Abstract: The relative content of strengthening element tantalum (Ta) and oxidation-resistant element chromium (Cr) is an essential value for superalloys to obtain an excellent combination of oxidation resistance and mechanical properties. In the present paper, the isothermal oxidation behavior of several single crystal Ni-base superalloys with different Ta/Cr (wt. %, similarly hereinafter) ratios at 1000 °C in static air has been systematically investigated to explore the optimal Ta/Cr for excellent oxidation resistance. A detailed microstructure study using X-ray diffraction (XRD), scanning electron microscopy (SEM) with energy dispersive spectroscopy (EDS) and an electro-probe microanalyzer (EPMA) was performed to reveal the oxidation products and mechanisms. For all alloys, a three-layer structured scale consisting of an outer (Cr, Al, Ti, Ni, Ta)-O layer, an inner Al\textsubscript{2}O\textsubscript{3} layer and an inner nitride layer was formed. As Ta/Cr increased, the amounts of Ta-containing products, cracks, holes and inner nitride increased. Meanwhile, the completeness of the Al\textsubscript{2}O\textsubscript{3} layer got worse. It was shown that if Ta/Cr ≤ 0.5, Ta increased the growth rate of Cr\textsubscript{2}O\textsubscript{3} via the doping effect induced by Ta cations. If Ta/Cr > 0.5, Ta reduced the completeness of Cr\textsubscript{2}O\textsubscript{3} through competitive growth of Ta\textsubscript{2}O\textsubscript{5} and Cr\textsubscript{2}O\textsubscript{3}. A good oxidation performance can be expected with the value Ta/Cr ≤ 0.5.

Keywords: superalloy; high-temperature oxidation; tantalum oxide

1. Introduction

Nickel-base single crystal superalloys are widely used in large industrial gas turbines (IGTs) because of their outstanding combination of mechanical properties and environmental performance [1]. As IGTs developed, the operating temperature of the turbine blades became higher and higher. It was a new challenge to the hot end component materials because of severe oxidation and hot corrosion damage. During oxidation and hot corrosion, elements that assume the role of strengthening are consumed [2]. In addition, the inner corrosion destroys the microstructure of the blades [3]. Therefore, environmental damage severely deteriorates the service life of the blades. In recent years, approaches such as more effective thermal barrier coatings and cooling systems for blades have been used to improve this situation [4]. At the same time, efforts in developing superalloys with better temperature capability have been carried out. Researchers have been trying to increase the concentration of refractory elements (Re and Ta), which exhibited excellent performance in improving mechanical properties [5–7]. Consequently, the mechanisms by which Re and Ta affect hot corrosion resistance and oxidation behavior are very important for further optimization of their contents in superalloys for IGTs.

The authors have systematically investigated the effect of Re on hot corrosion and oxidation. It was shown that the Re addition improved both oxidation and hot corrosion...
resistance remarkably. The mechanisms, however, are different [8,9]. The effect of Ta on hot corrosion was systematically investigated. A quantization parameter Ta/Cr (wt. %, similarly hereinafter) was proposed to describe the effect of Ta [3,10]. Considering the different mechanisms of Re on hot corrosion and oxidation, it is of interest to reveal the effect of Ta on oxidation.

The effect of Ta addition on oxidation behavior is presently controversial. According to Yang et al. [11], a small amount of Ta improved the oxidation resistance of the alloy, while too much Ta content reduced the stability of the Al2O3 scale and deteriorated the oxidation resistance of the alloy. However, Barrett et al. [12] showed that high Ta concentrations can promote oxidation resistance, although the mechanism is not clear. Wu et al. [13] have studied the effect of Ir-Ta-Al coating on the cyclic oxidation behavior of alloys and declared that the Ta addition effectively inhibited the precipitation of the TCP phase in the coating, stabilized the coating structure and retarded the diffusion of elements. However, the results acquired by Kuppusami [14] are just the opposite. The results showed that increasing Ta content in Ir-Ta coating worsened the protection capability because the substitution of Al3+ by Ta5+ increased the concentration of Al vacancy, which led to increased consumption of Al. Thus, it is necessary to investigate the effect of Ta addition on oxidation behavior.

Cr is one of the most important elements for superalloys to resist oxidation attacks. Sufficient Cr promotes the formation of a continuous and adherent Cr2O3 scale on the surface of the alloy [15,16]. The diffusion rate of ions through the Cr2O3 scale is extremely slow. A complete Cr2O3 scale can protect the substrate from being damaged by oxygen or other corrosive media. Thus, describing the effect of Ta by introducing a parameter including Cr was necessary and practical. In the present paper, the isothermal oxidation behavior of several Ni-base single crystal superalloys with different Cr and Ta contents was investigated at 1000 °C. The oxidation process and mechanism of interaction of Ta and Cr on oxidation behavior were discussed. An optimal Ta/Cr ratio for good oxidation resistance was proposed.

2. Materials and Methods

The chemical compositions of the experimental alloys are listed in Table 1. The four alloys were designed with different Ta/Cr (wt. %) varying from 0.34 to 0.69, with the compositions of other alloying elements nearly unchanged.

| Alloys | Cr  | Ta  | Re  | Mo  | W   | Ti  | Al  | Co  | C   | Ni  | Ta/Cr |
|--------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-------|
| E7     | 12.00 | 4.09 | 1.97 | 1.91 | 4.00 | 3.99 | 3.41 | 9.02 | 0.05 | Bal. | 0.34  |
| E71    | 12.00 | 6.15 | 2.03 | 1.93 | 4.01 | 3.94 | 3.39 | 9.04 | 0.04 | Bal. | 0.51  |
| E73    | 9.84  | 6.04 | 2.04 | 2.01 | 3.99 | 3.94 | 3.30 | 9.01 | 0.04 | Bal. | 0.61  |
| E75    | 9.89  | 6.87 | 2.00 | 1.03 | 3.82 | 3.96 | 3.46 | 8.94 | 0.06 | Bal. | 0.69  |

The single crystal rods measuring Φ16 × 220 mm of the alloys were fabricated using the high rate solidification (HRS) method. A subsequent heat treatment was conducted for each alloy. Samples used in this experiment were slices measuring 20 × 10 × 1.5 mm. They were cut from the heat-treated single crystal rods using electro-spark wire-electrode cutting, with the 20 mm side parallel to the (001)-direction of the rods and the other sides left random. The surfaces of the specimens were ground to #800 and ultrasonically cleaned with acetone and alcohol before the test.

The isothermal oxidation test was conducted according to the China Aviation industry standard HB5258-2000. Samples were put into separated corundum crucibles and then placed in a furnace, the temperature of which was set at 1000 °C. The crucibles were taken out of the furnace at 1, 4, 7, 10, 13, 25, 50, 75, 100, 200 and 500 h for morphology inspection and weight monitoring. As soon as the crucibles were taken out, they were covered with special corundum lids, avoiding any weight loss caused by splashing oxide scales. Weight measurement was carried out after 1 h of cooling. Each data point of mass change
was the average of three samples. XRD (Shimadzu, Kyoto, Japan), SEM (FEI, Hillsboro, USA) equipped with EDS and EPMA (Shimadzu, Kyoto, Japan) were employed for the sample analysis.

3. Results

Heat-treated microstructures of the experimental alloys are illustrated in Figure 1. After standard heat treatment, all of the alloys exhibited similar microstructure. The eutectics were almost completely eliminated.

![SEM micrographs of the experimental alloys after standard heat treatment.](image)

Figure 1. SEM micrographs of the experimental alloys after standard heat treatment. (a,e): E7. (b,f): E71. (c,g): E73. (d,h): E75.

3.1. Macroscopic Morphologies

The macroscopic morphologies of the E7, E71, E73 and E75 alloys after isothermal oxidation at 1000 °C are shown in Figure 2. After only 4 h of oxidation, all samples were covered by gray, complete and dense scale. As the oxidation proceeded to 500 h, however, dispersed dot spallation was observed on the E7, E73 and E75 specimens, while the E71 samples remained complete and dense.

![Macroscopic morphologies of the E7, E71, E73 and E75 alloys after isothermal oxidation at 1000 °C.](image)

Figure 2. Macroscopic morphologies of the E7, E71, E73 and E75 alloys after isothermal oxidation at 1000 °C. (a) E7—4 h. (b) E71—4 h. (c) E73—4 h. (d) E75—4 h. (e) E7—500 h. (f) E71—500 h. (g) E73—500 h. (h) E75—500 h.

3.2. Oxidation Kinetics

Weight gain versus time curves for the E7, E71, E73 and E75 alloys after isothermal oxidation at 1000 °C are shown in Figure 3. All alloys followed the logarithm law. In the initial 100 h, the mass changes of alloys increased rapidly. After 100 h, the increase in sample weights of the E7, E71 and E75 slowed down while E73 still presented obvious weight addition. The final mass gain of the four alloys decreased in this order: E75 > (E71, E73) > E7.
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Figure 3. Oxidation kinetic curves of the E7, E71, E73 and E75 alloys after isothermal oxidation at 1000 °C.

3.3. XRD Results

XRD patterns of the oxidation products are provided in Figures 4 and 5 and Table 2. After 100 h of oxidation, common products TiO$_2$, Cr$_2$O$_3$ and (Ni, Co)-oxides were detected on all samples. In addition, Al$_2$O$_3$ was detected on E7, E71 and E73; NiAl$_2$O$_4$ was detected on E73 and E75; (Ni, Co)Ta$_2$O$_6$ was formed on E75. After 500 h of oxidation, NiCr$_2$O$_4$ was formed on E7 and E71. Al$_2$O$_3$ and NiAl$_2$O$_4$ could not be detected on E73. The composition of the scale on E75 was unchanged.

Figure 4. XRD patterns of samples exposed for 100 h at 1000 °C.
3.4. Surface and Cross-Section Morphology

After 4 h of oxidation, samples were analyzed to reveal the initial stages of the oxidation process. After 4 h of oxidation, all of the alloys revealed a similar microstructure. The representative cross-section microstructure and element distribution are shown in Figures 6 and 7. A layer with a three-layer structure was formed. The outer layer was composed of mixed (Cr,Al,Co,Ni,Ti)-oxides. The middle layer was an Al₂O₃ layer. The inner layer was a nitride layer. The thickness of the scale was about 2 μm.

Table 2. XRD results of the E7, E71, E73 and E75 alloys after isothermal oxidation at 1000 °C.

| Time  | Alloys | Phase Constituents                                      |
|-------|--------|--------------------------------------------------------|
| 100 h | E7     | TiO₂, Co₃O₄, Al₂O₃, Cr₂O₃, γ + γ′                     |
|       | E71    | TiO₂, Co₃O₄, Al₂O₃, Cr₂O₃, γ + γ′                     |
|       | E73    | TiO₂, Co₃O₄, Al₂O₃, Cr₂O₃, NiAl₂O₄, γ + γ′            |
|       | E75    | TiO₂, NiAl₂O₄, (Ni,Co)Ta₂O₆, Cr₂O₃, (Ni,Co)O₂, γ + γ′ |
| 500 h | E7     | TiO₂, Co₃O₄, Cr₂O₃, Al₂O₃, NiCr₂O₄, γ + γ′            |
|       | E71    | TiO₂, Co₃O₄, Cr₂O₃, Al₂O₃, NiCr₂O₄, γ + γ′            |
|       | E73    | TiO₂, Co₃O₄, Cr₂O₃, γ + γ′                            |
|       | E75    | TiO₂, NiAl₂O₄, (Ni,Co)Ta₂O₆, Cr₂O₃, (Ni,Co)O₂, γ + γ′ |

Figure 5. XRD patterns of samples exposed for 500 h at 1000 °C.

Figure 6. Cross-section morphologies of the oxide scale formed on the E7 alloy after oxidation at 1000 °C for 4 h.
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Figure 7. Secondary electron micrograph and corresponding X-ray micrograph of the E7 alloy after oxidation at 1000 °C for 4 h.

The surface morphologies of the experimental alloys after isothermal oxidation at 1000 °C are illustrated in Figure 8. Two types of regions with different morphologies and constituents were formed on the samples: one is the oxides cluster made up of small granular Ti-enriched oxides; the other is the plain oxides scale composed of fine oxides enriched in Al, Ti, Cr and Ni. The four alloys differed by the relative proportion of the two regions. After 100 h of oxidation, the surface morphologies of E7 (Figure 8a) and E73 (Figure 8e) were similar. The morphologies of E71 (Figure 8c) and E75 (Figure 8g) were alike. The amount of Ti-rich oxides on their surfaces was less than that of E7 and E73. The surface morphologies of E7, E71 and E73 alloys did not change significantly after 500 h of oxidation. For E75, however, spallation occurred, exposing the bare substrate.

Cross-section morphologies of the oxide scales formed on E7, E71, E73 and E75 alloys after oxidation at 1000 °C for 100 h are illustrated in Figures 9 and 10. There are two types of regions in the cross-section: the raised surface region and the plain surface region, in correspondence with the two regions in Figure 8. For the E7 alloy oxidized for 100 h, the plain surface region consisted of three layers. The outer layer was a compact oxide layer enriched in Cr, Al and Ni. A small amount of white Ta-enriched oxides appeared in this layer. The inner layer was a continuous Al₂O₃ layer. In addition, small amounts of TiN particles were formed in the front of the oxidation layer (Figure 10a and Table 3). The raised surface region indicated the cross-section of the oxide clusters. The outer layer of this region was an oxide protrusion enriched in Ti and Cr. The inner layer was a smooth Al₂O₃ layer. Small amounts of TiN can be observed on the substrate. The scale morphologies of E71 were similar to that of E7. The main difference was that the amounts of Ta-enriched oxides and TiN were higher on E71 (Figure 9c). The amounts of Ta-enriched oxides and TiN in E73 increased further.
Figure 8. Morphologies and EDS analyses of the oxide scales formed on E7, E71, E73 and E75 alloys after isothermal oxidation at 1000 °C. (a,c,e,g): E7, E71, E73 and E75 for 100 h, respectively. (b,d,f,h): E7, E71, E73 and E75 for 500 h, respectively.
raised surface region indicated the cross-section of the oxide clusters. The outer layer of this region was an oxide protrusion enriched in Ti and Cr. The inner layer was a smooth Al₂O₃ layer. Small amounts of TiN can be observed on the substrate. The scale morphologies of E71 were similar to that of E7. The main difference was that the amounts of Ta-enriched oxides and TiN were higher on E71 (Figure 9c). The amounts of Ta-enriched oxides and TiN in E73 increased further.

Figure 9. Cross-section morphologies of the oxide scales formed on the E7, E71, E73 and E75 alloys after oxidation at 1000 °C for 100 h. (a,c,e,g): Flat surface regions on the E7, E71, E73 and E75 alloys at 100 h, respectively. (b,d,f,h): Oxides clusters on the E7, E71, E73 and E75 alloys at 100 h, respectively. “A” “B” “C” represent the products at which it is labeled. The products information corresponding to “A” “B” “C” is listed in Table 3.
Figure 10. Secondary electron micrograph and corresponding X-ray micrograph of the E7 (a) and E75 (b) alloys after oxidation at 1000 °C for 100 h.

| Table 3. EDS results of the regions “A”, “B” and “C” in Figure 9 (at. %). |
|---|---|---|---|---|---|---|
| Al | Cr | Ti | Ni | Ta | O |
| A  | 6.31 | 28.26 | 3.59 | 1.07 | - | 60.76 |
| B  | 2.53 | 3.87 | 23.22 | 0.36 | - | 69.45 |
| C  | -   | 1.15 | 20.73 | 2.36 | 13.11 | 62.65 |

The cross-section morphology of E75 was significantly different from the other three alloys. Oxide clusters were almost invisible. In the plain surface region, an almost continuous Ta-enriched layer can be seen at the middle of the oxide layer, with a lot of holes dispersing between this layer and adjacent oxides. The Al$_2$O$_3$ layer was continuous, with a large number of needle-like TiN structures dispersed on the substrate (Figure 10b). While in the raised surface region, the Ta-enriched layer and Al$_2$O$_3$ layer are discontinuous. The amount of TiN increased further.
After 500 h oxidation (Figure 11), the composition and morphology of the oxide scale on the surface of E7 and E71 were almost unvaried. Cracks and large holes were formed on E75. The scale of E75 became porous and fragile. The EPMA results of the scale formed on E75 are illustrated in Figure 12. The Ta-enriched layer and Al$_2$O$_3$ layer are discontinuous. The scale on E73 at 500 h was similar to that of E75 at 100 h. A continuous Ta-rich layer was accompanied by the formation of a large number of porosities (Table 4). The number of TiN particles increased. The scale became thick and loose, with visible cracks and holes.

Figure 11. Cross-section morphologies of the oxide scales formed on the E7, E71, E73 and E75 alloys after oxidation at 1000 °C for 500 h. (a,c,e,g): Flat surface regions on the E7, E71, E73 and E75 alloys at 500 h, respectively. (b,d,f,h): Oxides clusters on the E7, E71, E73 and E75 alloys at 500 h, respectively. “A” and “B” represent the products at which it is labeled. The products information corresponding to “A” and “B” is listed in Table 3.
E75 alloys at 500 h, respectively. “A” and “B” represent the products at which it is labeled. The products information corresponding to “A” and “B” is listed in Table 3.

Figure 12. Secondary electron micrograph and the corresponding X-ray micrograph of the E75 alloy after oxidation at 1000 °C for 500 h.

Table 4. EDS results of the regions “A” and “B” in Figure 11 (at. %).

|     | Al  | Cr  | Ti  | Ni  | Ta  | Co  | O    |
|-----|-----|-----|-----|-----|-----|-----|------|
| A   | 9.71| 19.63| 5.22| 3.85| -   | 0.62| 60.97|
| B   | -   | 9.56| 7.99| 4.54| 11.94| 0.63| 65.33|

4. Discussion

4.1. Oxidation Process

As soon as the alloy was exposed to a high-temperature oxidation atmosphere, the surface of the alloy was oxidized. Alloying elements diffused outward to participate in the oxidation process and resulted in a thin initial oxide layer above the sample surface. At the same time, the inward diffusion of oxygen and nitrogen was also underway. Inner oxidation and nitrogenization took place simultaneously.

Cr$_2$O$_3$ layers and Al$_2$O$_3$ layers are effective protective outer scales formed on superalloys under the oxidation atmosphere [15]. Due to the moderate Cr and Al contents in the alloys, neither a complete outer Cr$_2$O$_3$ nor Al$_2$O$_3$ layer was formed after the initial transition period. Instead, a Cr$_2$O$_3$-dominant layer accompanied by mixed oxides of Ni, Co, Ti, Al and their spinel was formed on the sample surface [15]. Ti has a high affinity for oxygen and a high diffusion rate in the alloy [17]. Therefore, TiO$_2$ was formed above the previously formed outer layer by outward diffusion of Ti.

Because of the establishment of an outer layer, the oxygen partial pressure at the interface of the outer layer and substrate was reduced. Consequently, the oxide that has the highest thermodynamic stability was formed here [15]. Thus, an inner Al$_2$O$_3$ layer was raised beneath the outer oxide layer. In the front of oxidation where the oxygen partial pressure was low, nitride was formed [18]. Since Al$_2$O$_3$ was extremely dense and the diffusion rate of ions passing through it was very low, it played an important role in
hindering the inward diffusion of nitrogen/oxygen and the outward diffusion of alloying elements [18]. Thus, for samples that had complete and dense outer oxide layers and inner Al₂O₃ layers, small amounts of nitrides appeared. For samples that had incomplete layers, large amounts of nitrides were generated [18].

In the incubation period, all of the alloys underwent rapid oxidation and their masses changed rapidly. The slope of the kinetics curve was large. At the end of the incubation period, a stable three-layer oxide scale consisting of an outer Cr₂O₃-dominant layer, an inner Al₂O₃ layer and an inner nitride layer was formed on the sample. From then on, the inward or outward diffusion of ions was significantly retarded, and the samples’ masses changed slowly. Therefore, all alloys followed the logarithm law [19].

4.2. Interaction of Ta and Cr

During the oxidation process, the structure of the scale on E7, E71, E73 and E75 was similar. However, differences in microstructure and constituents exist. Ta-containing products were formed on all alloys, with the least on E7 and the most on E75. The Al₂O₃ layer on E7 and E71 was continuous, while that on the E73 and E75 alloys was discontinuous. The amount of TiN on the E7 and E71 alloy was much less than that on E73 and E75. Holes and cracks appeared on E73 and E75. According to the results, increasing Ta content has limited benefit for improving the oxidation performance. Excessive Ta content destroyed the compactness of the oxide scale and deteriorated the isothermal oxidation property. Based upon the previous research [10], it is believed that the influence of Ta on the oxidation performance was due to the competitive growth of Ta₂O₅ and Cr₂O₃.

As the oxidation proceeded, the structure and constituents of the scale were affected by thermodynamic factors, such as the thermodynamic stabilities of oxides, and kinetic factors, such as the concentration of elements and their diffusivities and mobilities [10]. The element which has the higher affinity for oxygen will be oxidized first. It is the most stable component during the whole process. However, the element with the highest concentration and diffusivities will be oxidized faster. The relevant oxides are the most abundant and were prone to deriving the corresponding spinels [20]. The constituents of the final scale were the result of the competitive growth of oxides.

The standard free energy of the formation of oxides is illustrated in Figure 13. As is revealed, the ΔG⁰ of Ta₂O₅ is more negative than that of Cr₂O₃. Ta₂O₅ was thermodynamically favored. In addition, the diffusivity and mobility of Ta were faster than Cr [17,21]. Thus, another kinetic factor, the concentration of the two elements, is an important factor in affecting the competitive growth of Ta₂O₅ and Cr₂O₃ and further affecting the scale composition and the corrosion behavior. In the initial hours of oxidation, a variety of oxides can form, while the final composition of the scale was determined by thermodynamic factors as well as kinetic factors.

Because of the content advantage of Cr over Ta in E7, the dominant oxide of the outer layer was Cr₂O₃. A small amount of Ta₂O₃ and Ta-containing spinels dispersed among the Cr₂O₃, playing the role of dopant. When Cr₂O₃ is doped with Ta₂O₅, higher valence Ta ions, Ta⁵⁺, will increase the concentration of cation vacancies in Cr₂O₃ scale and further accelerate the growth rate of Cr₂O₃ [10,22]:

\[
3\text{Ta}_2\text{O}_5 = 6\text{Ta}^{5+}_\text{Cr} + 15\text{O}_2 + 4\text{V}^{\text{Cr}}''
\]  

(1)

Cr₂O₃ is an effective protective oxide under high-temperature oxidation because it is compact and dense to suppress the inward diffusion of oxygen [23–26]. The improved proportion of Cr₂O₃ in the oxide scale reduced the oxygen partial pressure at the outer layer/substrate interface, which then accelerated the selective oxidation of Al [15]. Therefore, the Al₂O₃ layer in E7 is continuous and thick. Few internal nitrides were formed, even after 500 h of oxidation. Because of the superior protection of the scales formed in the incubation period, the sample was well protected afterwards. The rate of mass change was reduced to an extremely low level after 100 h. The schematic diagram of the oxidation mechanism of E7 is illustrated in Figure 14a.
Figure 13. Standard free energy of formation of oxides. The diagram was plotted using data from HSC Chemistry® version 6.0.

Figure 14. Schematic diagrams of the oxidation mechanisms of the alloys: (a) alloys with Ta/Cr ≤ 0.5; (b) alloys with Ta/Cr > 0.5.

For E71, the increase in the Ta/Cr value in the alloy enhanced the kinetic advantage of Ta. More Ta-containing spinel TiTaO₄ (Table 3 “C”) was formed in E71. Spinel can reduce the diffusion rate of ions [27] and thereby improve the isothermal oxidation performance of the alloy. Small amounts of nitrides were formed. Thus, the rate of mass change was reduced to an extremely low level after 100 h. The oxidation mechanism of E71 can be described by the schematic diagram illustrated in Figure 14a.
The reduction in Cr and promotion of Ta in E73 and E75 narrowed the content difference between Ta and Cr, which reinforced the kinetic advantage of Ta. Competitive growth of Ta$_2$O$_5$ and Cr$_2$O$_3$ took place. The percentage of Cr$_2$O$_3$ in the outer layer was lowered. The reduction in Cr$_2$O$_3$ content retarded the selective oxidation of Al. Therefore, the Al$_2$O$_3$ layer in E73 and E75 was incomplete, and many more internal nitrides were produced. After long-term oxidation, the layers became loose and porous, with cracks and holes in them. The sample could not be protected by the scale. The rate of mass change remained high after 100h. The oxidation mechanism of E73 and E75 can be described by the schematic diagram illustrated in Figure 14b.

5. Conclusions

The isothermal oxidation behavior of a single crystal Ni-based superalloy with varying Ta/Cr ratio at 1000 °C was investigated. It was shown that excessive Ta content reduced the isothermal oxidation. Ta/Cr ratio affected the composition of the alloy oxide scale.

1. If Ta/Cr ≤ 0.5, Ta improved the growth rate of Cr$_2$O$_3$ and promoted the selective oxidation of Al. The oxidation resistance of the alloy was excellent. The mechanism was the doping effect induced by Ta cation.
2. If Ta/Cr > 0.5, Ta reduced the growth rate of Cr$_2$O$_3$ and suppressed the formation of Al$_2$O$_3$. The oxidation property of the alloy deteriorated. The mechanism was the competitive growth of Ta$_2$O$_5$ and Cr$_2$O$_3$.
3. A good oxidation performance can be expected with the value Ta/Cr ≤ 0.5.

Author Contributions: Conceptualization, D.W.; Data curation, W.F. and W.Z.; Funding acquisition, J.C. and J.D.; Investigation, J.C.; Methodology, J.C.; Resources, D.W.; Supervision, J.C. and D.W.; Writing—original draft, J.C.; Writing—review & editing, H.J., Y.L. and S.L. All authors have read and agreed to the published version of the manuscript.

Funding: This work was funded by the National Natural Science Foundation of China under grant No. 51901179, the National Science and Technology Major Project under grant No. 2017-VI-0019-0091 and the National Science Foundation of Shaanxi Province of China under grant No. 2018JQ5198.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: Not applicable.

Conflicts of Interest: The authors declare no conflict of interest.

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