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THESIS

An X-Ray Diffraction investigation of α -Al₂O₃ addition to yttria stabilized zirconia (YSZ) thermal barrier coatings subject to destabilizing vanadium pentoxide (V₂O₅) exposure

by

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September, 1993

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An X-ray Diffraction investigation of α -Al₂O₃ addition to yttria stabilized zirconia (YSZ) thermal barrier coatings subject to destabilizing vanadium pentoxide (∇_2 O₅) exposure

by

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ABSTRACT

Since the mid 1970's the U.S. Navy has used Yttria-stabilized zirconia (YSZ) as thermal barrier coatings for hot stage gas turbine components. Use of low cost, high contaminant, fuels has led to shortened component life from failure of YSZ coatings due to corrosive attack by vanadium and other combustion oxides. The object of this investigation was to determine the reactivity of adding α -Al₂O₃ to current YSZ ceramics for creation of a ceramic composite which could improve mechanical properties and show improved durability to corrosive chemical attack. Ten powder samples of ZrO_28mol Y_2O_3 , α -Al₂O₃, and V_2O_5 were annealed at 900°C for 100 hours. X-Ray Diffraction analysis utilizing a standard 'search and match' method was used to determine the phases present in the reacted powder samples. Peak intensity comparisons between reacted and un-reacted samples allowed for a quantitative determination for the reactivity of α -Al₂O₃ with the YSZ system exposed to V_2O_5 . This investigation indicated that α -Al₂O₃ is non-reactive in all YSZ samples exposed to

V₂O₅ at 900°C.

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I. INTRODUCTION

Since the mid 1970's the U.S. Navy has been utilizing thermal barrier coatings (TBC) on hot stage gas turbine superalloy components. Yttria stabilized zirconia (YSZ) has been effective in providing thermal barrier protection, allowing for realized operating temperatures from increases in 50-220°C, thereby obtaining increases in engine efficiencies from 10 to 15 percent [Ref. 1]. Many researchers have found that YSZ becomes highly unstable and prematurely fails when exposed to inorganic fuel contaminants such as vanadium and sulphur which are found in low cost fuels of non-aviation quality. Vanadium can be found in fuels in the form of vanadium porphyrin which is transformed during combustion into VO(OH), (gas) and V₂O₅ (solid) [Ref. 2]. Vanadium pentoxide (V_2O_5) is in liquid form at 672°C and reacts with the stabilizing agent Y_2O_3 to form yttria vanadate (YVO₄). This reaction leaves behind an unstable tetragonal (t) zirconia structure which transforms to monoclinic (m) zirconia while cooling The t \rightarrow m transformation results in a 4-5 to room temperature. percent volume increase within the matrix leading to spalling and eventual premature failure of the coating [Ref. 3]. The emphasis of the current research is to investigate alpha alumina $(\alpha - Al_2O_3)$ as a possible additive to the current YSZ ceramic for the creation of a composite which could show improved immunity to corrosive attack and improved durability to thermal cycling.

II. BACKGROUND

In order to interpret data on composites containing ZrO_2 , V_2O_3 , Al_2O_3 , and Y_2O_3 , it is necessary to understand all the binary and ternary phase diagrams of these systems and the crystallography of the phases present. The first section of this background will therefore be devoted to summarizing these fundamentals.

A. CRYSTALLOGRAPHY OF ZIRCONIA

Pure zirconia exhibits three different crystal structures at atmospheric pressure as shown in Figure 1 [Ref. 4].

1. Cubic

The cubic phase is the first polymorph to form from the liquidus state at $2680 \pm 15^{\circ}$ C. Cubic zirconia has the fluorite crystal structure which is FCC. Eight anions (oxygen) are located equidistantly at the eight tetrahedral positions. Each anion is tetrahedrally surrounded by four cations (zirconia) as seen in Figure 2 [Ref. 1]. Cubic zirconia is stable down to 2370° C.









Figure 2. Schematic diagrams of fluorite structure of cubic ZrO_2 emphasizing (A) face-centered cubic packing of cation sublattice and (B) simple cubic packing of anion sublattice.

2. Tetragonal

The tetragonal (t) polymorph forms at 2370° C and is stable through 1170° C. The cation is surrounded by eight anions, four equidistantly at 0.2465 nm and the other four at 0.2065 nm. This structure is classified as simple tetragonal and is shown in Figure 3 [Ref. 5:p. 96].



Figure 3. Crystal structures of monoclinic and tetragonal zirconia with lattice parameters at 950°C for both phases.

3. Monoclinic

The monoclinic phase is stable at all temperatures below 1170°C. The cation has a coordination number of seven which indicates a departure from pure ionic bonding, suggesting that covalent bonding is present [Ref. 6]. The simple monoclinic bravais lattice is shown in Figure 3.

B. CRYSTALLOGRAPHY OF ALUMINA

Alpha alumina (tradename corundum) is an excellent candidate for addition to the YSZ system due to it's properties of high elastic moduli, low density, and most importantly it's stability at extremely high temperatures [Ref. 7]. Corundum has the hexagonal close packed structure as shown in Figure 4 [Ref. 7].



Figure 4. Hexagonal closed-packed structure of α -alumina. A & B layers contain oxygen atoms. C₁, C₂, and C₃ contain aluminum atoms. The C layers are only two-thirds full.

C. MECHANICAL PROPERTIES OF ZIRCONIA SYSTEMS

As a ceramic, zirconia presents an attractive package of properties; high strength at extremely high temperatures, low density, and chemical inertness due to its ionic nature. The achilles heel for this and other ceramics is the absence of toughness. Ceramics are prone to catastrophic failures from minute internal or external flaws. TABLE I presents very promising possibilities for the creation of ceramic matrix composites (CMC) [Ref. 8 pg. 24].

TABLE I.

ROOM TEMPERATURE TOUGHINESS K_{ic} and strength of df ZrO₂-toughened Ceramics

Ceramic Material	Matrix alone		Matrix + ErO ₂	
	K_{ze} (MPa $n^{1/2}$)	σ _f (MPa)	K _z , (MPa m ^{1/2})	σ _ε (MPa)
c-ZrO ₂	2.4	180	2 - 3	200-300
PSZ			6 - 8	600-800
TZP			7 - 12	1000-2500
Al ₂ O ₃	4	500	5 - 8	500-1300

PSZ - Yttria Stabilized Zirconia

TZP - Tetragonal Stabilized Zirconia

The effect of 'alloying' ceramic particles or fibers are clearly evident. The fracture toughness values show an increase of nearly 300 percent, likewise rupture strengths have been shown to produce similar increases for ZrO_2 toughened ceramics. Contributions to the increased toughness and strength values are from different mechanisms, e.g., the stress-induced t \rightarrow m transformation (process zone mechanism), microcracking, crack deflection and bridging (from whisker composites), and inhibition of grain growth (grain boundary pinning).

D. BINARY PHASE DIAGRAMS OF ZrO2-AL203-Y203-V205 SYSTEMS

The use of binary phase diagrams is sufficient as a starting point when evaluating multiple compound formations. Unfortunately, the availability of ternary phase diagrams for the $ZrO_2-Al_2O_3-Y_2O_3$ system was extremely limited. One was obtained, but for a temperature of 1450°C [Ref. 9], well above the temperature of interest for this research.

1. Zr0,-Y,0, Phase Diagram

The zirconia-yttria system is one of the most well known systems due to the extensive research in the field of thermal barrier coatings. Figure 5 shows clearly that with an increasing amount of Y_2O_3 the cubic solid solution phase greatly expands, producing a stable cubic phase to about 300°C (at 25mol Y_2O_3) [Ref. 10]. A clearer view of the low-yttria region is depicted in Figure 6 [Ref. 11]. Yttria content below 4-5 percent creates the monoclinic polymorph upon cooling. Yttria percentages above 15

percent produce a fully stabilized zirconia (FSZ) at room temperatures.



Figure 5. System $ZrO_2-Y_2O_3$. F = fluorite type of ZrO_2 ; Y = Y₂O₃; Cub = cubic; Tet = tetragonal; Mon = monoclinic; Hex = hexagonal.





FSZ are produced for their stability and strength at room temperature and widely used in costume jewelry as diamond replacements. The phase desired near room temperature has a yttria content of between 6-12 mole percent. The tetragonal phase is desired because it is metastable and does not spontaneously transform into monoclinic upon cooling.

The monoclinic structure is not desired due to the 4-5 percent volume increase during cooling, this would create undue stresses in the matrix and cause coating failure. The FSZ structure is undesirable because it does not have the microstructural defects, specifically twinning of the tetragonal grains and production of anti-phase boundaries (APB), or show coherent precipitates at high annealing temperatures like the tetragonal phase. [Ref. 11] These microstructural defects of tetragonal zirconia allow for the dissipation of crack propagation energy through a crack deviation mechanism not found in the other two polymorphs. These phenomena explain why researchers have settled on eight (8) mole percent yttria as the optimum content for zirconia stabilization.

a. Monoclinic - Tetragonal Transformation

This transformation was determined by high temperature X-Ray Diffraction in 1929. Wolten was the first to suggest that this was a diffusionless 'martensitic like' transformation. A hysterisis effect exists with a forward transition temperature of 950°C and a reverse transformation temperature of 1170°C. This martensitic transformation can occur athermally or by stress-inducement (as found in the region of a

stress field ahead of a crack tip). [Ref. 12]

b. Cubic - Tetragonal Transformation

This displacive transformation is composition invariant and occurs with a change in crystal structure relative to the parent cubic phase (resulting in a strain free, undisturbed habit plane) [Ref. 11] and [Ref. 12]. Other characteristics of this transformation are a homogenous shape change due to the shear action which is reflected by surface relief where the product regions form. Twinning also occurs with this transformation in polycrystalline materials.

2. Zro,-V,O, Phase Diagram

The $ZrO_2-V_2O_5$ system shown in Figure 7 indicates low liquidus temperatures for high concentrations of V_2O_5 with a eutectic and peritectic based on pure ZrO_2 and ZrV_2O_7 [Ref. 13]. The stoichiometric structure ZrV_2O_7 (zirconium vanadate - cubic structure) will be shown later to develop over extended periods of time and is therefore unlikely to be as detrimental to a YSZ coating as the formation of YVO_4 .



Figure 7. System V₂O₃-ZrO₂.

3. AL₂0,-ErO, Phase System

One of the most promising aspects of corundum as an additive for ceramic matrix composites is its high temperature stability as indicated in Figure 8 [Ref. 14]. The eutectic point for the $\lambda l_2 O_3$ -ZrO₂ system is nearly 1000°C above the typical gas turbine operating temperature. Whether as a polycrystalline powder or fabricated whisker, α -alumina and zirconia remain in solidus form up to 1845°C, and appear to be immiscible in the solid state.

4. Al₂O₃-Y₂O₃ Phase Diagram

The $Al_2O_3-Y_2O_3$ phase diagram indicates that the various phases which can form are all solid at temperatures of interest in the present work as seen in Figure 9 [Ref. 15]. Of interest are the garnet structure (3:5 ratio) and the perovskite structure (1:1 ratio). Perovskite is a complex structure which is found in several electrical ceramics.



Figure 8. System Al₂O₃-ZrO₂ Phase Diagram.

5. Al₂O₃-V₂O₃ Phase Diagram

The $Al_2O_3-V_2O_5$ system shown in Figure 10 exhibits no propensity for stoichiometric compound formation, and remains in the solidus phase below the 660°C eutectic temperature. [Ref. 17]



Figure 9. System $Al_2O_3-Y_2O_3$, tentative. $3Y_2O_3*5Al_2O_3$ (3:5) = garnet structure type; $Y_2O_3*Al_2O_3$ (1:1) = perovskite; $2Y_2O_3*Al_2O_3$ (2:1) = monoclinic symmetry.



Figure 10. System V₂O₅-Al₂O₃.

6. Y.O.-V.O. Phase Diagram

The $Y_2O_3-V_2O_3$ phase diagram is one of the most important diagrams in helping to explain the deleterious effects of vanadium pentoxide on yttria stabilized zirconia. The formation of yttria vanadate (YVO₄) is possible for all percentages of V_2O_3 above 20 percent for operating temperatures of interest. Since V_2O_3 is a combustion byproduct it is obvious that cheaper fuels (less processing) will contain higher amounts of vanadium porphyrin. The $Y_2O_3-V_2O_3$ phase diagram is shown in Figure 11 [Ref. 18].



Figure 11. System $Y_2O_3-V_2O_5$. The $Y_2O_3-YVO_4$ subsystem is probably pseudobinary because of oxygen losses from the 4:1 and 5:1 phases. L and H = low and high forms respectively.

7. Additional Stabilising Oxides for Sixoonia

Oxides of calcia, magnesia, india, and scandia have also been seen to demonstrate a stabilizing effect on sirconia because of their high solubility in zirconia and ability to form the fluorite type structure. Current research with india (I_2O_3) and scandia (Sc_2O_3) have shown improvements over yttria from the destabilization of zirconia due to YVO₄ production and associated leaching of the zirconia matrix [Ref. 19]. Three of these common zirconia stabilizers, MgO, CaO, and Sc₂O₃ are shown below in Figure 12 [Ref. 20], Figure 13 [Ref. 10], and Figure 14 [Ref. 21], respectively.



Figure 12. System ZrO₂-MgO showing ZrO₂-rich side.



Figure 13. Zirconia-calcia binary system.



Figure 14. System $ZrO_2-Sc_2O_3$, partial subsolidus. Fl ss = flouritetype cubic ss: Mon = monoclinic; Tet = tetragonal. $Sc_2Zr_7O_{17}$, $Sc_2Zr_5O_{13}$, and $Sc_4Zr_3O_{12}$ show rhombohedral symmetry.

E. DESTABILIZATION OF YSZ BY CHEMICAL ATTACK

Fuel contaminants that are known to cause corrosive attack on ceramics are vanadium, sulphur, sodium, and phosphorous. All of these contaminants form oxide species during combustion (V_2O_5 , SO_3 , Na_2O_5 , and P_2O_5) [Ref. 22]. Of specific interest to this research is V_2O_5 .

1. Reaction of Vanadium Compounds With Ceramic Oxides

Ceramic oxides have particular acid or base characteristics. Lewis acid-base theory is used to explain the tendency for oxides to react with each other. R. L. Jones describes Lewis acid-base theory as:

Basicity is defined as the ability to donate electronic charge and acidity as the ability to accept electronic charge. For oxides, the oxide anion is the charge donor, and it's behavior is modified by the cation, where the cation acidity increases both as the cation positive charge increases and the cationic radii decreases... Thus, each oxide species has a characteristic acidity (or basicity), and the tendency for reaction between oxides will depend, to a first approximation, on the difference between their relative acid or base strengths. [Ref. 22 pg. 371]

Reactions of some vanadium compounds with ceramic oxides and their products are shown in Figure 15 [Ref. 22]. This figure clearly indicates the notion that based on Lewis acid-based theory, acids will react with bases, but not acids with acids nor bases with bases.



Figure 15. Reaction behavior and products of ceramic oxides with vanadium compounds.

2. Reaction of V.O. With ZrO.

Based on the phase diagram in Figure 7, the proposed reaction for the V_2O_5 -ZrO₂ system would be:

$$V_2O_5 + ZrO_2 = ZrV_2O_7$$
 (1)

As shown by Figure 15, ZrV_2O_7 forms but only after a long period of time and/or high temperature ZrO_2 exposure to vanadium pentoxide.

3. Reaction of V₂O₅ With Y₂O₅

Assuming that V_2O_3 and Y_2O_3 react on an equimolar basis, the creation of yttria vanadate is:

$$V_2O_3 + Y_2O_3 = 2YVO_4$$
 (2)

As shown in the phase diagram (Figure 10), YVO_4 can form with V_2O_3 concentrations above 20 percent. It is not surprising that from Lewis acid-base theory that YVO₄ rapidly forms as Y_2O_3 is very basic relative to V_2O_5 acidity. This important principle is a major reason why researchers are viewing alternative stabilizers such as scandia (Sc₂O₃) and india (I₂O₃) as alternative zirconia stabilizers. Both of these oxides, on a relative basis, are more acidic than yttria.

4. Reactions of YSZ With Vanadium Pentoxide

The reaction between V_2O_5 and Y_2O_3 occurs within the zirconia matrix. Rewriting equation (2):

 Y_2O_3 (base) + V_2O_5 (acid) = 2YVO₄ (salt) (3)

 Y_2O_3 (a<1) + V_2O_5 (a<1) = 2YVO₄ (s,a=1) (4)

$$Y_2O_3$$
 (ZrO₂) + V_2O_5 (melt) = 2YVO₄ (ppt.) (5)

Equation (3) is written to emphasize the Lewis acid-base nature specific to corrosive chemical attack on ceramics. [Ref. 23]. Equation (4) emphasizes that the activities of yttrium oxide in PSZ and vanadium pentoxide in melt are less than one (a<1), and the yttria vanadate is formed as a pure solid with activity equal to one (s, a=1) [Ref. 23].

Equation (5) indicates the physical nature of the corrosive attack. Molten V_2O_5 attaches to the YSZ surface. The zirconia matrix (as well as other oxide ceramics) surface is porous, allowing the molten salt to thoroughly penetrate the matrix surface [Ref. 23].

Diffusion of vanadium into the matrix (near the surface) precipitates the YVO_4 formation on the surface, creating a depletion zone within the interior of the surface. Depletion of

 Y_2O_3 from the zirconia matrix creates the unstable condition for the low temperature transformation of the desirable polymorph of ZrO_2 (tetragonal and cubic) to monoclinic ZrO_2 on the surface. Cracks initiate due to the cubic \rightarrow tetragonal \rightarrow monoclinic transformations associated with the thermal cycling from continued transformations. This repeated cycling will allow the cracks to grow and branch into the coating interior eventually leading to spalling and final coating failure. [Ref. 22]

Work by Patton et al on single crystal yttria stabilized zirconia exposed to vanadium pentoxide melts are presented in Figure 16 and illustrates this behavior [Ref. 22].



(b)

(a)



Figure 16. A comparison of (a) the concentration variations for yttrium, vanadium and zirconium existing in a YSZ specimen afterexposure to V_2O_5 for 100 hours using EPMA compared to (b) a schematic illustrating the orientation of the different regions.

III. FABRICATION PROCESS FOR THERMAL BARRIER COATINGS

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One important feature of any potential coating is its method of application. Current researchers have recently looked at the fabrication process of TBCs as a way of improving the durability and mechanical properties of these coatings. Recent work with Al_2O_3 -ZrO₂-SiC_(w) whisker composites utilizing the Surface Induced Coating (SIC) method have shown promising results in mechanical properties strengthening [Ref. 24]. The increased strength arises from the fabrication of a uniform spatial distribution of ZrO₂ and $SiC_{(w)}$ phases throughout the Al_2O_3 matrix. This homogenous fabrication method also shows the promise of minimizing the t \rightarrow m zirconia transformation due to the suppression of tetragonal grain growth within the matrix not enabling the tetragonal grain to reach the critical grain diameter (d_c) [Ref. 8], [Ref. 24], and [Ref. 25]. Colloidal processing techniques should be another aspect to consider with the addition of α -Al₂O₃ to the ZrO₂8mol Y_2O_3 matrix.

A. PLASMA SPRAY COATING

The plasma spray deposition method is widely used in the fabrication/application of thermal barrier coatings [Ref. 26]. Plasma spray deposition can evenly distribute ultra thin coatings of material and join dissimilar materials which are difficult to bond. A low-pressure plasma deposition (LPPD) system is depicted in Figure 17 [Ref. 7].



Figure 17. Schematic of low-pressure plasma deposition (LPPD) method.

Due to the large difference in the coefficients of thermal expansions (C.T.E.) between ceramic oxides and metal.ic superalloys, an intermediate bondcoat is necessary to minimize the surface/interface shear strains due to expansion and contraction of the interface due to thermal cycling. Figure 18 shows a typical cross-section of a partially stabilized zirconia thermal barrier coating on a superalloy substrate [Ref. 23].



Figure 18. Cross section of a plasma-sprayed zirconia thermal barrier coating under corrosion test conditions.

Probably the most significant aspect of the plasma sprayed deposition method are the high cooling rates associated with the molten deposit hitting the substantially cooler substrate. Van Valzah and Eaton have recently completed work on the effect of cooling rates on the tetragonal \rightarrow monoclinic transformation, thermal expansion behavior, and flexure strength of the plasma sprayed thermal barrier coatings [Ref. 27]. Their research indicates that t-ZrO₂ actually consists of two phases of ZrO₂. Metastable tetragonal (t') phase and another phase consisting of cubic and monoclinic ZrO₂. The metastable t' phase forms when the atomized ZrO₂ strikes the much cooler substrate and transforms rapidly from cubic to t'. The t' phase is termed 'nontransformable' due to a higher concentration of Y_2O_3 in t' than

tetragonal (t) phase. The t' phase undergoes phase separation via diffusion at temperatures greater than 1200°C [Ref. 27]. This phase separation will allow for a greater amount of tetragonal to monoclinic transformation upon cooling, leading to excessive microcracking and eventual coating failure.

The above findings occur for aged samples above 1200°C, the same principle can apply to lower temperature coatings. Although the plasma spray deposition method can lay down an even thin coating of material, the composition of the coating is harder to control. Phase segregation can occur during processing (laying a coating down). The dispersion of Y₂O, can not be controlled. The plasma spray method thus has some problem controlling precipitate Van Valzah and Eaton along with Lange also report that size. grain size of a critical diameter will also effect the t \rightarrow m transformation [Ref. 25] and [Ref. 27]. Precipitate tetragonal grains larger than a d will transform spontaneously upon cooling. Tetragonal precipitates smaller than d but larger than a minimum diameter (d_), are metastable at room temperature, but may transform in the front of a propagating crack. Grains smaller than d_m are stable at room temperature [Ref. 25] and [Ref. 27].

It is easy to understand that the method of fabrication can play an important role on the durability of thermal barrier coatings. Methods of application/fabrication which can produce a homogeneous spatial distribution would appear to have the ability to control the aforementioned problems of a phase segregation and control of critical grain size.
B. SURFACE INDUCED COATING METHOD

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The fabrication/processing of ceramic composites can show improved mechanical properties with improved spatial distribution of toughening agents within the ceramic matrix. Recent studies by Jang and Moon on zirconia toughened alumina (ZTA), and $Al_2O_3-ZrO_2-$ SiC_(w) whisker composites utilizing colloidal science with interfacial electrochemistry has developed superior microstructures with uniform size and spatial distributions of both matrix and dispersed phases in these two systems [Ref. 24] and [Ref. 28].

The keys to the success of creating a ZTA composite powder by the surface induced coating (SIC) method are: (1) the determination of a pH range for a kinetically stable Al₂O₃ dispersion (a pH range where Al₂O₃ particles do not coagulate); (2) the determination of a pH range which will enhance the interfacial concentration of ionic species used for selective formation of the ZrO₂ precursor only at the interface (a pH range which enhances ZrO₂ formation on the Al₂O₃ particle); (3) the pH range which optimizes the first two conditions [Ref. 28]. The reader is encouraged to read reference 24, and reference 28 for a precise presentation and theoretical analysis of the SIC method. A flow diagram for the processing of Al₂O₃-ZrO₂-SiC_(w) composite is shown in Figure 19 [Ref. 24].



Figure 19. Schematic representation of homogeneous fabrication of $Al_2O_3-ZrO_2-SiC_{(W)}$ whisker composite by surface-induced coating.

Homogeneously coated ZTA powder samples (10 vol. % ZrO₂) were pressurelessly sintered at 1500°C for six hours. XRD analysis indicated that nearly 95 percent of the ZrO₂ grains remained tetragonal down to room temperature. Mechanically mixed samples were also prepared, sintered and cooled for comparison. Figure 20 shows outstanding results of scanning electron microscopy (SEM) photographs for both the SIC method (20(a)) and mechanically mixed microstructures (20(b)) [Ref.28].



Figure 20. Scanning electron micrographs showing polished microstructures of zirconia-toughened alumina prepared by (a) surface-induced coating and (b) conventional mixing. As shown by Figure 20(a), the microstructure is characterized by a uniform spatial distribution of the dispersed tetragonal ZrO_2 phase throughout the composite and by the absence of large ZrO_2 grains formed from hard ZrO_2 agglomerates. Figure 20(b) indicates that there are large cracklike voids produced by large hard ZrO_2 agglomerates, and is the main fracture origin for ZTA composites. These voids tend to propagate to a ZrO_2 deficient region [Ref. 28].

Returning to the idea of critical grain sizes, the surface induced coating sample intergranular $3rO_2$ grain sizes were predominately under 1 µm (micro meter) in diameter, indicating that the martensitic t \rightarrow m transformation was suppressed [Ref. 24]. Whereas the mechanically mixed sample had a wider range of grain sizes, making the sample much more susceptible to unfavorable t \rightarrow m transformation. Figure 21 shows a comparison of matrix and dispersed phase grain sizes for both prepared samples [Ref. 24]. Ultrafine ZrO_2 grains (~ 0.1 µm or less) are unable to give transformation toughening due to the transformation resistance to monoclinic symmetry [Ref. 28].

The creation of a uniform spatial distribution of dispersed phase within the matrix phase is the result of the uniform coating of the ZrO_2 particles on Al_2O_3 particles, effectively pinning the Al_2O_3 grain boundaries and limiting the abnormal growth of Al_2O_3 grains.



Figure 21. Grain size distribution of Al₂O₃ (matrix phase) and ZrO₂ (dispersed phase) in ZTA samples processed by two different methods, surface-induced coating and conventional mixing.

. The addition of $SiC_{(w)}$ also provides additional strengthening mechanisms to the matrix by crack deflection, crack bridging, and The control of the t \rightarrow m transformation can whisker pullout. enhance mechanical properties via the process zone mechanism [Ref. When the t \rightarrow m transformation is not uniform and excessive, 24]. degradation of the thermal barrier coating will be accelerated. The $Al_2O_3-ZrO_2-SiC_{(w)}$ composite shows a three to four fold increase in rupture strength and fracture toughness, and just as important, the ability to control the microstructure of the ceramic matrix and avoid the degrading behavior of excessive $t \rightarrow m$ transformation. For these reasons, it is emphasized that the surface [Ref. 8] induced coating method be investigated as a possible alternative for the processing of yttria stabilized zirconia.

IV. SCOPE OF PRESENT WORK

Based upon previous research and literature introduced here, a summary of important conclusions is presented:

- ZrO₂ reacts slowly to form ZrV₂O₇ from Lewis-acid base theory.
- Y_2O_3 reacts on an equimolar basis with V_2O_5 to form YVO_4 very quickly which destabilizes the zirconia matrix, thereby increasing the chances of the t \rightarrow m transformation.
- Application/fabrication processes for CMCs can directly impact the mechanical characteristics of ceramic composites.
- Addition of strengthening agents such as corundum and SiC $_{(W)}$ have been shown to improve both fracture toughness and strength values of CMCs.

The present work will involve mixing alpha alumina in varying concentrations to $ZrO_28mol \$Y_2O_3$ and exposing these samples to vanadium pentoxide for 100 hours at 900°C. Analysis by X-Ray Diffraction (XRD) and Scanning Electron Microscopy (SEM) will be used to:

- Re-investigate the reaction between V_2O_5 and $ZrO_28moltY_2O_3$ in air at 900°C to verify the destabilization of tetragonal zirconia to the monoclinic polymorph.
- Determination of the degree of reactivity of α -Al₂O₃ in the pseudo ternary ZrO₂8mol Y_2O_3 -Al₂O₃-V₂O₅ system.

V. EXPERIMENTAL PROCEDURE

A. SAMPLE PREPARATION

Fine powder of yttria stabilized zirconia (TOSOH-Zirconia TZ-8Y ZrO₂8mol%Y₂O₃) were provided by TOSOH Corp., Tokyo, Japan. Vanadium pentoxide (V₂O₃ - 99.6% pure, Chlorine 0.15%, and Carbon Dioxide 0.09%), was provided by J.T. Baker Chemical Co., Phillipsburg, N.J., and alpha alumina (1.0 micron diameter), provided by BUEHLER LTD., Evanston, Illinois.

An accurate balance (Sartorius-Werke AG), was used in preparing ten 5.00 gram samples of varying concentrations of PSZ, V_2O_5 , and Al_2O_3 . Samples are presented in weight percent form in TABLE II. Each sample was mechanically mixed and placed in a Platinum (Pt) crucible and annealed at $900\pm15^{\circ}$ for 100 hours.

B. X-RAY DIFFRACTION

Each sample prior to annealing (mechanically mixed state), was run through the XRD to obtain a baseline value for comparison with a post-fired sample. In both samples, the powder sample was passed through a U.S. Standard #400 sieve mesh (38µm) and then mounted into a standard specimen holder and held in place by applying drops of acetone to 'cement' the sample in place. Excess powder and acetone residue was

removed by packing and smoothing with a razor blade to obtain a flat smooth surface.

X-Ray Diffraction was performed utilizing a Phillips XRG 3100 X-Ray generator, a PW 1710 Diffractometer controller, and directed by a Digital 3100 VAX work station. The X-Ray generator utilizes a copper target (Cu alpha 1,2 wavelengths - 1.54060, 1.54439 A respectively), with powel settings of 30 kV and 35 mA. The gonio scan program was set with a 20 angle range of 10-140 degrees, step size of 0.05 degrees, and scan rate of 10.0 seconds (for a total run time of 7:13:20). This scanning program and power setting was suitable for collecting all the raw data necessary for analysis.

Raw data files for each sample (pre and post annealed samples) were analyzed by using the VAX work station and the Phillips APD1700 software package. Peak positions and integrated intensities were calculated and recorded with each peak curve fitted for each sample. A matching program provides a probability that the constructed intensity curves 'match' other known intensity profiles on file within the The match and analysis program was too software database. vague in many instances and did not correlate well with small percentage (by weight percent) compounds that were expected in The Phillips PW 1891 Total Access the analyzed sample. Diffraction Databa*s*e (TADD), PDF-2 1-41, sets was

indispensable in using the 'search and match' method necessary in determining compounds within the powder sample.

C. SCANNING ELECTRON MICROSCOPY (SEM)

There was one reacted sample that formed an extremely hard and brittle mass in the Pt crucible. With this sample it was not possible to obtain enough of the sample to conduct a proper XRD analysis (post annealed gonio scan). After successive attempts to clean the Pt crucible with concentrated HCL, enough of a sample was stained from the bottom of the crucible to conduct an SEM analysis. The sample remnants were cold mounted, sanded and polished, and carbon coated to enhance conductivity.

The microstructure of the mounted sample was studied on a Cambridge Model S200 Scanning Electron Microscope utilizing a LaB₆ gun, and equipped with a Kevex x-ray energy dispersive spectrometer (EDX). Chemical analysis was also measured by EDX to confirm observations resulting from X-Ray Diffraction analysis. The SEM backscattered imaging method was used to improve detection of the phases present using atomic number contrast.

TABLE II.

PONDER SAMPLES OF THE ErO28mol. 8Y203-Al203-V205 SYSTEM BY WEIGHT PERCENT

Sample No.	Zr0,8mol. 8Y203	AL203	₹203
1	80	10	10
2	90	5	5
3	70	20	10
4	60	30	10
5	50	40	10
6	10	80	10
7	20	70	10
8	30	30	40
9	40	40	20
10	10	10	80

VI. RESULTS AND DISCUSSION

A. X-RAY DIFFRACTION RESULTS

1. YSZ-Al₂O₃-V₂O₅ Unreacted Powder Samples

All ten samples were analyzed by XRD analysis prior to annealing at 900°C for 100 hours. Samples analyzed indicated that all expected peak intensity and line positions accurately reflected the Hanawalt JCDP diffraction patterns verifying the phases present. Figure 22 shows sample number four plotted against each compound's diffraction pattern expected in the sample. Each compound's maximum intensity line is marked on the appropriate raw data peak. SCHERBINAITE (V₂O₅) has a peak intensity at the (001) reflecting plane, CORUNDUM (α -Al₂O₃) peak intensity at (113), and tetragonal ZrO₂ at (101).

2. YSZ-Al₂O₃-V₂O₅ Annealed Powder Samples

Assuming that the reaction between V_2O_5 and Y_2O_3 occurs on an equimolar basis, as shown by equation (2), and that ZrV_2O_7 forms after long periods of time, the expected phases present in the reacted samples can be calculated by converting the weight percentage of mixed constituents and converting them to atomic percent. Calculation of atomic percent is conducted by dividing each compound's gram weight percentage by the molecular formula for that compound. The atomic weights are then added to obtain the total molecular weight for each

sample. Atomic percentages can then be obtained by dividing each compound's atomic weight by the total molecular weight for that sample. TABLE III presents the expected atomic percentages for phases present in each reacted sample.

a. Destabilization of ZrO,

Determination of ZrO_2 destabilization is rather straightforward with a formula which is simply the ratio of monoclinic peak intensity (111) plane, over the summation of monoclinic plus tetragonal peak intensity (101) plane [Ref. 23]:

 $\text{B} Destabilization = M/[M + T] \times 100$

Figure 23 shows a comparison of a reacted and un-reacted sample (sample number one - 80%YSZ-10%Al₂O₃-10%V₂O₅), with marked peak heights for the tetragonal and monoclinic phases, along with the diffraction patterns for tetragonal $2rO_2$, BADDELEYITE (monoclinic $2rO_2$), and cubic $2rO_2$.

An expanded view of Figure 23 is shown in Figure 24. Analysis of all other samples with ten percent V_2O_5 , samples one, three, four, five, six, and seven) indicates that the average destabilization from tetragonal to monoclinic ZrO_2 is nearly 95 percent complete.



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TABLE III.

ANNHALED POWDER SAMPLES OF THE Zro, 8mol. %Y,0,-A1,0,-V,0,

SYSTEM

BY	ATCHIC	PERCENT	

Sample #	žr0,	A1,0,	YVO,	₹208	¥203
1	73.15	12.87	12.78	0.82	
2	82.75	6.49	7.22		3.58
3	62.51	25.43	10.87	1.58	
4	52.12	36.51	9.06	2.29	
5	42.28	47.39	7.35	2.96	
6	7.64	85.68	1.32	5.34	
7	15.66	76.81	2.72	4.79	
8	28.28	39.62	4.91	27.16	
9	34.71	48.63	6.03	10.61	
10	11.39	15.97	1.98	70.64	



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Peak intensity values for monoclinic ZrO_2 from all reacted samples are listed in TABLE IV by increasing atomic percent ZrO_2 . A plot of peak intensity versus atomic percent ZrO_2 is shown in Figure 25. Of significant interest in Figure 25 is the sharp increase in intensity at 28.3 atomic percent, and sharp decrease in intensity at 82.7 atomic percent ZrO_2 . One possibility for the major jump in intensity is the formation of an amorphous glassy phase.

Sample number eight had an abundance of V_2O_5 by weight and atomic percent. It is possible that with a glass phase formation that some phases were "hidden" from XRD analysis and were under-counted, while other phases were overcounted giving the impression that more percent phase is actually present. The second peculiarity on this graph is from sample number two (82.7 atomic percent ZrO_2). This vast drop can be explained by equation (2), and viewing the expected results as shown in TABLE III. The results show that there is a significant drop in monoclinic ZrO₂, however it is noteworthy to say that sample number two is the only sample that calculates excess Y_2O_3 , thereby indicating that all the V_2O_5 that could be consumed was used. A plot of reacted and un-reacted sample number Two raw data files are shown in Figure 26, emphasizing that the monoclinic and tetragonal peaks co-exist after 100 hours at 900°C.

TABLE IV.

PEAK INTENSITY VALUES FOR (111) MONOCLINIC Ero, BY INCREASING ATOMIC PERCENT Ero,

Sample Number	Atomic Percent ErO ₂	Peak Intensity for (111) ErO ₂
6	7.64	894
10	11.39	894
7	15.64	1,574
8	28.28	10,427
9	34.71	3,302
5	42.28	7,156
4	52.12	9,941
3	62.51	11,378
1	73.15	20,173
2	82.75	7,721









b. Ytria Vanadate (YVO,) Formation

As expected YVO₄ formation appears to occur rapidly and correlates well with the destabilization of ZrO_2 . Figure 27 shows the tetragonal peak for YVO₄ at the (200) reflecting plane formed in sample number one. TABLE V presents the peak intensities for all samples by increasing atomic percent YVO₄. A plot of peak intensity versus atomic percent YVO₄ is displayed in Figure 28.

c. $\nabla_{g}O_{g}$ Intensity

Vanadium pentoxide intensity readings support the YVO₄ formation reaction. The values obtained are somewhat sketchy in that two competing factors that impact the V_2O_5 intensity are at work. YVO₄ formation will deplete the amount of V_2O_5 available for diffraction. On the other hand, the remaining V_2O_5 will become more dense and sharpen the intensity peaks observed by the XRD. This analysis is more complicated than the formation transformation from tetragonal to monoclinic ZrO_2 . An example (sample number eight) of the V_2O_5 peak intensity plane (001) is shown in Figure 29.

d. a-11,0, Intensity

Peak intensity calculations for alpha alumina were conducted on all reacted and un-reacted samples at the (113) reflecting plane as shown in Figure 30. Peak intensity values for the (113) reflecting plane are displayed in TABLE VI by increasing atomic percent α -Al₂O₃. A plot of peak intensity

for the un-reacted and reacted samples versus atomic percent α -Al₂O₃ is presented in Figure 31 and presents some interesting results.

In all samples with the exception of two, samples numbers eight and ten intensity values increased after 100 hours of annealing at 900°C. The significance of sample number eight (40%V₂O₅ by weight) and sample number ten (80%V₂O₅ by weight) reporting sharply reduced intensity values corresponds well with the excessive monoclinic ZrO₂ peaks seen in Figure 25, and one of the excessive peaks of tetragonal YVO₄ seen in Figure 27. The logical conclusion would be the formation of an amorphous glassy phase in both of these samples, α -Al₂O₃ being less dense and being hidden by the glassy phase was under-recorded by the XRD.

The three plots $(\alpha-Al_2O_3, ZrO_2, and YVO_4)$ all share one thing in common and that is, the general shape of the plot with increasing atomic percent. The quasi-linear, exponential growth curve arises from the densification due to annealing (sharpening of the beam broadening), and increase in the mass absorption coefficient for the calculated relative intensity due to increasing atomic weight percent. [Ref. 29]

With the exception of samples number eight and ten, by relative measure it appears that α -Al₂O₃ does not react nor interfere with the ZrO₂8mol. Y_2O_3 -V₂O₅ system and corresponding reactions.



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, Plot of un-reacted and reacted sample No.One showing YVO, formation and peak intensity line for YVO, at the (200) reflection plane. Figure 27.

TABLE V.

PEAK INTENSITY VALUES FOR (200) TETRAGONAL YVO, BY INCREASING ATOMIC PERCENT YVO,

Sample Number	Atomic Percent	Peak Intensity
	XA0*	(200) YVO4
6	1.32	850
10	1.98	850
7	2.72	1,081
8	4.91	3,332
9	6.03	1,346
2	7.22	3,786
5	7.35	2,727
4	9.06	4,171
3	10.87	4,456
1	12.78	6,334







. Plot of sample No. Eight V_2O_5 peak intensity (001) reflecting plane. Figure 29.



Figure 30. Comparison of un-reacted and reacted sample No.Three peak intensity values for $(113) \alpha$ -Al₂O₃ reflecting plane.

TABLE VI.

PEAK INTENSITY VALUES FOR (113) α -Al₂O₃ UNREACTED AND REACTED SAMPLES BY INCREASING ATOMIC PERCENT α -Al₂O₃

Sample	Atomic Percent	Unreacted	Reacted
Number	a-Al ₂ 03	Intensity	Intensity
2	6.49	1,819	2,445
1	12.87	2,346	3,595
10	15.97	1,346	605
3	25.43	4,556	12,352
4	36.51	5,920	14,617
8	39.62	5,074	3,248
5	47.39	7,316	19,412
9	48.63	7,236	14,690
7	76.81	18,695	28,599
6	85.68	19,543	22,819



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Figure 31. Plot of peak intensity (113) α -Al₂O₃ for un-reacted and reacted samples versus atomic percent α -Al₂O₃.

B. SEM RESULTS

Two samples were cold mounted from sample number ten $(10\&2rO_2-10\&Al_2O_3-80\&V_2O_3)$. Samples were obtained from the bottom of the Pt crucible after repeated saturation cleanings with concentrated HCl acid. Only one of the two specimens survived final polishing with a 1.0 micron diamond compound. SEM photographs and EDX analysis prove the formation of a glassy phase in samples number eight and ten, thereby substantiating previous explanations for variations in intensity for the ZrO_2 , YVO_4 , and α -Al₂O₃ plots. Figure numbers 32, 33, and 34 show a sequence of pictures taken of the same region and the chemical analysis by EDX for each respective view.

All three pictures show varying concentrations of oxides based upon location of the EDX probe. Figure 32 indicates the presence of all expected reactants with the exception of YVO₄. In fact, YVO₄ does not appear in any of these three pictures. Figure 33 is nearly 100 percent ZrO_2 (view of ZrO_2 agglomerate). Figure 34 taken to the left of the ZrO_2 agglomerate appears amorphous in nature and contains nearly 62 percent V₂O₅ (as compared to expected value by atomic percent calculation of 70.6 percent).



Figure 32. Wide field view of sample number ten (X528). EDX analysis by oxide percent: 22.75% ZrO₂, 15.35% Y₂O₃, 30.7% V₂O₅, and 31.83% Al₂O₃; Analysis by weight percent: 16.84% Al, 16.84%V, 12.09% Y, 16.84% Zr, and 37.83% O.



Figure 33. ZrO₂ agglomerate from sample number ten (close up view of Figure 32), (X1, 32K). EDX analysis by oxide percent: 95.17% ZrO₂, and 4.83% Y₂O₃; Analysis by weight percent: 3.8%Y, 70.46% Zr, and 25.74% O.



Figure 34. View of EDX probe on the matrix surface to the left of the ZrO₂ agglomerate ("1,35K). EDX analysis by oxide percent: 34.13% ZrO₂, 4.32% Y₂O₃, and 61.55% Al₂O₃; Analysis by weight percent: 34.48% V, 3.4% Y, 25.26% Zr, and 36.85% O.

Two very contrasting examples of the crystalline and amorphous nature of sample number ten are presented in Figures 35 and 36 respectively. In particular, Figure 36 shows a wave-like nature to the amorphous phase with the hint of α -Al₂O₃ just below and breaching the surface of the glass.

All of the SEM photographs taken for this sample clearly indicate partial glass formation due primarily to the high concentrations of V_2O_5 in this sample, explaining the behavior of peak intensity fluctuations versus atomic percent of compound in the ZrO_2 , YVO_4 , and α -Al₂O₃ plots (Figures 25, 28, and 31).



Figure 35. Broad view of crystalline structure for sample number ten (X462). EDX analysis by oxide percent: 24.93% ZrO_2 , 7.06% Y_2O_3 , 33.04% V_2O_5 , and 34.97% Al_2O_3 ; Analysis by weight percent: 18.51% Al, 18.51% V, 5.56% Y, 18.46% Zr, and 38.97% O.



Figure 36. Amorphous glassy phase of sample number ten (X236). EDX analysis by oxide percent: 26.88% ZrO_2 , 35.52% V_2O_5 , and 37.6% Al_2O_3 ; Analysis by weight percent: 19.9% Al, 19.9% V, 19.9% Zr, and 40.3%O.
VII. CONCLUSIONS

Investigation of the pseudo-ternary $ZrO_28mol.$ $&Y_2O_3-Al_2O_3-V_2O_5$ system using X-ray diffraction and SEM analysis indicates the following:

- The exposure of powder samples to V_2O_5 at 900°C for 100 hours was sufficient to destabilize the Yttria stabilized zirconia.
- YVO₄ formation occurs rapidly and corresponds in relative intensity measurement to the increase in monoclinic ZrO₂.
- In all the samples analyzed there was no supportive evidence for the formation of zirconium divanadate (ZrV_2O_7) .
- Comparison of α -Al₂O₃ peak intensities at the (113) reflecting plane for unreacted and reacted powder samples indicates that α -Al₂O₃ is non-reactive with the ZrO₂8mol. Y_2O_3 -V₂O₅ compounds. Samples eight and ten showed decreased (113) intensity values due to the formation of an amorphous glassy phase.

VIII. RECOMMENDATIONS

With the appearance that α -Al₂O₃ is non-reactive with the $ZrO_28mol.&Y_2O_3-V_2O_5$ system, the next logical step is to test the strength and fracture toughness values of this composite by fabricating test coupons and subjecting them to operational type conditions, i.e., corrosive environment and thermal cycling to and from operating temperatures.

To address the desire of increased service life from corrosive destabilization, the fabrication process should be one which can best produce a homogeneous microstructure with spatially distributed reinforcements. The surface induced coating method should be thoroughly investigated as this process holds the promise of minimizing the destabilization of tetragonal to monoclinic ZrO_2 from YVO₄ formation.

During the literature review it was apparent that researchers are currently looking at alternative stabilizers to replace yttria oxide. Based on Lewis acid-base theory other stabilizers (which are more acidic than Y_2O_3) like Sc_2O_3 and I_2O_3 have shown promising results in the reduction of VO_4 formation. Again however, even the use of these stabilizers should be looked at with the idea of utilizing a fabrication

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process which could produce the much desired homogeneous microstructure.

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