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As cast and heat-treated alloys of the Pt-Al-V system at the Pt-rich corner

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Synopsis

The Pt-based alloys have a high potential for replacing some Ni-based superalloys (NBSAs) used in the highest temperature and most aggressive environments, and vanadium would be a beneficial alloying element.

Six alloys of average compositions Pt-26.6Al-9.1V, Pt-23.1Al-17.8V, Pt-9.6Al-21.1V, Pt-17.4Al-6.4V, Pt-22.3Al-7.9V, and Pt-6.0Al-10.1V (all in at.%) were manufactured by arc-melting and examined in the ascast condition using scanning electron microscopy and X-ray diffraction (XRD). Five samples were heat treated at 1 000°C for 1 500 hours and water quenched, then examined. The phases were identified using energy-dispersive X-ray spectroscopy and the identities were confirmed by XRD. In most of the samples, the phases existing in the as-cast condition were different from those at 1 000°C after ~1500 hours heat treatment. A solidification projection, an isothermal section and a liquidus surface projection were determined for the Pt-rich corner.

Keywords

Phase diagram, Pt-Al-V system, scanning electron microscopy, X-ray diffraction.

Introduction

Nickel-based superalloys (NBSAs) have been successfully used in turbine components since the late 1940s. Approximately 70 percent of the weight of a modern jet turbine comprises NBSAs, and their success is due to their high vield stress and excellent resistance to environmental attack at high temperatures¹. Although the NBSAs have excellent properties, they are hindered by their maximum application temperature, which is limited by the melting point of the nickel solid-solution matrix. Currently, the maximum temperature at which NBSAs operate is ~1 100°C, which is approximately 85 percent of their melting temperature¹. If the operating temperatures could be increased, there would be a number of advantages. Higher temperatures improve the efficiency of the turbine engines, and this enables greater thrust, improved fuel efficiency, and reduced pollution. Although thermal-barrier coatings can be used to increase the application temperature, the component is still restricted by the melting point of the substrate, so the maximum attainable temperatures are still limited if catastropic failure is to be avoided².

There is increasing interest in using a different alloy system with a much higher melting point. Ceramics and selected intermetallic phases have been discussed because of their high melting points and high strength; however, both suffer from reduced toughness compared with NBSAs, and their processing costs are high2. One solution would be to base the new materials on alloys with high melting points and use the naturally occurring precipitates of that system. This should also help keep the processing cost to a minimum. Thus, the systems developed would be similar in structure to the NBSAs, with a matrix and a fine dispersion of small, preferably coherent and stable precipitates. Face-centered cubic (fcc) structures (like nickel) are advantageous because, being closepacked, they are more creep resistant. Refractory metals (niobium, molybdenum, and tungsten) have been considered because of their high melting points (2 469°C, 2 623°C, and 3 020°C respectively), but their more open body-centred cubic (bcc) structures are more susceptible to creep. Additionally, they are prone to rapid oxidation, even at relatively low temperatures³. Platinum-group metals (platinum, iridium, rhodium, and palladium) were targeted for investigation because, with the exception of palladium, they have high melting points, good environmental resistance, and a mostly fcc structure4.

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Pt is similar to Ni in crystal structure (fcc) and chemistry. The important differences are the higher melting point (1769°C for Pt compared to 1 455°C for Ni) and improved corrosion resistance⁵. These facts inspired Wolff's and Hill's research into the development of Pt-based analogues to Ni-based superalloys that could potentially serve in the most critical and demanding of the high-temperature applications⁴. Thus, phases similar to $L1_2$ Ni₃Al could be used to obtain properties similar to those found in the NBSAs. Although platinum-based alloys are unlikely to replace all NBSAs on account of both higher price and density, it is likely that they can be used for the highest application temperature components.

Alloys based on Pt-Al have the highest potential, not only because of possible precipitation strengthening through Pt₃Al, but also due to high oxidation and corrosion resistance. The resistance to oxidation and corrosion is attributed to the formation of a protective Al oxide scale^{6,7}. At least 11 at.% Al is needed to form a thin protective oxide coating on the ternary and quartenary alloys at high temperatures⁸. It is generally accepted that the L1₂ structure of the γ' phase is a prerequisite for the success of the Ni-based superalloys. Due to the close crystallographic relationship, precipitates with ordered fcc, L1₂ have coherent interfaces with the surrounding fcc matrix. Coherent γ/γ' interfaces with small lattice misfit lead to low interfacial energies, thus prohibiting fast coarsening of γ'^8 .

Under the auspices of the Platinum Development Initiative, platinum-based superalloys were investigated for high-temperature and special applications requiring good corrosion and oxidation resistance⁹. Ternary alloys based on Pt-Al, where the ternary additions comprised chromium, iridium, molybdenum, nickel, rhenium, ruthenium, tantalum, titanium, and tungsten were tested. As well as phase characterization work, mechanical and oxidation tests were undertaken. The best systems were found to be Pt-Al-Cr and Pt-Al-Ru. The microstructures were similar to NBSAs, and comprised ~Pt₃Al precipitates in a Pt-based matrix. However, the volume fraction of ~Pt₃Al was only ~40% instead of the ~70% found in the NBSAs⁹.

Chromium was found to stabilize the cubic form of the ~Pt₃Al phase, whereas ruthenium acted as a solid-solution strengthener¹⁰. In order to exploit the additional benefits from the two ternary additions (Cr and Ru), quaternary alloys were made containing both. A number of these quaternary alloys have been investigated extensively. They exhibited a two-phase γ/γ' structures analogous to the NBSAs ¹¹.

Much work has been done on the quarternary alloys of the Pt-Al-Cr-Ru system in an attempt to optimize the composition with respect to the precipitate volume fraction and hardness. The Vickers hardness of the Pt-Al-Cr-Ru alloys within the composition ranges selected was relatively independent of the chemical composition and the number of annealing stages and fell within the range HV₁₀ ~400 to ~430. The volume fraction of the precipitate, using image analysis, was estimated at approximately 25 to 30%¹².

There is a need to increase the hardness and consequently the strength, melting temperature, and volume fraction of the precipitates. It is also desirable to reduce density and cost. Although vanadium may not meet all these requirements, it is expected to reduce cost and density by replacing some Pt in a chosen Pt-Al-Cr-Ru alloy. It is also expected to act as a solid

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solution strengthener by going into solution in (Pt), as well as increase the melting temperature. The binary phase diagram of Pt-V shows that ~20.5% V will go into solution in (Pt) at 700°C, increasing to ~57% at 1 720°C, and the melting temperature of (Pt) has a maximum of ~1 805°C¹³. Addition of Nb, Ta, and Ti to Pt-based alloys resulted in elevated strengths at high temperatures through precipitation-hardening. The polycrystalline Pt-based alloys containing 5 at.% Nb or Ta are stronger than the single-crystal NBSA CMSX-4 in (001) orientation above 1 200°C¹⁴. V is near Nb, Ta, and Ti in the periodic table and is smaller in size compared to Nb and Ta. Therefore, V may act as a precipitation-hardener in addition to solid-solution strengthening.

Experimental procedure

Six alloy buttons (Samples 1 to 6) weighing ~2 g each were prepared from Pt and Al of 99.9 percent purity and V of 99.6 percent purity. Each of the six alloys is named with respect to its average overall composition in at.% in the as-cast condition. The samples were manufactured by arc-melting under argon in a water-cooled copper hearth. Each button was turned over and re-melted three times to ensure thorough mixing of the elements.

Each of the samples was halved using a ceramic blade mounted in a Struers Secotom-10 cutting machine. The spindle speed was 3 500 r/min and a feed of 0.5 m/s was used. One half were prepared for metallographic examination, in the as-cast condition, by grinding on silicon carbide down to 1 200 grit, diamond polishing down to 1 μ m and then final polishing by an oxide polishing system (OP-S).

Five of the other halves (samples 1H, 2H, 3H, 5H and 6H, where the number refers to the original as-cast sample and 'H' refers to the 1 000°C heat treatment) were placed in silica glass tubes 16 mm in diameter. The glass tubes were then made into vacuum ampoules by evacuating air and sealing while the vacuum pump was running. The ampoules with the samples were placed in a tube furnace and annealed at 1 000°C for 1 500 hours. Quenching was done by removing the glass ampoules from the furnace, dropping them in water and breaking the glass, as quickly as possible in order to retain the phases existing at 1 000°C. The samples were then prepared for metallographic examination in the same way as the as-cast samples.

The microstructures of all the alloys were analysed using a scanning electron microscope (SEM) model HR-NovaNano SEM200. Imaging in all the analyses was done using the backscattered electron (BSE) mode, while overall and phase compositions were determined by energy dispersive X-ray spectroscopy (EDX). Often it was difficult to discern between some of the phases, even in BSE mode because of the similarity in average atomic number of these phases. There was also the concern that orientation and different electron channelling would also change the contrast¹⁵, therefore the morphology of the phases was also used to differentiate them. Since the interaction volume generating the X-rays can be as much as 3 µm across and deep (especially for higher atomic number elements), the analyses for smaller regions than this are an indication only. Usually, an accuracy of ± 1 at.% would be expected for this technique. Spot analyses were used on single phases and areal analyses on two-phase regions. The compositions of each phase and overall

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compositions were determined by taking EDX readings from at least five different areas and reporting the average and the standard deviation. X-ray diffraction (XRD) was used to confirm the phase identities.

Results

As-cast alloys

As-cast Alloy 1, average overall composition Pt-26.6Al-9.1V (at.%)

Figure 1 shows the microstructure of the as-cast Alloy 1. It consists of medium grey dendrites surrounded by a dark thin layer, light needles, and a eutectic comprising the light phase and a barely resolveable two-phase dark region. The compositions of these phases are shown in Table I. The medium grey dendrites were identified as ~Pt₃Al and the light needles as ~Pt₅Al₃. The thin dark layer on the dendrites was identified as ~Pt₃V. XRD confirmed the identities of these phases. Solidification started with the ~Pt₃Al dendrites, followed by peritectic formation of ~Pt₅Al₃ which then solidified independently as needles. The ~Pt₃Al dendrites and the ~Pt₅Al₃ needles reacted peritectoidally to form ~Pt₂Al. The next reaction was eutectic, forming β + ~Pt₅Al₃ and the β decomposed eutectoidally to form \sim PtAl + \sim Pt₅Al₃¹³. The final reaction was peritectoid formation of a thin layer of ~Pt₃V between the ~Pt₃Al dendrites and the ~Pt₅Al₃ needles.



Figure 1—SEM-BSE image of Alloy Pt-26.6Al-9.1V (at.%) showing medium grey dendrites (~Pt₃Al) with a dark thin layer (~Pt₃V), light needles (~Pt₅Al₃), and a eutectic. Scale bar is 30 μ m

As-cast Alloy 2, average overall composition Pt-23.1Al-17.8V (at.%)

Figure 2 shows the microstructure of the as-cast Alloy 2 at different magnifications. It consisted of medium grey dendrites surrounded by a light phase, also appearing dendritic, and a complex eutectic. When the compositions of these phases were plotted (Table II), the medium grey dendrites were identified as ~Pt₂V, the light phase as ~Pt₅Al₃, while the dark component of the eutectic was identified as PtAl, although the areas are likely to be affected by the nearby ~Pt₂V and ~Pt₅Al₃. XRD confirmed the identities of ~Pt₂V, ~Pt₅Al₃, and PtAl. Solidification started with (Pt). This was followed by a peritectic reaction between the (Pt) and the liquid, resulting in ~Pt₅Al. There then followed a eutectic reaction with the liquid forming $\sim Pt_5Al_3$ and β , with β decomposing at high temperature to form $PtAl + -Pt_5Al_3^{13}$. The (Pt) dendrites ordered to $-Pt_2V$ (identified by unique peaks of those phases amongst the common peaks). There was also a solid-state cellular precipitation of $\sim Pt_2V / (Pt)$ in the ~Pt₅Al₃ phase, and these phases were too fine to analyse using SEM-EDX.

As-cast Alloy 3, average overall composition Pt-9.6Al-21.1V (at.%)

Figure 3 shows the microstructure of as-cast Alloy 3, average overall composition Pt-9.6Al-21.1V (at.%), at two different magnifications. It consists of dendrites, a light phase, and a dark phase between the dendrites. When the compositions of the phases in Table III were plotted, the dendrites were identified as ~Pt₃V, and the light phase as ~Pt₃Al. The dark phase was identified as ~Pt₂Al, which formed peritectoidally at high temperature from ~Pt₅Al₃ and ~Pt₃Al. Here, any ~Pt₅Al₃ that formed was consumed. Solidification started with (Pt), followed by a weak peritectic reaction between the liquid and (Pt) resulting in a sparse eutectic with the dark component ~Pt₂Al in the light ~Pt₃Al. The final reaction was (Pt) ordering to ~Pt₃V. XRD confirmed the phase identifications.

As-cast Alloy 4, average overall composition Pt-17.4Al-6.4V (at.%)

Figure 4 shows that the as-cast Alloy 4, average overall composition Pt-17.4Al-6.4V (at.%), consisted of a single phase having grains at different orientations. The gain was increased to enhance the contrast. The compositions in Table IV are for different grains i.e. orientations, and hence different contrast. The plot of these compositions confirmed

Overall and phase compositions of Alloy 1, average overall composition Pt-26.6Al-9.1V (at.%) as determined by EDX

Component	Pt	AI	v	Phases
Overall	64.3±0.7	26.6±0.8	9.1±0.4	-
Medium dendrites	67.4±0.6	18.7±0.6	14.0±0.5	~Pt ₃ Al
Dark	65.3±0.4	29.4±0.8	5.3±0.5	~Pt ₂ Al
Light needles	59.0±1.2	30.7±1.1	10.3±0.4	~Pt ₅ Al ₃
Thin dark layer on dendrites	65.5±0.9	13.6±1.1	20.9±1.2	~Pt ₃ V
Eutectic overall	63.0±0.3	29.1±0.7	8.0±0.5	\sim PtAI + \sim Pt ₅ AI ₃

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Figure 2—(a) SEM-BSE image of Alloy 2 showing medium grey dendrites (\sim Pt2V), a light phase \sim Pt₅Al₃), and a eutectic (β + \sim Pt₅Al₃). Scale bar is 30 µm. (b) SEM-BSE image of Alloy 2 showing medium grey dendrites (\sim Pt₂V), a light phase (\sim Pt₅Al₃) and solid state cellular precipitation. Scale bar is 5 µm

Table II					
Overall and phase compositions of Alloy 2, average overall composition $Pt_{59.1}$:Al _{23.1} :V _{17.8} (at.%) as determined by EDX					
Component	Pt	AI	v	Phases	
Overall	59.1±0.3	23.1±0.4	17.8±0.1	-	
Medium dendrites	60.5±0.4	8.3±0.4	31.2±0.4	~Pt ₂ V	
Light phase	62.0±0.3	32.3±0.3	5.7±0.5	~Pt ₅ Al ₃	
Eutectic	58.2±0.4	25.7±1.2	16.1±1.0	$\sim Pt_5Al_3 + \beta$	
Dark eutectic component (2-phase)	54.4±1.7	32.8±1.9	12.9±0.9	PtAI + ~Pt ₅ Al ₃	



Figure 3—(a) SEM-BSE image of as-cast Alloy Pt-9.6AI-21.1V (at.%) showing dendrites (~Pt₃V) and a light phase (~Pt₃AI). Scale bar is 50 µm. (b) SEM-BSE image of as-cast Alloy Pt-9.6AI-21.1V (at.%) dendrites (~Pt₃V), a light phase (~Pt₃AI) and a dark phase (~Pt₂AI) between the dendrites. Scale bar is 10 µm

Table III Overall and phase compositions of Alloy 3, average overall composition Pt-9.6AI-21.1V (at.%) as determined by EDX

Component	Pt	AI	v	Phases	
Overall	69.3±1.2	9.6±1.0	21.1±1.1	-	
Dendrites	69.7±1.3	4.9±1.3	25.2±0.6	~Pt ₃ V	
Light phase	69.8±0.4	25.5±0.6	4.7±0.6	~Pt ₃ Al	
Dark	62.8±1.3	27.9±1.2	9.3±0.4	~Pt ₂ Al	



Figure 4—SEM-BSE image of as-cast Alloy Pt_{76.2}Al_{17.4}V_{6.4} (at.%) showing a single phase (~Pt_3Al) with grains at different orientations. Scale bar is 400 μm

that the alloy was single phase ~Pt₃Al, and XRD also confirmed the phase identity.

As-cast Alloy 5, average overall composition Pt-22.3Al-7.9V (at.%)

Figure 5 is a SEM-BSE image of the microstructure of as-cast Alloy 5, showing light- and medium-contrast dendrites with a thin dark contrast phase. Since the average atomic numbers of the phases were similar, the contrast between them was low, giving a poor image. Both the light and medium grey phases were identified as \Pt_3Al , from the close compositions and the same dendritic morphology, while the thin dark grey phase was identified as \Pt_2Al from the composition. Solidification started with \Pt_3Al , followed by a peritectic formation of a thin layer of \Pt_5Al_3 which subsequently



Figure 5—SEM-BSE image of as-cast Alloy Pt-22.3Al-7.9V showing a light grey phase (~Pt₃Al), a medium grey phase (~Pt₃Al) and a thin dark grey phase (~Pt₂Al). Scale bar is 30 μ m

reacted peritectoidally with ~Pt₃Al, to form ~Pt₂Al.

As-cast Alloy 6, average overall composition Pt-6.0Al-10.1V (at.%)

Figure 6 shows the microstructure of as-cast Alloy 6, average overall composition Pt-6.0Al-10.1V (at.%), with a medium-contrast phase which was identified as (Pt) and a fine eutectic. The existence of the eutectic was confirmed after it coarsened during heat treatment for 1 500 hours at 1 000°C (described later). The eutectic and its components were too fine to be analysed accurately by EDX. Solidification started with (Pt), followed by the eutectic reaction producing (Pt) and ~Pt₃Al. XRD confirmed the presence of (Pt) and ~Pt₃Al (Figure 7).

Table IV					
Compositions of Alloy 4A, average overall composition Pt-17.4AI-6.4V (at.%) as determined by EDX					
Component	Pt	AI	v	Phases	
Overall	76.2	17.4	6.4	-	
Dark	75.3±0.1	18.9±3.7	5.8±2.7	~Pt ₃ Al	
Medium	76.4±0.3	16.6±1.0	7.0±0.9	~Pt ₃ Al	
Light phase	75.7±0.1	17.6±1.5	6.8±1.6	~Pt ₃ Al	

Table V Compositions of Alloy 5, average overall composition Pt-22.3AI-7.9V (at.%) as determined by EDX				
Component	Pt	AI	v	Phases
Overall	69.8±0.5	22.3±0.5	7.9±0.5	-
Light	70.6±0.3	21.2±0.1	8.2±0.2	~Pt ₃ Al
Medium	72.2±1.1	18.7±0.9	9.1±0.4	~Pt ₃ Al
Thin dark	65.8±0.2	30.4±0.2	3.8±0.3	~Pt ₂ Al

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Figure 6—SEM-BSE image of as-cast Pt_{83.9}:Al₆:V_{10.1} (at.%) showing a medium grey phase (Pt), a fine eutectic ((Pt) + ~Pt₃Al) and porosity (dark spots). Scale bar is 30 μ m



Figure 7—XRD pattern of as-cast Alloy 6, average overall composition Pt-6.0Al-10.1V (at.%). Key: (Pt) – 1, ~Pt₃Al – 2, and CaCO₃ - 3 in the resin mounting

Heat-treated alloys

Annealed Alloy 1H, average overall composition Pt-26.2AI-9.1V (at.%)

Figure 8 depicts the microstructure of annealed Alloy 1H, average overall composition Pt-26.2Al-9.1V (at.%), showing a medium-contrast phase, a light phase, a dark phase and a eutectic. In some cases, the medium contrast phase had a thin dark layer around it, indicating that these were the dendrites in the as-cast alloy although their composition had changed.



Figure 8—SEM-BSE image of annealed alloy $Pt_{64,7}$: $Al_{26,2}$: $V_{9,1}$ (at.%) showing a medium grey phase, a light phase, a dark phase, a eutectic, and dark precipitate the within medium grey phase. Scale bar is 10 μ m

The original dendrites showed precipitates of a darker phase, presumed to be PtV considering the contrast and the morphology, which varied between coarse (easily visible) and fine (very difficult to resolve). The plot of the compositions identified the light phase as \sim Pt₅Al₃, the dark phase as either \sim PtV or \sim Pt₂V, while the medium grey phase was identified as \sim Pt₃Al. The individual components of the eutectic were too small to be analysed accurately by EDX. XRD confirmed the identity of the light phase as \sim Pt₅Al₃ and the dark phase as \sim PtV (Figure 9).

Annealed Alloy 2H, average overall composition Pt-21.1AI-17.1V (at.%)

Figure 10 shows images of annealed alloy 2H, average overall composition Pt-21.1Al-17.1V (at.%), at different magnifications. Figure 10(b) shows a dark phase with light solid-state precipitates and a light phase with dark solid-state precipitates. Plotting identified the dark phase as ~PtV and the light phase as ~Pt₅Al₃. The precipitates were too small to be analysed accurately by EDX. XRD confirmed the presence of ~PtV and ~Pt₅Al₃. Compared to the as-cast sample, the ~PtV dendrites had reduced in volume with precipitates of ~PtV.

Annealed Alloy 3H, average overall composition Pt-10.8AI-25.5V (at.%)

Figure 11 shows Alloy 3H, average overall composition Pt-10.8Al-25.5V (at.%), at different magnifications. Figure

Table VI					
Compositions of as-cast Alloy 6, average overall composition Pt-6.0Al-10.1V (at.%) as determined by EDX					
Component	Pt	AI	v	Phases	
Overall	83.9±0.3	6.0±0.3	10.1±0.3	-	
Medium	88.5±0.4	3.2±0.51	8.3±0.3	(Pt)	
Eutectic	78.1±7.2	12.1±2.1	9.8±5.6	$(Pt) + Pt_3AI$	

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Figure 9—XRD pattern of Alloy 1H, average overall composition Pt-26.2AI-9.1V (at.%). Key: ~Pt₃AI - 1, ~Pt₂AI - 2, ~PtV - 3, ~Pt₅AI₃ - 4, CaCO₃ - 5

11 (a) shows a medium grey phase, a light phase, and a dark phase. Comparison with the as-cast structure (Figure 3) shows that the amount of \sim Pt₂V dendrites had been reduced, and a lighter phase, \sim Pt₃Al, had precipitated within the dendrites (Figure 11 (b)). The phase was too small to analyse accurately, although larger areas of the medium phase were analysed. Plotting identified the medium phase as \sim Pt₂V, the dark phase as \sim PtV, while the light phase was identified as \sim Pt₃Al. XRD confirmed these identities.

Annealed Alloy 5H, average overall composition Pt-23.2AI-7.1V (at.%)

Alloy 5H, average overall composition Pt-23.2Al-7.1V (at.%), contained a dark and a medium contrast phase of the same morphology and a small amount of a light phase. EDX compositions of the phases showed that the dark and

Table VII

Overall and phase compositions of annealed Alloy 1H, average overall composition Pt-26.2AI-9.1V (at.%) as determined by EDX

Component	Pt	AI	v	Phases
Overall	64.7±0.3	26.2±0.2	9.1±0.2	-
Medium dendrites	65.3±0.4	17.5±0.8	17.2±0.7	~Pt ₃ AI
Light phase	65.4±0.9	30.5±1.5	4.1±0.8	~ Pt ₅ Al ₃
Dark phase	57.9±1.0	5.3±2.0	36.6±2.5	~PtV
Eutectic	63.8±1.2	24.5±2.5	11.8±1.5	?
Two-phase dendrites	66.0±0.5	20.4±1.1	13.6±1.3	~PtV pptn. within ~Pt ₃ Al



Figure 10—(a) SEM-BSE image of annealed Alloy Pt-21.1Al-17.1V (at.%) showing a dark phase (~PtV), a light phase (~Pt₂Al) and dark precipitates in the light phase. Scale bar is 30 µm. (b) SEM-BSE image of annealed Alloy Pt-21.1Al-17.1V (at.%) showing a dark phase, a light phase, a eutectic, and a possible light precipitates in the dark phase. Scale bar is 10 µm

Table VIII

Overall and phase compositions of annealed Alloy 2H, average overall composition Pt-21.1Al-17.1V (at.%) as determined by EDX

Component	Pt	AI	v	Phases
Overall	61.8±0.3	21.1±0.6	17.1±0.3	-
Dark	57.6±0.4	1.8±0.2	40.6±0.2	~PtV
Light	64.4±0.3	33.8±0.4	1.8±0.2	~Pt ₅ Al ₃
Dark ppt. within the light matrix	63.8±0.9	28.2±2.0	7.9±1.2	~PtV pptn. within ~Pt ₂ Al
Light ppt. within the dark phase	59.1±0.9	8.9±0.9	32.4±1.6	~ Pt_5Al_3 pptn. within ~ PtV

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Figure 11—(a) SEM-BSE image of annealed alloy Pt-10.8AI-25.5V (at.%) showing a medium grey phase (\sim Pt₂V), a light phase (\sim Pt₃AI), and a dark phase (\sim PtV). Scale bar is 30 μ m. (b) Figure 11(b). SEM-BSE image of annealed alloy Pt-10.8AI-25.5V (at.%) showing a medium grey phase (\sim Pt₂V), a light phase (\sim Pt₃AI), a dark phase (\sim Pt₃AI), a dark phase dendrites. Scale bar is 10 μ m

Table IX

Overall and phase compositions of annealed Alloy 3H, average overall composition Pt-10.8AI-25.5V (at.%) as determined by EDX

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Component	Pt	AI	v	Phases	
Overall	63.7±0.5	10.8±0.6	25.5±0.3	-	
Medium	64.0±0.4	3.2±0.5	32.8±0.6	~Pt ₂ V	
Light	65.3±0.8	15.9±1.4	18.8±0.9	~Pt ₃ Al	
Dark	58.4±0.6	0	41.5±0.6	~PtV	
2-phase dendrites	64.2±0.6	7.4±0.7	28.3±0.3	~Pt ₃ Al pptn. within ~Pt ₂ V	

Table X

Overall and phase compositions of annealed Alloy 5H, average overall composition Pt-23.2AI-7.1V (at.%) as determined by EDX

Component	Pt	AI	v	Phases
Overall	69.7±0.1	23.2±0.3	7.1±0.1	-
Medium	70.6±0.3 70.4±0.6	21.0±0.2 21.6±1.0	8.4±0.2 8.1±0.7	~Pt ₃ AI ~Pt ₃ AI
Light	65.0±0.7	32.7±0.7	2.2±0.2	~Pt ₂ Al

medium contrast phases were the same phase, ~Pt₃Al, and the appearance was likely due to different orientations. The light phase was identified as ~Pt₂Al. XRD confirmed the presence of both phases (Figure 11).

Annealed Alloy 6H, average overall composition Pt-5.8Al-10.3V (at.%)

Figure 13 shows annealed Alloy 6H, average overall composition, Pt-5.8Al-10.3V (at.%), at different magnifications. Both have poor contrast owing to the close average atomic numbers of the phases. Figure 13(a) shows a light contrast phase and a dark phase. From the plotting, the light phase was identified as (Pt), and the dark phase as ~ Pt_2V , which was not observed in the as-cast sample. The coarsened eutectic could just be discerned in Figure 13(a) and was more



Figure 12—XRD pattern of alloy 5H, average overall composition Pt-23.2Al-7.1V (at.%). Key: \sim Pt_3Al – 1 and \sim Pt_2Al – 2

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As cast and heat-treated alloys of the Pt-Al-V system at the Pt-rich corner



Figure 13—(a) SEM-BSE image of annealed alloy Pt-5.8AI-10.3V (at.%) showing a light grey phase (Pt), a dark phase (\sim Pt₂V), and a eutectic. Scale bar is 10 μ m. (b) SEM-BSE image of annealed alloy Pt-5.8AI-10.3V (at.%) showing a light grey phase, and a eutectic at a higher magnification (x 10 000). Scale bar is 5 μ m

Table XI

Overall and phase compositions of annealed Alloy 6H, average overall composition Pt-5.8AI-10.3V (at.%) as determined by EDX

Component	Pt	AI	v	Phases
Overall	83.9±0.9	5.8±0.6	10.3±0.5	-
Light	87.8±0.5	3.7±0.3	8.6±0.5	(Pt)
Dark	70.1±0.7	-	29.9±0.7	~Pt ₂ V
Eutectic	78.5±1.0	6.8±0.2	14.8±1.2	(Pt) + ~Pt ₃ Al

clearly resolved in Figure 13 (b). The components of the eutectic were too small to be analysed accurately by EDX. Extrapolation indicated that the eutectic is between (Pt) and ~Pt₃Al. XRD confirmed the presence of ~Pt₂V, ~Pt₃Al, and (Pt).

Discussion

Although some of the phases were very difficult to resolve, mainly because they had very similar average atomic numbers, the morphologies, analyses, and XRD results were consistent, even if the alloys were not fully homogeneous.

The solidification projection, obtained from the as-cast samples, at the Pt-rich corner is given in Figure 14, although changes to the phase field boundaries are expected as more alloys are studied. All the phases identified in the as-cast alloys were extensions of the binary phases. The phase \sim Pt₃Al extended the most, particularly in the annealed condition (Figure 15).

The (Pt) phase had a greater solubility for Al above ~20 at.% V than between 10–20 at.% V. The annealed results were used to obtain the isothermal section in Figure 15. There were similar shapes for the phase field boundaries in the solidification projection and the isothermal section, particularly for (Pt) between 10–20 at.% V. It is not yet possible to discern the shape of the ~Pt₃Al phase field boundary on the Pt-rich side, but new samples are in preparation.

Alloy 1 had four visible phases, \sim Pt₃Al, \sim Pt₂Al, \sim Pt₅Al₃, and \sim Pt₃V, in the as-cast condition, while a fifth phase, PtAl,

could not be resolved. In the annealed condition Alloy 1 contained ~PtV, ~Pt₅Al₃ and ~Pt₃Al. The light phase was deduced to be to be ~Pt₅Al₃ on the basis of not showing the



Figure 14—Solidification projection of the Pt-Al-V system at the Pt-rich corner





Figure 15—Isothermal section at 1 000°C of the Pt-Al-V system at the Pt-rich corner

laths as found by Tshawe *et al.*¹⁶ in the Pt-Al alloys as well as the XRD pattern. The major change is that in the as-cast condition the alloy contained ~Pt₃V, while in the annealed condition it contained ~PtV. The change can be attributed to diffusion as the phase compositions change, or a reaction in which ~Pt₃V changed to ~PtV (although the involvement of the other phases cannot be identified yet). The microstructure of the annealed alloy was also coarser than that of the ascast, which is expected.

Alloy 2 contained three phases in the as-cast condition, namely ~Pt₂V, ~Pt₅Al₃, and PtAl. There was a lamellar structure which consisted of a solid-state cellular precipitation of ~Pt₂V /(Pt) in ~Pt₂Al, which shows that the ~Pt₅Al₅ solvus retreats at lower temperatures. The complex eutectic was originally ~Pt₅Al₃ + β , and the β decomposed eutectoidally¹³ to give almost unresolveable ~Pt₅Al₃ + ~PtAl. The annealed Alloy 2H contained two phases, ~PtV, ~Pt₅Al₃, and the remnants of a eutectic. The precipitation of ~PtV within the ~Pt₂Al matrix (Figure 9 (b)) shows that the ~Pt₅Al₃ solvus range decreases markedly with lower temperatures, as seen by the cellular precipitation in the as-cast alloy (Figure 2 (b)). Additionally, there was precipitation of ~PtV solvus retreats substantially with decreasing temperature.

Alloy 3 contained three phases, $\sim Pt_3V$, $\sim Pt_3Al$, and $\sim Pt_2Al$ (in a small proportion), in the as-cast condition. In the annealed condition, there was $\sim Pt_2V$, $\sim PtV$, and $\sim Pt_3Al$, and precipitation of $\sim Pt_3Al$ in the $\sim Pt_2V$ phase.

Alloy 4, which had an average overall composition Pt-17.4Al-6.4V, was single-phase ~Pt₃Al in the as-cast condition. This alloy has not yet been heat treated. Alloy 5 had an average overall composition Pt-22.3Al-7.9V, with two phases, ~Pt₃Al and ~Pt₂Al in the as-cast condition and the same two phases after annealing.

Alloy 6 contained (Pt) dendrites and a (Pt) + \sim Pt₃Al eutectic in the as-cast condition. After annealing, the alloy had the same (Pt) and the (Pt) + \sim Pt₃Al eutectic as well as a third phase, \sim Pt₂V which must have formed as a result of a reaction during annealing.

Conclusion

The solidification projection, an isothermal section, and a liquidus surface projection for the Pt-rich corner of the Pt-Al-V were found to be consistent with each other. Using the solidification sequences from the phases and their morphologies, a tentative liquidus surface was derived. The \sim Pt₂Al phase forms only in the solid state, and so does not appear on the liquidus surface.

Figure 16 (b) shows the solidification reactions for each of the as-cast alloys. These are schematic, since the analysed primary phase is assumed not to change on further solidification. Solidification starts with the composition moving directly away from the primary solid (on that primary



Figure 16—(a) Liquidus surface projection for Pt-AI-V at the Pt-rich corner. (b) Solidification reaction on the liquidus surface for Pt-AI-V at the Pt-rich corner

P

a

P

e

P²

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liquidus surface) until a boundary with another liquidus surface is met, and the liquid composition then follows the boundary between the two liquidus surfaces for either a eutectic or a peritectic reaction (ignoring the fact that on nonequilibrium cooling, the liquid composition can sometimes run across the liquidus surface of the peritectic component. For Alloy 1, the second phase was also used to derive the direction of the liquid composition as the ~Pt₅Al₃ formed peritectically. Unfortunately the extrapolation of this falls on the direction of the liquidus for Alloy 5, but this is of no significance and is coincidental. Of the reactions observed, all had been reported in the Pt-Al binary, except for the peritectoid reaction ~Pt₃Al + ~Pt₅Al₃ \rightarrow ~Pt₃V, which formed as a thin layer.

Several phases showed solid state-precipitation, indicating that the solvus of the original phase retreats noticeably with decreasing temperature. These included ~Pt₃Al, ~Pt₅Al₃, ~Pt₂V, and ~PtV. The solidification projection and the isothermal section were consistent with each other.

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