Abstract: Stainless steel-based WC composite layers fabricated by a laser cladding technique, have strong mechanical strength. However, the wear resistance of WC composite layers is not sufficient for use in severe friction and wear environments, and the corrosion resistance is significantly reduced by the formation of secondary carbides. Low-temperature plasma nitriding and carburizing of austenitic stainless steels, treated at temperatures of less than 450 \(^\circ\)C, can produce a supersaturated solid solution of nitrogen or carbon, known as the S-phase. The combined treatment of nitriding and carburizing can form a nitrocarburizing S-phase, which is characterized by a thick layer and superior cross-sectional hardness distribution. During the laser cladding process, free carbon was produced by the decomposition of WC particles. To achieve excellent wear and corrosion resistance, we attempted to use this free carbon to form a nitrocarburizing S-phase on AISI 316 L stainless steel-based WC composite layers by plasma nitriding alone. As a result, the thick nitrocarburizing S-phase was formed. The Vickers hardness of the S-phase ranged from 1200 to 1400 HV, and the hardness depth distribution became smoother. The corrosion resistance was also improved through increasing the pitting resistance equivalent numbers due to the nitrogen that dissolved in the AISI 316 L steel matrix.

Keywords: plasma nitriding; laser cladding; stainless steel; tungsten carbide; hardness; corrosion resistance

1. Introduction

Laser cladding is an effective method for surface modification, such as improving the wear and corrosion resistance of metal substrates, and has many advantages, such as high deposition speed with rapid cooling, dense metallography, strong bonding strength between the cladding layer and substrate, low heat input, and distortion of the substrate [1]. In recent years, the fabrication of three-dimensional objects by additive manufacturing (AM) using laser as a heat source has attracted much attention. There are many metal AM technologies, for example, laser-based powder bed fusion (PBF) methods have two types of selective laser melting (SLM) and selective laser sintering (SLS), in which a laser is irradiated on top of the stacked powders [2–4]. Meanwhile, directed energy deposition (DED) is based on laser cladding with a coaxial nozzle system for feeding powder in the axial direction of the laser beam and has been developed not only as a coating for surface modification, but also as a manufacturing process for various engineering materials and objects. DED achieves a near-perfect net-like shape appearance with faster processing speed and without mold tooling, which can be applied for manufacturing, part repair, and rapid prototyping. DED also enables the reduction in manufacturing costs and has been increasingly used in recent years [5–9].

Among the classes of austenitic stainless steels for laser cladding, AISI 316 L stainless steel is heavily used owing to its excellent corrosion property, high versatility, and the easy of obtaining steel powder [10]. However, deposited AISI 316 L stainless steel layers have
a low hardness and poor wear resistance properties, and therefore are difficult to use in severe wear conditions. Several researchers have reported methods to improve the wear resistance and strength of deposited steel layers, and it is concluded that the addition of carbide particles as a reinforcement is highly effective. When the hard carbide particles were dispersed in the steel matrix, the steel structure was modified to one akin to sintered metal matrix composite materials. Consequently, the wear resistance is significantly improved [11–13].

It is required that reinforcement carbides have high hardness, chemical affinity to steel, and high melting temperature [14,15]. Tungsten carbide (WC) is a hard ceramic that can be uniformly dispersed in a steel matrix with good wettability; therefore, WC particles have been considered as excellent candidates for reinforcement. Steel-WC composite layers are currently the most widely applied for hardness and strength characteristics [16,17]. As for AISI 316 L stainless steel layers, it has been reported that the properties of the cladding layers were improved by compositing with SiC, TiC and titanium carbonitride [18–20]. As a matter of course, the WC particles composite for AISI 316 L stainless steel layers obtained by the laser cladding process also leads to improved mechanical properties, in particular, surface hardness and wear resistance [21–23]. During the heating of the laser cladding process, the WC particles decompose and solidify in the AISI 316 L steel matrix, and consequently, the steel matrix structure modifies to form carbide networks as a dendritic structure [21,22,24]. In addition, WC, W₂C phases, M₂₃C₆, and (Fe,W)₃C complex carbide phases form, and tungsten and carbon elements solidify [25].

In this way, the wear resistance of WC composite layers can be improved by the hardness of WC particles themselves and also by the formation of carbides in the stainless-steel matrix. However, the hardness of stainless-steel matrix is not sufficient compared with that of hard materials such as ceramics. In addition, when a higher amount of WC particles is added, cracks will appear in the composite layer [26]. Therefore, there is a limit to the amount of WC particles, and the effect of improving wear resistance is also limited. Recently, the fabrication of coatings and objects by laser cladding and DED has attracted considerable attention, especially because of its high functionality for use in severe environments. Accordingly, in addition to the composite of WC particles, further enhancement of wear resistance by other methods should be investigated.

Low-temperature plasma nitriding and carburizing at temperatures less than 450 °C for austenitic stainless steels can improve their wear resistance. Rather than producing a nitride or carbide, this treatment produces supersaturated solid solution of nitrogen or carbon in the face-centered cubic lattice, which is known as the S-phase (or the expanded austenite) [27–33]. Accordingly, the corrosion resistance does not degrade because the ability to form a passive film is maintained after plasma treatment [34,35]. In addition, a combined treatment of nitriding and carburizing has advantages of thickening and improving the toughness of the S-phase compared with the individually nitriding and carburizing [36–39].

These plasma treatments have been applied not only to steel plates but also to thermal spray coatings and laser cladding of AISI 316 L stainless steel, and their effectiveness has been recognized by the authors [27–30]. In addition, the effects of alloy elements in stainless steel plates on the formation and properties of the S-phase have also been examined, and several papers have reported the role of chromium, nickel and molybdenum [32,40–42]. The increase or decrease in these alloy elements causes a variation in the thickness of the S-phase layer and influences the wear and corrosion properties. Furthermore, the nickel element of stainless-steel layers fabricated by laser cladding was reported by the authors [30]. However, the effects of tungsten and tungsten carbides on the S-phase have not yet been reported for both steel plates and cladding layers.

The WC composite layers contain free carbon produced by the decomposition of the WC particles during laser cladding process. In general, a nitrocarburizing S-phase requires two processes of nitriding and carburizing continuously or simultaneously. Through the use of this free carbon, there is a possibility that a carburizing S-phase is formed
by nitriding process alone. In this study, AISI 316 L stainless steel-based WC particle composite layers were fabricated using laser cladding, subsequently performed by low-temperature plasma nitriding. Consequently, a thick nitrocarburizing S-phase had formed. Therefore, the metallurgical changes and carbide formation of the WC composite layers during laser cladding process were examined in detail. The formation mechanism of the nitrocarburizing S-phase by low-temperature plasma nitriding were investigated, and the properties of the hardness and corrosion resistance with the WC content were discussed.

2. Experiments

AISI 316L stainless steel-based WC composite layers were obtained using direct laser metal deposition. The mixed powder of AISI 316 L stainless-steel powder with 20 wt.% or 40 wt.% of tungsten carbide (WC) powder was used as a material for the laser cladding. The powder diameter was \(-212/\pm 63 \mu m\) of AISI 316 L stainless-steel powder, and \(-180/\pm 53 \mu m\) of WC powder.

The laser cladding system consisted of a continuous-wave diode laser beam machine of LDM-2000-60 (Laserline, GmbH, Mulheim-Karlich, Germany) with an attached a coaxial powder feed head of COAX12 (Fraunhofer, Munchen, Germany). The condition of the laser beam was a wavelength of 940 nm and an output power of 1.4 kW. The laser head was scanned one-layer deposition on AISI 304 stainless steel plates (size of \(65 \text{ mm} \times 65 \text{ mm} \times 10 \text{ mm}\)) with a traverse speed of 4 mm s\(^{-1}\) and length of 50 mm at 3 mm interval. The mixed powder was delivered in the coaxial direction of the laser beam through the COAX12 laser head with mass flow rates of 17–19 g min\(^{-1}\), and using argon as the carrier gas.

To remove the surface oxidation film and flatten the surface profile such as a mirror, the deposited AISI 316 L stainless steel-based WC composite layers were polished, and the final polishing was treated using a 1 \(\mu m\) diamond paste. After the polishing, the thickness of the deposited layers was over 1 mm.

DC plasma ion treatment machine of FECH-1N (Fuji Electronics Industry, Osaka, Japan) was used in low-temperature plasma nitriding. The AISI 316L stainless steel-based WC composite layers on the AISI 304 stainless steel plate were connected to a cathode electrode in the glass chamber, and the copper plate was connected to an anode electrode. DC current was applied between the anode and cathode electrode at a voltage of 300 V. The plasma gas with a composition ratio of \(\text{N}_2:\text{H}_2 = 80:20\) flowed at a rate of 1 L min\(^{-1}\) in the chamber to keep the pressure at 667 Pa. The nitriding temperatures were at 400 \(^\circ\text{C}\), 425 \(^\circ\text{C}\) \(\text{and}\) 450 \(^\circ\text{C}\), and the processing time was 4 h after reaching the nitriding temperatures.

The crystal structure of the WC composite layers was examined using an X-ray diffraction measurement of SmartLab (Rigaku, Tokyo, Japan) in the conventional \(\theta-2\theta\) scan of a Cu-K\(\alpha\) radiation with 40 kV and 150 mA.

Cross-sectional micrographs of the WC composite layers were obtained using an optical microscope of ECLIPSE MA100N (Nikon, Tokyo, Japan). The thickness of the nitride layers was measured from the cross-sectional micrographs at approximately 60 points for each sample.

The compositions of the WC composite layers were investigated by SEM with an energy-dispersive X-ray (EDX) analysis, SEM of ERA-8900FE (Elionix, Tokyo, Japan), and EDX of ApolloX (AMETEK, Berwyn, IL, USA), and the operating SEM voltage was 20 kV. In addition, the nitrogen and carbon depth distribution profiles were obtained using a glow-discharge optical emission spectroscopy (GDOES) of GDA750 system (Rigaku, Tokyo, Japan).

The hardness depth profiles in the cross-section of the nitrided layers were measured using a nanoindentation tester of ENT-1100a (Elionix, Tokyo, Japan) with a test load of 1.5 mN. The measurements for each sample were repeated thrice. The surface hardness of the WC composite layers was measured using a Vickers tester of HM-220D (Mitutoyo, Tokyo, Japan) with a test load of 0.245 N. The Vickers tests were repeated for each sample at least eight times, and the maximum and minimum values were rejected.
The corrosion resistance of the WC composite layers was estimated from anodic polarization curves. The anodic polarization curves were obtained using a potentiostat of HSV-110 (Hokuto Denko Corporation, Tokyo, Japan). The reference and counter electrode were an Ag/AgCl and a platinum plate, respectively. The measurement voltage was applied at a range of $-0.6 \text{ V} \text{ to } 1 \text{ V}$ with a scan rate of $1 \text{ mV s}^{-1}$. The experiments were carried out in a 3.5 wt.% NaCl solution at a temperature of 30 °C. Before the measurements, the dissolved oxygen in the NaCl solution was degassed by bubbling with N$_2$ gas for at least 0.5 h. After the measurements, the corrosion morphologies of the surfaces were obtained using a digital microscope of Dino-Lite (AnMo Electronics Corporation, Taipei, Taiwan).

3. Results and Discussion

3.1. Carbide Formation of AISI 316 L Stainless Steel with WC Particles during Laser Cladding

Figure 1 shows an overall view of the cross-section of 20 wt.% and 40 wt.% of WC particle composite layers. Note that the WC compositions in the text are the content of the feeding powder material used for the laser cladding, and the compositions of the deposited layers should be varied during the laser cladding process. The WC particles were dispersed in the AISI 316 L steel matrix, similarly to the metal matrix composite materials. However, cracks perpendicular to the substrate were observed in some areas of the WC 40 wt.% layers.

![Figure 1. Cross-sectional micrographs of the as-deposited AISI 316 L steel-based WC composite layers with various WC compositions: (a) WC 20 wt.%, and (b) WC 40 wt.%.](image)

The XRD patterns in Figure 2 present that the as-deposited WC composite layers contained the austenite phase of AISI 316 L stainless steel, the carbides of WC, W$_2$C and M$_6$C, and M$_{12}$C in only the 40 wt.% layer. It has been reported that WC, W$_2$C phases, M$_{23}$C$_6$, and (Fe,W)$_3$C complex phases are formed simultaneously in Fe/WC metal matrix composite coatings produced using the Yb: YAG laser [25].

Furthermore, W$_2$C carbide is produced by the decomposition of WC, the metastable $W_2C$ decomposes to M$_6$C of Fe$_5$W$_2$C [43,44], and the Eta-carbides of M$_6$C and M$_{12}$C are formed in the quaternary system Fe-W-C-Cr [45]. In this examination, the WC particles in the deposited layers were decomposed to W$_2$C; subsequently, W$_2$C decomposed and reacted with the AISI 316 L steel. Finally, the secondary carbides of M$_6$C and M$_{12}$C were synthesized. In Figure 3 of the cross-sectional microstructures after the etching treatment with Marble’s reagent (5 mL HCl, 5 mL H$_2$O$_2$, and 1 g CuSO$_4$), the gray contrasting areas with a net-like appearance in the AISI 316 L steel matrix indicate the precipitation of M$_6$C and M$_{12}$C as secondary carbides.

The magnified image by SEM in Figure 4 also showed the presence of eutectic carbides between the austenite in the WC 40 wt.% layer. In addition, elemental analysis of SEM-EDX showed that tungsten was dissolved in the austenite, and the chemical compositions of W ranged from 10 to 15 wt.%.
Figure 2. X-ray diffraction patterns of the AISI 316 L steel-based WC composite layers with various WC compositions and nitriding temperatures: (a) WC 20 wt.%, and (b) WC 40 wt.%.

Figure 3. Cross-sectional micrographs of the AISI 316 L steel-based WC composite layers with various WC compositions and nitriding temperatures: (a) WC 20 wt.% at 400 °C, (b) WC 20 wt.% at 425 °C, (c) WC 20 wt.% at 450 °C, (d) WC 40 wt.% at 400 °C, (e) WC 40 wt.% at 425 °C, and (f) WC 40 wt.% at 450 °C.

Figure 4. Cross-sectional SEM image of the AISI 316 L steel-based WC 40 wt.% composite layer.
3.2. Formation Mechanism of Nitrocarburizing S-Phase by Plasma Nitriding

The XRD of the nitrided WC composite layers in Figure 2 show that the (111), (200) and (220) planes of the γ-phase shifted towards the lower angles than those of the as-deposited layers. These shifts of the γ-phase peaks are caused by expanding the fcc lattice constants due to the dissolution of nitrogen. In addition, the cross-sectional microstructures with the etching treatment in Figure 3 indicate that the bright contrast layers are present on the surface. These XRD peak shifts and presence of the bright contrast layers suggest the formation of the S-phase.

Figure 5 shows the thickness of the S-phase measured from the bright contrast layers in the cross-sectional microstructures (Figure 3). The average thickness of the S-phase increased with increasing nitriding temperature. WC 20 wt.% layers were 6.7 µm at the nitriding temperature of 400 °C, 13.5 µm at 425 °C, and 20.6 µm at 450 °C. WC 40 wt.% layers were 5.6 µm at 400 °C, 9.8 µm at 425 °C, and 15.7 µm at 450 °C.

![Figure 5](image-url) Thickness of the S-phase layers with various WC compositions and nitriding temperatures.

The nitrogen depth distributions of the S-phase obtained by the GDOES measurements in Figure 6, show that nitrogen’s distribution is plateau and step-like, rather than a smooth decrease which would obey Fick’s law. Plateau and step-like distributions have been reported due to the dissolved chromium atoms binding to nitrogen, and it is considered that these distributions are unique to the S-phase [46–48]. Meanwhile, the carbon distributions show that the carbon peaks are present at a greater depth than where the nitrogen diffusion had completed. These distributions of carbon are very similar to that obtained by continuously or simultaneously nitriding and carburizing treatments [36–38]. Furthermore, the thickness of the S-phase, measured from the bright contrast layers in the cross-sectional microstructures (Figure 5), was thicker than the diffusion depth of nitrogen in GDOES. The thickness was almost equal to the sum of the diffusion depth of the nitrogen and the depth of the peak position of carbon. These results suggest that a carburizing S-phase presents where the carbon peaks exist.

The source of carbon in the carburizing S-phase seems to be free carbon produced by the decomposition of WC to W2C during the laser cladding process, since the possibility of decomposition of carbides by nitriding is low from the point of view of thermodynamics. Furthermore, some WC particles may have dissolved due to the local temperature increase caused by the laser scanning [49]. It is also possible that free carbon derived from the formation of the secondary carbides, M6C and M12C. It has been reported that the diffusion rate of carbon in austenitic stainless steels is fast compared with that of nitrogen [36]; therefore, the free carbon could penetrate ahead of nitrogen due to the nitrogen pushing the free carbon inward during the plasma nitriding, finally the carburizing S-phase would be formed. Consequently, the S-phase became a dual layer of nitrocarburizing; the nitriding S-phase sited at the outside, and the carburizing S-phase at the inside.
Figure 6. Nitrogen and carbon depth distributions of the AISI 316 L steel-based WC composite layers with various WC compositions and nitriding temperatures: (a) WC 20 wt.%, and (b) WC 40 wt%.

3.3. Effects of WC Content on Thickness of Nitrocarburizing S-Phase

The thickness of the S-phase of the WC composite layers was greater than that of 8.2 μm of the single AISI 316 L steel layer without WC at 450 °C [30]. The formation of the carburizing layer as well as the nitriding layer has resulted in the thickening of the S-phase. The GDS result for WC 20 wt.% at 450 °C in Figure 6 presents the nitrogen plateau region reaching up to 12 μm depth. Even excluding the carburizing S-phase, the nitriding S-phase was thicker than that of the single AISI 316 L steel layer. As shown in the previous section, the compositions of the solid solution of tungsten in the austenite ranged from 10 to 15 wt.%. Therefore, the solid solution of tungsten may promote nitrogen diffusion.

Meanwhile, comparing the nitrogen diffusion depth between the 20 wt.% and 40 wt.% WC layers in GDOES (Figure 6), the WC 40 wt.% layers diffused at a much shallower rate. The WC 40 wt.% layers had dense secondary carbide precipitations of a net-like appearance (Figure 3), resulting in the decrease in the crystal grain size. These dense carbide precipitations and small grain size would prevent the nitrogen diffusion. In contrast, the thickness of the carburizing S-phase (width of the carbon peaks in Figure 6) was almost the same in both layers. The WC 40 wt.% layers had a higher volume of the free carbon. Therefore, the shallow diffusion of nitrogen would not change the thickness of the carburizing S-phase layers.

3.4. Hardness Depth Profiles and Vickers Hardness of Nitrocarburizing S-Phase

Figure 7 shows the nanoindentation hardness-depth profiles of the cross-section at a nitriding temperature of 450 °C. The hardness gradually decreased from the surface to the inside, and hardening was observed up to a depth of approximately 20 μm for the 20 wt.% layer and 15 μm for the WC 40 wt.% layer. These hardening depths are almost identical to the thickness of the S-phase (Figure 5). In general, hardness depth profiles formed by nitriding alone have been reported to abruptly decrease at the boundary between the S-phase layer and the base metal. In contrast, a combined treatment of nitriding and carburizing decreases gradually from the surface to the inside [50]. Therefore, these hardness depth profiles also prove the formation of the carburizing S-phase. The variations in hardness in the deep region where the S-phase was not formed are likely due to the hardness of the secondary carbides. From these variations, the hardness of the secondary carbides was derived to be 10–20 GPa, which is lower than the hardness of the surfaces of the nitriding S-phase layers.
Figure 7. Hardness depth profiles of the S-phase layers at the nitriding temperature of 450 ºC with various WC compositions.

The Vickers hardness values of the surfaces of the WC composite layers are shown in Figure 8. The Vickers indentations were aimed at the surfaces of the AISI 316 L steel matrix, except for the WC particle location. The average hardness of the as-deposited 20 wt.% and 40 wt.% WC layers was 540 HV and 654 HV, respectively. Meanwhile, the single AISI 316 L steel layer without WC was 205 HV [30]. Therefore, the composite of WC particles improved the hardness of the as-deposited WC composite layers by more than 300 HV. The reason for this improvement owes to the formation of secondary carbides and the dissolved solid solution of tungsten. In addition, the hardness of the as-deposited 40 wt.% layer was 100 HV harder than that of the WC 20 wt.% layer because more secondary carbides existed.

Figure 8. Vickers hardness of the surfaces of the as-deposited and nitrided layers with various WC compositions and nitriding temperatures.

In contrast, the average hardness of the S-phase at a nitriding temperature of 450 ºC was 1384 HV for the 20 wt.% layer and 1308 HV for the WC 40 wt.% layer. This hardness was almost the same as that of the S-phase on the single AISI 316 L steel layer without WC [30]. As mentioned above, the hardness of the secondary carbides was lower than that of the nitriding S-phase. Therefore, after plasma nitriding, the secondary carbides and solid solution of tungsten are not effective in improving the hardness. On the contrary, the WC 40 wt.% layers had a low hardness compared to the WC 20 wt.% layers, suggesting that an increase in WC content decreases the S-phase hardness because the many secondary carbides with low hardness are present.

In consequence, the composite of WC particles only improved the hardness of as-deposited AISI 316 L steel matrix, not improved the S-phase hardness. However, the nitrocarburizing S-phase was formed by the composite of the WC particles, that has the
advantage of smoothing the hardness depth distribution, which is expected to restrict the delamination of the S-phase layer from the substrate when an external force is applied [50].

3.5. Corrosion Resistance of Nitrocarburizing S-Phase

Figure 9 shows the anodic polarization curves of the as-deposited and nitrided layers. The as-deposited WC 20 wt.% layer had a narrower passive region, and the WC 40 wt.% layer had no passive region.

![Anodic polarization curves](image)

**Figure 9.** Anodic polarization curves in 3.5 mass% NaCl solution of the as-deposited and nitrided layers with various WC compositions and nitriding temperatures: (a) of as-deposited layers, and (b) of nitrided layers.

The absence of a passive region implies a low corrosion resistance. These results indicate that the WC composite layers were inferior to the single AISI 316 L steel layer without WC [30]. In general, the effect of alloying elements on the pitting resistance of various types of stainless steels reported that Cr, Mo, W, and N strongly influence the pitting resistance equivalent numbers (PREN) as per Equation (1) [51].

\[
\text{PREN} = \%\text{Cr} + 3.3 \times (\%\text{Mo} + 0.5\%\text{W}) + 16 \times \%\text{N} 
\]

The solid solution of tungsten in the AISI 316 L steel matrix can improve the PREN. However, elemental analysis of SEM-EDX indicated that the compositions of chromium were approximately 11–13 wt.% of both the 20 wt.% and 40 wt.% WC layers, which showed a decrease compared to 17 wt.% of the feeding AISI 316 L steel powder. Passive films on austenitic stainless steels are composed of oxy-hydroxides of iron, chromium, and molybdenum species [52]; in particular, chromium oxy-hydroxides play an important role. Therefore, the decrease in the solid solution of chromium would degrade the passive film. Furthermore, secondary carbides should also prevent the uniform formation of the passive film on the AISI 316 L steel matrix. Therefore, the as-deposited WC composite layers exhibited a poor corrosion resistance.

In contrast, the nitrided WC composite layers had a passive region, and the corrosion current density was generally lower than that of the as-deposited layers. Figure 10 shows that the surfaces of the as-deposited layers exhibited pitting corrosion around the boundary of the tested area (5 mm × 5 mm), while the nitrided layers were discolored slightly, and no pitting corrosion was observed. Note that the black dots visible on the nitrided layers were not pitting corrosion, but WC particles. The dissolved nitrogen greatly improves the PREN; therefore, the formation of the S-phase improved the corrosion resistance.

The corrosion current density of the 40 wt.% layers was higher than that of the WC 20 wt.% layers because containing more secondary carbides would increase the corrosion current. In addition, the XRD pattern in Figure 2b indicates small CrN peaks in WC 40 wt.% layers treated at nitriding temperatures of 425 °C and 450 °C. The formation of CrN can also
decrease the solid solution of chromium to deteriorate the corrosion resistance. Therefore, increasing the WC content would deteriorate the corrosion resistance. For the same reason, the S-phase of the WC composite layers was inferior to that of the single AISI 316 L steel layer without WC [30].

![Graphs showing anodic polarization curves](image)

Figure 10. Surface morphologies after anodic polarization measurements of the as-deposited and nitrided layers with various WC compositions and nitriding temperatures: (a) WC 20 wt.% with as-deposited, (b) WC 20 wt.% at 400 °C, (c) WC 20 wt.% at 425 °C, (d) WC 20 wt.% at 450 °C, (e) WC 40 wt.% with as-deposited, (f) WC 40 wt.% at 400 °C, (g) WC 40 wt.% at 425 °C, and (h) WC 40 wt.% at 450 °C.

3.6. Summary and Perspectives

Nitriding as heat treatments has been applied to steel materials and has been highly effective in improving their properties. On the other hand, several papers about nitriding for stainless steel and other steel materials produced by laser cladding, selective laser melting and powder bed fusion, have been published on the improvement of wear and other properties [53–56]. In this study, it was found for the first time that the WC composite stainless steel layers have dendritic secondary carbides and dissolved tungsten, but the S-phase is formed by low-temperature plasma nitriding. In addition, not only a nitriding S-phase was formed, but also a carburizing S-phase was simultaneously formed by free carbon decomposed from WC particles. Owing to the nitrocarburizing, the S-phase thickened, and the hardness depth profiles were improved. This nitrocarburizing S-phase formed by nitriding alone is a unique phenomenon.

Recently, laser cladding and DED can be applied for an additive manufacturing as well as coatings. In these applications, there is a high demand for improved properties such as wear and corrosion resistance. As for steel materials, various heat treatment technologies have been developed and put into practical use to improve the characteristics. It is expected that heat treatment technologies will be applied to laser cladding and DED, contributing to the industry by expanding its applications.

4. Conclusions

In this study, we attempted to investigate whether low-temperature plasma nitriding applied to improving the hardness and corrosion resistance of AISI 316 L stainless steel-based WC composite layers fabricated by laser cladding. The effects of tungsten and tungsten carbides on the S-phase, which has not yet been clarified, were investigated. During laser cladding process, free carbon was produced by the decomposition of WC particles. Then, we attempted to form a nitrocarburizing S-phase by plasma nitriding alone using this free carbon. For these purposes, the metallurgical structure changes in the AISI 316 L stainless steel by WC particle dissolution during laser cladding, were examined in
detail. The formation mechanism of the S-phase was discussed, and the hardness and corrosion resistance were evaluated.

1. W\textsubscript{2}C, M\textsubscript{6}C, and M\textsubscript{12}C were synthesized in the as-deposited layers during the laser cladding process because the WC particles decomposed and reacted with the AISI 316 L steel matrix. The AISI 316 L steel matrix became a dendritic structure, and secondary carbides were distributed in a network. In addition, tungsten was dissolved in the austenite of the steel matrix;

2. Free carbon was also produced during the laser cladding process. By low-temperature plasma nitriding, this free carbon was pushed inward by the diffused nitrogen, forming a carburizing S-phase. Consequently, the S-phase became a dual layer, the nitriding layer sited at the outside, and the carburizing layer at the inside;

3. Owing to the nitrocarburizing, the thickness of the S-phase was greater than that of the single AISI 316 L steel layer without WC. In the WC 40 wt.% layers, secondary carbides densely precipitated, and the grain size was small, which would prevent the nitrogen diffusion. As a result, the WC 20 wt.% layers had thick S-phase compared with the WC 40 wt.% layers;

4. The Vickers hardness of the surfaces was improved to the range of 1200 to 1400 HV by low-temperature plasma nitriding. This hardness was almost the same as that of the S-phase on the single AISI 316 L steel layer without WC. The WC 40 wt.% layers were of a low hardness compared with the WC 20 wt.% layers, because the many secondary carbides with a low hardness were present. Meanwhile, the hardness depth profiles were improved to gradually decrease in the depth direction by the nitrocarburizing S-phase. This is expected to enhance the wear resistance and prevent the delamination of the S-phase layer from the substrate;

5. The corrosion resistance of the as-deposited layers was degraded due to the WC particle composite. In contrast, the nitried layers exhibited a low current density, and those surfaces after the anodic polarization measurements showed a slight color change, with no major damage such as pitting corrosion. Accordingly, low-temperature plasma nitriding of the WC composite layers improved the corrosion resistance to a generally excellent degree. The corrosion current density of the 40 wt.% layers was higher than that of the WC 20 wt.% layers because containing more secondary carbides and CrN.

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