APPLICATION OF Ir-BASE ALLOYS TO NOVEL OXIDATION RESISTANT BOND-COATINGS

H. Murakami1, A. Suzuki2, F. Wu3, P. Kuppusami4, and H. Harada2

1 Department of Materials Engineering, School of Engineering, The University of Tokyo, Tokyo, JP
2 Graduate School, Shibaura Institute of Technology, Tokyo, JP
4 Physical Metallurgy Section, Indira Gandhi Centre for Atomic Research, Kalpakkam, INDIA

Abstract

High temperature oxidation and hot corrosion properties of Ir-Ta, Ir-Pt coated and aluminized coatings are presented. An Ir-Ta binary alloy, proposed as a novel metallic bond coat material, was coated on a Ni-base single crystal superalloy TMS-75 using electron beam physical vapor deposition (EB-PVD), followed by a conventional Al pack cementation process. Cyclic oxidation tests and hot corrosion tests revealed that these Ir-Ta coated and aluminized specimens showed reasonably good oxidation and hot corrosion resistance. In addition, the formation of TCP phases is suppressed by the presence of the Ir-Ta enriched layer. These results indicated that the Ir-Ta alloy system is promising as a new metallic bond coat for high temperature structural materials. On the other hand, substitution of Pt with Ir in Pt-Al coatings is of interest for cost reduction and strengthening purposes. Cyclic oxidation properties of (Ir, Pt)-coatings, which were successfully electrodeposited on TMS-75, followed by aluminization, were also investigated. Cyclic oxidation tests at 1373 K in air revealed that materials having better thermal cyclic oxidation resistance can be arranged in the following order: Pt-Al = (36at%Ir, Pt-Al >> Ir-Al) simply aluminized. The presence of Ir and/or Pt may promote the formation of dense and adherent oxide scale and thus retard the growth of thermally grown oxides. Ir addition is thus promising, not just in terms of cost reduction, but also expected solid solution strengthening.

Introduction

Ni-base superalloys are used in the hottest section of gas turbines due to their good balance of adequate strength, high level of resistance to corrosion, oxidation and erosion at high temperatures. They are usually coated with a ceramic topcoat and an underlying metallic bond coat, such as NiCoCrAlY or aluminide in order to provide further oxidation protection to the superalloy substrate. However, recent demand in increasing the efficiency of turbines has led the components to be exposed at much higher operating temperatures. Under such circumstances, rapid interdiffusion of elements between substrates and bond-coats will promote the formation of detrimental TCP phases, resulting in the deterioration of mechanical properties of substrates. Therefore, there is a great demand for improving the protective performance of a bond coat material which also acts as a “diffusion barrier” to prevent the formation of detrimental phases. With this in mind, we propose Ir-base alloys as bond coat materials because Ir has excellent oxidation and corrosion resistance and has the highest melting temperature (2716K) among platinum group metals. It is also reported that the addition of Ir to Ni-base superalloys increase the melting temperature and thus may enhance the high temperature mechanical properties of the alloys [1]. In this study, we discuss the protective performance of several Ir-base alloys deposited on a Ni-base single crystal superalloy (TMS-75), followed by an Al-pack cementation process. Characteristics of Ir-Ta alloys using an EB-PVD method, and electrodeposited Ir-Pt alloys are discussed in terms of oxidation and corrosion resistance.

2. Experimental

In this paper, all the substrates were prepared from a heat-treated Ni-base single crystal superalloy TMS-75 (Ni:63.1, Co:12.6, Cr:3.5, Mo:1.3, W:2.0, Al:13.7, Ta:2.0, Hf:0.04, Re:1.7 in at%) [2,3]. The alloy was solution heat treated at 1573 K for 1 h followed by 1593 K for 5 h and gas fan cooled. A two-step aging treatment was then conducted, first at 1373 K for 4 h, second at 1143 K for 20 h, both followed by gas fan cooling to form coherent cuboidal γ precipitates.

For EB-PVD, the heat treated TMS-75 was cut into 17 x 17 x 2 mm plates, polished with emery paper up to #600, and buffed with the 0.25 µm diamond paste, and subsequently cleaned with acetone in an ultrasonic bath. An Ir-20at%Ta alloy, fabricated by the vacuum are melting method, was selected as a source material. The typical coating conditions were as follows, chamber pressure: 1.2-3.5x10⁻³ Pa and an electron beam acceleration voltage of 10 kV. Under these conditions, the typical deposition rate of approximately 10 µm/h was obtained for a single-sided coating. After that, the specimens were turned over and the other side of the specimens was coated. Entire surfaces including the edges were coated in this manner. For electrodepositing Ir-Pt alloys, The substrates were cut into disks 10 mm in diameter and 2 mm in thickness. The electrolytes were prepared from amidosulfuric acid, iridium chloride, chloroplatonic acid hexahydrate. Sodium nitrate was added to improve the conductivity. The nominal compositions of basic electrolyte are listed in Table 1.

Two platinum sheets were used as the counter electrodes with an area ratio of 2:1 to that of the specimens. The pretreatment procedure of the specimens prior to the electrodeposition consisted of following sequence: i) polishing with emery paper down to 2000 grit; ii) degreasing with acetone in an ultrasonic bath for 5 minutes; and iii) immersing the sample in a 30 wt% hydrochloric acid solution at 313 K for 5 minutes and then rinsing with distilled water. The pH value of the electrolyte was measured before and after every deposition ( HANNA pH meter 98108). In the event of a pH change, the value was corrected by adding 2N hydrochloric acid solution at 313 K for 5 minutes and then rinsing with distilled water. The pH value of the electrolyte was measured before and after every deposition ( HANNA pH meter 98108). In the event of a pH change, the value was corrected by adding 2N hydrochloric acid solution solution or 1N sodium hydroxide solution. In our study, all the experiments were carried out at controlled pH value of 2 ± 0.2. The electrolyte bath was stirred by a magnetic stirrer with a stirring speed of 60 rpm. The electrodeposition was conducted at the electrolyte temperature of 353 K and stirring with magnetic drum. The electrolyte temperature was controlled by constant temperature water bath with the accuracy of ± 2 K.
Typically coatings of 6 μm in thickness were obtained for 60 minutes of electrodeposition. The Ir-Ta and Ir-Pt-coated TMS-75 were then treated by a conventional pack-cementation aluminizing process, where these specimens were embedded in an Al2O3 retort containing a mixture of Al2O3, Al and Fe powder, and NH4Cl, under flowing Ar atmosphere. The pack-cementation aluminizing treatment was carried out at 1373 K for 5 h for Ir-Ta coatings and 1273 K for 5 h for Ir-Pt coatings, respectively. For comparison, uncoated TMS-75 was also aluminized under both conditions. Hereafter, these Ir-Ta coated, Ir-Pt-coated and uncoated TMS-75 samples with Al-pack cemented samples are denoted as (Ir, Ta)-Al-coated TMS-75, (Ir, Pt)-Al coated TMS-75, and simply aluminized TMS-75, respectively.

For the characterization of specimens at high temperatures, cyclic oxidation tests and hot corrosion tests were carried out in a programmable muffle furnace (Yamada Denki, MSFT-1500) at 1373 K in still air. The temperature variation in the furnace was controlled within ±2 K. Each thermal cycle consisted of 20 h at 1373 K and 4 h at ambient temperature and the tests were conducted up to 30 cycles. The oxidation resistance of the specimens was evaluated, in part, by the periodic weight changes of the specimens which were measured by a precision analytical balance with the accuracy of ±0.05 mg.

Hot corrosion tests were conducted for (Ir, Ta)-Al-coated and simply aluminized TMS-75. Specimens dipped with Na2SO4 molten salts were heated at 1173K for 20h in air, followed by air-cooling, which was defined as one heating cycle. This procedure was carried out up to 10 heating cycles. The crystal structure, surface morphology, and composition of the coated layer and oxides were analyzed by X-ray diffraction (XRD: RINT-2000/PC, Cu/Kα), scanning electron microscopy (SEM: Philips XL 30 with energy dispersive spectroscopy : EDS), and electron probe microanalysis (EPMA: JEOL JXA 9900R), respectively.

Table 1 Typical composition of Pt–Ir alloy electrolyte

<table>
<thead>
<tr>
<th>Component</th>
<th>Concentration (g/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iridium chloride (IrCl₃·4H₂O)</td>
<td>0 ~ 40.0 g/l</td>
</tr>
<tr>
<td>Chloroplatinic acid hexahydrate (H₂PtCl₆)</td>
<td>0 ~ 55.5 g/l</td>
</tr>
<tr>
<td>Amidosulfuric acid (HOSO₂NHS)</td>
<td>38.0 ~ 48.0 g/l</td>
</tr>
<tr>
<td>Sodium nitrate (NaNO₃)</td>
<td>40.0 ~ 50.0 g/l</td>
</tr>
</tbody>
</table>

pH (adjusted by 2N HCl or 1 N NaOH) 1.8 ~ 2.2
Temperature (T₆) 333 ~ 363 K
Current density (i₆) 0.5 ~ 4.0 A/dm²
Magnetic stirring 60 rpm
Deposition time 30 min

3. Results and discussions

3.1 Oxidation and hot corrosion properties of (Ir,Ta)-coated TMS-75

Oxidation resistance

The Ir-Ta alloy coatings, typically 8 μm in thickness were deposited on TMS-75 using the EB-PVD method. The coated layer is uniform, dense without any compositional variations. However, the average composition in the coating layer is Ir-70 at% Ta, while the evaporation source is composed of Ir-20at%Ta, indicating that the deposition rate of Ta is much greater than that of Ir. Though EB-PVD offers a possibility to deposit multi-component alloy coatings with different compositions, there are certain factors, which need to be taken into consideration. The composition variation of the coating is ascribed to the relative...
coated, and simply aluminized TMS-75, together with, shows the changes in weight of the (Ir, Ta)-Al-coated, Ta-Al-
investigated together with unaluminized specimens. Figure 3
The oxidation resistances of these aluminized specimens were
precipitation of other phases was not observed in either sample.
profile to that of the Ir-Ta-Al coated TMS-75, except for the
aluminized TMS-75 (Fig.2b) demonstrates a similar concentration
J
NiAl layer is enriched with Ta, and Cr, forming an Al-diffusion
E
Ir may have inward-diffused from the coating during the pack-
amount of Co, Ta, and Cr. A part of Ta and very small amount of
intermediate layer is mainly composed of β-NiAl-based outer
layer and ~20 μm Al-diffusion zone as the inner layer.
Figs 1b and 2b illustrate the concentration distribution of the
elements measured along the entire cross section of the sample
shown in Figs. 1a and 2a, respectively. For the (Ir, Ta)-Al-coated
TMS-75 (Fig. 1b), the outer Ir-Ta base alloy layer contains about
50 at% of Al together with a small amount of Ni (~5at%) and Co
(1at%) which may have outward-diffused from the substrate. The
intermediate layer is mainly composed of β-NiAl with a small
amount of Co, Ta, and Cr. A part of Ta and very small amount of
Ir may have inward-diffused from the coating during the pack-
cementation aluminizing process. The inner layer below the β-
NiAl layer is enriched with Ta, and Cr, forming an Al-diffusion
zone accompanied by a coarsened γ / γ’ structure. Although
having slightly thicker intermediate and inner layers, the simply
aluminized TMS-75 (Fig.2b) demonstrates a similar concentration profile to that of the Ir-Ta-Al coated TMS-75, except for the
presence of Ir-Ta-Al enriched layer in the latter. In addition,
precipitation of other phases was not observed in either sample.
The oxidation resistances of these aluminized specimens were
investigated together with unaluminized specimens. Figure 3
shows the changes in weight of the (Ir, Ta)-Al-coated, Ta-Al-
coated, and simply aluminized TMS-75, together with,
unaluminized, (Ir, Ta)-coated TMS-75 and as-heat-treated TMS-75
as a function of oxidation cycles. The sample weight gain is
mainly attributed to the oxidation of the surface, while weight loss
is attributed to the exfoliation of volatile oxide scales from the
specimen surface. Therefore, a sample demonstrating low weight

Figure 3: Kinetic oxidation curves of specimens as a function of oxidation cycle. Specimens were kept at 1373K for 20 hours and air-cooled for each cycle.

vapor pressure of Ir and Ta, the temperature of evaporation source, the distance between the specimen and the source, the position of the specimen in the vapor cloud, and other processing parameters [4]. Since the interaction among these factors mentioned above is complicated, and remains unclear especially in the case of Ir and Ta co-deposition, a new approach to precisely control the Ir-Ta composition in the coatings by EB-PVD should be developed near future.
The specimens are then treated by the conventional Al-pack cementation process at 1373K for 5 hours. Figures 1 and 2 show the cross-sectional microstructures and the concentration distribution of the elements for the (Ir, Ta)-Al-coated and simply aluminized TMS-75, respectively [5]. It can clearly be seen from Fig. 1a that the Ir-Ta-Al coated TMS-75 has a triplex layer structure, namely, an outer Ir-Ta-base alloy layer about 8-μm thick, an intermediate β-NiAl-based layer about 100 μm thick, and an inner Al-diffusion zone about 25 μm thick. On the other hand, for the uncoated simply aluminized TMS-75 (Fig.4a), only a duplex layer structure is observed, ~135 μm β-NiAl-based outer layer and ~20 μm Al-diffusion zone as the inner layer.
Figs 1b and 2b illustrate the concentration distribution of the elements measured along the entire cross section of the sample shown in Figs. 1a and 2a, respectively. For the (Ir, Ta)-Al-coated TMS-75 (Fig. 1b), the outer Ir-Ta base alloy layer contains about 50 at% of Al together with a small amount of Ni (~5at%) and Co (1at%) which may have outward-diffused from the substrate. The intermediate layer is mainly composed of β-NiAl with a small amount of Co, Ta, and Cr. A part of Ta and very small amount of Ir may have inward-diffused from the coating during the pack-
cementation aluminizing process. The inner layer below the β-
NiAl layer is enriched with Ta, and Cr, forming an Al-diffusion
zone accompanied by a coarsened γ / γ’ structure. Although
having slightly thicker intermediate and inner layers, the simply
aluminized TMS-75 (Fig.2b) demonstrates a similar concentration profile to that of the Ir-Ta-Al coated TMS-75, except for the
presence of Ir-Ta-Al enriched layer in the latter. In addition,
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Figure 4: Change in (a) microstructure and (b) corresponding chemical composition profile of (Ir, Ta)-Al-coated TMS-75 after cyclic oxidation tests.

Figure 5: Change in (a) microstructure and (b) corresponding chemical composition profile of simply aluminized TMS-75 after cyclic oxidation tests.
change, especially low weight loss during this oxidation test, can be regarded as having a good oxidation resistance. As for unaluminized specimens, sample weight linearly decreased with a rate of approximately 2.25 (mg/cm² per oxidation cycle) after several oxidation cycles, suggesting that these specimens had completely lost protective properties against oxidation. As for Ta-Al-coated and aluminized specimens, sudden decrease of weight after a few oxidation cycles is due to the spallation of surface layers. These results indicated that the (Ir,Ta)-Al-coated and simply aluminized TMS-75 have better oxidation resistance than the other specimens. However, for simply aluminized TMS-75, the sample weight increased after a few oxidation cycles and then became stable until 10th cycles. After that, the sample weight gradually decreased, suggesting that the spallation or exfoliation of surface oxidation layer started after 10 oxidation cycles. On the other hand, the weight change of (Ir,Ta)-Al-coated TMS-75 is much smaller than that of simply aluminized TMS-75, especially between the 10th and 30th oxidation cycles. For instance, the decrease in sample weight in (Ir,Ta)-Al-coated TMS-75 between the 10th and 30th cycles (2.38 mg/cm²) is about 4 times smaller than that in simply aluminized TMS-75 (13.0 mg/cm²), indicating that the former has better oxidation resistance at 1373 K.

Oxidation at high temperatures can involve microstructural evolutions of substrate materials. Figures 4 and 5 show the (a) cross-sectional microstructure of and (b) corresponding concentration profiles of solute elements in (Ir,Ta)-Al-coated (Fig. 4) and simply aluminized (Fig. 5) TMS-75 after 30 oxidation cycles.

It can clearly be seen from Fig.4a that an approximately 20-μm thick surface oxide layer was formed on the (Ir,Ta)-Al-coated TMS-75. The Ir-, Ta-, Ni-and Al-enriched layer of about 50-60 μm thick remains under the oxide layer. Under that, a β-NiAl-based layer of about 30 μm thick and an inner Al-diffusion zone being about 20 - 30 μm thick were formed. It should be noted that TCP phases precipitated were observed only within 300 μm in depth from the surface. For the simply aluminized TMS-75 (Fig.5a), an outer oxide layer of about 10-μm thickness was formed on the surface. Although the β-NiAl-based layer almost keeps its thickness of ~160 μm, there are a number of TCP precipitates spread not only in the β-NiAl-based layer, but throughout the observed region. Also, the Ni-content in the oxide layer is much higher in simply aluminized TMS-75 than in (Ir,Ta)-Al-coated TMS-75.

These microstructural changes occurring during oxidation tests can be explained in the following manner. When the aluminized specimens are exposed under hot air, the Al₂O₃ layer forms on the surface, which prevents the internal oxidation of the specimens. During oxidation tests, Al is continuously consumed in order to compensate for the spallation or evaporation of surface oxides. Simultaneously, interdiffusion between the Al-enriched layer and the substrate occurs for the uniform distribution of solute elements. Consequently, inward diffusion of Al in the Al-enriched surface layer to the substrate and alternative outward diffusion of solute elements in the substrate to the surface, such as Ni, Cr, and Co, occur during the oxidation tests. In addition, the concentration of Al in the surface area decreases with increasing oxidation cycles. These phenomena will lead to two detrimental effects, i) formation of brittle spinel oxides such as NiAl₂O₄ or volatile NiO and ii) precipitation of detrimental TCP phases in the substrate. Thermodynamic investigation confirmed that NiAl₂O₄ or NiO forms when the aluminum activity is low and those of oxygen and nickel are high [6]. Once a uniform and dense Al₂O₃ layer forms on the surface, such secondary oxides would never form because the permeability of nickel into the Al₂O₃ scale is very low. However, cyclic thermal stress during the oxidation test may initiate cracks in the oxide scale. If Ni is present right beneath the cracked oxide layer, it can react with oxygen in the gas and/or Al to form NiAl₂O₄ and/or NiO due to the low Al activity (Al concentration) at the reaction zone. The NiAl₂O₄ and/or NiO can then continue to grow by the transport (of oxygen gas) through the cracked Al₂O₃ layer. The slight decrease in weight for simply aluminized TMS-75 after 10 oxidation cycles can thus be explained by the spallation of brittle NiAl₂O₄ or exfoliation of volatile NiO. Indeed, in our previous investigation, X-ray analysis demonstrated the presence of NiAl₂O₄ and NiO in simply aluminized TMS-75 after 10 oxidation cycles [7]. In addition, Ni concentration in the oxide layer in (Ir, Ta)-Al-coated TMS-75 (less than 5 at%) is much lower than that in simply aluminized TMS-75 (typically 30 at%) after 30 oxidation cycles, as shown in Fig. 4(b) and Fig. 5(b), respectively. These results support the above discussion. On the other hand, (Ir, Ta)-Al-coated TMS-75 maintains stable Al₂O₃-based surface layer even after 30 oxidation cycles, resulting in the smaller mass change (2.38 mg/cm²) than that of simply aluminized TMS-75 (13.0 mg/cm²) from 10 to 30 oxidation cycles. Although further investigation would be required to understand the oxidation kinetics, it is speculated that the Ir-Ta-Al enriched oxide layer just beneath the surface, as shown in Fig.4(b), may hinder the outward diffusion of Ni, thus retarding the formation of NiAl₂O₄ and NiO.

Corrosion resistance
Having good hot corrosion resistance is another important requirement for coatings of high temperature structural materials. Comparisons of hot corrosion properties have been carried out in this section. Figure 6 shows the sample weight change as a function of hot corrosion cycles. Similar to the cyclic oxidation tests, the sample weight gain is mainly attributed to the formation of brittle and corrosive products such as sulfides or oxides while weight loss is mainly attributed to the evaporation of volatile oxides or spallation of such brittle and corrosive products as mentioned previously. Therefore, the significant variation in sample weight of the specimens except for (Ir, Ta)-Al-coated and simply aluminized TMS-75 indicates the repeated formation and

![Figure 6: Kinetic corrosion curves of specimens as a function of heating cycle. Samples were kept at 1173K for 20 hours and air-cooled for each cycle.](image)
spallation of brittle or corrosive products during corrosion tests, implying the poor corrosion resistance. On the other hand, sample weight changes of both (Ir, Ta)-Al-coated and simply aluminized TMS-75 are similar and smallest, suggesting that they have better corrosion resistance than the others. However, microstructural investigations revealed that they have different morphologies. Backscattered electron (BSE) images with concentration maps of these two specimens, derived from EPMA, are shown in Figure 7. In simply aluminized TMS-75, three kinds of layers are observed: an outer oxide layer with a thickness of about 5 μm (in Fig.7(a)), an intermediate Ni-Al based layer with a thickness of about 60 μm ((2) in Fig.7(a)) and an inner Al-diffusion layer with a thickness of about 30 μm ((3) in Fig.7(a)) from the surface. Note that in the BSE image, there are a number of dark spots corresponding to cracks or voids. The distribution of these dark spots agrees very well with that of oxygen enriched region. In addition, in the Ni-Al based layer, co-segregation of Cr and S is observed as red spots, which propagated to approximately 70 μm in depth from the surface. These results indicate that internal oxidation and sulfuration occurred in simply aluminized TMS-75. In the case of (Ir, Ta)-Al-coated TMS-75, on the other hand, an Ir-Ta-Al enriched layer lies in between the oxide layer and the Ni-Al enriched layer. The number of cracks and vacancies are much smaller than those in simply aluminized TMS-75. Neither the enrichment of O, nor the co-enrichment of Cr and S are observed in the (Ir, Ta)-Al-coated TMS-75 specimens.

Phase identification of the surfaces of the (a) simply aluminized TMS-75 and (b) (Ir, Ta)-Al-coated TMS-75 after the hot corrosion tests was conducted by X-ray diffraction analysis as in Figure 8. Na2SO4, which was used as the molten salt for the corrosion test, was not identified for either specimen, confirming that the molten salt was thoroughly removed from the surface when the samples were cleaned. NiAl and Al2O3 phases were identified on both surfaces of the specimens. In addition, the CrS phase was identified in simply aluminized TMS-75 whilst not sulfides but Ta2O5 and NaTaO3 were identified in (Ir, Ta)-Al-coated TMS-75. These results are consistent with the EPMA analysis described above.

From these investigations, it is revealed that (Ir, Ta)-Al-coated TMS-75 has higher corrosion resistance than any other specimens at 1173 K. There may be two major roles of the Ir-Ta layer for improving hot corrosion resistance: i) a diffusion barrier of solute elements as described in the previous section and ii) formation of tantalum oxides to protect the corrosive attacks by S.

<table>
<thead>
<tr>
<th>Component</th>
<th>Intensity ( Arbitrary Unit )</th>
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<tbody>
<tr>
<td>CrS</td>
<td>High</td>
</tr>
<tr>
<td>Ta2O5</td>
<td>High</td>
</tr>
<tr>
<td>NaTaO3</td>
<td>Low</td>
</tr>
</tbody>
</table>

It is well established that S and O in the molten salts diffuse very quickly at intergranular sites [8]. Especially, S diffuses extremely quickly and attacks grain boundaries, which gives rise to the formation of cracks. Also, when small voids are formed in a coating layer, O and S segregate at these small voids to form oxides and sulfides, which may become crack propagation sites. In simply aluminized TMS-75 as explained in the previous section, Ni, Co and Cr easily diffuse outwards. The outward diffused Cr and S thus formed CrS as shown in Fig. 7, which may have enhanced the initiation and propagation of cracks. Although further investigation is required to understand the detailed mechanism of CrS formation and the reason why only CrS is detected, it is concluded that many cracks and voids generated in simply aluminized TMS-75 are attributed to the formation of CrS which would degrade the mechanical properties of the coatings. In (Ir, Ta)-Al-coated TMS-75, on the other hand, α-Al2O3, NiAl, Ta2O5 and NaTaO3 peaks were identified from X-ray analysis. Lee et al. [9] reported that NaTaO3 is formed from Na2SO4 and Ta2O5 according to the following reactions:
Since the melting point of NaTaO$_3$ is 2083 K, NaTaO$_3$ will not be melted during the hot-corrosion tests and thus will not work as the corrosive agents. In addition, Ta$_2$O$_5$, a melting point of 2163 K, is known as the stable oxide. In 'Ir-Ta+Al', the presence of both α-Al$_2$O$_3$ and Ta$_2$O$_5$ on the specimen surface may have protected the surface from so-called "sulfur and oxygen attack" and prevented the formation of sulfides such as CrS and voids. It is also suggested that the Ir-Ta-Al enriched layer hinders the outward diffusion of Cr to the surface, thus avoiding the reaction between Cr and S. These results can explain the better corrosion resistance of (Ir, Ta)-Al-coated TMS-75.

3.2 Oxidation properties of (Ir, Pt)-coated TMS-75

In the previous section, we have proposed the Ir-Ta coatings as a candidate for the next-generation bond-coat materials. However, practical turbine blade industries will seek low-cost coating technologies. While proposing new materials, we are trying to develop new technologies to deposit Ir-Ta coatings with lower cost. On the other hand, addition of Ir to Pt-modified coatings is of interest because Ir-Pt alloys can be deposited using the electroplating process which we have recently developed [10]. Another advantage is the cost of the materials. Since Ir is cheaper than Pt, Ir deposition would reduce the cost of airfoils, compared to Pt-modified aluminate coatings [11].

Microstructure of as-aluminized specimens

Ir-Pt alloys with various compositions were deposited on TMS-75 followed by the al-pack cementation at 1273 K for 5 h. Figures 9 and 10 show the cross-sectional microstructure and the concentration distribution of the elements for the Pt–Al coated TMS-75 (Fig. 9), Ir-Al coated (Fig. 10) and simply aluminized TMS-75 (Fig. 3), respectively. It can clearly be seen from Fig. 9a and Fig. 10a that the Pt–Al coated and Ir-Al coated TMS-75 have triplex layer structure. XRD and EDS analyses revealed that both the coatings have B2 ordered structure, consisting of three continuous layers: a dense and uniform outer of β-(Pt,Ni)Al- or β-(Ir,Ni)Al-based layer ~14 μm thick, an intermediate β-NiAl-based layer ~50 μm thick, and an inner Al-diffusion zone ~5 μm thick. It should be noted that (Ir, Pt)-Al coated specimens also have a similar structure to that of Pt-Al-coated and Ir-Al coated TMS-75. On the other hand, simply aluminized TMS-75 at 1273 K showed a very similar structure to that aluminized at 1373 K, except for the thickness of β-NiAl-based layer (~60 μm at 1273 K while ~135 μm at 1373 K.)

Figures 9b and 10b illustrate the concentration distribution of the elements measured along the entire cross-section of the sample shown in Fig. 9a and 10a, respectively. For the Ir-Al coated TMS-75 (Fig. 10b), the outer β-(Ir,Ni)Al-based layer contains approximately 56 at.% of Al, 32 at.% of Ir together with a small amount of Ni (~10 at.%) and Co (1 at.%) which may be attributed to the outward-diffusion from the substrate. The intermediate layer is mainly composed of β-NiAl with a small amount of Co, Ta and Cr. Some Ir may diffuse inwardly from the coating during the pack-cementation aluminizing process. The inner layer below the β-NiAl layer is enriched with Ta, and Cr, forming an Al-diffusion zone accompanied by a coarsened γ’ structure. Although having slightly higher Al concentration and lower Ni concentration in the β-(Pt,Ni)Al layer, the Pt-Al coated TMS-75 (Fig. 10a) demonstrates a similar concentration profile to that of the Ir-Al coated TMS-75. The simply aluminized TMS-75
also demonstrates a similar behavior, except for the presence of Ir–Al or Pt-Al enriched layers in the former specimens.

**Oxidation behavior**

In order to evaluate the stability of oxides formed on the coated specimens under cyclic thermal stresses, a cyclic oxidation test was performed in exactly the same manner as described in the previous section. Specific mass gain as a function of the number of cycles for Pt-Al coated, (36at%Ir, Pt)-Al coated, Ir–Al coated and simply aluminized TMS-75 are given in Figure 11. The mass gains of the Pt-Al coated, (36at%Ir, Pt)-Al coated and Ir–Al coated TMS-75 increase monotonically with the increase of oxidation cycles. This indicates that the oxide scales formed on the modified TMS-75 are dense and have good adherence to the coating. On the other hand, in the case of simply aluminized TMS-75, the mass gain increases up to 7 oxidation cycles and then slightly decreases. This indicates that the spallation of the oxide scale from the surface has started after 7 oxidation cycles. In addition, it was found that for each oxidation cycle, the simply aluminized TMS-75 shows larger mass gain than the other specimens, confirming that Ir-Pt alloy deposition retards the growth rate of thermally grown oxides (TGO). Among the coated specimens, the Ir-Al coated one exhibits a slightly higher weight gain rate than Pt-Al coated or (36at%Ir, Pt)-Al coated specimens, yet the maximum mass gain of the Ir–Al coated TMS-75 is about half as large as that of the simply aluminized TMS-75. Note that Pt-Al and (36at%Ir, Pt)-Al showed almost identical oxidation kinetics, suggesting that the growth rate of TGO in Ir–Al coated TMS-75 is lower than that in the simply aluminized TMS-75.

Figure 12 shows SEM images of the surface morphologies and cross-sectional microstructures of the specimens after ten oxidation cycles. It can be seen that the surface of simply aluminized TMS-75 is rougher (Fig. 12g) and the oxide scale formed on the surface is thicker (~10 μm) and more porous (Fig. 12h) than the others, the oxide scale of which are typically ~3 μm thick. In addition, the amount of γ’-phase formed under the oxide scale in the simply aluminized TMS-75 is much larger than the other aluminized TMS-75, suggesting that the transformation from the β-NiAl phase to the γ’-phase was decelerated due to the presence of (Pt, Ir, Ni)Al enriched layer. The deceleration of the γ’-phase formation is beneficial in improving oxidation resistance of coated specimens, because the γ’-phase does not have good oxidation resistance. When comparing the Pt-Al coated (Fig. 12a&b), and Ir-Al coated (Fig. 12c&d) TMS-75, Pt-Al coated TMS-75 seems to show a smoother surface and smaller amounts of γ’-phase precipitates just beneath the oxide scale, indicating that Pt-Al coated TMS-75 has slightly better oxidation resistance. However, it is noteworthy that surface roughness and γ’-phase precipitates are almost identical between, Pt-Al coated and (36at%Ir, Pt)-Al coated specimens, suggesting that appropriate substitution of Ir for Pt may not affect the oxidation properties of coated materials.

Improved oxidation resistance at high temperature is essential for the next-generation high temperature structural materials and their coatings. They must protect the substrates by forming a dense and adherent oxide scale. The lower mass gain for the Pt-Al-coated, simply aluminized TMS-75 and Ir-Al coated shows a smooth surface and little amount of γ’-phase precipitation (Fig. 12a&b&c&d). Figure 13 shows SEM images of the cross-sectional microstructures of the specimens after ten oxidation cycles. It can be seen that the surface of simply aluminized TMS-75 is rougher (Fig. 12g) and the oxide scale formed on the surface is thicker (~10 μm) and more porous (Fig. 12h) than the others, the oxide scale of which are typically ~3 μm thick. In addition, the amount of γ’-phase formed under the oxide scale in the simply aluminized TMS-75 is much larger than the other aluminized TMS-75, suggesting that the transformation from the β-NiAl phase to the γ’-phase was decelerated due to the presence of (Pt, Ir, Ni)Al enriched layer. The deceleration of the γ’-phase formation is beneficial in improving oxidation resistance of coated specimens, because the γ’-phase does not have good oxidation resistance. When comparing the Pt-Al coated (Fig. 12a&b), and Ir-Al coated (Fig. 12c&d) TMS-75, Pt-Al coated TMS-75 seems to show a smoother surface and smaller amounts of γ’-phase precipitates just beneath the oxide scale, indicating that Pt-Al coated TMS-75 has slightly better oxidation resistance. However, it is noteworthy that surface roughness and γ’-phase precipitates are almost identical between, Pt-Al coated and (36at%Ir, Pt)-Al coated specimens, suggesting that appropriate substitution of Ir for Pt may not affect the oxidation properties of coated materials.
(36at%Ir, Pt)-Al-coated and Ir–Al-coated TMS-75 observed in the kinetic curves of the cyclic oxidation tests suggests the formation of a more dense and adherent oxide scale than that in the simply aluminized TMS-75, which retards the degradation of microstructure. Protective oxide scales kept for longer periods on the Pt-Al-coated, (36at%Ir, Pt)-Al-coated and Ir–Al-coated TMS-75 compared to the simply aluminized TMS-75 lead to higher oxidation resistance.

Although this study revealed that oxidation resistance of an Ir-Al coated TMS-75 is not as good as that of Pt-Al coated TMS-75, Ir-modified aluminide coating can be regarded as a promising material for the next-generation turbines because i) addition of Pt to Ir in the aluminized coating may act as a solid solution strengthener of the coatings, ii) up to 36at% of Ir addition may not deteriorate the oxidation resistance of the coated materials and iii) Ir is known to increase the melting point of substrate Ni-base superalloys [1], thus an Ir coating may not drastically affect the creep properties of substrates, as has been reported in Pt-modified aluminized coatings.

3. 3 Summary
Through this study, use of Ir for the novel bond-coat material has been proposed, and found that i) Ir-Ta coatings deposited using EB-PVD, followed by aluminization showed better oxidation and hot corrosion resistance than simply aluminized coatings, ii) electrodeposited Ir behaves similarly to Pt, and up to 36at% of Ir addition to Pt may not deteriorate the oxidation resistance of the coated materials. In addition, Ir-based alloy coatings in general seem to retard the interdiffusion of solute elements between substrates and coatings. These results confirmed that Ir-based alloys are promising for use as next-generation bond-coat materials. Ir-Ta coatings may be used under high temperature and corrosive atmosphere, and Ir-Pt modified aluminized coatings can straightforwardly replace current Pt-modified aluminized coatings. In order to realize practical usage for these coatings, further investigation would be required. Optimization of the coating parameters and characteristics, such as, composition of Ir-Ta and/or Ir-Pt, and the optimization of aluminizing parameters, in accordance with the characteristics of Ir-based coatings and substrate superalloys, would be utmost importance. Detailed investigations on the distribution of Ir, Ta and Pt presented in the coated materials after cyclic oxidation tests, the diffusion kinetics of solute elements such as Ir, Pt, Ta, Ni, and Al, and the mechanical properties such as creep and fatigue properties of the coatings, should be carried out in the future.

4. Conclusions
The results and the subsequent discussions described above support the following conclusions.

1. The Ir-Ta coated TMS-75 with Al-pack cementation showed better oxidation resistance than simply aluminized TMS-75. Moreover, the precipitation of the third phase was suppressed.

2. The hot corrosion tests and the following investigations revealed that in the Ir-Ta coated TMS-75 with Al-pack cementation, an Ir-Ta-Ni-Al enriched layer prevented the infiltration of oxygen and sulfur, and retarded the formation of voids and cracks in the inner layer, confirming that the Ir-Ta coated superalloys with Al-pack cementation also has better corrosion resistance.

3. Cyclic oxidation tests revealed that coated materials having better thermal cyclic oxidation resistance can be arranged in the following order: Pt-Al = (36at%Ir, Pt)Al > Ir-Al >> simply aluminized. This can be attributed to the presence of Pt and/or Ir which not only improved the adherence of oxide scale but also retarded the transformation of the β-(Ir,Ni)Al phase to the γ’ phase during oxidation.

4. (Pt, Ir)-Al coatings are promising for replacing conventional Pt-Al ones because Ir may play a role as solid solution strengthener and will not deteriorate the creep properties of substrates, although oxidation resistance of Ir-Al is not as good as that of Pt-Al. Thermal cyclic oxidation resistance of the Ir-Al coated TMS-75 is higher than that of simply aluminized TMS-75. This can be attributed to the presence of Ir, which not only improved the adherence of oxide scale but also retarded the transformation of the β-(Ir,Ni)Al phase to the γ’ phase during oxidation.

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