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### Electro-codeposition of MCrAIY Coatings for Advanced Gas Turbine Applications

by

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#### Summary

MCrAlY coatings electrodeposited in a Watts bath typically contain a trace amount of sulfur, which can adversely affect their high-temperature oxidation resistance. Sulfur-free, all-chloride plating solutions were utilized in the electro-codeposition process to lower the sulfur content in the Ni-Co coating matrix. Three solutions were prepared. One solution included no wetting agent, while the other two solutions contained sodium lauryl sulfate (SLS) and Triton X-100 (TX100), respectively. After plating, the NiCo-CrAlY composite coatings were heat treated at 1080°C in vacuum and were converted to the NiCoCrAlY coatings that consisted of phases such as  $\gamma$ -(Ni,Co),  $\beta$ -NiAl, etc. The cyclic oxidation resistance of the coating plated in the all-chloride solutions were evaluated in air at 1100°C for 200 hr (1-hr per cycle with a total of 200 cycles). The coating plated in the solution with SLS wetting agent showed the best oxidation performance. This coating also demonstrated much better oxidation resistance that the bare CMSX-4 alloy. The results suggest that the all-chloride solution can be used to reduce the sulfur content in the MCrAlY coatings and satisfactory oxidation performance could be achieved. Further evaluation is needed to elucidate the effects of wetting agent on both the coating quality and oxidation behavior.

### Technical report

### I. Introduction

To improve high-temperature oxidation and corrosion resistance of critical superalloy components in gas turbine engines, metallic coatings such as diffusion aluminides or MCrAIY overlays (where M = Ni, Co or Ni+Co) have been employed, which form a protective oxide scale during service.<sup>1</sup> The state-of-the-art techniques for depositing MCrAIY coatings include electron beamphysical vapor deposition (EB-PVD) and thermal spray processes.<sup>1</sup> Despite the flexibility they permit, these techniques remain line-of-sight which can be a real drawback for depositing coatings on complex-shaped components. Further, high costs are involved with of the EB-PVD process.<sup>2</sup> Several alternative methods of making MCrAIY coatings have been reported in the literature, among which electro-codeposition appears to be a more promising coating process.

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Electrolytic codeposition (also called "composite electroplating") is a process in which fine powders dispersed in an electroplating solution are codeposited with the metal onto the cathode (specimen) to form a multiphase composite coating.<sup>3,4</sup> The process for fabrication of MCrAIY coatings involves two steps. In the first step, pre-alloyed particles containing elements such as chromium, aluminum and yttrium are codeposited with the metal matrix of nickel, cobalt or (Ni,Co) to form a (Ni,Co)-CrAIY composite coating. In the second step, a diffusion heat treatment is applied to convert the composite coating to the desired MCrAIY coating microstructure with multiple phases of  $\beta$ -NiAl,  $\gamma$ -Ni, etc.<sup>5</sup>

Compared to conventional electroplating, electro-codeposition is a more complicated process because of the particle involvement in metal deposition. It is generally believed that five consecutive steps are engaged:<sup>3,4</sup> (i) formation of charged particles due to ions and surfactants adsorbed on particle surface, (ii) physical transport of particles through a convection layer, (iii) diffusion through a hydrodynamic boundary layer, (iv) migration through an electrical double layer and (v) adsorption at the cathode where the particles are entrapped within the metal deposit. The quality of the electro-codeposited coatings depends upon many interrelated parameters, including the type of electrolyte, current density, pH, concentration of particles in the plating solution (particle loading), particle characteristics (composition, surface charge, shape, size), hydrodynamics inside the electroplating cell, cathode (specimen) position and post-deposition heat treatment, if necessary.<sup>3-6</sup>

There are several factors that can significantly affect the oxidation and corrosion performance of the electrodeposited MCrAIY coatings, including: (i) the volume percentage of the CrAIY powder in the as-deposited composite coating, (ii) the CrAIY particle size/distribution and (iii) the sulfur level introduced into the coating from the electroplating solution. This three-year project aims to optimize the electro-codeposition process for improved oxidation/corrosion performance of the MCrAIY coatings. The three main tasks are as follows:

- Task 1 (Year 1): Effects of current density and particle loading on CrAIY particle incorporation.
- Task 2 (Year 2): Effect of CrAIY particle size on CrAIY particle incorporation.
- Task 3 (Year 3): Effect of electroplating solution on the coating sulfur level.

### II. Background

A typical MCrAIY coating consists of 8–12% AI, 18–22% Cr, and up to 0.5% Y (in wt%). Other more complicated compositions of MCrAIYs contain additional elements such as hafnium, silicon and tantalum.<sup>7,8</sup> The concentrations of some minor elements (*e.g.*, sulfur, yttrium and hafnium) play an important role in affecting the growth and adhesion of the oxide scale. The detrimental effect of sulfur on oxide scale adherence of MCrAIY alloys has been well documented.<sup>9</sup> Small amounts of sulfur can segregate to the alumina-metal interface and weaken the interface.<sup>10</sup>

The electrolytes used to deposit the nickel or cobalt metal matrix for forming the MCrAIY coating are typically sulfate- or sulfamate-based solutions.<sup>11,12</sup> Approximately 0.006-0.013 wt% (60-130 ppm) of sulfur has been reported in electroplated nickel coatings using these solutions.<sup>13,14</sup> Table 1 lists the five commercial nickel plating solutions and their deposition parameters.<sup>15</sup> According to our previous study results, the fluoborate bath is not suitable for codeposition of CrAIY-based particles, due to the reaction between the solution and the powder at 50°C, leading to the formation of a dark powdery coating. The all-chloride solution appeared more promising. In this study, the oxidation performance of the MCrAIY coatings electrodeposited in the all-chloride solution was evaluated.

### III. Experimental Procedure

The substrate material used in this study was Ni-based superalloy CMSX-4 (60.7 Ni-5.9 Al-6.3 Cr-9.6 Co-6.5 W-0.6 Mo-2.9 Re-6.5 Ta-1.0 Ti, in wt%, and 1100 Hf-17 C-1 S in ppmw). Disc specimens (1.6 mm thick, ~17 mm in diameter) were ground to #600 grit using SiC grinding papers, followed by grit blasting with #220 Al<sub>2</sub>O<sub>3</sub> grit. The spherical CrAIY powder used in this research was manufactured via gas atomization by Sandvik. The powder had a mean particle diameter of 9.5  $\mu$ m and it was sieved through a 20- $\mu$ m screen (625 mesh) prior to usage.





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Solution ID	Nickel chloride (g/L)	Cobalt chloride (g/L)	Boric acid (g/L)	Wetting agent type and amount (g/L)		
Solution 1 (All-chloride)	225	25	33	0		
Solution 2 (All-chloride, with SLS)	225	25	33	Sodium lauryl sulfate (SLS): 0.02 Triton X-100 (T100): 0.02		
Solution 3 (All-chloride, with T100)	225	25	33			
Current density	pH: 2.0-3.0					
Temperatu	Plating time: 5 h					

Table 1 - Composition of all-chloride solutions and plating conditions

A rotating barrel system similar to the one illustrated in Fig. 1 was used in the electro-codeposition experiments.<sup>6</sup> Three allchloride solutions were prepared. Their compositions and the electro-codeposition parameters are given in Table 1. Two types of wetting agents were utilized. The sulfur-containing wetting agent was sodium lauryl sulfate (SLS), and the sulfurfree one was Triton X-100 (T100).<sup>16</sup>

After plating, the coated specimens were heat-treated in vacuum for 6 hr at 1080°C. The purpose of the heat treatment was to convert the as-deposited Ni/Co-CrAIY composite coating to the NiCoCrAIY coatings that consisted of phases such as  $\gamma$ -(Ni,Co),  $\beta$ -NiAI, etc. The post-deposition heat treatment was carried out in a horizontal alumina tube furnace. A vacuum of at least 10<sup>-6</sup> torr was maintained. The heating rate was 20°C/min. After completion, the samples were furnace cooled to room temperature. The coatings were examined by optical microscopy and scanning electron microscopy (SEM) equipped with energy dispersive spectroscopy (EDS) (Fig. 2).



Figure 1 - Schematic of the barrel system.

The oxidation performance of the MCrAIY coatings was evaluated in cyclic oxidation testing at Oak Ridge National Laboratory (ORNL). The test was carried out in laboratory air with 1-hr cycles at 1100°C with a 10-minute cooling period. The specimens were removed and weighed every 20 cycles for the first 100 hours and every 50 cycles thereafter. The mass changes ( $\pm 0.02$  mg/cm<sup>2</sup> accuracy) were recorded.

The coating surface compositions determined by EDS before and after heat treatment are summarized in Table 2. For all three coatings, a decrease in Cr content was observed, as a result of Cr evaporation in vacuum at 1080°C, similar to what was reported for MCrAIY coatings made by thermal spraying and EB-PVD.<sup>17,18</sup>

Coating #2 (plated in the solution with SLS) had higher Co, Cr and Al levels. Coating #1 (plated in the solution without wetting agent) had acceptable levels of Cr and Al, whereas the Co content was the lowest (*i.e.*, 3.4 wt%). This was possibly due to a lower cathode efficiency, since there was no wetting agent in this solution and the specimen was plated at a lower pH value. The Co, Cr and Al levels were lower for Coating #3 (plated in the solution with T100). This was likely due to a lower particle incorporation in the coating.





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Figure 2 - SEM surface images of the specimens for oxidation tests before and after heat treatment.

- (a) and (b): sample plated in the all-chloride solution (Coating #1, without wetting agent);
- (c) and (d): sample plated in the all-chloride solution with SLS wetting agent (Coating #2);
- (e) and (f): sample plated in the all-chloride solution with TX100 wetting agent (Coating #3).

Table 2 - Coating surface compositions before and after heat treatment, as determined by EDS (wt%).





Specimen - Solution Used	Before Heat Treatment			After Heat Treatment				
	Ni	Со	Cr	AI	Ni	Со	Cr	AI
Coating #1 (no wetting agent)	57.9	3.1	24.9	14.1	72.1	3.4	13.8	10.7
Coating #2 (SLS)	62.9	10.8	17.6	8.7	62.9	12.0	11.2	13.9
Coating #3 (TX100)	69.9	4.3	15.7	10.1	77.9	5.3	9.9	6.9

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The specific mass changes after 200 1-hr cycles are presented in Fig. 3. The uncoated CMSX-4 substrate and a previous coating plated in the Watts bath were also included for comparison. A coating with good oxidation resistance should have a slow and steady growth in weight, due to the formation of a thin and adherent Al<sub>2</sub>O<sub>3</sub> scale on the surface. Coating #1 had a slow steady mass gain in the first 100 cycles, but started to lose weight after 100 hr, indicating spallation of the oxide scale. Coating #3 (T100), on the other hand, showed a very rapid mass gain, as a result of fast growth of the oxide scale. Coating #2 (SLS) had the best oxidation performance, exhibiting a steady mass gain and only started to show signs of mass loss near the end of the 200-hr test. This coating plated in the Watts bath. In addition, the oxidation results were in good agreement with the coating surface composition.



Figure 3 – Specimen mass changes during 1-hr cyclic oxidation in laboratory air at 1100°C.

The current oxidation results suggest that with the proper wetting agent, the all-chloride solution can be used to reduce the sulfur content in the MCrAIY coatings and satisfactory oxidation performance could be achieved. Further evaluation is needed to elucidate the effects of wetting agent on both coating quality and oxidation behavior.

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#### Past project reports

- 1. Quarter 1 (January-March 2018): Summary: *NASF Report* in *Products Finishing*, *NASF Surface Technology White Papers*, 82 (12), 13 (September 2018); Full paper: <u>http://short.pfonline.com/NASF18Sep1</u>.
- Quarter 2 (April-June 2018): Summary: NASF Report in Products Finishing, NASF Surface Technology White Papers, 83 (1), 13 (October 2018); Full paper: <u>http://short.pfonline.com/NASF18Oct1</u>.
- 3. Quarter 3 (July-September 2018): Summary: *NASF Report* in *Products Finishing*, *NASF Surface Technology White Papers*, 83 (3), 15 (December 2018); Full paper: <u>http://short.pfonline.com/NASF18Dec1</u>.
- 4. Quarter 4 (October-December 2018): Summary: *NASF Report* in *Products Finishing*, *NASF Surface Technology White Papers*, 83 (7), 11 (April 2019); Full paper: <u>http://short.pfonline.com/NASF19Apr1</u>.
- 5. Quarter 5 (January-March): Summary: *NASF Report* in *Products Finishing*, *NASF Surface Technology White Papers*, **83** (10), 11 (July 2019); Full paper: <u>http://short.pfonline.com/NASF19Jul1</u>.
- 6. Quarter 6 (April-June): Summary: *NASF Report* in *Products Finishing*, *NASF Surface Technology White Papers*, **84** (1), 17 (October 2019); Full paper: <u>http://short.pfonline.com/NASF19Oct2</u>.
- 7. Quarter 7 (July-September): Summary: *NASF Report* in *Products Finishing*, *NASF Surface Technology White Papers*, **84** (3), 16 (December 2019); Full paper: <u>http://short.pfonline.com/NASF19Dec1</u>.
- Quarter 8 (October-December): Summary: NASF Report in Products Finishing, NASF Surface Technology White Papers, 84 (6), 11 (March 2020); Full paper: <u>http://short.pfonline.com/NASF20Mar3</u>.
- 9. Quarter 9 (January-March): Summary: *NASF Report* in *Products Finishing*, *NASF Surface Technology White Papers*, 84 (9), 9 (June 2020); Full paper: <u>http://short.pfonline.com/NASF20Jun2</u>.
- 10. Quarter 10 (April-June): Summary: NASF Report in Products Finishing, NASF Surface Technology White Papers, 85 (1), 15 (October 2020); Full paper: http://short.pfonline.com/NASF20Oct1.
- Quarter 11 (July-September): Summary: NASF Report in Products Finishing, NASF Surface Technology White Papers, 85 (4), 14 (January 2021); Full paper: <u>http://short.pfonline.com/NASF21Jan2</u>.

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