoshouni Diffusion bonding aluminium al-loys and composites: new approaches and modeling (doctoral thesis) University of Cambridge

43. Oliveira JP, Cavaleiro AJ, Schell N, Stark A, Braz Fernandes FM (2018) Effects of laser processing on the transformation characteristics of TiNi: a contribute to additive manufacturing. Scr Mater 152:122–126

44. Marattukalam JJ, Balla VK, Das M, Bontha S, Kalpathy SK (2018) Effect of heat treatment on microstructure, corrosion, and shape memory characteristics of laser deposited TiNi alloy. J Alloys Compd 744:337–346

45. Beres J, Polak A, Indacocha J (2006) Joining YSZ to 444 SS by in situ alloying in a N-Ti filler metal for SOFC applications. In: Stephens JJ, Weil KS (eds) Proceeding of the 3rd international brazing and soldering conference. American Welding Society, San Antonio, pp 125–132

46. Massalski TB, Okamoto H, Subramanian PR, KaprzkBinary L (1990) Alloy phase diagrams, 2nd edn. ASM Intl, OH

47. Deng YQ, Sheng GM, Xu C (2013) Evaluation of the microstruc-ture and mechanical properties of TiAl to 444 SS by in situ alloying in a N-Ti filler metal for SOFC applications. In: Stephens JJ, Weil KS (eds) Proceeding of the 3rd international brazing and soldering conference. American Welding Society, San Antonio, pp 125–132

48. Batalovč K, Koteski V, Stojić D (2013) Hydrogen storage in mar-tensite Ti-Zr-Ni alloy: a density functional theory study. J Phys Chem C 117(51):26914–26920

49. Toprek D, Belosevic-cavor J, Koteski V (2015) Ab TiNio studies of the structural, elastic, electronic and thermal properties of TiNi2 intermetallic. J Phys Chem Solids 85:197–205

50. Chen Q, Huang Z, Zhao Z, Hu C (2013) First-principles study on the structural, elastic, and thermodynamics properties of Ni3X (X:
Al, Mo, Ti, Pt, Si, Nb, V, and Zr) intermetallic compounds. Appl Phys A Mater Sci Process 116(3):1–12
51. Li B, Rong L, Li Y, Guinter VE (2000) Fabrication of cellular TiNi intermetallic compounds. J Mater Res 15(1):10–13
52. Battezzati L, Pappalepore P, Durbiano F, Gallino I (1999) Solid state reactions in Al/Ni alternate foils induced by cold rolling and annealing. Acta Mater 47(6):1901–1914
53. Shao X, Guo XL, Han YF, Lin ZJ, Qin J, Lu WJ, Zhang D (2017) Preparation of TiNi films by diffusion technology and the study of the formation sequence of the intermetallics in Ti-Ni system. J Mater Res 29(22):2707–2716
54. Zhang Y, Cheng X, Cai H (2016) Fabrication, characterization and tensile property of a novel Ti2Ni/TiNi micro-laminated composite. Mater Des 92:486–493
55. Brandes EA (ed) (1983) Smithells metals reference book. Butterworth, London
56. Otsuka K, Ren X (2005) Physical metallurgy of Ti-Ni-based shape memory alloys. Prog Mater Sci 50:511–678
57. Ye DL, Hu JH (2002) Utility inorganic materials thermodynamics data handbook, 2nd edn. Metallurgy Industry Press, Beijing
58. Villars P, Prince A, Okamoto H (1995) Hand book of ternary phase alloys, vol 7. ASM Int., Mater. Park, Ohio, pp 8903–8928
59. Massalski TB (ed) (1986) Binary phase diagrams. American Society for Metals, Metals Park, OH
60. Raghavan V, Phase J (1970) Fe-Ni-Ti (Iron-Nickel-Titanium). Equilib Diff 31:186–189
61. Orhan N, Khan TI, Eroglu M (2001) Diffusion bonding of a microduplex stainless steel to Ti-6Al-4V. Ser Metall 45:441–446
62. Ma Y, Li H, Yang L, Hu A (2018) Reaction-assisted diffusion bonding of Ti6Al4V alloys with Ti/Ni nanostructured multilayers. J Mater Sci Technol 262:204–209
63. Muralimohan CH, Ashfaq M, Ashiri R, Muthupandi V, Sivaprasad K (2016) Analysis and characterization of the role of Ni interlayer in the friction welding of titanium and 304 austenitic stainless steel. Mater Trans A 47:347–359
64. Yildiz A, Kaya Y, Kahraman N (2016) Joint properties and microstructure of diffusion-bonded grade 2 titanium to AISI 430 ferritic stainless steel using pure Ni interlayer. Int J Adv Manuf Technol 86:1287–1298
65. Liu J, Cao J, Song X, Wang Y, Feng J (2014) Evaluation on diffusion bonded joints of TiAl alloy to Ti3SiC2 ceramic with and without Ni interlayer: interfacial microstructure and mechanical properties. Mater Des 57:592–597
66. Hinotani S, Ohmori Y (1988) The microstructure of diffusion-bonded Ti/Ni interface. Trans Jpn Inst Mater 29:116–124
67. Benedictus R, Böttger A, Mittemeijer E (1996) Thermodynamic model for solid-state amorphization in binary systems at interfaces and grain boundaries. Phys Rev B 54(13):9109–9125
68. Li Y, Cui L, Shi P, Yang D (2001) Phase transformation behaviors of prestrained TiNi shape memory alloy fibers under the constraint of a hard substrate. Mater Lett 49(3):224–227
69. De Boer F, Boom R, Mattens W, Miedema A, Niessen A (1989) Cohesion in metals. Transition Metal Alloys, Cohesion and Structure (North-Holland, Amsterdam Chap. 2
70. Miedema A, De Chatel P, De Boer F (1980) Cohesion in alloys-fundamentals of a semi-empirical model. Physica B 1 C 100(1):1
71. Miedema A (1976) On the heat of formation of solid alloys. II. J Less-Common Met 46(1):67
72. Liu J, Su Y, Xu Y, Luo L, Guo J, Fu H (2011) First phase selection in solid Ti/Al diffusion couple. Rare Metal Mater Eng 40(5):753
73. Kundu S, Chatterjee S (2007) Structure and properties of diffusion bonded transition joints between commercially pure titanium and type 304 stainless steel using a nickel interlayer. J Mater Sci 42:7906–7912
74. Kundu S, Chatterjee S, Olson D, Mishra B (2007) Effects of intermetallic phases on the bond strength of diffusion-bonded joints between titanium and 304 stainless steel using nickel interlayer. Mater Trans A 38:2053–2060
75. Chawla N, Chawla KK (2006) Metal Matrix Composites. Springer Science and Business Media Inc., Berlin

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