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UPDATE ON THE DEVELOPMENT OF PLATINUM-BASED ALLOYS FOR POTENTIAL HIGH-TEMPERATURE APPLICATIONS

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Abstract

Pt-based alloys for high-temperature applications in aggressive environments have been under development for over 10 years, and are targeted to be used as either as bulk, or as coatings. The alloys comprise Pt, Al, Cr, and Ru, and the microstructure has been improved by composition to a best possible analogue of the nickel-based superalloys, which these alloys could partially replace. This was necessary because the previous best alloy had a strengthening precipitate volume proportion of only approximately 40 vol.%, whereas the nickel-based superalloys have around 70 per cent, and a lower volume would mean that the strength would not be the best that could be obtained. The microstructures were assessed using electron microscopy, and have been related to the alloys' hardness values. The current microstructures are much more like those of NBSAs, with a high proportion of the strengthening $^{\rm Pt_3Al}$ precipitates. Since the samples have to be small (platinum is expensive), hardness has been used as an indication of strength. Nano-indentation studies showed that the hardness and Young's modulus were higher for $^{\rm Pt_3Al}$ than the (Pt) matrix.

More extensive oxidation studies have been undertaken on the previous optimum sample, and the effect of cooling rate after heat treatment has also been ascertained. In addition, the samples were studied after different heat treatment times, and cross-sections were made in order to characterize the alumina scale that formed. The oxide scales of Pt-11Al-3Cr-2Ru (at.%) up to 100 h exposure did not spall, and were at least as good as those of the ternary alloys. There was no discernable Al depletion zone in the substrate, although it could have been at a greater depth than thickness of the samples studied.

Further additions to the Pt-Al-Cr-Ru alloys have been studied, to increase the melting temperature as well as to reduce the platinum content without compromising the properties. Reduced platinum content would have the benefits of reduced density and cost. The target elements include vanadium and niobium, of which vanadium is of special interest, since it is a South African product. Before these additions could be made, phase equilibria studies were undertaken so that the maximum addition could be ascertained, and also to check any possible ternary phases, which could be deleterious. Work on the Pt-Al-V system revealed a ternary phase, $V_{27}Pt_{54}Al_{19}$ (at.%), which has a eutectic reaction with (Pt). The maximum V addition is likely to be around 15 at.%, otherwise Pt_3V or the ternary phase will form, and not the required (Pt)/ Pt_3Al phases.

Introduction

The Pt-based project has been running for many years, and was initially funded by the Platinum Development Initiative which comprised Anglo American Platinum, Lonmin, andImpala Platinum, and two LEAD grants from the Government¹⁻⁷. Most of the work was done either at Mintek or the University of the Witwatersrand, with involvement from other South African universities such as the universities of Cape Town, Limpopo, and the Free State.

There were also parallel studies at Fachhochschule Jena and the University of Bayreuth, Germany⁸⁻¹², with some collaborations¹³⁻¹⁵, as well as with the National Institute for Materials Science (NIMS), Japan¹⁶⁻¹⁷.

The rationale behind the work is that Ni-based superalloys (NBSAs) are reaching their temperature limit, and despite improvements with the higher-generation alloys, the improvements are slowing down, and the last 24 years have produced an increase of approximately 100°C¹⁸. There have already been improvements with design, using forced aircooling, and huge benefits from using coatings, which have an insulating outer layer. Both of these techniques are expensive and can have disastrous effects if they fail, for example, if the coating fails locally. Thus, there has been a push to develop a totally new suite of materials, and platinum-based alloys are one possible solution. Platinum was selected because it has the same fcc structure as Ni, but a higher melting point¹⁹, similar chemistry, and better oxidation and corrosion resistance²⁰. The earlier work has already shown that the same microstructure as the NBSAs could be attained, although the volume of the strengthening precipitate was much less¹⁻⁷, and that the corrosion resistance was much better²¹. The work soon identified Pt-Al as the best system both for beneficial microstructures and for formation of the protecting alumina film, and ternary additions were studied. The best ternary alloys were Pt-Al-Cr and Pt-Al-Ru, and these were then combined^{5,22}. The target temperature of the Pt-based alloys is 1300°C, which is about 200°C higher than the currently-used NBSAs, and the optimum composition was Pt-11Al-3Cr-2Ru (at.%). The effect of heat treatment was studied³⁻⁷. However, the problems of the high expense and density remained.

In 2007, the study split, with work on coatings ongoing at Mintek²³, and the more fundamental investigations continuing at the University of the Witwatersrand. This is an update on that work, and shows that there has been progress on both alloying to potentially reduce the proportion of expensive and dense platinum, and improving the microstructure to increase the proportion of the strengthening precipitates. Also, more work has been done on the oxidation of the alloys at high temperature, showing that the platinum-based alloys have good potential in aggressive environments through the formation of a stable and tenacious alumina film.

Improving the microstructure

The NBSAs obtain their strength from the high proportion of $\sim Ni_3Al$ precipitates in a (Ni) matrix. The precipitates have an ordered fcc structure and so the interface between the precipitates and the matrix has low interfacial energy, meaning that the driving force for coarsening is low, and hence the fine structure can be retained to relatively high temperatures⁴.

The Pt-based alloys have also been shown to have nearly the same structure, although there is a complication, because unlike in the Ni-Al system, the Pt-Al system has at least three different structures for the ^{Pt_3}Al precipitates²⁵⁻²⁶, and the cubic high-temperature form is the preferred one. It can be stabilized by additions of Ti, Cr, and Ta^{16, 27}.

The worst problem was that all the studies indicated that despite attempted to optimize the eat treatment, the highest precipitate proportion attained was around 30 per cent. Although the mechanical properties were promising²⁸⁻³⁰, obviously with such a small proportion of the strengthening precipitates, the alloys would not achieve their potential strength. The ruthenium additions were for both increased corrosion resistance as well as for solid-solution strengthening.

Studies of six alloy compositions in the as-cast and heat-treated states, and hardness tests, identified a new optimum composition: Pt-11Al-6Cr-5Ru (at.%), on the basis of a higher volume of the fine precipitates and increased hardness³¹⁻³³. The casting was done in a button-arc melter under an argon atmosphere, with titanium used as an oxygen getter, and the heat treatment conditions were taken from the work on Pt-Al-Cr-Ni alloys^{8, 13}: heat treated in air at 1500°C for 18 hours, then water-quenched, followed by 1100°C for 120 hours and air cooling. Furthermore, this particular heat treatment is more similar to current industrial practices for NBSAs, compared to the much longer times used in earlier work¹⁻⁷. A study of the misfit, obtained from XRD, showed that the degree of misfit was related to shape of the precipitates³⁴, just as in NBSAs²⁴.

Using alloys around the new optimum composition of Pt-11Al-6Cr-5Ru (at.%), samples of approximately 3 g were manufactured by button-arc melting, then heat treated as above. Sections of approximately 250 μ m were cut from the centre of the button, and 3 μ m diameter cores were cut from these and then prepared as transmission electron microscope (TEM) samples³⁵, with the final thinning being done on a Gatan 691 Precision Ion Polishing System (PIPS), with a final milling done at a lower energy and for a short time to reduce the ion damage. However, the ion damage was not sufficiently removed, and the method of Witcomb had to be utilized³⁶. The samples were immersed in a solution made up of equal amounts of phosphoric, nitric and sulphuric acids at 2.8V AC at 20°C, with the sample positioned between two Pt electrodes. The cleaning time was about 30s³⁶, and afterwards, the microstructures of ~Pt₃Al precipitates in the (Pt) matrix were clearly seen.

The microstructure had much higher proportions of the 2 Pt₃Al precipitates³⁵ than in earlier studies^{1-7, 8, 13}, and were comparable to the NBSAs (Figure 1). The average size of the precipitates was 200 nm. The morphology of the current precipitates was more cubic, and thus more similar to NBSAs than those in earlier work. The matrix was less easily removed and protruded more, showing that it was more resilient to the thinning. Electron diffraction patterns were obtained from several areas of the precipitates and the matrix by tilting, and the angles and distances were measured between the spots. The results were compared to the XRD results from the same samples to deduce the structures^{35, 37-39}, by comparing different sets of *hkl* values. The different comparisons showed that the precipitates were more likely to be cubic than tetragonal, although some precipitates were more difficult to identify conclusively.



Figure 1–TEM micrograph for nominal Pt-12Al-Cr4-Ru2 (at.%) showing ~Pt₃Al (marked A) with channels of (Pt) matrix [35]

The nanohardness and the elastic (Young's) modulus were analysed on the separate phases of nominal Pt-12Al-Cr4-Ru2 (at.%) using a CSM nano-indentor with a maximum load of 2.5 mN. Ten individual indentations were averaged for determining the hardness and modulus of elasticity of the matrix and precipitates. The nanohardenss results were 11.4 \pm 0.9 GPa for ~Pt₃Al, and 7.3 \pm 0.6 GPa for (Pt). Similarly, the elastic (Young's) modulus values were 259.7 \pm 23.7 for ~Pt₃Al and 233.3 \pm 21.8 for (Pt). The ~Pt₃Al values were higher than the matrix, as expected. Durst *et al.*⁴⁰ performed nanoindentation on the third-generation CMSX-10 NBSAs, and found the nanohardness to be 9.4 GPa for the γ' -precipitates and 7.2 GPa for the matrix, with an elastic modulus of 255 GPa for γ' and 231 GPa for γ . The γ' -phase in CMSX-10 is softer compared to that of Pt-12Al-Cr4-Ru2, while the matrix nanohardnesses are similar. Similar elastic modulus values were observed for both phases in CMSX-10 and Pt-12Al-Cr4-Ru2. The γ' and γ nanohardnesses of Pt-12Al-Cr4-Ru2 were similar to that of Pt-12Al-Gcr-5Ni-2Re⁴¹, which were 11.5 GPa for γ' and 7.5 GPa for γ . The elastic modulus for γ' was slightly lower compared to the 282 GPa of Pt-12Al-6Cr-5Ni-2Re, while the γ -phase elastic modulus of Pt-12Al-Cr4-Ru2 was slightly higher than that of Pt-12Al-6Cr-5Ni-2Re.

Oxidation studies

Obviously, if alloys are targeted to be utilized in a fuel-burning environment, the reactions with oxygen and other deleterious components in the fuel have to be known.

Earlier studies had already shown the Pt-based alloy's superior corrosion resistance over NBSAs²¹, as well as good alumina scale-forming capabilities of the Pt-Al-Cr and Pt-Al-Ru ternary alloys between 1150°C and 1350°C⁴²⁻⁴³, after the formation of a discontinuous oxide layer during an initial transient stage of up to 10 hours. A continuous α -Al₂O₃ scale with no zone of discontinuous oxide or any internal oxidation was formed on Pt-14Al-3Cr-3Ru isothermally oxidized in air at 1350°C up to 500 hours⁴⁴, but there were concerns that the scale formed too quickly, which might not be sustainable in long-term service. Wenderoth *et al.*¹² also observed the parabolic formation of alumina scale on a Pt-12Al-6Cr-5Ni (at.%) alloy when isothermally oxidized in air at temperature between 1100°C and 1300°C for up to 400 hours, and the Al depletion zone in the substrate increased with oxidation time and temperature. A more indepth study was necessary to study how the alumina formed over time.

As-cast (button-arc melted) Pt-11Al-3Cr-2Ru (at.%) samples were heat-treated at 1350°C for 96 hours in a muffle furnace, then cooled in either water or air⁴⁵. Approximately 1 mm thick slices were cut and prepared metallographically on both sides to a 1 μ m diamond finish. Isothermal oxidation tests were done in air at 1350°C for up to 100 hours. Specimens were removed after specific time intervals from 1 hour to 100 hours, and weighed after cooling. Specimens were prepared metallographically for cross-sectional examination using a SEM⁴⁵.

The specific mass gain for each specimen in both quenching media increased parabolically (Figure 2) with exposure time, with the oxidation rate of the air-cooled specimens increasing faster⁴⁵. There was no mass loss over the entire 100 hours' exposure at 1350°C, indicating good oxidation resistance, and no scale spallation. Although the higher oxidation rate of the aircooled samples could be due partially to the additional oxidation that occurred while the samples were cooling, there is also the effect of available aluminium. The scale on the aircooled specimens had potentially better adhesion than that on the water-quenched specimens, based on deeper protrusions of the scales into the substrate. This was a result of faster scale growth rates, leading to increased scale thicknesses. Both the mass- and thickness-related parabolic constants of the air-cooled specimens had higher scale growth rate constants than those of Süss et al.⁴³, which may be due to a slightly higher Al content of the quaternary alloy, as well as beneficial effects of other alloying elements. Vorberg et al.⁹ showed that Pt-12.5Al-3Cr-6Ni (at.%) quenched in water after heat-treatment at 1500°C contained small y' precipitates with a volume fraction of about 3 per cent, whereas air-cooled samples were found to contain evenly-distributed cuboid-shaped γ' precipitates, with a volume fraction of approximately 30 per cent. The faster scale growth kinetics of the air-cooled specimens may be due to a higher volume fraction of γ' in the as-cast alloy.

Here, the microstructure of the air-cooled specimens could not be resolved, but the Al content waslower, which would be expected to result in a lower volume fraction of γ' . In both studies, the air-cooled specimens had thicker scales. One reason for thicker scale could be more free Al, but this was not true for the current alloys.

Another reason could be more γ' (~Pt₃Al) precipitates, which would give more available Al for the scale. Vorberg *et al.*⁹ reported the depletion of Al near the surface was due to the depletion of γ' precipitates.

This showed that the Al in the scale originated from the γ' precipitates, so it is likely that the aircooled specimens had more γ' precipitates, especially since Pt-7Al-5Cr-3Ru (at.%)⁴ had approximately 30 per cent precipitates by volume. These alloys have four components, so the proportion of γ' (~Pt₃Al) precipitates might not be directly proportional to the Al content, since Al is present in both ~Pt₃Al and (Pt) matrix. The faster scale kinetics of the air-cooled specimens, resulting in thicker scales, might have resulted from more ~Pt₃Al in those samples. Mass-parabolic rate constants, k_p , for both the air-and water-quenched samples were derived from the classical parabolic equation⁴⁶ $\Delta m^2 = k_p t + C$ (where Δm = specific mass gain, t = exposure time, and *C* is the intercept from the graph). The values were compared with those obtained from Pt-10Al-4Cr and Pt-10Al-4Ru alloys⁴³ in Table I.



Figure 2–Specific mass gain for water-quenched and air-cooled Pt-11Al-3Cr-2Ru (at.%) specimens after isothermal oxidation in air at 1350°C [45]

Composition (at.%)	Exposure range (h)	<i>k_p</i> (mg ² .cm ⁻ ⁴ .h)	Behaviour	<i>k_{ps}</i> (μm ² .h)	Behaviour
Pt-10Al-4Cr ⁴³	1-100	0.005	Parabolic	1.3467	Parabolic
Pt-10Al-4Ru ⁴³	1-100	0.006	Linear	0.7159 *	Parabolic
Pt-11Al-3Cr-2Ru (w)	1-100	0.006 ± 0.0004	Parabolic	1.55 ± 0.01	Parabolic
Pt-11Al-3Cr-2Ru (a)	1-100	0.1 ± 0.0014	Parabolic	2.10 ± 0.007	Parabolic

Table I-Properties of the different alloys (w = water quenched; a = air quenched; * after 1000 h)-⁴⁵

The surface oxide grain size increased with oxidation time (Figure 3), with the pores becoming less discernable, as more and larger flakes formed⁴⁵. The pores were deduced to arise from the way the oxide grains grew together unevenly. Oxide grains of approximately 1.63 \pm 0.6 μ m had grown after 100 hours. The oxide grains in the air-cooled samples were more rounded and joined (Figure 4).



Figure 3–SEM-BSE images of the surface of water-quenched Pt-11Al-3Cr-2Ru (at.%) specimens after isothermal oxidation in air at 1350°C for: (a) 1 h, (b) 10 h, (c) 50 h and (d) 100 h⁴⁵



Figure 4–SEM-SE image of the surface of air-cooled Pt-11Al-3Cr-2Ru (at.%) specimen after 100 h oxidation in air at 1350°C⁴⁵

The SEM-SE cross-sections (Figure 5) showed that the scales after oxidation at 1350°C were continuous with limited porosity⁴⁵. X-ray elemental mapping of the scale after 100 hours' oxidation showed that the scale comprised mainly Al and O, which was confirmed by XRD. Scale thickness increased with oxidation time for both quenching media. Air-cooled specimens formed thicker scales than the water-quenched specimens, although high errors showed the differences were minor. The scales were also non-uniform in thickness, with irregular scale-substrate and scale-gas interfaces, which had no apparent relationship to each other (Figure 5). No zones of discontinuous oxides or any internal oxidation were observed, unlike similar alloys¹⁷. Beneficially, there was no void formation at the scale-alloy substrate interface, indicating that the scales were potentially protective.

The ridges or intrusions of oxide scale increased with exposure time. The oxide scales were also observed to protrude outwards at the scale-gas interface. The presence of substrate in the scale was deduced to be due to continuous lateral growth of the oxide after protrusion.



Figure 5–SEM-BSE images of cross sections of Pt-11Al-3Cr-2Ru (at.%) specimens after 100 h oxidation in air at 1350°C (a) water-quenched (b) air-cooled⁴⁵

The irregular oxide-alloy interfaces improved the adhesion of the scale, due partially to mechanical keying⁴⁷, although deleterious cavity formation at ridges associated with Pt-Al grain boundaries has been reported⁴⁸⁻⁴⁹. The high Pt content of the quaternary alloy may have improved the scale adherence, due to various mechanisms discussed by Reed *et al.*⁵⁰.

Future work to be reported will include the measurement of stress in the scales, using Raman spectroscopy. The stress is important because it is one of the determining factors for the durability of the scales.

Alloying with vanadium

Although platinum has many beneficial properties, its high price and density are disadvantageous, and if the same structure and reasonable chemical properties could be retained, alloying could reduce the Pt content and hence the cost and density. A beneficial addition is vanadium which has a solubility in (Pt) of up to approximately 20.5 at.% V at 700°C which increases with increasing temperature, reaching approximately 7 at.% V at 1720°C⁵¹. Thus, as well as decreasing cost and density, V has potential both as a solid-solution strengthener and to increase the melting temperature¹⁹. Except for this ongoing study⁵²⁻⁵⁴, there is no reported data for Pt-Al-V, and phase diagram work on the system was necessary to check the extents of the phases and existence of any ternary phases that might be detrimental. Knowledge of the phases will help in defining the compositions of developmental alloys in the Pt-Al-Cr-Ru-V system.

Nine different Pt-Al-V composition alloys were manufactured by arc-melting under argon, with a titanium oxygen-getter, and sectioned. One half was prepared metallographically, then characterized using SEM with EDX, with XRD used for phase confirmation. The second half was annealed and will be reported in a later paper.

As expected, the alloys had very different microstructures, even from those reported earlier⁵⁴. For example, Alloy 1⁵⁴, average overall composition Pt-27Al-9V (at.%), had ~Pt₃Al dendrites, ~Pt₅Al₃ needles, a thin ~Pt₃V layer on the dendrites, and a complex eutectic-like microstructure. This consisted of ~PtAl + Pt₅Al₃ + ~Pt₅Al₃, and resulted from the eutectic reaction L $\rightarrow \beta$ + ~Pt₅Al₃ followed by the eutectoid decomposition $\beta \rightarrow$ ~PtAl + ~Pt₅Al₃. Some of the later alloys had a ternary phase (Figure 6). An interesting example was Alloy 13, average overall composition Pt-9Al-38V (at.%), which had ~PtV dendrites in a sparse eutectic comprising ~V₃₀Pt₅₆Al₁₄ (a ternary phase) + ~PtV (Figure 7).



A solidification projection of all the samples to date was plotted (Figure 7), and the liquidus surface⁵⁴ was extended to show the liquidus surface for the ternary phase. These figures are very important because they identified the likely limit of the addition of V at which the (Pt)/~Pt₃Al microstructure will be retained. Thus, Figure 8 shows that the maximum V addition is probably around 15 at.%, otherwise the ~Pt₃V phase will be obtained, and the liquidus⁵⁴ shows a slightly lower value. A series of quinary alloys based on Pt-12Al-Cr4-Ru2 (at.%) have been manufactured, and the as-cast alloys show mainly dendrites with little or no eutectic, which agrees with the liquidus surface⁵⁴. Additionally, the ternary phase needs to be avoided, but this phase only appears at V contents higher than 17 at.%.

Although the structure has yet to be determined, it is likely to be brittle. The next stage is to conduct hardness tests on the remaining alloys, and also to heat treat and characterize all the Pt-Al-Cr-Ru-V alloys to ascertain the effect of different amounts of vanadium.



Figure 8–Solidification projection of the Pt-Al-V system above 50 at.% Pt

Conclusions

Ongoing work on the Pt-based alloys has shown that microstructures much more like those of the NBSAs can be obtained with comparably high proportions of the strengthening Pt_3Al precipitates, and the majority were the preferred cubic allotrope. Nano-indentation studies confirmed the higher hardness and Young's modulus of the Pt_3Al compared to the (Pt) matrix. Recent work has shown that the oxide scale of Pt-11Al-3Cr-2Ru (at.%) up to 100 hours' exposure has potential for protection, and did not spall. There was no discernable Al depletion zone in the substrate, probably because it was greater than the depths of samples studied. The scales were at least as good as those of the ternary alloys.

Phase diagram work on the Pt-Al-V system has shown that there is a ternary phase of ${}^{\sim}V_{27}Pt_{54}Al_{19}$ (at.%) composition which has a eutectic reaction with (Pt). The maximum useful V addition is likely to be around 15 at.%, otherwise ${}^{\sim}Pt_{3}V$ or the ternary phase will form, and not the required (Pt)/ ${}^{\sim}Pt_{3}Al$ phases.

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