Role of carbon nanotube dispersion in fracture toughening of plasma sprayed aluminum oxide - carbon nanotube nanocomposite coating

Kantesh Balani
Florida International University

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ROLE OF CARBON NANOTUBE DISPERSION IN FRACTURE TOUGHENING OF PLASMA SPRAYED ALUMINUM OXIDE – CARBON NANOTUBE NANOCOMPOSITE COATING

A dissertation submitted in partial fulfillment of the requirements for the degree of

DOCTOR OF PHILOSOPHY

in

MECHANICAL ENGINEERING

by

Kantesh Balani

2007
To: Dean Vish Prasad  
College of Engineering and Computing  

This dissertation, written by Kantesh Balani, and entitled Role of Carbon Nanotube Dispersion in Fracture Toughening of Plasma Sprayed Aluminum Oxide – Carbon Nanotube Nanocomposite Coating, having been approved in respect to style and intellectual content, is referred to you for judgment.

We have read this dissertation and recommend that it be approved.

______________________________________________  
Kinzy Jones  

______________________________________________  
Norman Munroe  

______________________________________________  
Wenzhi Li  

_________________________  
Sind Agarwal, Major Professor  

Date of Defense: May 10, 2007  

The dissertation of Kantesh Balani is approved.

______________________________________________  
Dean Vish Prasad  
College of Engineering and Computing  

______________________________________________  
Dean George Walker  
University Graduate School  

Florida International University, 2007
DEDICATION

This dissertation is dedicated to everloving memory of my father. He would have been extremely proud of this moment. I dedicate this dissertation to my parents and my maternal uncle.
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ABSTRACT OF THE DISSERTATION

ROLE OF CARBON NANOTUBE DISPERSION IN FRACTURE TOUGHENING OF PLASMA SPRAYED ALUMINUM OXIDE – CARBON NANOTUBE NANOCOMPOSITE COATING

by

Kantesh Balani

Florida International University, 2007

Miami, Florida

Professor Arvind Agarwal, Major Professor

Aluminum oxide (Al₂O₃, or alumina) is a conventional ceramic known for applications such as wear resistant coatings, thermal liners, heaters, crucibles, dielectric systems, etc. However applications of Al₂O₃ are limited owing to its inherent brittleness. Due to its excellent mechanical properties and bending strength, carbon nanotubes (CNT) is an ideal reinforcement for Al₂O₃ matrix to improve its fracture toughness.

The role of CNT dispersion in the fracture toughening of the plasma sprayed Al₂O₃-CNT nanocomposite coating is discussed in the current work. Pretreatment of powder feedstock is required for dispersing CNTs in the matrix. Four coatings namely spray dried Al₂O₃ (A-SD), Al₂O₃ blended with 4wt.% CNT (A4C-B), composite spray dried Al₂O₃-4wt.% CNT (A4C-SD) and composite spray dried Al₂O₃-8wt.% CNT (A8C-SD), are synthesized by plasma spraying. Owing to extreme temperatures and velocities involved in the plasma spraying of ceramics, retention of CNTs in the resulting coatings necessitates optimizing plasma processing parameters using an inflight particle diagnostic sensor. A bimodal microstructure was obtained in the matrix that consists of fully melted
and resolidified structure and solid state sintered structure. CNTs are retained both in the fully melted region and solid-state sintered regions of processed coatings.

Fracture toughness of A-SD, A4C-B, A4C-SD and A8C-SD coatings was 3.22, 3.86, 4.60 and 5.04 MPa m$^{1/2}$ respectively. This affirms the improvement of fracture toughness from 20% (in A4C-B coating) to 43% (in A4C-SD coating) when compared to the A-SD coating because of the CNT dispersion. Fracture toughness improvement from 43% (in A4C-SD) to 57% (in A8C-SD) coating is evinced because of the CNT content. Reinforcement by CNTs is described by its bridging, anchoring, hook formation, impact alignment, fusion with splat, and mesh formation.

The Al$_2$O$_3$/CNT interface is critical in assisting the stress transfer and utilizing excellent mechanical properties of CNTs. Mathematical and computational modeling using ab-initio principle is applied to understand the wetting behavior at the Al$_2$O$_3$/CNT interface. Contrasting storage modulus was obtained by nanoindentation (~ 210, 250, 250-350 and 325-420 GPa in A-SD, A4C-B, A4C-SD, and A8C-SD coatings respectively) depicting the toughening associated with CNT content and dispersion.
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Fig. 4.71: Plasma sprayed A8C-SD coating showing, a) Surface topography, b) Modulus mapping, c) Line scan distribution of storage modulus with distance, and d) Histogram of modulus distribution.
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<table>
<thead>
<tr>
<th>SYMBOL</th>
<th>EXPANSION</th>
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<tbody>
<tr>
<td>$c$</td>
<td>Radial Crack Length</td>
</tr>
<tr>
<td>$f_c$</td>
<td>Capillarity Force</td>
</tr>
<tr>
<td>$h_m$</td>
<td>Meniscus Height</td>
</tr>
<tr>
<td>$k$</td>
<td>Stiffness</td>
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<tr>
<td>$m$</td>
<td>Mass of Sensor</td>
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<tr>
<td>$p$</td>
<td>Pressure</td>
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<tr>
<td>$r$</td>
<td>Radius</td>
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<td>Radial Distance</td>
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<tr>
<td>$t$</td>
<td>Time</td>
</tr>
<tr>
<td>$t$</td>
<td>CNT Wall Thickness</td>
</tr>
<tr>
<td>$t$</td>
<td>Crystallite Size</td>
</tr>
<tr>
<td>$v$</td>
<td>Velocity</td>
</tr>
<tr>
<td>$w$</td>
<td>Displacement Along Axial Direction</td>
</tr>
<tr>
<td>$x$</td>
<td>Displacement</td>
</tr>
<tr>
<td>$\dot{x}$</td>
<td>Rate of Change of Displacement</td>
</tr>
<tr>
<td>$\ddot{x}$</td>
<td>Acceleration</td>
</tr>
</tbody>
</table>

| $A$    | Area |
| $A_c$  | Contact Area of Nanoindenter Tip |
| $B$    | Full Width Half Maximum Broadening (radians) |
| $C$    | Damping of System |
| $C_2$  | constant (1.4388 cm.K) |
| $D$    | Distance |
| $E$    | Young’s Modulus (390 GPa for Al$_2$O$_3$) |
| $E'$   | Storage Modulus |
| $E''$  | Loss Modulus |
| $F_0$  | Sinusoidal Force |
$G$: Free Energy

$G$: Shear Modulus

$H_v$: Vickers’ Hardness

$I_v$: Nucleation Frequency

$K$: Fracture Toughness (MPa m$^{1/2}$)

$K_v$: Kinetic Parameter

$L$: Length of CNT

$N$: Number of Walls in CNT

$P$: Applied Load

$R$: Ratio of Radiant Energy

$T$: Surface Temperature of Radiating Body

$T_s$: Surface Tension

$V$: Volume

$\alpha$: Rhombohedral Al$_2$O$_3$ (hcp)

$\gamma$: Cubic Al$_2$O$_3$ (fcc)

$\gamma$: Surface Energy

$\gamma \cos \alpha$: Vertical Component of Surface Tension

$\delta$: Tetragonal/Orthorhombic Al$_2$O$_3$ (fcc)

$\varepsilon$: Spectral Emissivity

$\eta$: Cubic Al$_2$O$_3$ (fcc)

$\theta$: Monoclinic Al$_2$O$_3$ (fcc)

$\theta$: Bragg Angle

$\theta$: Torsion Angle

$\theta$: Wetting Angle

$\theta'$, $\theta''$: Monoclinic Al$_2$O$_3$ (hcp)

$\kappa$: Orthorhombic Al$_2$O$_3$ (hcp)

$\lambda$: Wavelength

$\sigma$: Axial Stress

$\tau$: Shear Stress
\( v \) : Poisson’s Ratio

\( \chi \) : Hexagonal Al\(_2\)O\(_3\) (hcp)

\( \phi \) : Phase Shift

\( \chi \) : Material independent constant (\(=0.016\))

\( \omega \) : Frequency

\( \Gamma \) : Surface Energy

\( \Omega \) : Surface Projection Height
1. INTRODUCTION

The goal of the proposed research is to improve fracture toughness of nanocrystalline aluminum oxide by addition and dispersion of multiwalled carbon nanotube (CNT) reinforcements using plasma spray technique. Fig 1.1 summarizes the overall research plan.
Aluminum oxide (commercial name alumina, chemical formula \( \text{Al}_2\text{O}_3 \)) is viewed as an ideal material for high temperatures application. However, the extremely brittle nature of \( \text{Al}_2\text{O}_3 \) ceramic limits its applications [1-14]. Hence there is a need to toughen \( \text{Al}_2\text{O}_3 \) for its use in structural applications. Due to their excellent mechanical properties, carbon nanotubes (CNT) has been identified as an ideal reinforcement to toughen the ceramics, including \( \text{Al}_2\text{O}_3 \) [9, 15-31]. Thermal spray processing of ceramic has accomplished superior coatings with improved mechanical properties such as indentation crack resistance, spallation resistance and enhanced fracture toughness [3, 32-36]. Therefore, the plasma spraying technique has been adopted in synthesizing \( \text{Al}_2\text{O}_3 \)-CNT nanocomposite coating with enhanced fracture toughness. But processing and synthesis of CNT reinforced ceramic nanocomposite coating involves several challenges as summarized below:

1.1. Challenges in Fabricating CNT Reinforced Ceramic Nanocomposites

1.1.1. Dispersion of CNTs in the Ceramic Matrix

CNTs have a strong tendency to agglomerate owing to the high surface-area/volume ratio. Dispersion of CNTs offer advantages of excellent mechanical properties since individual CNTs possess such as very high Young’s modulus (~ 1 TPa), and fracture strength (~ 50 GPa) in enhancing the fracture toughness of ceramic nanocomposites [11, 22, 25, 37, 38]. Owing to layer by layer deposition in the plasma spraying, CNT homogeneity in the coatings can be achieved by dispersion of CNTs in the powder feedstock and subsequent coating process. Hence, powder feedstock pretreatment becomes a requirement for dispersing CNTs in the \( \text{Al}_2\text{O}_3 \) matrix. In the present study, the
challenge of dispersing CNTs has been addressed by spray drying of composite ceramic and CNT powder. Consequently, plasma spraying should be optimized in order to retain CNTs in the sprayed coatings.

1.1.2. Retention of Undamaged CNTs in Plasma Sprayed Deposit

Particles passing through thermal plasma plume experience very high temperatures in excess of 10,000 Kelvin [1, 39-41]. In addition, the exiting powder particles attain high velocity from sonic to supersonic regime [1, 39, 41]. Hence retention of undamaged CNTs at such extreme temperatures and impact is quite a challenging problem. Though earlier works have demonstrated successful retention of CNTs in the plasma sprayed metal matrix nanocomposite, ceramic matrix requires enhanced enthalpy plasma parameters which might damage CNTs during processing [42-44]. This challenge has been met by monitoring and controlling plasma parameters of processing via in-flight diagnostics. Owing to rapid solidification behavior of the plasma spraying and short flight time, undamaged CNTs can be retained in the nanocomposite coating.

1.1.3. Grain Growth during Consolidation of Nanocrystalline Ceramic Matrix:

Nanostructure enhances the strength and fracture toughness of the material. However, conventional consolidation techniques result in the grain growth of the nanoparticles [2, 5, 8-10, 17, 21-31, 45-52]. Hence, retention of nanocrystalline structure after consolidation remains a hurdle to overcome. This has been achieved by controlled melting/heating of powder particles that require suitable processing protocol and
optimized consolidation mechanism towards retaining nanostructure. High cooling rates which are inherent to the plasma spraying also aids in retention of nanostructure.

1.2. Research Objectives

The overall objective of current research is to elicit the role of CNT dispersion in the fracture toughening of plasma-sprayed aluminum oxide – carbon nanotube ceramic nanocomposite coating.

The specific research objectives of this work can be summarized as:

- Incorporation and retention of CNT to improve toughness of Al₂O₃-CNT nanocomposite structure via plasma spray forming.
- Dispersion of undamaged CNT in the Al₂O₃ matrix.
- Retention/refinement of nanostructure in Al₂O₃ matrix towards improving fracture toughness.
- Evaluation of bulk mechanical properties such as fracture toughness and elastic modulus of plasma sprayed coating.
- In-depth analysis of interfacial phenomenon and wetting characteristics between Al₂O₃ matrix and CNT reinforcement using experimental technique and molecular modeling of interface via SIESTA 1.3 simulation software.

This dissertation is systematically structured in separate chapters to thoroughly present the background, execution of ideas, assembling of results, scientific interpretation and future scope of the work. Chapter 2 describes the systematic progression of work done by various researchers in extending the scientific knowledge. Adoption of various
techniques and ideas for achieving enhanced fracture toughness is reviewed in this chapter. Chapter 3 describes the methodologies of experiments and analysis adopted in the current work. The heart of the dissertation is the results and discussion, chapter 4, where the toughening enhancement of Al2O3 by CNTs is presented and discussed in detail. Conclusions of the research are stated in chapter 5 summarizing the key findings and important feats. Chapter 6 states the recommendation for future work, which seem potential measures in continued improvement of the current research.
2. LITERATURE REVIEW

2.1. Need of Developing Tough Structural Ceramics

Whether it be structural ceramics or ultra high temperature ceramics for reentry space vehicles, the very competence of ceramics requires: high refractoriness, chemical inertness, resistance to wear, good oxidation resistance, low coefficient of thermal expansion, high thermal conductivity, and good creep properties [1]. But, inherent brittleness of ceramics have created a need for enhancing the fracture toughness and flexural bend strength of the ceramics [1, 3, 4, 12, 49, 53, 54]. The structural ceramics can be broadly classified into non-oxide and oxide ceramics.

2.1.1. Non-Oxide Ceramics

Non-oxide ceramics constitute the borides, carbides, nitrides, and silicides. Ultrahigh temperature ceramics such as HfC, TaC, ZrB$_2$, BN, HfN, TiN/TiB$_2$ and their composites have been used for rocket science and engineering applications that require structural integrity at temperature $> 2100$ K [55]. Toughening in the non-oxide ceramics has been achieved by grain refinement, densification by HIPping (hot-isostatic pressing) and crack-healing agents (such as glassy phase, or presence of microporosity) [8, 48]. A summary of strengthening and toughening mechanisms in non-oxide structural ceramics is presented in Table 2.1. It can be observed that most of the research work and
commercial development of the non-oxide ceramics has been limited to SiC and Si$_3$N$_4$ [45, 48, 50, 56].

Table 2.1: Fracture toughening of non-oxide structural ceramics

<table>
<thead>
<tr>
<th>Composition</th>
<th>Toughening Mechanism</th>
<th>Fracture Toughness</th>
<th>Other features/Comments</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiAlON-SiC</td>
<td>Liquid phase sintering</td>
<td>$\sim 6.0$ MPa $m^{1/2}$</td>
<td>Fine equiaxed grains and dense structure</td>
<td>[45]</td>
</tr>
<tr>
<td>SiC + AlN + Rare earths</td>
<td>Smaller rare earth cations resulting clean boundaries</td>
<td>$\sim 6.5$ MPa $m^{1/2}$</td>
<td>Composition and microstructure of sintering additives affect toughness</td>
<td>[56]</td>
</tr>
<tr>
<td>WC- 0.5 wt % Co – 0.25 wt % VC</td>
<td>Abnormal grain growth reinforcement</td>
<td>7.34 MPa $m^{1/2}$</td>
<td>Toughening by crack-cutting elongated grains</td>
<td>[5]</td>
</tr>
<tr>
<td>Si$_3$N$_4$/SiC (5 wt% Y$_2$O$_3$ with upto 13 wt.% SiC)</td>
<td>Inter and intra SiC nano inclusions</td>
<td>5.8 MPa $m^{1/2}$ (at 8 wt. % SiC)</td>
<td>Hardness increased with increase in SiC content</td>
<td>[50]</td>
</tr>
<tr>
<td>ZrB$_2$–MoSi$_2$</td>
<td>Uniform and fine microstructure</td>
<td>$\sim 2.6$ MPa $m^{1/2}$</td>
<td>Flexural strength $\sim 655$ MPa and 500 MPa at 1200 and 1500 $^0$C respectively.</td>
<td>[10]</td>
</tr>
<tr>
<td>C/SiC</td>
<td>Presence of micro cracks, carbon fiber pullout, homogeneity and non-decomposition of matrix</td>
<td>--</td>
<td>Flexural strength $\sim 124$–287 MPa</td>
<td>[57, 58]</td>
</tr>
</tbody>
</table>

Composite of Si$_3$N$_4$ with inter and intra SiC inclusions in wood-cutting ceramics provided enhancement of mechanical properties (fracture toughness increase from 5.1 to 5.8 MPa $m^{1/2}$ and microhardness increase from 16 GPa to 19 GPa) [50]. In addition,
pressing and sintering techniques (thermal/pressureless/liquid-phase) to manipulate microstructure in ZrB$_2$-MoSi$_2$, SiC, SiAlON-SiC ceramics have proved to be significant in enunciating toughening of ceramic composites [10, 45, 56]. The role of utilizing abnormal grain growth in WC, and development of tough ternary carbide phase ceramics were next big feats in the field of non-oxide ceramic [2, 5, 8, 10, 45-50]. C/SiC composites fabricated by chemical vapor deposition/infiltration and sol-gel technique have emerged for applications such as aircraft brakes, and re-entry shields, rocket nozzles etc [57, 58].

Figure 2.1 demonstrates change of fracture toughness during sintering with addition of rare earth oxides in SiC matrix [56]. Clean interphase boundaries without amorphous inter granular phase, as observed in Fig. 2.2, is resulted by the addition of smaller rare-earth cations [56]. Composition and microstructure of the sintering additives strongly aid the toughening and strengthening of SiC ceramics. Figure 2.3a features increase of hardness of Si$_3$N$_4$/SiC nanocomposite with increase in SiC content [50]. Apart from increasing SiC content, microstructure refinement hinders dislocation movement within Si$_3$N$_4$ matrix to enhance hardness. Liquid phase sintering resulted percolation of SiC nano inclusions in the intergranular regions enhancing the fracture toughness, Fig. 2.3b.
Fig. 2.1: Fracture toughness of SiC ceramics sintered with Re$_2$O$_3$ and AlN [56].

Fig. 2.2: HRTEM micrographs revealing clean interfaces without the amorphous intergranular phase in SiC-AlN-Sc$_2$O$_3$: (a) SiC–SiC boundary and (b) SiC-junction phase boundary [56].
Fig. 2.3: Comparison of a) nano- and macro-hardness and b) fracture toughness of Si$_3$N$_4$/SiC nanocomposites [50].

Figure 2.4 shows Vicker indentation generating radial crack in cobalt bonded WC. Vicker-crack generate due to residual stress relief upon unloading and the balancing relation of stress intensity ahead of crack tip with the crack-termination can indirectly help evaluating the fracture toughness of the material. When the Young’s modulus for ceramics is known (taken as 390 GPa for bulk Al$_2$O$_3$), indentation fracture toughness ($K$) can be calculated by the semi-empirical formula given by Antis, equation 2.1 [59]:

\[
K = \chi \left( \frac{E}{H_V} \right)^{1/2} \left( \frac{P}{c} \right)^{3/2}
\]

Equation 2.1

where \(\chi=0.016\) is material independent constant, \(E\) is the Young’s modulus, \(H_V\) is the Vicker’s hardness, \(P\) is the applied load and \(c\) is the crack length.
Fig. 2.4: Vicker Indentation on WC ceramic depicting origination of radial cracks [5].

Oxide ceramics and their toughening mechanism are reviewed in the next section.

2.1.2. Oxide Ceramics

Among ceramics, the oxides are most useful and common owing to ease of fabrication and stability at high temperatures than nitrides, carbides, sulfides and borides [60, 61]. Oxide ceramics, such as Al$_2$O$_3$, ZrO$_2$, TiO$_2$, Cr$_2$O$_3$, SiO$_2$, and Y$_2$O$_3$ offer advanced technological applications owing to high hardness, resistance to corrosion, and high refractoriness apart from high wear-, fretting-, cavitation- and erosion-resistance, and high dynamic modulus [3, 13, 33, 34, 62]. Though non-oxide ceramics display unique properties as potential candidates for extreme environments, they often require a protective oxide layer to create diffusion barrier for oxidation protection [60, 63, 64].
Oxidation resistance becomes a natural requirement for the materials operating at elevated temperature in air. Aptly oxide ceramics emerge as oxidation-resistant structural materials [60, 64, 65]. High temperature applications introduce large volumetric phase transformations leading to the structural instability. And, inherent brittleness of the ceramic oxides makes them more susceptible to thermal shock failures. Thereby, toughening of the oxide ceramics becomes a prerequisite in their structural applications.

Conventionally, fracture toughening enhancement in oxides has been achieved through phase transformation or by introducing controlled/graded microstructure [18, 66-68]. Incorporation of secondary phases in restricting crack propagation and nanocrystalline structure in enhancing grain sliding has also been prominent in improving the fracture toughness of ceramic nanocomposites [6, 36, 69, 70]. Table 2.2 features fracture toughening mechanisms in oxide ceramics.

Specific examples of Y-ZrO₂ and TiO₂ are considered to elicit toughening mechanisms in oxide ceramics. Initial particle size of Y-ZrO₂ was ~50-200 nm, whereas addition of fine TiO₂ was selected in range of 100-300 nm. Fig. 2.5a shows pressed and sintered (at 1400°C for 4h) Y-ZrO₂ composite [66]. Due to agglomeration tendency of TiO₂ particles, addition of 10 vol.% TiO₂, Fig. 2.5b, resulted increase in inhomogeneity and porosity of the nanocomposite. Increase in fracture toughness with addition of 10 vol.
%TiO$_2$ was attributed to nucleation of Y-ZrO$_2$-ZrTiO$_4$ with more pronounced transformation toughening when compared to without the TiO$_2$ addition [66].

Table 2.2: Fracture toughness and toughening mechanisms for oxide ceramics

<table>
<thead>
<tr>
<th>Ceramic Oxide</th>
<th>Toughening Mechanism</th>
<th>Fracture Toughness</th>
<th>Other features/Comments</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>YSZ (Yttria stabilized</td>
<td>Transformation toughening (tetragonal to monoclinic ZrO$_2$)</td>
<td>4.5 MPa m$^{1/2}$</td>
<td>Additional toughening by ZrTiO$_4$ phase</td>
<td>[66]</td>
</tr>
<tr>
<td>tetragonal zirconia)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nano-TiO$_2$</td>
<td>Crack arrest by nanostructured zones</td>
<td>~ 27 MPa m$^{1/2}$</td>
<td>Isotropic crack propagation, and distribution of agglomerated nanoparticles</td>
<td>[6]</td>
</tr>
<tr>
<td>SiO$_2$</td>
<td>Restricting cristobalite by AlN particles</td>
<td>Upto 2.96 MPa m$^{1/2}$ at 1400 °C</td>
<td>Microcrack deflection and divergence</td>
<td>[71]</td>
</tr>
<tr>
<td>MgAl$_2$O$_4$</td>
<td>Restraining the grain growth by nano Al$_2$O$_3$</td>
<td>7.79 MPa m$^{1/2}$</td>
<td>Low temperature diffusional creep, thermally activated deformation</td>
<td>[69]</td>
</tr>
<tr>
<td>ZrO$_2$</td>
<td>Transformation toughening</td>
<td>10.1 MPa m$^{1/2}$</td>
<td>Addition of secondary toughening by WC, crack deflection</td>
<td>[67]</td>
</tr>
</tbody>
</table>
Fig. 2.5: SEM micrographs of the thermally etched surfaces of the Y–ZrO₂–ZrTiO₄ composites sintered at 1400°C with a) 0 vol.%, and b) 10 vol.% TiO₂ [66].

Fig. 2.6 shows hardness and crack propagation resistance of HVOF (high velocity oxy-fuel sprayed) and APS (air plasma sprayed) TiO₂ coatings. Though hardness of all three processed coatings (viz. air plasma and HVOF sprayed TiO₂, and HVOF sprayed nano TiO₂) is similar, nano TiO₂ demonstrated extremely high fracture toughness when compared to conventional processing, Fig. 2.6 (also see Table 2.2) [6]. Enhanced fracture toughening was attributed to arrest of cracks by nano-structured zones as observed in Fig. 2.7 [6]. Nanozones impede the crack path and restrict the crack extension by absorbing the crack-propagation energy and arresting the crack to result enhanced fracture toughness.
Fig. 2.6: Vickers hardness and crack propagation resistance for coatings made from nanostructured and conventional TiO$_2$ feedstock sprayed via HVOF and APS [6].

Fig. 2.7: Vickers indentation impression (1 kgf) in the cross-section of the HVOF-sprayed nanostructured TiO$_2$ coating (a) and the indentation crack tip being arrested by a zone of nanostructured particles (b) [6].
2.2. Nature and Properties of Aluminum Oxide

Al₂O₃ is white and odorless oxide ceramic (also known as alumina or aluminum oxide), with a hexagonal crystal structure shown in Fig. 2.8. In nature, Al₂O₃ is the hardest mineral after diamond with a hardness of 18-20 GPa. Due to its high hardness and refractory nature, Al₂O₃ is widely used as thermal liners, thermal barrier installations, high temperature insulating systems, crucibles, ceramic boards and brackets, heaters, etc. Crystal structure, physical, thermal, and mechanical properties of Al₂O₃ are presented in Table 2.3.

Fig. 2.8: Crystal Structure of Al₂O₃ (a = b = 4.7564 Å, c = 12.9894 Å, and α = β = 90°, γ = 120°).
Table 2.3: Crystal Structure, physical, thermal and mechanical properties of Al₂O₃

<table>
<thead>
<tr>
<th>Properties of Al₂O₃</th>
<th>Description</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Crystal Structure</td>
<td>Hexagonal</td>
<td>lattice parameters $a = b = 4.7564 \text{ Å}$, $c = 12.9894 \text{ Å}$, and $\alpha = \beta = 90^\circ$, $\gamma = 120^\circ$</td>
</tr>
<tr>
<td>2. Physical Properties</td>
<td>Melting Point</td>
<td>2327 K</td>
</tr>
<tr>
<td></td>
<td>Density</td>
<td>3.99 kg/m³</td>
</tr>
<tr>
<td></td>
<td>Molar Volume</td>
<td>25.554 cm³/mol</td>
</tr>
<tr>
<td></td>
<td>Molecular Weight</td>
<td>101.96 g/mol</td>
</tr>
<tr>
<td>3. Thermal Properties</td>
<td>Coeff. Of Therm. Expansion</td>
<td>$6.76 \times 10^{-6} \text{ K}^{-1}$</td>
</tr>
<tr>
<td></td>
<td>Specific Heat</td>
<td>78.7 J/mol/K</td>
</tr>
<tr>
<td></td>
<td>Thermal Conductivity</td>
<td>36.16 W/m/K</td>
</tr>
<tr>
<td>4. Mechanical Properties</td>
<td>Young’s Modulus</td>
<td>389.5 GPa</td>
</tr>
<tr>
<td></td>
<td>Poisson’s Ratio</td>
<td>0.22</td>
</tr>
<tr>
<td></td>
<td>Hardness</td>
<td>18-20 GPa</td>
</tr>
<tr>
<td></td>
<td>Fracture Toughness</td>
<td>~ 3.2 MPa m¹/²</td>
</tr>
<tr>
<td></td>
<td>- Monolithic</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Plasma Sprayed</td>
<td>~ 2.0 MPa m¹/²</td>
</tr>
</tbody>
</table>

Aluminum oxide has several polymorphs. Among many polymorphs of aluminum oxide, $\alpha$-Al₂O₃ (with space group $R\bar{3}c$) is the most thermodynamically stable form. Other metastable phases can be classified into: (i) face-centered closed (fcc) packing as: cubic ($\gamma, \eta$), monoclinic ($\theta$), tetragonal/orthorhombic ($\delta$), (ii) hexagonal closed packing (hcp) as rhombohedral ($\alpha$), orthorhombic ($\kappa$), and hexagonal ($\chi$). Other monoclinic phases are identified as $\theta'$, $\theta''$, and $\lambda$ [14].

Though atomic stacking for oxygen ion is similar in amorphous and $\gamma$- Al₂O₃, transformation from $\gamma$- to $\alpha$-Al₂O₃ is primarily associated to rearrangement of oxygen sublattice as shown in Fig. 2.9. Difference in the activation energy of amorphous $\rightarrow$ $\gamma$ transition (4.5 eV) compared to the activation energy of $\gamma$ $\rightarrow$ $\alpha$ transformation (5.2 eV) suggests that the mechanism of atomic rearrangements controlling these
Fig. 2.9: (a) Change of stacking of \{111\} spinel oxygen layers from fcc to hcp. (b) Shear of an oxygen layer having aluminum cations in the interstices. (c) Cooperative migration of the cations around regularly distributed vacancies, resulting in a "honey comb" lattice for \(\alpha\)-Al\(_2\)O\(_3\) [14].
transformations is not the same [14]. No experimental confirmations are available till
date to clearly explain direct $\gamma \rightarrow \alpha$ transformation mechanism. But following orientation
relationships are presented between $\gamma$ and $\alpha$ Al$_2$O$_3$: $\{001\}_\gamma \parallel \{0001\}_\alpha$, $\{440\}_\gamma \parallel \{30 \bar{3} 0\}_\alpha$, and $\{310\}_\gamma \parallel \{2110\}_\alpha$ determined from electron diffraction [14].

Transformation concepts are important in understanding the grain growth and
phase evolution of Al$_2$O$_3$ during processing. These aspects become highly important
especially in rapid processing techniques that can form metastable phases, such as
thermal spraying, rapid melt quenching, vapor deposition, laser processing, etc [72-74].
Correspondingly, understanding of solid-state transformations associated with Al$_2$O$_3$
processing become beneficial in attaining required properties via optimizing the
microstructure.

2.3. Ceramics Processing via Thermal Spraying

Thermal spraying was practiced since early 1900s where flame was utilized as
heat source for melting the material [41]. Since the development of plasma spray torch by
Thermal Dynamic Corp. in 1957, *plasma spraying* has established itself in depositing
thick ceramic coatings (>50 $\mu$m). High velocity oxy-fuel (HVOF) spraying, vacuum
plasma spraying (VPS), detonation-gun (D-gun) spraying and cold spraying are other
processes in the family of thermal spray. HVOF utilizes high velocity of carrier gas (3-5
mach) and uses combustion as source of thermal energy. Deposition of coatings in HVOF and D-gun is achieved through plastic deformation with secondary assistance from thermal energy. Hence microstructure is usually observed with minimal porosity (and high density) in HVOF and D-gun sprayed coatings. Deposition of coating in cold spray is achieved primarily through plastic deformation. Hence, cold spray process is limited to deposition of soft metallic coatings. Plasma spraying utilizes thermal energy as the primary source to melt the powders and deposit consolidated coatings. Thermal spraying has evolved as effective processing tool to synthesize ceramic coatings with improved properties such as fracture toughness, indentation crack resistance, spallation resistance against bend and cup test, adhesion strength, abrasive wear resistance and sliding wear resistance [75-84]. Table 2.4 shows a list of ceramics processed by thermal spray processes. Conventional thermal spray processes such as wire arc spraying and flame oxy-fuel spraying, etc are not considered here.

Since plasma spraying of Al₂O₃ based coatings is the focus of current research, HVOF, VPS and D-gun processing of ceramics is not discussed here. Owing to ease of attaining high temperatures in plasma plume (excess of 10000 K) and being able to melt any known material, processing of ceramics has made plasma spraying the first choice [101]. Its ease of use and low cost associated with spraying has helped its wide commercialization. Plasma spraying has emerged as material processing techniques in
Despite of its inception of specifically depositing coatings [41]. Plasma spraying involves several processing parameters such as plasma power, primary and secondary gases types and their flow rates, powder feedstock (size, shape and morphology), feed rate of powder.

Table 2.4: Fabrication of ceramics by thermal spraying

<table>
<thead>
<tr>
<th>Ceramics</th>
<th>Thermal Spraying Processing</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-Oxides</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MoSi$_2$-Si$_3$N$_4$</td>
<td>VPS</td>
<td>[85]</td>
</tr>
<tr>
<td>TiC-Ni</td>
<td>HVOF</td>
<td>[86]</td>
</tr>
<tr>
<td>TaC</td>
<td>VPS</td>
<td>[87]</td>
</tr>
<tr>
<td>SiC/ ZrB$_2$</td>
<td>Controlled APS</td>
<td>[88]</td>
</tr>
<tr>
<td>WC</td>
<td>APS</td>
<td>[89]</td>
</tr>
<tr>
<td>Oxides</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>APS</td>
<td>[90]</td>
</tr>
<tr>
<td>Al$_2$O$_3$/SiC</td>
<td>HVOF, APS</td>
<td>[91]</td>
</tr>
<tr>
<td>Mo-MoO$_2$</td>
<td>APS</td>
<td>[92]</td>
</tr>
<tr>
<td>ZrO$_2$-Al$_2$O$_3$</td>
<td>APS</td>
<td>[93]</td>
</tr>
<tr>
<td>Al$_2$O$_3$-TiO$_2$</td>
<td>APS</td>
<td>[94]</td>
</tr>
<tr>
<td>TiO$_2$</td>
<td>HVOF, VPS</td>
<td>[6, 95-97]</td>
</tr>
<tr>
<td>ZrO$_2$</td>
<td>APS</td>
<td>[98]</td>
</tr>
<tr>
<td>Al$_2$O$_3$-Ni</td>
<td>HVOF</td>
<td>[12]</td>
</tr>
<tr>
<td>HAP*- ZrO$_2$</td>
<td>HVOF</td>
<td>[99]</td>
</tr>
<tr>
<td>HAP*-CNT</td>
<td>APS</td>
<td>[54]</td>
</tr>
<tr>
<td>HAP*</td>
<td>VPS, D-Gun</td>
<td>[100]</td>
</tr>
</tbody>
</table>

(and its flowability), external cooling, coating/formed shape thickness and many more
[41, 101]. In addition secondary parameters such as thermal conductivity of powder
particles, application of external cooling, power fluctuations, inconsistent flow of
powders due to gravity, etc also add to the complexity of process. Following section
discusses plasma spraying in detail.

2.3.1. Plasma Spraying

Plasma is the fourth state of matter, namely ionized gas. An arc is struck between
tungsten cathode and copper anode to generate plasma, where the temperature reaches in
excess of 10,000 K. Plasma spraying involves feeding powder particles in the plasma
stream with pressurized gases creating velocities upto Mach 1-3 [102-104]. A schematic
of plasma spraying process is presented in Fig. 2.10. Powders experience differential heat
(temperature) and velocity owing to their location, size and morphology. Molten/semi-
molten particles exiting from the plasma plume get resolidified and impact on the
substrate to form a coating. Successive deposition lead to increasing thickness of the
layer, which can be subsequently obtained as a free standing structure in case the
substrate is separated from the deposit [87].
2.3.1.1. Plasma Sprayed Nanocrystalline Ceramic Coatings

Plasma spraying has been utilized by several researchers to synthesize nanocrystalline ceramic coatings [33, 73, 75]. With stringent material requirements, ceramic nanocomposites (such as WC-Co, Mo-Si-B, Hydroxyapatite, FeAl, Y₂O₃-ZrO₂, ZrO₂-Al₂O₃, Al₂O₃-TiO₂, etc) have been plasma sprayed for improved friction, wear, biocompatibility, oxidation resistance etc. [36, 73, 81, 90, 105-110]. Nanostructured coatings also provide a solution to improving the fracture toughness of the ceramics. Grain size refinement provides the required strength and improved toughness due to Hall-Petch relationship [51, 52].

Synthesis of nanocrystalline ceramic coating by plasma spraying is associated with two major challenges: (i) grain growth and (ii) lack of flowability of nanosize
powders. Plasma spraying involves rapid cooling rates (~ 10^6 K/s) and formation of fine grain size is inherent to the process [43]. Additional cooling can further prevent grain growth during plasma spraying. However, low mass of nanoparticles and subsequent inability to be carried in gas stream for deposition onto substrate is a major challenge. Flowability of such fine powders during plasma spraying is limited owing to high surface/volume ratio. In order to reduce interparticle friction and enhance flowability, nanoparticles must be constituted into spherical micron sized agglomerates via powder treatment. Thereby, spray drying as necessary powder pretreatment in atomizing nanoparticles as spherical micron sized agglomerates becomes a requirement. Also, the limitation of flowing nanoparticles can be obviated by passing powders as liquid powder feed. Liquid evaporates and leaves the nanoparticles as coating on the substrate. Reduced deposition rates associated with solution precursor plasma spraying makes the process less attractive. Current research has adopted spray drying for achieving good flowability of nanoparticles, which is explained in the following section.

2.3.1.2. Spray Drying of Ceramic Nanopowders

Spray drying is a process utilizing spraying of fluid powder-feed (powder dispersed in solvent- (generally alcohol) or water-based suspensions), and consequent drying (with nitrogen or air) to result spherical agglomerated mass of dry powder [111].
Organic binder is added to keep the solid particles combined with one another. Characteristics of the spray-dried powder are influenced by: type of atomizer, composition and viscosity of suspension, feed rate, drying temperature and atomizing pressure. Spray-dried powders possess high flowability and thereby its coherent flow results high densities and precise thickness in the plasma sprayed structures [112, 113].

Spray drying is a complex process involving (i) droplet formation, (ii) evaporation and ballooning, (iii) droplet explosion, and (iv) particle formation, Fig. 2.11 [111]. When suspension is pumped into the atomizing chamber, it fragments into small droplets by compressed feed gas. Consequent evaporation of solvent causes shrinkage of the droplet in the second stage. During the process, liquid takes along solid particles from core to the outside of the droplet creating a void when evaporation rate inside the droplet exceeds the diffusion rate through the droplet. When pressure exceeds a certain limit, thickness of droplet is not sufficient to hold the ballooning by expansion of gases in its core, causing explosion in some cases. Final stage is recognized by shrinkage from surface tension until it completely dries. Spray dried powders are spherical and free flowing, which are consequently sieved to the ideal powder particle size range of 15-60 μm required for plasma spraying. High quality thermal spraying of micro and nanostructured coatings such as WC-Co, TaC, and Al₂O₃-TiO₂ are described in literature [33, 87, 98, 114].
Plasma spraying of spray-dried powders may reduce interlamellar boundaries/cracks and may give increased thermal corrosion and wear resistance [115-117].

Fig. 2.11: Spray drying of ceramics [111].
Plasma sprayed coatings nano-ZrO₂ exhibited excellent dynamic modulus (~ 170 GPa) and enhanced damping capacity as observed in Fig. 2.12 [13]. TEM micrograph of nano ZrO₂ coating in Fig. 2.13 revealed fine grains in the order of 70-100 nm. Improved damping capacity was attributed to enhanced grain-boundary relaxations.

Fig. 2.12: Damping capacity vs dynamic modulus map of various thermally sprayed samples [13].
Another study by Goberman and Dell focused on plasma spraying of nano Al₂O₃-13 wt% TiO₂ [73, 94]. Secondary electron micrograph (Fig. 2.14a) and backscattered electron micrograph (Fig. 2.14b) elicited bimodal microstructure in nano Al₂O₃-13 wt% TiO₂ coatings. Presence of bimodal microstructure assisted toughening of the coatings. In order to define the processing parameter, a critical plasma spray parameter (CPSP) was defined, as equation 2.2, for the plasma spraying with keeping powder feed rate, gun speed, carrier gas flow rate, spray distance and flow ratio of Ar to H₂ as constant [73]:
Fig. 2.14: a) Secondary and b) Backscattered electron micrographs of plasma sprayed nano-Al$_2$O$_3$-13wt.%TiO$_2$ with additives. (a) Regions of splats (region A) and small particulates (region C), and (b) The backscattered electron image illustrates that the embedded particles appear darker than in the matrix [94].
CPSP = \textit{voltage x current / primary gas (Ar) flow rate} \quad \text{Equation 2.2}

CPSP was related directly to plasma torch/particle temperature. It was evident that metastable phases were inherent in thermal spray processes owing to non-equilibrium nature of fabrication technique [73, 94, 118-120]. Generation of metastable nanocrystalline $\gamma$-Al$_2$O$_3$ phase was observed to vary with varying CPSP, Fig. 2.15 [94].

![X-ray diffraction patterns](image)

Fig. 2.15: X-ray diffraction patterns from (113) $\alpha$-Al$_2$O$_3$ and (400) $\gamma$-Al$_2$O$_3$ peaks for reconstituted Al$_2$O$_3$-13wt.%TiO$_2$ with (modified) additives. The relative integrated intensity of these peaks are observed to vary as a function of CPSP [94].
2.4. Toughening of Aluminum Oxide

Toughening of Al₂O₃ is worked upon by using secondary phases as additives, consolidating nanocrystalline Al₂O₃ powder particles via pressing and sintering, and reinforcing with particulates, fibers or whiskers [35, 121-136]. Following example elicit specific examples of toughened Al₂O₃ ceramic.

Ultrafine grained (1-50 nm) bulk Al₂O₃ ceramic was pressed at 1120 MPa and sintered at 1150°C to evince enhanced resistance to mechanical and thermal shock [121]. Moreover, metastable phases of Al₂O₃ such as gamma and theta nucleate as secondary precipitates and assist in precipitation strengthening of Al₂O₃ ceramic[122-124]. Li et al demonstrated that nano α-Al₂O₃ can be obtained using two-step sintering, Fig. 2.16 [121]. Along with sintering additives, sintering time and temperature have a strong effect on the generated microstructure making sintering an exciting technique to alter the microstructure and attain enhanced structural properties in nanoceramics [125, 126]. Since fracture toughness of Al₂O₃ is lower than stress required to initiate plastic deformation, role of additives in enhancing the energy absorption (by generating banded structure, Fig. 2.17) becomes more prominent [127, 128]. Table 2.5 lists role of various additives in enhancing the fracture toughness of Al₂O₃ ceramic composites. Though introduction of fibers, particulates and whiskers was highly initiated, the recent trend is using CNTs as an effective reinforcement, and is discussed in detail in the next section.
Fig. 2.16: The relative density of the Al₂O₃ nanoceramics pressed at 1120 MPa and sintered at different temperatures for 5 h as a function of mean particle size of the starting α-Al₂O₃ nanopowders [121].

Fig. 2.17: TEM micrograph of an Al₂O₃–1 mol% TiO₂–1 mol% MgO solid solution sintered at 1500°C for 5 h in air [127].
Table 2.5: Role of additives in enhancing fracture toughness of Al$_2$O$_3$

<table>
<thead>
<tr>
<th>Ceramic/Metal</th>
<th>Additive (P/F/W)*</th>
<th>Fracture Toughness</th>
<th>Toughening Mechanism</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al$_2$O$_3$</td>
<td>Yttria Stabilized Zirconia (P)</td>
<td>8.9 MPa.m$^{1/2}$</td>
<td>Ferroelastic domain switching and transformation toughening</td>
<td>[125]</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>SiC (W)</td>
<td>3.8 MPa.m$^{1/2}$</td>
<td>Crack healing by SiC whisker</td>
<td>[35]</td>
</tr>
<tr>
<td>MgAl$_2$O$_4$</td>
<td>Al$_2$O$_3$ (P)</td>
<td>2.5-5.82 MPa.m$^{1/2}$</td>
<td>Grain boundary sliding and creep diffusion</td>
<td>[129]</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>Yttrium Aluminum Garnet (P)</td>
<td>4.54 MPa.m$^{1/2}$</td>
<td>YAG inclusion leading to residual stress. Reduction of grain and flaw size.</td>
<td>[34]</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>MgSiO$_3$</td>
<td>3.3 MPa.m$^{1/2}$</td>
<td>Stepped and micro-faceted cleavage</td>
<td>[130]</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>Monoclinic Zirconia (P)</td>
<td>11.8 MPa.m$^{1/2}$</td>
<td>Phase transformation toughening</td>
<td>[131]</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>CeO$_2$ (P)</td>
<td>4.7 MPa.m$^{1/2}$</td>
<td>Particulate toughening</td>
<td>[132]</td>
</tr>
<tr>
<td>Ni</td>
<td>Al$_2$O$_3$ (F)</td>
<td>10-35 MPa.m$^{1/2}$</td>
<td>Metallic Bridging layer</td>
<td>[133]</td>
</tr>
<tr>
<td>Al, Al-Cu</td>
<td>Al$_2$O$_3$ (P)</td>
<td>6.5-30 MPa.m$^{1/2}$</td>
<td>High stress triaxiality between narrowly spaced ceramic particles, ductile matrix, and high Young’s modulus</td>
<td>[134, 135]</td>
</tr>
<tr>
<td>Al</td>
<td>Al$_2$O$_3$ (F)</td>
<td>~4.8 MPa.m$^{1/2}$</td>
<td>Plastic deformation of the metal, testing at 1200 °C</td>
<td>[136]</td>
</tr>
</tbody>
</table>

2.5. Carbon Nanotubes as Reinforcement for Nanocomposite for Improving Mechanical Properties

Nanocomposite is a novel class of materials that combines two or more materials, physically and/or chemically distinct, in creating a resultant material in which size of one of the phases is less than 100 nanometer [137]. The advantage of using nanocomposites is achieving the best qualities of the constituted material and often some, which are even superior to those of constituent materials possess.

Carbon nanotube (CNT) is a new class of material that has superior mechanical properties with theoretical tensile strength of 200 GPa and Young's modulus reaching ~1 TPa [11, 22, 25, 37, 38]. CNTs have also displayed superplasticity despite their sp\(^2\) bonding [138]. Superplastic deformation of upto 280% in Fig. 2.18, hints CNT as an ideal reinforcement for toughening ceramics. Such high deformation is attributed to kink nucleation and movement apart from atomic movement to heal vacancies [138]. These properties of CNTs make them suitable candidate for as reinforcement for creating nanocomposites with improved mechanical properties.

Wavelike distortions were observed in repeated bending of the CNTs with local strains of upto 16% without failure, Fig. 2.19 [139]. The bending modulus of CNTs decreased with the increasing diameter of CNTs. Sustained integrity at large strain with no sign of brittleness, plasticity or atomic arrangement inferred good resilience [140]. Hence CNTs serve as potential reinforcements for improving ductility in the ceramic nanocomposites. In addition, excellent mechanical properties of CNTs do not restrict their participation in enhancing the strength of polymer-CNT and metal-CNT nanocomposites as discussed in the consequent sections.
2.5.1. CNT Reinforced Polymer Matrix Nanocomposites

Most of the research on CNT reinforced nanocomposites has been carried out on polymer matrix. CNT reinforcement in matrices such as UHMWPE (ultra high molecular weight polyethylene), PMMA (polymethylmethacrylate), PS (polystyrene), PAN (polyacrylonitrile), PMEMA (Methyl-ethyl methacrylate copolymer), etc have been utilized [38, 141-146]. The main purpose of adding CNTs in polymer matrix had been to improve the elastic modulus and tensile strength via effective load transfer between polymer and CNT. Table 2.6 summarizes CNT reinforced polymer nanocomposites and
improvement in mechanical properties [142-146]. CNT dispersion, polymer-CNT bonding, and CNT pullout are dictated as the main reasons of strengthening the nanocomposite. Chemical interaction at the CNT-polymer interface are critical and affect the toughening of polymer-CNT nanocomposite.

Fig. 2.19: TEM image of bent CNT with A and B) radius of curvature ~ 40 nm, and C) amplitude of ripples increase at the outer layer [140].

Polymer-CNT reinforcement has shown enhanced role of CNT in load transfer, as seen in Fig. 2.20 via CNT pulling and bridging [141]. Modulus and strength improvements of
Table 2.6: CNT-reinforced polymer nanocomposites

<table>
<thead>
<tr>
<th>Polymer</th>
<th>CNT Loading</th>
<th>Microstructural Features</th>
<th>Mechanical Property</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epoxy</td>
<td>0.5 wt. %</td>
<td>CNT pullout. Intimate CNT contact with matrix. Ductile deformation in fractured surface.</td>
<td>Increase in Young’s modulus by 200% (upto 10.9 MPa). Tensile strength increase by 140% (upto 0.44 GPa) (softer epoxy matrix)</td>
<td>[142]</td>
</tr>
<tr>
<td>Polypropylene</td>
<td>1 wt. %</td>
<td>Well dispersed CNTs. Good wetting observed.</td>
<td>Enhancement of Young’s Modulus (from 5.2 to 8.0 GPa) and tensile strength increased from 7.9 to 9.8 GPa.</td>
<td>[143]</td>
</tr>
<tr>
<td>PMMA (poly methyl methacrylate)</td>
<td>4 wt.%</td>
<td>Lack of chemical interaction between PMMA and CNT. Weaker interfacial adhesion resulted higher toughness.</td>
<td>No improvement in Young’s Modulus. Considerable improvement in impact resistance (from 40 to 160 KJ/m²)</td>
<td>[144]</td>
</tr>
<tr>
<td>Epoxy</td>
<td>5 wt.%</td>
<td>CNT aggregation limited reaching theoretical predicted values. CNT bundles easily slide pass one another.</td>
<td>Elastic Modulus increase by 75 % (~ 7 GPa) and hardness increase by 30 % (~0.42 GPa)</td>
<td>[145]</td>
</tr>
<tr>
<td>Polystyrene</td>
<td>1 wt. %</td>
<td>Good CNT dispersion. Good interfacial bonding</td>
<td>~ 42 % increase in elastic modulus (from 1.19 to 1.69 GPa) and ~25% increase in break stress (from 12.8 to 16 MPa)</td>
<td>[146]</td>
</tr>
</tbody>
</table>
upto 200% and toughness improvements of upto 150 % have been reported in epoxy-
CNT nanocomposites [141]. Since matrix provides a secondary role in bearing the stress,
stress transfer to the reinforcing CNT fiber is highly critical [147, 148]. Importance of
CNT incorporation as a dispersed phase is impressed upon by various research works,
with dispersion being achieved by sol-gel technique, surface activation, inorganic
coating, heterocoagulation, ball milling, solvent incorporation, etc [28, 149-155].

Fig. 2.20: CNT ropes are observed bridging a fatigue fracture surface in an epoxy matrix
[141].

Though literature has continually reported enhancement of tensile strength and
modulus experimentally, these values are far below the theoretical predictions. The
difference between experimental observations and theoretical predictions has been
attributed to poor dispersion of CNT in the matrix and lack of understanding in engineering CNT/matrix interface [38, 141, 144]. Next class of nanocomposite is CNT reinforced metal-matrix composite as explained in section 2.5.2.

2.5.2. CNT Reinforced Metal Matrix Nanocomposites

Comparatively limited research has been performed on metal matrix-CNT nanocomposites [156-158]. Basic challenge arising in the metal-CNT nanocomposite is associated with damage to CNTs at high temperature processing, as required for metal processing [43, 44, 118]. Secondary challenge includes low interfacial bonding strength between CNT and metal interface [156-158].

CNT addition in Al and Ti matrix has shown improvement in elastic modulus and hardness [43]. Spark plasma sintered and isostatically pressed Cu-CNT nanocomposites depicted enhanced wear resistance upto three times whereas hardness increased two times [159, 160]. A summary of improvement in mechanical properties of metal-CNT nanocomposites is presented in Table 2.7.

CNT dispersion in the metal matrix is critical to impart stress transfer towards improving strength, and elastic modulus [164]. Dangling CNTs in the fractured surface of the plasma sprayed Al-Si/CNT nanocomposite is presented in Fig. 2.21a [44] indicating strong anchoring by CNTs. Smooth CNT surface indicated no physical damage during plasma spraying. Survival of CNTs was possible because of rapid kinetics inherent to the plasma spray processing. Laha et al has investigated CNT/metal interface in Al-Si/CNT system [43, 44, 118]. It was concluded that a thin layer of SiC on CNT is formed (Fig. 2.21b) suggesting interfacial reaction which improved interface wettability required for
Table 2.7: Mechanical property enhancement in metal-CNT nanocomposite

<table>
<thead>
<tr>
<th>Metal</th>
<th>CNT content</th>
<th>Microstructure</th>
<th>Property</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu</td>
<td>1 vol. %</td>
<td>Severe plastic deformation (Equi channel angular pressing). Homogeneous deformation</td>
<td>Increase in indentation load by upto 20 % (from 56 to 75 kgf for 300 μm deformation.</td>
<td>[156]</td>
</tr>
<tr>
<td>Sn-Ag-Cu</td>
<td>0.07 wt.%</td>
<td>Extruded. Addition of CNTs enhanced the wetting behavior. Large difference in CTE induced multidirection thermal stress.</td>
<td>Improvement in yield strength from 31 to 36 MPa, microhardness increase from 16.3 to 17.0 Hv</td>
<td>[157]</td>
</tr>
<tr>
<td>Ti</td>
<td>20 vol. %</td>
<td>Hot pressed. Dispersion strengthening, refinement of grain size, and TiC formation</td>
<td>Hardness increase from 221 to 1216 Hv, and Young’s Modulus increase from 120 to 198 GPa.</td>
<td>[158]</td>
</tr>
<tr>
<td>Mg</td>
<td>2 wt. %</td>
<td>Hot pressed. Uniform CNT dispersion in Mg matrix. Improved interface bonding and effective load transfer. Upto 98 % dense.</td>
<td>Young’s modulus improvement by 9% (from 35.3 to 38.6 GPa)</td>
<td>[161]</td>
</tr>
<tr>
<td>Ag</td>
<td>8 vol. %</td>
<td>Pressing and sintering. Uniform distribution of CNTs. Agglomeration observed for higher CNT content (&gt;10%). CNT pullout, CNT bridging, and CNT alignment.</td>
<td>9% increase in bend strength (from 427 to 465 MPa)</td>
<td>[162]</td>
</tr>
<tr>
<td>Al-Si</td>
<td>10 wt.%</td>
<td>CNT retention after plasma spraying. SiC layer over CNT surface</td>
<td>Elastic modulus increased from 79 to 114.5 GPa. UTS of ~ 234 MPa</td>
<td>[43, 44, 163]</td>
</tr>
</tbody>
</table>
effective load transfer between metal/CNT interface. CNT reinforced ceramic nanocomposites are discussed in the next section.

2.5.3. CNT Reinforced Ceramic Nanocomposites:

Incorporation of fibers and CNTs in the ceramic matrix has shown promising results in improving the fracture toughness of ceramic matrix. [9, 17, 21-31]. Toughening behavior of CNT reinforced matrix is mainly linked to CNT crack bridging, CNT dispersion and crack deflection [17, 165, 166]. It has been theoretically dictated that spherical particles do not contribute to toughness of ceramic composites [18]. Hence research directed to using non-spherical particles/fibers/nanotubes in enhancing the fracture toughness of ceramic nanocomposites is well justified. Temporal evolution of CNT reinforced ceramics by various processing techniques is presented in Table 2.8.
Hot pressing of Fe-Al₂O₃-CNT composite depicted continuous network of CNTs between grains, a pronounced effect of CNT straightening before fracture was expected to show enhanced fracture toughening [9]. Study on Si₃N₄-Al₂O₃-Y₂O₃-CNT nanocomposite showed improvement in modulus of elasticity by 15%, whereas bending strength increased by 37% [21]. Peigney et al studied hot pressing of MgAl₂O₄-CNT composite with improper CNT dispersion and reinforcement effect was not to the order that was expected out of the system [31]. Ning et al stressed on the requirement of improvement in CNT dispersion to enhance the strengthening by CNTs [28].

Mechanical aspects of load transfer at CNT-matrix interface have been relatively poorly studied in these studies without detailed experimental evidence. Computational tools have been utilized in grasping a better idea of interfacial mechanism in CNT/ceramic system [17, 147, 165, 181-183]. Table 2.9 is extracted from Table 2.8 as a subset presenting details of investigations on some of the CNT-ceramic composite system. Table 2.9 explains the processes involving high temperature sintering consolidation of various ceramic nanocomposites. Different degree of strengthening and fracture toughening were elicited such as poor cohesion between matrix and CNTs, straightening of CNTs and improved CNT dispersion. Since the direct attention is given to the Al₂O₃-CNT nanocomposites in the current work, detailed review on this system follows in the consequent section.

2.6. Aluminum Oxide-CNT Nanocomposite

Aluminum oxide-CNT nanocomposites for improved fracture toughness have been the focus of recent investigations [31, 37, 184]. Hot pressing, spark plasma
Table 2.8: Evolution and development of CNT-reinforced ceramics by various processing techniques

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<tbody>
<tr>
<td>Surfactant assisted processing</td>
<td>UC Dev</td>
<td>Ning,</td>
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<td>Sol Gel</td>
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<td>Molecular level mixing</td>
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<td>Thermal CVD/Spray Pyrolysis</td>
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<td>Liquid Precursor</td>
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sintering, sol-gel synthesis, surfactant assisted processing, Catalytic CVD, hot isostatic pressing, molecular level mixing, spray pyrolysis, high temperature extrusion, liquid precursor spraying, colloidal processing, etc, have been used to fabricate Al₂O₃-CNT nanocomposite [16, 17, 37, 128, 154, 155, 166-168, 171, 172, 177, 178, 180]. Siegel et al were the pioneers in processing Al₂O₃-CNT nanocomposite and utilized hot pressing as consolidation technique [16]. Improvement of upto 24% in fracture toughness was achieved in Al₂O₃-CNT nanocomposite [16]. Zhan et al. adopted spark plasma sintering and evinced tremendous improvement in the electrical conductivity (from 10⁻¹² to 10⁵0 S/m) and fracture toughness (by three times) of Al₂O₃-CNT nanocomposite [185]. Wang et al adopted the similar methodology as that of Zhan et al., but did not observe the similar enhancement in fracture toughness [172]. Damage tolerance of Al₂O₃-CNT nanocomposite led to absence of Vicker indentation cracks [172]. Disparity of results were attributed to dissimilar densities and residual compressive stresses in the Al₂O₃-CNT nanocomposite [186]. Mo et al observed toughening of Al₂O₃-CNT nanocomposite by crack bridging [176]. An outlook and description of Al₂O₃-CNT nanocomposite by various researchers is presented in Table 2.10.

Effort of CNT dispersion by colloidal processing and consolidation by hot pressing was studied by Sun et al [11]. CNT pullout was observed in the fractured surface of the Al₂O₃-CNT composite, Fig. 2.22. Hot pressed Al₂O₃-CNT nanocomposite displayed 10% improvement (from 496 to 554 MPa) in the bend
Table 2.9: Description of ceramic-CNT nanocomposites research

<table>
<thead>
<tr>
<th>Peigney (Univsite Paul-Sabatier, France) [9]</th>
<th>Balazsi (Hungarian Academy of Sciences, Hungary) [21]</th>
<th>Peigney (Univsite Paul-Sabatier, France) [31]</th>
<th>Ning (Chinese Academy of Sciences, China) [28]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ceramic</td>
<td>Fe-(\text{Al}_2\text{O}_3)</td>
<td>Si(_3\text{N}_4) 90%-(\text{Al}_2\text{O}_3) 4%-(\text{Y}_2\text{O}_3)-6 %</td>
<td>Mg (\text{Al}_2\text{O}_4)</td>
</tr>
<tr>
<td>CNT</td>
<td>Arc Discharge (SWNT + MWNT) 1.5-15 nm. Mostly 2-3 walls CNT (Grown in-situ)</td>
<td>Catalytic decomposition of acetylene on aluminum oxide supported Co/Fe catalyst</td>
<td>SWNT (CCVD): In-situ</td>
</tr>
<tr>
<td>Mixing/Processing Method</td>
<td>Hot Pressing (1475°C) in vacuum (Thermal treatment optimization)</td>
<td>Ball-milled in ethanol (several hours), ultrasonicated, Pressing: 220 MPa, Sinter 1700°C</td>
<td>Hot Pressing (1475°C)</td>
</tr>
<tr>
<td>Measurement</td>
<td>Surface Area to calculate CNT content</td>
<td>Modulus of Elasticity, Bending Strength</td>
<td>conductivity studies</td>
</tr>
<tr>
<td>Fracture Toughness</td>
<td>1.8x1.8x16 mm specimen</td>
<td>Not done (Sample: 3.5x5x50 mm)</td>
<td>Not done</td>
</tr>
<tr>
<td>Key Results</td>
<td>1050°C, 6 min thermal treatment found optimum cohesion between CNT and matrix is poor.</td>
<td>Deterioration of carbon fibers (ref. Sample) was observed 15-37% improvement in mech. properties. Pressure, sintering time and temperature resulted disappearance of MWNT</td>
<td>continuous network of CNT between oxide grains</td>
</tr>
<tr>
<td>Other</td>
<td>Quality and Quantity defined by author for defining CNT</td>
<td>Better to shorten sintering time and pressure.</td>
<td>Increased electrical conductivity with CNT</td>
</tr>
</tbody>
</table>
Table 2.10: Description of Al₂O₃-CNT nanocomposites research

<table>
<thead>
<tr>
<th></th>
<th>Amiya (UC Davis) [186]</th>
<th>Siegel (RPI, New York) [16]</th>
<th>Padture, Tanaka (U.Connecticut, CT) (NIMS, Japan) [172]</th>
<th>Mo (KAIST, Korea) [176]</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Vendor</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td><strong>Al₂O₃ Powder</strong></td>
<td>(80%α + 20%γ) Al₂O₃</td>
<td>Nanophase Tech. (39 nm average diameter)</td>
<td>Baikowski Intl (80%α + 20%γ) Al₂O₃</td>
<td>Al-tri-sec-butoxide</td>
</tr>
<tr>
<td><strong>Morphology</strong></td>
<td>300 nm (40nm Crystallite) and 20 nm: Irregular</td>
<td></td>
<td>300 nm (40nm Crystallite) and 20 nm: Irregular</td>
<td></td>
</tr>
<tr>
<td><strong>Vendor</strong></td>
<td>Carbon Nanotechnology (HIPco process)</td>
<td>MWNT: Electric Discharge method/ Oxidation ~640°C</td>
<td>Carbon Nanotechnology (Same as Zhan)</td>
<td>MWNT:10-30 nm; 10-50μm CVD:aluminum oxide supported</td>
</tr>
<tr>
<td><strong>Processing Method</strong></td>
<td>Ethanol + ultrasonic agitation Ball Milling (ZrO₂ ball/ 24h)/ SPS</td>
<td>Ethanol + ultrasonic agitation, dried and crushed/ Hot Pressed at T: 1300°C, 1h, 60 MPa</td>
<td>Methanol + ultrasonic agitation / Ball Milling (ZrO₂ ball/ 24h)/ SPS</td>
<td>CNT dispersed in Ethanol + Aluminum oxide sol (Yolda’s Process) Hydrolysis and Peptization of Al-hydroxide (AIOOH). SPS (1650°C, 5min); sintered at 1000C, 6h</td>
</tr>
<tr>
<td><strong>Electrical Conductivity (S/m)/ Density (g/cc)</strong></td>
<td>Al₂O₃: 10⁻¹⁰ - 10⁻¹² 10 vol. % SWNT: 1510 S/m</td>
<td>Al₂O₃ + 4 vol. % CNT : Best Wear Resistance</td>
<td>Al₂O₃: 3.898 g/cc 10 vol. % SWNT: 3.568 g/cc</td>
<td>Drying 350°C, 6h; calcination 1250°C, 1h</td>
</tr>
<tr>
<td><strong>Strength/Fr. Tough. (MPa m⁻²)</strong></td>
<td>Al₂O₃ : 3.33 10 vol % SWNT: 9.7</td>
<td>Al₂O₃ : 3.4 10 vol% MWNT: 4.2 (5 kg)</td>
<td>Al₂O₃ : 3.22 10 vol.% SWNT: 3.32</td>
<td>Al₂O₃: 1.0 (Normalized) 1.5 vol% MWNT : 1.1</td>
</tr>
<tr>
<td><strong>Key Results</strong></td>
<td>No Vicker cracking Highest Indentation toughness at Al₂O₃ / 10 vol. % CNT Significant inc. in Electrical conductivity</td>
<td>24% Increase in Fr. Toughness (Al₂O₃ + 10 vol % MWNT) Damage delocalization (well dispersed MWNT)</td>
<td>-Enhanced contact damage resistance</td>
<td>Toughening of CNT reinforced aluminum oxide crack bridging effect of CNT</td>
</tr>
<tr>
<td><strong>Other</strong></td>
<td>Did not present toughening mechanism</td>
<td>No discussion on interfacial properties</td>
<td>Clear explanation for the result is missing</td>
<td>Does not includes discussion on the bonding of CNT with matrix</td>
</tr>
</tbody>
</table>

46
strength [11]. Only marginal improvement from 3.9 to 4.0 MPa m$^{1/2}$ was observed in the fracture toughness of the nanocomposite [11]. Marginal improvement in the toughness is attributed to damage of CNTs at high temperature processing as intimated by Peigney et al [9]. In addition, low relative density (~ 85-95 %) possibly reduced the fracture toughness of hot-pressed nanocomposites.

High temperature extrusion (1500$^\circ$C at 43 MPa) elicited retention of undamaged CNTs in the CNT–Fe–Al$_2$O$_3$ and CNT–Fe/Co–MgAl$_2$O$_4$ composite, respectively (Fig. 2.23a and b) [9]. This effort was directed for adjusting the electrical conductivity value in the composite [9]. Higher electrical conductivity (~ 20 S/m) was observed along extrusion direction, whereas it dropped to 0.6 S/m along transverse direction. Alignment of CNTs was responsible towards imparting such a difference in the electrical conductivity [9]. Presence of CNTs also assisted in restricting grain growth and serving as lubricating agent during hot extrusion. No mechanical property data was discussed in the paper, but retention of undamaged CNTs and CNT alignment via extrusion insinuates possible anisotropic property enhancement [9].

With many publications in processing nanocrystalline aluminum oxide, Zhan et al, and Duan et al demonstrated improved fracture toughness, of upto three times than that of aluminum oxide, using spark plasma sintering consolidation of CNT-Al$_2$O$_3$ composite, Fig 2.24 [125, 166, 171, 186-191]. However, the primary interest of Zhan et
al was also concentrated on enhancing electrical properties using CNTs in the aluminum oxide matrix [166]. Electrical conductivity enhancement from $10^{-10}$ S/m to 3345 S/m was observed in 15 vol. % CNT-Al$_2$O$_3$ nanocomposite. This conductivity increase was because of CNT addition, and formation of CNT-ropes along grain boundaries creating network of electrically conductive paths [166]. Dispersion of CNTs along Al$_2$O$_3$ grains is shown in Fig. 2.25 [166]. Later, Wang et al repeated the methodology of the Zhan et al and showed that toughness of CNT-Al$_2$O$_3$ matrix was not improved [172]. The
Fig. 2.23: SEM images of fracture profiles of the CNT–Fe–Al$_2$O$_3$ (a) and CNT–Fe/Co–MgAl$_2$O$_4$ (b) composites prepared by extrusion at 1500°C. Note the alignment of the CNTs [9].
discrepancy in data was because Zhan et al had used indentation methods to calculate fracture toughness, whereas the Al₂O₃-CNT composite shows contact-damage resistance, which was tracked through Hertzian indentation [172]. This unusual mechanical behavior was attributed to shear-deformable nature of CNTs in redistributing stress field upon indentation. Also, different densification achieved by Zhan (> 99%) and Wang (~ 95%) induced discrepancy [185]. Generation of crack was limited by high porosity in Wang’s

![Fracture toughness versus carbon nanotube volume content in aluminum oxide based composites as reported in literature[166]. (SWCNT: Single walled CNT).](image)

Fig. 2.24: Fracture toughness versus carbon nanotube volume content in aluminum oxide based composites as reported in literature[166]. (SWCNT: Single walled CNT).
Fig. 2.25: TEM Micrographs of 5.7 vol.% SWCN/ Al₂O₃ nanocomposite. (a) Bright-field TEM image and (b) high-resolution TEM image of specimens in the fully dense 5.7 vol.% SWCN/Al₂O₃ nanocomposite. The arrows indicate the SWCN phase [166]. (SWCNT: Single walled CNT).
work, since indentation initiated collapse of crack before the induction of plastic deformation in the material. On the other hand, Vicker indentation cracks were clearly observed in Zhan’s paper. Moreover, high residual compressive stress (2.0 GPa) in the alumina matrix, crack bridging, fiber pull-out and crack deflection were responsible for improved fracture toughness of spark plasma sintered CNT-Al₂O₃ nanocomposite.

Surface assisted processing of CNTs followed by hot pressing (25 MPa at 1300°C) helped increasing the bending strength (from 51.5 to 97.0 MPa) and fracture toughness (from 1.0 to 2.46 MPa m¹/²) of SiO₂ ceramic [28]. CVD processing of CNT-Al₂O₃ composite was elucidated by Xia et al along with computational modeling of CNT-Al₂O₃ interface demonstrating CNT pullout, Fig. 2.26a, and crack deflection at CNT-Al₂O₃ interface, Fig. 2.26b [17, 178]. Defects, residual stresses, alignment of crack path, and deflection of CNTs were stressed upon by researchers towards estimating the fracture toughness of the composite [17].

High aspect ratio of CNT diameter caused severe agglomeration. Hence a new technique for CNT dispersion was required. Catalytic CVD method was adopted by Rul et al for dispersing CNTs in MgAl₂O₄ matrix [29]. Insitu CCVD synthesis followed by hot pressing at 1300°C (43 MPa in vacuum of 10⁻⁴ torr) retained the CNT distribution in the nanocomposite. CNTs inhibited grain growth and improved the electrical conductivity by seven orders of magnitude (from 10⁻¹⁰ to 4 x 10⁻³ S/cm) [29]. Variety of
Fig. 2.26: a) SEM photograph of a fracture surface of the 20 μm-thick sample demonstrating CNT pullout, longitudinal delamination, and residual holes; (b) crack deflection at CNT along interface [17].
samples were produced, out of which the nomenclature of the samples CMH, C5, C10, CM2, and CM12 is presented in Table 2.11 A very well distributed network of CNTs was observed in the composite powders. Increased CNT content resulted restricted full densification, which was attributed to grain growth inhibition by CNTs. Fine grained matrix and widely interconnected CNT network was depicted in the fractured surface of nanocomposite. Fractured surface illustrated straightening of CNTs before breaking during the fracture, Fig. 2.27, which might show enhanced fracture toughness because of CNT pullout and CNT bridging [29].

Spray pyrolysis emerged as in-situ technique for fabrication of aluminum oxide-CNT composites [179]. Here crack deflection and nanotube debonding served as site for fracture energy dissipation [179]. Molecular level mixing utilized functionalization of CNTs reacting with metal ions in solution [177]. Hardness of the Al₂O₃-CNT nanocomposite improved from 1500 Hv to 1700 Hv. Addition and dispersion of functionalized CNTs was responsible for strengthening and toughening enhancement of upto 25% and 15% respectively for CNT-Al₂O₃ nanocomposite [177]. CNT pullout and CNT bridging was also observed in the fractured surface with most of CNTs residing at grain boundary. Hence, CNT reinforced the matrix via load sharing and bridging mechanism [177]. Chemical bonding was observed between CNT and amorphous Al₂O₃ matrix, which imparted such increase in mechanical properties.
Table 2.11: Nomenclature for powders and hot-pressed MgAl$_2$O$_4$ samples

<table>
<thead>
<tr>
<th>Sample</th>
<th>Co (wt.%</th>
<th>Mo (wt.%</th>
<th>Preparation Mode</th>
<th>CNT content (wt.%)</th>
<th>Density (%)</th>
<th>Conductivity (S/cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMH</td>
<td>20</td>
<td>10</td>
<td>Powder (Urea)</td>
<td>0</td>
<td>97.3</td>
<td>$10^{-10}$</td>
</tr>
<tr>
<td>C5</td>
<td>1</td>
<td>0</td>
<td>Powder (Urea)</td>
<td>0.55</td>
<td>100</td>
<td>0.0072</td>
</tr>
<tr>
<td>C10</td>
<td>20</td>
<td>0</td>
<td>Gelcasting Foam (Urea)</td>
<td>3.70</td>
<td>92.7</td>
<td>0.530</td>
</tr>
<tr>
<td>CM2</td>
<td>20</td>
<td>10</td>
<td>Powder (Urea)</td>
<td>5.10</td>
<td>83.8</td>
<td>1.00</td>
</tr>
<tr>
<td>CM12</td>
<td>10</td>
<td>5</td>
<td>Gelcasting Foam (with Citric Acid)</td>
<td>12.20</td>
<td>68.0</td>
<td>8.53</td>
</tr>
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</table>

Sol-gel technique was utilized to counteract the van der Waals attractive force between CNTs in the gel network. Sol-gel techniques depicted homogeneous dispersion in the aluminum oxide matrix without agglomeration, Fig. 2.28 [176]. The process was followed by spark plasma sintering to consolidate the Al$_2$O$_3$-CNT composite. Strong CNT bonding with matrix as CNT pullout was observed in the fractured surface indicating significant stress transfer from matrix to CNT. Enhanced density (> 99.5 %), significant load transfer between matrix to CNTs, and bridging effect of CNTs during crack propagation were reasoned for enhanced fracture toughness (by 10% for 1.5 vol. % CNT) of the Al$_2$O$_3$-CNT nanocomposite [176].
Fig. 2.27: High resolution SEM images of the fracture surface of the hot-pressed composites: (a) Sample without CNT showing mainly transgranular fracture. (b)–(e) Samples C5, C10, CM2 and CM5, respectively, showing CNT emerging out of the surface fracture. C10 and CM2 show restricted grain-growth effect (and intergranular fracture). (f) Sample CM12 showing many CNT emerging out of large pores and straightened before breaking during the fracture [29].
Fig. 2.28: The micrographs of carbon nanotube reinforced aluminum oxide composite powders. (a) The optical micrographs of dried gel and (b) the SEM micrographs of calcinated carbon nanotube/aluminum oxide composite powders [176].
2.7. What’s Missing and What’s the Challenge?

A review of the Al₂O₃-CNT nanocomposite in the previous section leads to the following conclusions:

- Improvement in the fracture toughness has been achieved by addition of CNTs to the Al₂O₃ matrix.
- Uniform dispersion of CNTs in the Al₂O₃ matrix still remains a challenge.
- Al₂O₃-CNT nanocomposite processing/consolidation techniques are limited to solid state processing (e.g. HIP, sintering, spark plasma sintering, and extrusion).
- The interface between Al₂O₃/CNT is relatively less understood. A few works are limited to computational studies without sufficient experimental evidence.

Based on observations and limitations from other researchers’ work, the present work addresses those specific challenges by synthesizing Al₂O₃-CNT nanocomposite coating by plasma spraying. In addition to the above listed challenges, this work also focuses on the specific challenges which are unique to plasma spraying of Al₂O₃-CNT system. Some of these challenges are listed below:

- Improved CNT Dispersion: Achieving CNT dispersion in the powder makes powder treatment becomes a necessity in exercising them as required feedstock.
• Dispersed CNTs allow interfacial interactions and load-transfers towards attaining best properties in ceramic nanocomposites. Consequently, *layered deposition typical in plasma spraying will further improve CNT dispersion* in the processed coating. Hence plasma spraying of pretreated (spray-dried) powder feedstock is incorporated in the current research to obtain coatings with improved fracture toughness.

• Bimodal Matrix Structure: Presence of second phases act as strengthening medium, thereby bimodal phase/grain generation can further enhance the fracture toughness of the nanocomposite. Consequently, *controlling the bimodal microstructure* becomes an immense challenge to overcome. Bimodal grain/phase can be obtained by *imminent control of plasma processing parameters* to result *surface melting followed by resolidification and consolidation of core by solid state sintering*. Further, extreme temperature (> 10000 K) and impact (~ 1-3 Mach velocity) associated with the plasma spraying creates an additional challenge of *retaining CNTs in the processed coating* [1, 39-41].

• Al$_2$O$_3$/CNT interface: Most of the ceramic-CNT processes involve “solid state processing” wherein aluminum oxide and CNT are both in solid state. But plasma spraying involves alumina in molten state, so wetting between alumina and CNT becomes highly important. This very novel concept of wettability of Al$_2$O$_3$/CNT
interface is to be conceptualized, but currently there is no work dictating the interfacial mechanism occurring at the Al\textsubscript{2}O\textsubscript{3}-CNT interface. And limited research on the aluminum oxide-graphite interface elicits non-wetting of the interface [90, 192-197]. Hence aspects like wettability, interfacial bonding, load transfer mechanism, etc. remain unanswered. Herein, ab-initio computer modeling of Al\textsubscript{2}O\textsubscript{3}-CNT interface has been incorporated towards eliciting interactive response at the Al\textsubscript{2}O\textsubscript{3}/CNT interface.

Figure 2.29 summarizes the Al\textsubscript{2}O\textsubscript{3}/CNT nanocomposite research efforts by various groups, their conclusions and remaining challenges.
Fig. 2.29: A bird’s view of Al₂O₃/CNT nanocomposite research efforts by various groups, their conclusions and remaining challenges.
3. EXPERIMENTAL PROCEDURE

The aim of the current work is to fabricate Al$_2$O$_3$-CNT nanocomposite coating via plasma spraying with uniform dispersion of CNTs in the matrix for enhanced fracture toughness. Various methodologies of powder treatment are adopted for dispersion of CNTs in the powder feedstock.

3.1. Pretreatment of Powder Feedstock

Since nano size of the powders tend to clog the nozzle during plasma spraying, as-received nano Al$_2$O$_3$ and CNTs have been pretreated to result powder feedstock ideal for plasma spraying. Powder feedstock treatments prior to plasma spraying is explained in this section.

3.1.1. As Received Al$_2$O$_3$ Powder and CNTs

Nano α-Al$_2$O$_3$ powder (99.8 % pure, 150 nm particle size, and 40 nm crystallite size) was obtained from Inframat Corporation, Farmington, CT. Smooth and irregular faceted morphology of the as-received Al$_2$O$_3$ powder is presented in Fig. 3.1. Owing to increased surface area of the faceted nano particles, increased friction causes clogging of the nozzle during plasma spraying. Hence to reduce the clogging tendency of fine particles, powder treatment becomes necessary to reduce the contact area by consolidating them as spherical agglomerates.

Multiwall carbon nanotubes (95%+ purity, OD 40-70 nm, 0.5-2.0 μm in length) were obtained from Nanostructured and Amorphous Materials Inc., Houston, TX. CNT
diameter of about 70 nm is observed in the TEM image, Fig. 3.2, with a wall thickness of \(~25-30\) nm. In order to tap superior mechanical properties of CNTs, it becomes imperative to uniformly disperse CNTs in the matrix. But strong agglomerating tendency of CNTs require special treatments such as functionalization, molecular level dispersion, ultrasonication etc.

![Fig. 3.1: As-received Al₂O₃ powder particles.](image)

3.1.2. Spray Drying of As-received Al₂O₃ Powder (A-SD):

Commercial spray drying process was utilized to obtain spherical agglomerates of Al₂O₃ particles (or mixture of Al₂O₃ and CNTs). Nanosize powder particles are dispersed in an aqueous organic binder to form slurry. The slurry is passed through an atomizing orifice, which mechanically binds the fine particles as spherical agglomerates. Resulting spherical agglomerates cause reduced interparticle friction, improved
flowability and avoid clogging of powders in the nozzle during plasma spraying. Sieved spherical agglomerates in the size range of 15-60 μm were obtained that are ideal for plasma spraying. Three powder treatment methodologies have been adopted to prepare the powder feedstock for consequent plasma spraying.

![Fig. 3.2: TEM of Multiwalled CNTs.](image)

Pure α-Al₂O₃ (99.8%) was spray dried to obtain spherical agglomerates for powder feedstock (referred to as A-SD). Fig. 3.3a shows the typical size range of 15-60 μm spherical agglomerates, and an enlarged image, Fig. 3.3b elicits the mechanically bounded porous cake of fine powders.
3.1.3. Blending of Spray Dried Al₂O₃ with 4 wt.% CNT (A4C-B):

Spray dried Al₂O₃ agglomerates were blended with 4 wt.% CNT in a jar mill (US Stoneware, East Palestine, OH) for 24 hrs to obtain dispersion of CNTs in Al₂O₃ matrix (referred to as A4C-B powder). Schematic and SEM morphology of as-received Al₂O₃, spray-dried Al₂O₃ agglomerate and resulting blended AC-B powder has been shown in Fig. 3.4. CNTs are observed to disperse onto (i) surface of agglomerate spray dried Al₂O₃ powder and (ii) dome-cavity regions of spray dried powder.

3.1.4. Spray Drying of Composite Al₂O₃ and CNT:

Further refinement of CNT dispersion to what obtained in A4C-B powder can be achieved by spray drying of composite Al₂O₃ and CNT powder in a single step. Two powder compositions, namely Al₂O₃- with 4 wt. % CNT and 8 wt. % CNT were used for composite spray drying.
Fig. 3.4: Process schematic and SEM images of a) nano Al₂O₃, b) spray dried Al₂O₃ (A-SD) agglomerates, and c) spray dried Al₂O₃ blended with 4 wt. % CNT for 24 h to result nanocomposite A4C-B powder feedstock.

Al₂O₃ powder and 4 wt.% CNTs (A4C-SD):

Spray drying of composite Al₂O₃ powder and 4 wt.% CNTs (referred to as A4C-SD) produced spherical agglomerates of 15-60 μm, Fig. 3.5a. In contrast to surface distribution of blended powder (Fig. 3.4), CNTs are also dispersed both at the powder
agglomerate surface and the inside of the powder. No agglomeration of CNTs was observed along the powder surface, Fig. 3.5b. CNTs were also distributed uniformly inside the powder agglomerate, as seen in the fractured powder Fig. 3.5c.

**Al₂O₃ powder and 8 wt.% CNTs (A8C-SD):**

Spray dried composite Al₂O₃ and 8 wt.% CNTs (A8C-SD) powder agglomerates show spherical morphology with size range of 15-60 μm, Fig. 3.6. Alike A4C-SD powder, uniform CNT dispersion is achieved in the powder agglomerate. Increase in the CNT content of the spray dried powder (from 4 wt. % to 8 wt. %) is an extra parameter to analyze effect of CNT content in improving the fracture toughness of the plasma sprayed structure (apart from its different dispersion powder treatment when compared to A4C-B powder). Fracture surface of A8C-SD powder, Fig. 3.7, illustrates the non-agglomeration tendency of the spray-dried powder inside the powder agglomerate. This becomes significant since CNTs are dispersed non-preferentially everywhere uniformly in the powder agglomerate.

**3.2. Plasma Spraying of Al₂O₃-CNT Nanocomposite Coating**

Atmospheric plasma spraying, Fig. 3.8, of various powders is done using SG 100° plasma gun (Praxair Surface Technologies, Indianapolis, IN). Overall features of plasma spraying are listed in Table 3.1. Plasma processing parameters were optimized in terms of coating microstructure (partially melted, fully melted, and porosity), CNT distribution and fracture toughness. A representative set of plasma processing parameters is listed in Table 3.2. Consequently, A-SD, A4C-B, A4C-SD, A8C-SD powders were
Fig. 3.5: SEM image of A4C-SD powder showing a) powder agglomerates, b) dispersion of individual CNTs, and c) fractured surface showing inside of the powder.
Fig. 3.6: Spray dried A8C-SD powder showing a) spherical agglomerates, and b) uniform CNT dispersion on surface.
Fig. 3.7: Fracture surface of A8C-SD powder indicating uniform CNT dispersion in the core of the powder agglomerate.

Fig. 3.8: Schematic of plasma spraying process.
Table 3.1: Features of plasma spraying

<table>
<thead>
<tr>
<th>Features of Plasma Spraying</th>
<th>Description of the Feature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heat source</td>
<td>Plasma arc with gun voltage and current 35 V and 800 A, respectively</td>
</tr>
<tr>
<td>Spray gun</td>
<td>SG-100® plasma gun, Praxair Surface Technologies, Indianapolis, IN</td>
</tr>
<tr>
<td>Particle velocity</td>
<td>75-750 m/s [198-201]</td>
</tr>
<tr>
<td>Temperature</td>
<td>10,000-15,000 K [198, 200-202]</td>
</tr>
<tr>
<td>Gases used</td>
<td>Primary gas: Ar Secondary gas: He Carrier gas: Ar</td>
</tr>
</tbody>
</table>

Table 3.2: Representative processing parameters for plasma spraying

<table>
<thead>
<tr>
<th>Plasma Spraying</th>
<th>Volts (V)</th>
<th>Current (Amp)</th>
<th>Power (kW)</th>
<th>Feed Rate (rpm)</th>
<th>Primary Gas (slm')</th>
<th>Secondary Gas (slm')</th>
<th>Carrier Gas (slm')</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al₂O₃ with CNT coatings</td>
<td>35-40</td>
<td>600-780</td>
<td>18.5-32.3</td>
<td>2.5-3.0</td>
<td>Ar (32.1)</td>
<td>He (59.5)</td>
<td>Ar (19.8-21.7)</td>
</tr>
</tbody>
</table>

*slm: standard liters per minute

plasma sprayed onto AISI 1020 steel substrate (100 x 20 x 3 mm³ coupon) for detailed evaluation. A typical plasma sprayed coating is presented in Fig. 3.9.

Control of plasma parameters can be attained by inflight monitoring of powder particles in plasma plume using AccuraSpray (Tecnar Automation Ltée, QC, Canada), Fig. 3.10. This allows extra leverage in understanding the relation of plasma parameters responsible for generating resulting coating and microstructure. Optical fiber measures the particle velocity and temperature by controlled distance traveled by particle and the differential wavelength emission respectively. Particle velocity is indicative of degree of flattening of particles whereas particle temperature suggests its heated/melted state.
Degree of melting and flattening of particles directly constitute the microstructure of the coating that affects its properties.

Fig. 3.9: Typical plasma sprayed coating showing uniform coating thickness.

Fig. 3.10: AccuraSpray: Inflight particle diagnostic sensor: (a) Controller with display monitor, (b) the sensor head, and (c) a screenshot showing temperature, velocity and plume profile.
The particle velocity is measured based on a time of flight technique. The sensor head images two optical fibers in space. As the particles pass through these images incandescent light is gathered and transmitted to photo multiplier tubes that generate an electronic pulse that corresponds to the particle passing through the fiber images, Fig. 3.11. The figure below shows a representation of the two pulses generated as the particle passes in front of the first and second fiber. A trigger level is set on the slope of each pulse to start and stop a timing clock. The velocity, $v$, is given by,

$$v = \frac{D}{t_2 - t_1}$$  \hspace{1cm} Equation 3.1

where $D$ is the physical spacing of the two fibers imaged at the measurement volume and $t_2 - t_1$ is the measured time it takes for the particle to fly past the two fibers.

Fig. 3.11: Detection of particle velocity from the time traveled within fixed distance between two optical fibers.
The basic premise behind all radiation thermometry is Plank's Law, which describes the emissive power of a radiating body as a function of wavelength, emissivity and temperature. Dual-wavelength (ratio or two color) pyrometry involves the measurement of the spectral energy in two different wavelength bands. Using Wien’s approximation to Planck's law the ratio of radiant energy, R, in two different wavelength bands, λ₁ and λ₂, is given by:

\[
\frac{I_{\lambda_1}}{I_{\lambda_2}} = R = \frac{\varepsilon_{(\lambda_1,T)}}{\varepsilon_{(\lambda_2,T)}} \left(\frac{\lambda_2}{\lambda_1}\right)^5 \exp \left[\frac{C_2(1/T) - (\lambda_1/\lambda_2)}{C_2}\right] \quad \text{Equation 3.2}
\]

where \(T\) is the surface temperature of the radiating body, \(C_2\) is a constant (= 1.4388 cm.K), and \(\varepsilon_{(\lambda,T)}\) is the spectral emissivity. AccuraSpray utilizes a CCD camera in capturing the plume intensity of plasma plume.

And solving for \(T\) gives:

\[
\frac{I}{T} = \ln \left[\frac{\varepsilon_{(\lambda_1,T)}}{\varepsilon_{(\lambda_2,T)}} \left(\frac{\lambda_2}{\lambda_1}\right)^5 \frac{I_{\lambda_2}}{I_{\lambda_1}}\right] = \frac{C_2}{\left(\frac{1}{\lambda_1} - \frac{1}{\lambda_2}\right)} \quad \text{Equation 3.3}
\]

Temperature and velocity of the powder particles is major factor in deciding the development of microstructure in a coating. Density, degree of melting, cooling rate, degree of impact, etc are deciding factors in deposition of a coating. Hence access to the inflight particle diagnostic data is beneficial in relating the microstructure in terms of experienced thermal history.
3.3. Microstructural and Phase Characterization

3.3.1. Optical Microscopy and Quantitative Analysis of CNT Dispersion

Quantitative and morphological analysis of plasma sprayed coatings is performed using an optical microscope (Versamet 3, Buehler Ltd, IL, USA) with an attached CCD camera (Leica DFC 320). CNT dispersion in the powder feedstock was measured using Image-Pro ® Plus, Version 5.1 imaging software (2004 Media Cybernetics Inc.) on SEM images. Development of microstructures in terms of fully melted, partially melted and porosity in the plasma sprayed coatings were quantified using imaging software. Partially melted region is recognized with solid state sintering of the particles showing necking/sintering at the particle interface. Fully melted region is characterized by melting and resolidification of the powder particles. Porosity is the region of voids (inter-splat, intra-splat or unfilled regions). Quantification of such features helps in assimilating the role of regions in enhancing the fracture toughness and optimizing the plasma processing parameters.

SEM images were used in analyzing CNT dispersion through Image-Pro imaging software. Quantitative image analysis illuminates the nature of CNT dispersion in the various treated powders. Angular alignment of CNTs in the dome cavity or surface-dispersion of CNTs in the powder agglomerate allows significant inference to the powder behavior during plasma spraying and consequent coating microstructure. Moreover, uniform CNT dispersion also becomes obvious through the quantitative analysis depicting uniform spreading of CNTs homogeneously in the matrix.
3.3.2. Phase Analysis in Powder Feedstock and Plasma Sprayed Coatings

Phase analysis of the powders and coatings was carried out using a Siemens 500D X-ray Diffractometer (XRD) with CuK\textsubscript{\alpha} radiation (of 1.54 Å) operating at 40 kV and 20 mA. A graphic software DIFFRAC\textsuperscript{plus} EVA (Version Rev 0, Bruker axs, Madison, WI, USA) was used to analyze the XRD spectra. Crystallite size ($t$) measurement was carried out using Scherrer equation, equation 3.4. Standard NIST Si powder sample was employed to determine the instrumental broadening and the calibration of the XRD patterns of powder samples using Gaussian profile.

\[
t = \frac{0.9 \lambda}{B \cos \theta}
\]

Equation 3.4

where $\lambda$ is the incident wavelength, and $\theta$ is the Bragg angle, and $B = \sqrt{B_m^2 - B_s^2}$, where $B_m$ and $B_s$ are full width half maximum (FWHM) broadening of specimen and standard sample respectively.

3.3.3. Confirmation of CNT Retention by Raman Spectroscopy

Micro-Raman spectroscopy of the powder feedstock and plasma sprayed nanocomposites was carried out to validate the carbon nanotube structure. Ti-sapphire crystal target with a laser wavelength of 785 nm was used for this purpose. The laser was produced using a laser source from Spectra Physics (Model 3900S, California, USA) and the detector was from Kaiser Optical Systems, Inc. (Michigan, USA).
3.3.4. **Microstructural Characterization of Al\textsubscript{2}O\textsubscript{3}-CNT Nanocomposite**

Coating morphology and CNT dispersion in the powders and plasma sprayed deposits was investigated using FESEM JEOL JSM 6330 F scanning electron microscope. Samples were metallographically polished to 0.5 µm using diamond slurry for preparing sample surface. No sample preparation was done for imaging powders and fractured coating surfaces. These were consequently gold coated (using Pelco SC-7 Auto Gold Sputter coater) for 30 seconds to allow earthing of electronic charge. SEM was operated at 15 kV and 12 µA for microstructure observation. Elemental analysis of the composite was determined by performing energy dispersive spectroscopy (EDS) integrated with the FESEM.

Transmission electron microscope (TEM) was used to analyze the CNT distribution and observe Al\textsubscript{2}O\textsubscript{3}-CNT interface in the plasma sprayed coatings. Two different types of TEMs were used in characterizing the coatings, viz. (i) Philips PW 6061 TEM system (model CM 200, Eindhoven, Netherlands), and (ii) FEI Technai F30 high-resolution transmission electron microscope operated at 300 kV. Samples for TEM observation were fabricated by two methods: (i) crushing and dispersion in ethanol followed by ultrasonification (Branson 2510, Danbury, CT), and (ii) metallographic thinning, dimpling (Gatan, Inc. Model 656 Mk3, California, USA) and ion milling (Gatan Precision Ion Polishing, Model 691, California, USA).
3.4. Mechanical Testing

3.4.1. Vicker Indentation Toughness

Zhongguo HXD-100 TMC Shanghai Taiming Optical Instruments microhardness tester (200g load and 15s dwell time) was used for estimating hardness and fracture toughness of the plasma sprayed samples using indentation technique. Student t-test was utilized, with better than 95% confidence level, towards comparing the statistical difference between the fracture toughness values for eight indentations on each sample. Fracture toughness was calculated from the Anti’s semi-empirical relationship based on radial crack generation during Vicker indentation [203].

3.4.2. Nanoindentation

The elastic modulus values of the as-sprayed nanocomposites in modulus mapping mode were evaluated by the nanoindentation technique. Indentation experiments were conducted using Hysitron TriboIndenter® (Hystrion Inc., Minneapolis, MN). Three-sided Berkovich diamond indenter was used for indentation. The load and displacement data obtained from the tests were analyzed using the methods reported by Oliver and Pharr [204]. The initial calibration of the instrument was done using a standard aluminum and fused silica sample provided by Hysitron.

Modulus mapping is performed using dynamic mechanical analysis approach of applying cyclic stress in compression. Quasi-static load is applied to the probe tip with smaller dynamic load (at a predescribed frequency (200 Hz). Dynamic load is then analyzed to measure amplitude and phase shift of the original signal. The equation of motion for sinusoidal force ($F_0$) is given as:
\[
F_0 \sin(\omega t) = m \ddot{x} + C \dot{x} + k x 
\]

where \( m \) is the mass of sensor, \( k \) is the stiffness, and \( C \) is the damping of the system. And displacement response is given as:

\[
x = X \sin(\omega t - \phi) 
\]

Solution to the differential equations are:

\[
X = \frac{F_0}{\sqrt{(k - m \omega^2)^2 + (C \omega)^2}},
\]

and

\[
\phi = \tan^{-1} \left( \frac{\omega C}{k - m \omega^2} \right) 
\]

rearranging the equations, \( k \) and \( C \) can be calculated by:

\[
k = \frac{F_0}{X} \frac{1}{\sqrt{1 + \tan^2 \phi}} + m \omega^2
\]

and

\[
C = \sqrt{\frac{(F_0/X)^2 \tan^2 \phi}{1 + \tan^2 \phi} \frac{1}{\omega}}
\]

During instrument set up, stiffness, damping and mass of sensor are evaluated by frequency sweep of sensor in air. Stiffness and damping of the instrument can be directly subtracted to obtain actual stiffness and damping of the tested sample. Correspondingly, storage modulus \( (E') \) and loss modulus \( (E'') \) can be calculated as following:

\[
E' = \frac{k \sqrt{\pi}}{2 \sqrt{A_v}}
\]

Equation 3.11
\[ E'' = \frac{\omega C_s \sqrt{\pi}}{2 \sqrt{A_c}} \]  

Equation 3.12

where \( A_c \) is contact area, which is dependent on the contact depth. Contact depth of the indenter is described through tip area function during instrument calibration.

3.5. Ab-initio Molecular Modeling of Al₂O₃-CNT Interface

SIESTA (Spanish Initiative for Electronic Simulations with Thousands of Atoms) modeling scales \( \text{Al}_2\text{O}_3\)-CNT interface was created with 1 x 1 x 1 crystal lattice of alpha-alumina interfacing 2 x 2 x 2 crystal layers of graphite. Plane wave basis set was used for the interfacial system, limiting in z-direction and periodicity in x-y direction with cell size of 4.928 x 4.928 x 26.4114 Å³ and \( \alpha=90^0, \beta=90^0, \) and \( \gamma=120^0 \). Standard Kohn Sham self-consistent density functional was utilized with LCAO (Linear combination of atomic orbitals) basis set in local density approximations. Spin polarized Ceperly-Adler scheme (Perdew and Zunger) was used for defining Al, O and C exchange correlation functional. Improved Troullier Martins pseudo potential generation was employed to describe nonlocal, and normconserving interaction between core and valence electrons. First principle pseudopotentials were generated from spin polarized non relativistic ground state components of Kleinman and Bylander projectors. Al was defined with ground state \( 3s^2 \) \( 3p^1 \) \( 3d^0 \) with cutoff 1.86, 2.25 and 3.07 bohrs respectively [192]. Oxygen was stated as \( 2s^2 \) \( 2p^4 \) \( 3d^0 \) \( 4f^0 \) with cutoff 1.15, 1.15, 1.15, and 1.15 bohrs respectively, whereas Carbon was described through \( 2s^2 \) \( 2p^2 \) ground state with cutoff 1.50 and 1.54 bohrs respectively [192, 205].
Localized spin density (LSD) Hamiltonian was calculated by matrix diagonalization to generate self-consistent Kohn Sham solution. Conjugate gradient (CG) method was used for coordinate optimization with limiting force of 0.05 eV/Å or 50 iterations whichever came first. Maximum displacement during CG optimization run was limited to 0.2 bohr.
4. RESULTS AND DISCUSSION

Effect of CNT dispersion in enhancing the fracture toughness of Al₂O₃-CNT nanocomposite has been conferred to in this section. Role of powder pretreatment and optimized plasma spraying parameters in retaining and distributing CNTs are elucidated with underlying consolidation mechanism. Microstructural effect on mechanical property is undertaken to confirm the fracture toughness enhancement. Wettability studies and computational modeling is pursued for understanding the interfacial behavior between Al₂O₃ matrix and CNT reinforcement. Subsequent sections enunciate the overall behavior of plasma sprayed Al₂O₃-CNT nanocomposite interlinking the CNT dispersion with generated microstructure and toughening mechanisms.

4.1. Powder Treatment of n-Al₂O₃ as Ideal Feedstock for Plasma Spraying

Nano-Al₂O₃ powder was spray dried to result A-SD powder (explained in section 3.1.2). Though the powder particles are agglomerated as spherical cake in the size range of 15-60 μm, there is no change in the phase of the treated powder particles, and no chemical reaction occurred during the spray drying. Identity of initial powder particles is maintained during the spray drying as confirmed by XRD spectrum, Fig. 4.1 and Raman Spectrum, Fig. 4.2. This powder (A-SD) is used as a feedstock for optimizing the plasma spraying parameters.

Spray dried aluminum oxide powder is referred to as A-SD from now onwards, and details of initial particle size, spray drying treatment, and adopted nomenclature are presented in Table 4.1.
Fig. 4.1: XRD spectrum of as-received and spray dried n-Al2O3 powder indicating no phase change and chemical reaction.

Fig. 4.2: Raman spectrum showing similar peaks in the as-received and spray dried n-Al2O3 powder.
Nano scale surfaces are known to be unstable because of their high surface to volume ratio, and have strong tendency towards agglomeration [206]. During mechanical blending of the A-SD and CNT powder mixture, agglomeration and settling of CNTs in the dome cavity of A-SD powder is observed in the scattergram, Fig. 4.3a, via Image Pro quantitative analysis. Agglomeration of CNTs in the histogram, Fig. 4.3b, endorses the higher CNT content in the dome cavity when compared to the CNTs adhering at the surface of the powder agglomerate. Though CNTs are dispersed in A4C-B powder to some extent, further improvisation of CNT dispersion can have tremendous impact on improving mechanical properties of the nanocomposite. This is achieved by composite spray drying of the n-Al₂O₃ and CNT. Laplace equation (of free energy change with particle radius) clearly elicits the instability of nanoparticles with the decreasing radius of particles. But, increasing the surface area by contacting two unstable surfaces reduces the overall energy of system to a lower value. CNTs, when mixed with nano-Al₂O₃ powder particles (during composite spray drying), reduce the surface charges by physically attaching to open surfaces[207]. Because of high surface area of nano-Al₂O₃ particles, CNTs do not find difference in associating itself with nano-Al₂O₃ [207]. Surface energy of CNTs range about 0.2 J/m², whereas Al₂O₃ particles have surface energy of ~ 1.59 J/m² [194]. It has been shown that Al-graphite interface can bring down the energy in the 0.02-0.4 J/m² range [194]. Nano-Al₂O₃ powder particles thereby aid dispersion of CNTs in the solid state mixing of nanocomposite powder [208]. Random surface availability therefore results in the uniform dispersion of CNTs. Distribution of CNTs in the A4C-SD and A8C-SD powder feedstock as a function of their aspect ratio is presented in Fig. 4.4 and Fig. 4.5 respectively. Figure 4.4 and Fig. 4.5 show non-preferential distribution of
Table 4.1: Initial powder size, spray drying treatment and adopted nomenclature

<table>
<thead>
<tr>
<th>Initial Powders</th>
<th>Powder Treatment -Step 1</th>
<th>Powder Treatment -Step 2</th>
<th>Resulting Powder</th>
<th>CNT Dispersion</th>
<th>Plasma Sprayed Coating Nomenclature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al₂O₃ (150 nm particle size)</td>
<td>Spray Drying of Al₂O₃ (agglomerate ~15-60 μm)</td>
<td>---</td>
<td>A-SD (Al₂O₃ agglomerate ~ 15-60 μm)</td>
<td>---</td>
<td>A-SD Coating (Al₂O₃)</td>
</tr>
<tr>
<td>CNTs (Multi walled, OD 40-70 nm, 0.5-2.0 μm long)</td>
<td>Spray Drying of Al₂O₃ (agglomerate ~15-60 μm)</td>
<td>Blending with 4 wt.% CNTs for 24 h</td>
<td>A4C-B (Al₂O₃ - 4 wt.% CNT agglomerate ~ 15-60 μm)</td>
<td>Onto surface and dome cavity</td>
<td>A4C-B Coating (Al₂O₃ - 4 wt.% CNT)</td>
</tr>
<tr>
<td></td>
<td>Spray Drying of Al₂O₃ and 4 wt.% CNTs (agglomerate ~15-60 μm)</td>
<td>---</td>
<td>A4C-SD (Al₂O₃ - 4 wt.% CNT agglomerate ~ 15-60 μm)</td>
<td>Throughout the powder agglomerate</td>
<td>A4C-SD Coating (Al₂O₃ - 4 wt.% CNT)</td>
</tr>
<tr>
<td></td>
<td>Spray Drying of Al₂O₃ and 8 wt.% CNTs (agglomerate ~15-60 μm)</td>
<td>---</td>
<td>A8C-SD (Al₂O₃ - 8 wt.% CNT agglomerate ~ 15-60 μm)</td>
<td>Throughout the powder agglomerate</td>
<td>A8C-SD Coating (Al₂O₃ - 8 wt.% CNT)</td>
</tr>
</tbody>
</table>
Fig. 4.3: a) Scattergram and representative micrograph describing CNT dispersion angle with respect to aspect ratio, and b) Histogram of A4C-B powder depicting agglomeration of CNTs in the dome cavity of the powder agglomerate.
Fig. 4.4: a) Scattergram and representative micrograph describing CNT dispersion angle with respect to aspect ratio, and b) Histogram of A4C-SD powder depicting dispersion angle with respect to the number of CNTs in the powder agglomerate.
Fig. 4.5: a) Scattergram and representative micrograph describing CNT dispersion angle with respect to aspect ratio, and b) Histogram of A8C-SD powder depicting dispersion angle with respect to the number of CNTs in the powder agglomerate.
CNTs throughout the powder agglomerate. Visually no bundles/agglomerates of CNTs were observed in the spray dried powder validating the non-preferred sticking sites for CNTs. Fractured A4C-SD and A8C-SD powders displayed excellent CNT dispersion in the core of spray dried agglomerate. Aspect ratio is helpful in understanding the pinning effect of CNTs, and corresponding dispersion angle suffices the reinforcement in the subsequent direction.

An initial signature of Raman spectrum was acquired for the powder feedstock, Fig. 4.6, to be able to compare the retention of CNT structure in the plasma sprayed powders.

![Raman Spectrum of Powders](image)

Fig. 4.6: Raman spectrum of the initial powder feedstock (before plasma spraying). D and G peaks correspond to defect and graphitic structure of CNT.
coatings. D and G peaks correspond to defect and graphitic peaks of CNT [42, 178]. The presence of G peaks in the plasma sprayed coatings will confirm the presence of CNT structure. Since high temperature and impact associated with plasma spraying might damage the CNTs during the processing, an initial Raman spectrum assists in realizing importance of optimized plasma spray parameters in retaining CNTs in the plasma sprayed coatings.

4.2. Optimization of Plasma Processing Parameters

First step is to optimize the plasma spraying parameters in order to achieve uniform coating thickness and required microstructure for improved fracture toughness. Optimized processing of spray dried composite powder was required to result controlled melting of the surface and solid state sintering of core, Fig. 4.7.

![Optimized plasma spraying process](image)

Fig. 4.7: Optimized plasma spraying to result surface melting and resolidification and core consolidation via solid state sintering.
Surface melting and resolidification results metallurgical consolidation of splats, whereas solid state sintering of core retains the nanocrystalline nature of the powder feedstock. Increasing the power-input (from the optimized condition) will increase the degree of melting and the nanocrystalline nature of the powder may not be retained in the processed coating. On the other hand reducing the power input will not result proper consolidation between splats and the integrity of the coating may not be achieved. Thereby, plasma-processing parameters are optimized in terms of bimodal microstructure in the matrix and fracture toughness of resultant coatings. Bimodal microstructure is characterized through fully melted structure (surface melting and resolidification) and partially melted structure (solid state sintering of the core), Fig. 4.7. Partially melted/solid state sintered (PM) grains provide ductility by grain shearing, whereas fully melted (FM) regions provide strength owing to dense structure.

Spray-dried nano-\(\text{Al}_2\text{O}_3\) (n-\(\text{Al}_2\text{O}_3\): ASD powder) was used as the starting powder to obtain optimized processing conditions. Purpose of using nano \(\text{Al}_2\text{O}_3\) is to take advantage of nanocrystalline grain with enhanced strength and ductility. Conventional micro ceramic grains are brittle in nature, whereas nanocrystalline ceramics show grain boundary shear sliding. Moreover crack propagation is more tortuous in nanocrystalline material owing to enhanced grain boundary area, which enhances the energy absorption during impact. Table 4.2 differentiates deformation mechanisms in micro and nanocrystalline ceramics.
Table 4.2: Toughening in micro- and nano-crystalline grains

<table>
<thead>
<tr>
<th>Toughening of Ceramic</th>
<th>Conventional Micro Grain</th>
<th>Nanocrystalline Grain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deformation Mechanism</td>
<td>Limited Plastic Deformation: Brittle</td>
<td>Deformation via Cracking and Shear Sliding along Grain Boundaries</td>
</tr>
<tr>
<td>Crack Path</td>
<td>Less Tortuous</td>
<td>More Tortuous</td>
</tr>
</tbody>
</table>

4.2.1. Optimization of Plasma Parameters on n-Al₂O₃ Coating

Overall plasma spraying parameters utilized in the optimization of depositing n-Al₂O₃ coating are listed in Table 4.3. Processing parameters for plasma spraying of A-SD powder were selected based on earlier experience and exiting literature [39, 118, 209-211]. Uniform coating thickness is observed in the plasma sprayed ASD coatings, as presented in Fig. 4.8. Gas flow rate is expressed in standard liters per minute (slm). Powder feed rate is expressed in rpm (rounds per minute) of the rotating wheel in the Praxair (model #1264) powder feeder. The conversion of rpm to the feed rate in g/min depends on the volume of powder being fed (in c.c.) per minute at specified speed of feeding wheel (in rpm).

It must be noted that slight tweaking of plasma parameters was required in order to obtain a uniform coating. Practical difficulties such as non-deposition of coating with lower plasma power required change in the carrier gas velocity and feed rate as observed in Table 4.3. Lower carrier gas flow rate increase the residence time of particles in plasma plume, so higher degree of melting occurs for powder particle. Similarly, lower feed rate converts to small amount of powder in same plasma plume volume. Thereby lower feed rate and lower carrier gas flow rate help coating deposition at low power. An
overall picture of the plasma parameters and the resultant microstructure is presented in Fig. 4.9.

Coating thicknesses of 300, 500, 400 and 120 μm were obtained for the samples A, B, C and D respectively as shown in Fig. 4.10 a-d. Coatings are denser with increasing power, whereas lower power processed coatings depict increasing solid-state sintered region (Fig. 4.9). Image Pro imaging software dictated dense coatings (porosity between 13-18 %) with varying degree of differential structure as explained in the consequent section.

Powder treatment (spray drying) and control of plasma parameters results in two distinct regions, viz. partially melted /solid-state sintered (PM), and fully melted and resolidified (FM) structure in the coating with varying plasma power. This bimodal structure is attributed to 30-45 percent porosity of spray dried powder feedstock [94] coupled with optimized plasma parameters. Velocity ($v$) and temperature ($T$) are the controlling parameters deciding the microstructure of the sprayed coatings. Velocity corresponds to the kinetic energy attained by inflight particle. This in turn decides the degree of flattening and subsequent densification of the powder particle upon impact. The degree of melting is decided by the temperature attained by inflight particles. Thus the interplay of kinetic- (through velocity) and thermal- energy (via temperature) is critical in the generation of the microstructure. Thereby, inflight sensor monitoring of velocity and temperature (of the inflight powders) serve as controlling factors in optimizing the plasma processing parameters.
Fig. 4.8: Plasma sprayed coatings (from top: Sample A, B, C, and D respectively) on an AISI 1020 steel coupon of size 100 x 19 x 3.1 mm$^3$. Coating thickness varies between 120-500 μm.
Table 4.3: Plasma spraying parameters used for spraying Al₂O₃

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Sample A</th>
<th>Sample B</th>
<th>Sample C</th>
<th>Sample D</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current (Amperes)</td>
<td>780</td>
<td>727</td>
<td>642</td>
<td>512</td>
</tr>
<tr>
<td>Voltage (Volts)</td>
<td>39.5</td>
<td>38.3</td>
<td>37.3</td>
<td>36.2</td>
</tr>
<tr>
<td>Power (kW)</td>
<td>30.81</td>
<td>27.84</td>
<td>23.95</td>
<td>18.53</td>
</tr>
<tr>
<td>Primary, Ar (slm*)</td>
<td>32.1</td>
<td>32.1</td>
<td>32.1</td>
<td>32.1</td>
</tr>
<tr>
<td>Secondary, He (slm*)</td>
<td>59.5</td>
<td>59.5</td>
<td>59.5</td>
<td>59.5</td>
</tr>
<tr>
<td>Carrier, Ar (slm*)</td>
<td>21.7</td>
<td>19.8</td>
<td>19.8</td>
<td>19.8</td>
</tr>
<tr>
<td>Feed Rate (rpm)</td>
<td>4.0</td>
<td>4.0</td>
<td>3.0</td>
<td>2.5</td>
</tr>
<tr>
<td>Standoff Distance (mm)</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Coating Thickness (μm)</td>
<td>300</td>
<td>500</td>
<td>400</td>
<td>120</td>
</tr>
</tbody>
</table>

*slm: standard liters per minute

Various plasma parameters required different powder feed and gas flow rates in order to deposit a coating under different plasma power conditions. Subsequent development of the bimodal phases is optimized by plasma spraying of spray dried nano Al₂O₃ powder feedstock. Consequently, the generated microstructure is characterized depending upon the bimodal structure and the mechanical properties (hardness and
fracture toughness) attained by the plasma sprayed coatings. The role of initial powder treatment and the subsequent role of plasma parameters are discussed in the current section.

Fig. 4.9: Plasma spray parameter optimization showing relation of microstructural features of sprayed coatings with differential plasma parameter settings.
Fig. 4.10: Plasma sprayed nAl₂O₃ coating: a) Sample A, and b) Sample B from A-SD powder.
Fig. 4.10: Plasma sprayed nAl$_2$O$_3$ coating: c) Sample C, and d) Sample D from A-SD powder.
Owing to poor thermal conductivity of porous spray dried agglomerate, partial melting is sustained in the core whereas fully melted region is obtained on the surface. The surface-melting of powder particles results fully melted and re-solidified regions whereas the core of the powder agglomerate gets sintered in solid state. Hence, solid state sintering of the Al$_2$O$_3$ nano-particles occurs in the core region without destroying the nano nature of the starting powders. Thus, the matrix results in bimodal grain structure, i.e. fully melted and resolidified outer region and partially melted/solid-state sintered core region. Partially melted/sintered region helps in distributing the shock energy experienced during an impact, and helps deflecting crack resisting its propagation. Fully melted regions imparts strength and binding integrity to the composite coating [94, 114].

Increasing power shows increased FM zone, and increased grain size in the PM region, Fig. 4.11 through Fig. 4.14. Grain size is highest (~ 1 μm) in the highest plasma power setting, Fig. 4.11, and decreases to 0.8 μm (Fig. 4.12), to 0.5 μm, (Fig. 4.13) and to 0.35 μm (Fig. 4.14) in Sample A, B, C and D respectively. Influence of plasma parameters on microstructural features is quantified in Table 4.4 and Fig. 4.9.

Fracture toughness of the coatings is measured using Anti’s equation (eq. 2.1) from Vicker indentation at 200g with a dwell time of 15 s, Table 4.4 [59]. Radial crack generate upon unloading, and the crack is restrained when the crack tip stress intensity is balanced by the fracture toughness of the material to restrict its propagation as shown in Fig 4.15. Uniform bimodal microstructure distribution and highest fracture toughness (~3.29 MPa m$^{1/2}$) were obtained for sample C (Voltage: 37.3 V, and Current: 642 A). Hence, processing parameters from sample C were selected as optimized condition to spray Al$_2$O$_3$-CNT coatings for improved fracture toughness.
Fig. 4.11: Plasma sprayed nAl₂O₃ coating (Sample A) showing a) PM, FM and porosity distribution, and b) solid-state sintering of powders. Note that grain size is around 1 µm.
Fig. 4.12: Plasma sprayed nAl₂O₃ coating (Sample B) showing a) PM, FM and porosity distribution, and b) solid-state sintering of powders. Note that grain size is decreased to around 0.8 μm.
Fig. 4.13: Plasma sprayed nAl₂O₃ coating (Sample C) showing a) PM, FM and porosity distribution, and b) solid-state sintering of powders. Note that grain size is further decreased to around 0.5 μm.
Fig. 4.14: Plasma sprayed nAl₂O₃ coating (Sample D) showing a) PM, FM and porosity distribution, and b) solid-state sintering of powders. Note that grain size is ~0.35 μm.
Fig. 4.15: Radial crack generation in plasma sprayed n-Al₂O₃ coating via Vicker indentation.

Table 4.4: Microstructural features and fracture toughness of plasma sprayed nAl₂O₃ coatings

<table>
<thead>
<tr>
<th>Features → Coatings ↓</th>
<th>FM Content (%)</th>
<th>PM Content (%)</th>
<th>Porosity (%)</th>
<th>Coating Thickness µm</th>
<th>Fracture Toughness MPa m¹/²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample A</td>
<td>65.3 ± 2.9</td>
<td>20.2 ± 2.4</td>
<td>14.3 ± 1.2</td>
<td>300</td>
<td>1.51-3.55</td>
</tr>
<tr>
<td>Sample B</td>
<td>64.5 ± 2.3</td>
<td>22.7 ± 1.9</td>
<td>13.3 ± 0.0</td>
<td>500</td>
<td>1.28-1.36</td>
</tr>
<tr>
<td>Sample C</td>
<td>56.5 ± 1.8</td>
<td>30.1 ± 3.8</td>
<td>15.0 ± 2.5</td>
<td>400</td>
<td>2.53-3.29</td>
</tr>
<tr>
<td>Sample D</td>
<td>56.9 ± 1.6</td>
<td>24.6 ± 1.7</td>
<td>17.7 ± 1.0</td>
<td>120</td>
<td>1.07-1.66</td>
</tr>
</tbody>
</table>
4.3. Plasma Spraying of n-Al₂O₃ with Addition of CNTs

A4C-B, A4C-SD, and A8C-SD powders were plasma sprayed with the plasma parameters listed in Table 4.5. It was observed that plasma parameters optimized for A-SD coating (Sample C) were not sufficient to deposit a uniform CNT reinforced coating. Since CNTs are uniformly distributed in spray dried powders, they extract heat from Al₂O₃ particles due to their high thermal conductivity. Thereby higher power is required to melt Al₂O₃ particles and get uniformly thick coatings. Accuraspray inflight sensor data clearly elicits such observation and thereby higher power plasma parameters were utilized in order to accommodate deposition of a uniform coating (Table 4.5). Comparative inflight particle diagnostic data (temperature and velocity of inflight powder particles) for all samples is discussed in later section.

Table 4.5: Plasma parameters for spraying A4C-B, A4C-SD and A8C-SD coating

<table>
<thead>
<tr>
<th>Coatings → Parameters ↓</th>
<th>A4C-B Coating</th>
<th>A4C-SD Coating</th>
<th>A8C-SD Coating</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current (Amperes)</td>
<td>630</td>
<td>778</td>
<td>761</td>
</tr>
<tr>
<td>Voltage (Volts)</td>
<td>40.8</td>
<td>39.6</td>
<td>42.5</td>
</tr>
<tr>
<td>Power (kW)</td>
<td>25.7</td>
<td>30.8</td>
<td>32.3</td>
</tr>
<tr>
<td>Primary, Ar (slm)</td>
<td>32.1</td>
<td>32.1</td>
<td>32.1</td>
</tr>
<tr>
<td>Secondary, He (slm)</td>
<td>59.5</td>
<td>59.5</td>
<td>59.5</td>
</tr>
<tr>
<td>Carrier, Ar (slm)</td>
<td>19.8</td>
<td>19.8</td>
<td>19.8</td>
</tr>
<tr>
<td>Feed Rate (rpm)</td>
<td>3.0</td>
<td>3.0</td>
<td>3.0</td>
</tr>
<tr>
<td>Standoff Distance (mm)</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Coating Thickness (μm)</td>
<td>350</td>
<td>450</td>
<td>500</td>
</tr>
</tbody>
</table>

* slm: standard liters per minute
In this section, comparable plasma parameters, Table 4.5, are used for studying the role of CNT dispersion in generating a contrasting microstructure and mechanical properties. Plasma sprayed A-SD coating under optimized conditions (Sample C with highest fracture toughness) is taken as a reference for comparing the CNT reinforced Al₂O₃ coatings. Steel samples coated with A-SD, A4C-B, A4C-SD and A8C-SD are presented in Fig 4.16 a-d. A-SD coating is whitish (Fig. 4.16 a), A4C-B coating is grayish (Fig. 4.16 b), A4C-SD coating is dark-grayish (Fig. 4.16 c), whereas A8C-SD coating is blackish (Fig. 4.16 d) in color. A-SD coating is white owing to the pure Al₂O₃ coating (Fig. 4.16a). Black specs and grayish color is observed in A4C-B coating (Fig. 4.16b) owing to agglomeration and dispersion of CNTs respectively. Uniform dark grayish nature of A4C-SD (Fig. 4.16c) is attributed to the uniform CNT dispersion in the coating. Uniformly black color of A8C-SD (Fig. 4.16d) is attributed to higher CNT content and their uniform dispersion. Overall microstructure generated in each CNT reinforced coating is discussed in following sections.

Table 4.6 shows the inflight particle diagnostic data for various plasma sprayed coatings. Temperature variation of different powders owing to CNT dispersion is clearly evinced in Table 4.6. Contrasting difference is attributed to role of high thermal conductivity of CNTs in the A4C-B powder (CNT agglomeration) and A4C-SD powder (CNT dispersion).

4.3.1. Plasma Sprayed A4C-B Coating

Fig. 4.17 shows plasma sprayed A4C-B nanocomposite coating on the AISI 1020 steel substrate. Uniform and homogeneous A4C-B nanocomposite coating of ~350 µm is
Fig. 4.16: Plasma sprayed a) A-SD, b) A4C-B, c) A4C-SD, and d) A8C-SD coating.
Table 4.6: Inflight particle diagnostic data for various plasma sprayed coatings

<table>
<thead>
<tr>
<th>Coating</th>
<th>Temperature (K)</th>
<th>Velocity (m/s)</th>
<th>Dwell Time (s)</th>
<th>Cooling Rate (K/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A-SD Coating</td>
<td>2512</td>
<td>289</td>
<td>$3.46 \times 10^{-4}$</td>
<td>$7.26 \times 10^{6}$</td>
</tr>
<tr>
<td>A4C-B Coating</td>
<td>2898</td>
<td>271</td>
<td>$3.69 \times 10^{-4}$</td>
<td>$7.85 \times 10^{6}$</td>
</tr>
<tr>
<td>A4C-SD Coating</td>
<td>2332</td>
<td>244</td>
<td>$4.10 \times 10^{-4}$</td>
<td>$5.69 \times 10^{6}$</td>
</tr>
<tr>
<td>A8C-SD Coating</td>
<td>2241</td>
<td>232</td>
<td>$4.31 \times 10^{-4}$</td>
<td>$5.20 \times 10^{6}$</td>
</tr>
</tbody>
</table>

obtained. The coating is free from cracks and adherent to the substrate. Coating was removed from the substrate to obtain as a freestanding structure. Density of 3.40 g/cc (87.2 % dense as compared to theoretical density of 3.90 for Al$_2$O$_3$-4 wt. % CNT composite) was obtained through Archimedes water immersion technique.

![A4C-B Coating](image.png)

Fig. 4.17: Cross-sectional image of A4C-B coating showing uniform and adherent coating.
At higher magnification, differential microstructure with fully melted and resolidified (FM) and partially-melted and solid state sintered (PM) regions is observed in Fig. 4.18. This microstructure is similar to that of optimized plasma sprayed A-SD coating (Fig. 4.13).

![Figure 4.18: Plasma sprayed A4C-B coating showing fully melted (FM) and solid state sintered (PM) region.](image)

To preserve the advantages of CNT reinforcement, it is necessary for CNTs to survive their sojourn in the harsh environment of plasma plume. As seen in Fig. 4.19a and 4.19b, interlinked-CNTs are distributed, undamaged and retained in the plasma sprayed structure. Survival and distribution of CNTs is attributed to: i) dispersion of CNTs along the surface and the dome-cavity region of the spray dried Al$_2$O$_3$ powder
Fig. 4.19: Fractured surface image of A4C-B coating depicting a) chain loop formation of CNTs, and b) fusion of CNTs in Al₂O₃.
(also seen in Fig. 4.3), and ii) controlled plasma processing parameters (2898 K, 271 m/s) to cause partial melting of core. Damage of CNTs from harsh plasma environment is also shrouded by poor thermal conductivity of powder-agglomerate porosity, and limited melting of the surface causing the core and tail regions of powder agglomerate to retain CNTs during successive deposition of splats. Survival of CNTs play an important role towards resulting microstructural ingenuity owing to their excellent mechanical properties [26, 31, 212]. CNT chain-loop formation, Fig. 4.19a, acts as anchors for providing improved fracture toughness. In addition, entangling of CNTs induce torsion, making torsion stiffness an important concept of providing toughness to the reinforced nanocomposite [213]. Fusion of CNTs along the Al₂O₃ melt, Fig. 4.19b, shows the good wettability observed in the plasma sprayed nanocomposite coating owing to high plasma parameters melting the Al₂O₃ surface and trapping the CNTs within.

4.3.2. Plasma Sprayed A4C-SD Coating

A uniform ~450 μm thick coating is obtained by plasma spraying A4C-SD powder, Fig. 4.20. Coating is dense (3.52 g/cc with 90.2 % theoretical density) and adherent to the substrate. Higher degree of PM region is observed in the plasma sprayed A4C-SD coating, and is compared and quantified with other coatings in the later section. Processing of this coating is aimed to correlate the role of CNT dispersion in enhancing fracture toughening in comparison to A4C-B coating, which also has similar composition.
It is clearly indicated that the powder particles have undergone reduced temperature (of 2332 K when compared to 2898 K for A4C-B coating) deposition owing to its high PM content, Fig. 4.21. Inflight particle temperature data indeed confirms the role of CNT dispersion in reducing thermal exposure to the powders during plasma spraying.

CNTs are retained in the plasma sprayed A4C-SD coating, Fig. 4.22a. CNTs appear thicker (~ 100-120 nm) when compared to starting CNTs (40-70 nm) implying that CNTs are coated with Al₂O₃ during processing. Spreading of CNTs throughout the matrix without agglomeration confirms good dispersion (Fig. 4.22b).
Fig. 4.21: Plasma sprayed A4C-SD coating depicting enhanced PM content.

CNT dispersion plays an important role in reducing thermal damage and consequently CNTs are retained in the matrix. CNTs are observed both near the PM and FM regions, indicating their survival in the plasma plume during their processing. Survival of CNTs in dispersed condition is critical towards uniform toughening of ceramic nanocomposite.

Enhanced wettability of CNTs is dictated by $\text{Al}_2\text{O}_3$ coating on CNT surface (indicated by increase in the CNT diameter), Fig. 4.22a, with good anchoring with the splats. On one hand where excellent wettability of CNTs anchors the splats, on the other hand, dispersed CNTs provide uniform reinforcements. Synergetic effect of dispersed CNTs depicting good wettability is expected to ensue enhanced fracture toughening of the A4C-SD coating.
Fig. 4.22: a) Cross-section and b) fractured surface of plasma sprayed A4C-SD coating showing CNT retention and distribution.
Most of the ceramic CNT composite processing is performed via solid state sintering techniques such as spark plasma sintering, hot pressing, extrusion, etc) consolidation [16, 172, 176, 186]. Absence of molten ceramic in conventional processing (and in most of advanced techniques) has obviated attention to aspects of CNT surface coating and anchoring in such structures. Hence wetting of CNT by alumina is a novel issue by itself that has been discussed later.

4.3.3. Plasma Sprayed A8C-SD Coating

Plasma sprayed A8C-SD shows a uniform and ~500 μm thick coating in Fig. 4.23. Coating is dense (3.53 g/cc with 94 % theoretical density) and adherent to the substrate. Magnified SEM image of the coating, Fig. 4.24, elicits similar features as observed for A4C-SD coating. Comparative solid state sintered region (PM), fully melted region (FM), and porosity is defined by the microstructural features, as discussed in section 4.5.1. Enhanced PM content is attributed to uniform CNT distribution and higher CNT content leading to reduced thermal exposure.

A typical solid state sintered region of A8C-SD coating is shown in Fig. 4.25a. Reduced thermal exposure (2241 K) to the powders is attributed to increased CNT content (from 4 wt. % to 8 wt.%), which extracts heat from surrounding Al2O3 particles. Consequent consolidation, in turn, entraps dispersed CNTs in the solid state sintered region, Fig. 4.25a. CNT dispersion is also observed at neck region between two adjoining splats, resolidified region and surface of Al2O3 particles, Fig. 4.25b. CNTs remain undamaged and are uniformly dispersed at various locations such as melted and resolidified region, high-impact area of adjoining neck region and also at surface.
Fig. 4.23: Cross-sectional SEM image of plasma sprayed A8C-SD coating.

Fig. 4.24: Plasma sprayed A8C-SD coating showing FM and PM regions.
Fig. 4.25: Solid state sintered region showing a) CNT retention in the plasma sprayed A8C-SD coating, and b) CNT distribution at neck, resolidified region and surface of Al$_2$O$_3$ particles.
Distribution of CNTs in the coatings is affirmed owing to retention of distributed CNTs as present in the starting A8C-SD powder.

The purpose of plasma spraying A8C-SD coating is to evaluate toughening properties with an increase in CNT content in comparison to A4C-SD coating. Hence it will be possible to capture fracture toughening enhancement both because of CNT dispersion and CNT content.

Density of plasma sprayed coatings (measured from water immersion technique) is summarized in Fig. 4.26. Minimum density of ~87% was achieved for A-SD coating, whereas addition of CNTs has resulted increased in the density of plasma sprayed coatings. This is attributed to dispersed CNTs promoting better heat transfer and thereby resulting enhanced densification. This is confirmed via enhanced PM region with

![Density of Plasma Sprayed Coatings](image)

Fig. 4.26: Density of plasma sprayed coatings measured from water immersion technique.
increase in CNT dispersion and CNT content. In addition, flexing nature of CNTs fill-in the gaps between n-Al₂O₃ splats and reduce the overall porosity during deposition.

4.4. Retention of CNTs in Plasma Sprayed of Al₂O₃ – CNT Coatings

Retention of CNT structure in the plasma sprayed coatings is confirmed by the D (\(\sim 1340 \text{ cm}^{-1}\)) and G peaks (between 1500 and 1600 cm\(^{-1}\)), Fig. 4.27 which are same as observed in the initial powder feedstock. D-peak corresponds to disordered graphite, whereas G-peak corresponds to stretching mode of graphite, a characteristic of

![Raman Spectrum of Plasma Sprayed Coatings](image)

Fig. 4.27: Raman spectrum of plasma sprayed coatings showing retention of CNT structure in the plasma sprayed coatings.
nanotubes. Hence, Raman spectroscopy dictates that optimized plasma spraying parameters contribute in the retention of undamaged CNTs in the plasma sprayed coatings. This effect is assisted by enhanced porosity of spray-dried powder, which blankets the damage to CNTs via its reduced thermal conductivity, and adds in retaining CNTs even under high power plasma parameters.

4.5. Effect of CNTs in Generating Microstructure and Phases in Plasma Sprayed Coatings

Bimodal grain size distribution, corresponding inflight particle diagnostic data, generation of secondary metastable phases is presented in the current section. Microstructural PM and FM regions observed in the A-SD, A4C-B, A4C-SD, and A8C-SD coatings are analyzed and influence of inflight particle diagnostic on generating such a microstructure is discussed. XRD analysis further elucidates the rapid kinetics inherent to the plasma spray processing.

4.5.1. Role of CNT Dispersion in Creating Bimodal Grain Distribution in Matrix

Distribution of the FM, PM and porosity in the plasma sprayed coatings is analyzed using ImagePro imaging software and presented in Table 4.7. Such a differential microstructure is obtained because of different thermal history experienced by powder particles during their transit through plasma plume.

Porosity measurement by quantitative microscopy considers only the provided image information, which in some cases might not be true representative. Different magnification images and selection of area for porosity calculation is user-dependent.
Hence a more concrete approach of evaluating porosity by water immersion technique is utilized in the current research. Water immersion technique evaluates both macro and micro porosity of the plasma sprayed coatings, and, hence, is more reliable.

Table 4.7: Volume percentage of microstructural features of the plasma sprayed coatings obtained from quantitative microscopy and image analysis

<table>
<thead>
<tr>
<th>Coating</th>
<th>Coating Thickness (μm)</th>
<th>Fully Melted (FM)</th>
<th>Partially Melted (PM)</th>
<th>Porosity (Image Analysis)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A-SD Coating</td>
<td>400</td>
<td>80.9 %</td>
<td>15.9 %</td>
<td>3.2 %</td>
</tr>
<tr>
<td>A4C-B Coating</td>
<td>350</td>
<td>82.1 %</td>
<td>12.1 %</td>
<td>5.8 %</td>
</tr>
<tr>
<td>A4C-SD Coating</td>
<td>450</td>
<td>71.4 %</td>
<td>24.7 %</td>
<td>3.9 %</td>
</tr>
<tr>
<td>A8C-SD Coating</td>
<td>500</td>
<td>52.6 %</td>
<td>46.3 %</td>
<td>1.1 %</td>
</tr>
</tbody>
</table>

Inflight particle diagnostic data was presented in Table 4.6. These contrasting velocities and temperature of inflight particles that form coatings indicate strong influence of CNT content and distribution in imparting reduced thermal exposure to the blended and spray dried powder particles. A4C-B coating experienced higher temperature when compared to A-SD coating, which is attributed to CNT distribution on the surface and the dome cavity (as explained in section 4.1). Owing to higher thermal conductivity of CNTs (~3000 W/mK) when compared to Al₂O₃ (~ 36 W/mK), high heat is absorbed on the surface leading to enhanced thermal exposure and higher attained temperatures in A4C-B coating. Inflight particle diagnostic data, therefore affirms the mechanism of the heat distribution in the A4C-B coating. But, the reduction of thermal exposure in spray
dried coatings (A4C-SD and A8C-SD coating) is attributed to uniform CNT dispersion both on the surface and inside of the powder particles (explained earlier in section 4.1). Directly stating, now less number of CNTs are present on the surface of spray dried powder (since content is same and CNTs are distributed uniformly also at inside of powder), thereby superheating of surface by CNTs is reduced. Additionally, CNT content in the inside of the powder now ‘soaks up’ the heat acquired by the surface CNTs towards reducing the temperature attained by powder particles. Hence, reduced temperature for A4C-SD coating was observed. Though addition of CNTs (in A8C-SD powder) marginally should increase the surface temperature, but uniform CNT distribution inside of the particle increases too, and the rate of soaking in the heat also increases. And owing to rapid kinetics inherent to the process, short residence times of ~ 4 x 10^4 s in the plasma plume do not suffice superheating of surface CNTs. This phenomenon further lowers the temperature acquired by the A8C-SD powder particles in the plasma plume. This offers two advantages: first that damage to CNTs is reduced due to reduced thermal exposure, and secondly that solid state sintered region can successfully retain CNTs.

It becomes clear now that increased temperature observed by A4C-B powder particles in plasma plume undergo higher degree of heating and induces higher FM content. Hence, it can be concluded that increasing plasma parameters and CNT content and dispersion are critical in defining temperature exposure experienced by powder particles and subsequent partially melted and fully melted structure.
4.5.2. Generation of Metastable $\gamma$-Al$_2$O$_3$

Owing to rapid kinetics of the plasma spraying (with cooling rates as high as $7.85 \times 10^6$ K/s), generation of metastable phases becomes inherent. XRD spectrum of the coatings is presented in Fig. 4.28. Since CNT peaks are not observed in the presence of Al$_2$O$_3$, only one XRD spectrum for powder feedstock is shown (instead of individual A-SD, A4C-B, A4C-SD, and A8C-SD powder spectra). Scherrer formula is used to calculate the crystallite size of the phases ($t$) present in powder feedstock and plasma sprayed coatings.

$$t = \frac{0.9\lambda}{B \cos \theta} \quad \text{Equation 4.1}$$

where $\lambda$ is the X-ray wavelength, $\theta$ is the diffraction angle, $B$ is the peak broadening measured at half of maximum intensity in radians (obtained as $\sqrt{B_m^2 - B_s^2}$ where $B_m$ is the measured broadening at peak intensity half maximum, and $B_s$ is the peak intensity half-maximum of the standard). Silicon (Si) from NIST is taken as standard for calculating peak broadening of machine.

For crystallite size measurement of $\alpha$-Al$_2$O$_3$, FWHM (full width at half maximum) at most intense peak of 2-theta ($\sim 35$ degrees) is considered, whereas for $\gamma$-Al$_2$O$_3$, FWHM (full width at half maximum) at most intense peak of 2-theta ($\sim 46$ degrees) is measured. Alpha and gamma Al$_2$O$_3$ content was measured with the ratio of area of corresponding peaks to the area under cumulative peaks. It must be noted that starting powder feedstock was 100 % $\alpha$-Al$_2$O$_3$, hence only the crystallite size is reported in Table 4.8.
Fig. 4.28: XRD spectrum of plasma sprayed coatings. Indicated peaks belong to $\gamma$-$\text{Al}_2\text{O}_3$ whereas all other peaks belong to $\alpha$-$\text{Al}_2\text{O}_3$.

Table 4.8: Crystallite size and content of powder and plasma sprayed coatings

<table>
<thead>
<tr>
<th>Crystallite Size and Phase Content</th>
<th>Powder*</th>
<th>Coating $\alpha$-$\text{Al}_2\text{O}_3$</th>
<th>Coating $\gamma$-$\text{Al}_2\text{O}_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Size (nm)</td>
<td>Size (nm)</td>
<td>Content (%)</td>
</tr>
<tr>
<td>A-SD</td>
<td>44</td>
<td>55</td>
<td>76.6</td>
</tr>
<tr>
<td>A4C-B</td>
<td>46</td>
<td>95</td>
<td>68.4</td>
</tr>
<tr>
<td>A4C-SD</td>
<td>45</td>
<td>73</td>
<td>81.7</td>
</tr>
<tr>
<td>A8C-SD</td>
<td>43</td>
<td>76</td>
<td>74.9</td>
</tr>
</tbody>
</table>

*Powder is 100% $\alpha$-$\text{Al}_2\text{O}_3$. 
Presence of CNTs affects the nucleation of γ-Al₂O₃ that occurs from the melting and rapid solidification of α-Al₂O₃. High thermal conductivity and superheating of CNTs create favorable sites for nucleation of γ-Al₂O₃. Generation of γ-Al₂O₃ phase is related to the CNT dispersion and location in the powder agglomerate. Nucleation of γ-Al₂O₃ phase is dependent on the content, location and dispersion behavior of CNTs.

4.5.2.1. Dependence of γ-Al₂O₃ on CNTs in the Matrix

CNTs on the surface of the powder particle are responsible for nucleating γ-Al₂O₃ by absorbing the plasma heat, assisting the melting on surrounding alpha, and leading to enhanced cooling rates in its vicinity to generate γ-Al₂O₃. Owing to their high thermal conductivity (∼3 x 10³ W/m/K) when compared to that of Al₂O₃ (∼36 W/m/K), CNTs heat up melting the surrounding Al₂O₃, and quickly lose heat to cooler surrounding Al₂O₃ in the core (because of optimized surface melting and solid state sintering of core). Rapid kinetics involved inherently with the processing adds to the γ-Al₂O₃ nucleation in the processed coating. Hence surface CNTs are responsible for enhancing the gamma content of the plasma spayed structure.

CNTs entrapped in the core absorb the heat from the surface. Since the travel time of powder particles is in order of ∼4 x 10⁻⁴ s, CNTs present in the core do not allow superheating of the CNTs located on the surface. Restriction to reach high temperature presents two benefits in the processed coatings. First it shrouds the damage to surface CNTs by restricting the CNT temperatures. This behavior is linked by the inflight particle diagnostic data observed for the spray dried coatings, where reduced temperature profiles were observed even for higher plasma parameters (Table 4.7). Secondly, it reduces the
gamma precipitation on CNTs in core by disallowing high cooling rates as experienced by surface CNTs in the absence of core CNTs.

Dome cavity CNT agglomeration (as in A4C-B coating) can achieve exceedingly high temperatures owing to agglomerated mass of CNTs. Reduced resistance path and high thermal conductivity of CNTs renders high cooling rates to the surrounding Al$_2$O$_3$. Absence of core-CNTs not only tender comparatively enhanced damage to surface-CNTs, but also is expected to impart enhanced $\gamma$-Al$_2$O$_3$ precipitation. Though $\gamma$-Al$_2$O$_3$ assists with second phase strengthening, the total potential of CNT-toughening goes untapped.

Dispersion of CNTs (as in A4C-SD and A8C-SD coatings) allows entrapment of CNTs both in the molten- and solid state sintered region. CNTs present in the core region are retained by solid state sintering and retain entrapped in the densified region. Surface CNTs disperse along intersplat region as successive impact of molten/semi-molten particles. Therefore, dispersion of CNTs potentially creates more sites for $\gamma$-Al$_2$O$_3$ nucleation. But two factors must be considered before making conclusions. First that since CNTs act as nucleating sites for $\gamma$-Al$_2$O$_3$, increasing CNT content overall increases the $\gamma$-Al$_2$O$_3$ content. Secondly that CNTs in the core act as ‘heat absorbing’ sites to reduce the surface temperature of the Al$_2$O$_3$, thereby reducing the degree of melting and kinetics of rapid solidification. Hence precipitation of $\gamma$-Al$_2$O$_3$ is a balance between CNT dispersion, surface distribution and total CNT content.

It can be reasoned from the observed date, Table 4.8, that addition of CNTs increased gamma content, but reduces size of gamma crystallite (A4C-B coating when compared to A-SD coating). This behavior is attributed to distribution of CNTs only on
the surface. Since CNTs are distributed onto the surface, rapid quenching due to thermal heat absorption enhances the kinetics increases the nucleation rate, but reduces the growth rate, thereby reduce the γ-Al\textsubscript{2}O\textsubscript{3} size and increase the γ-Al\textsubscript{2}O\textsubscript{3} content.

Contrasting feature of CNT dispersion is observed in the spray dried powder, where reduced γ-Al\textsubscript{2}O\textsubscript{3} size and content is observed (A4C-SD coating compared to A-SD coating). Since CNTs are dispersed throughout the powder, superheating of CNTs is restricted by CNTs present in the core of the powder agglomerate. On one hand, the overall content of CNTs on the surface of A4C-SD powder is less (when compared to A4C-B powder because of same CNT content with better dispersion even in the core of powder), and secondly core CNTs soak up the thermal heat of the surface CNTs. Hence overall sites for gamma nucleation are reduced and enhanced time for cooling (reduced cooling rate) allows only minimal precipitation of γ-Al\textsubscript{2}O\textsubscript{3}.

Now, with the A8C-SD powder, interesting feature of both enhanced γ-Al\textsubscript{2}O\textsubscript{3} size and content (when compared to A4C-SD powder) is observed. This is attributed to similar dispersion as that of A4C-SD powder, but enhanced CNT content on the surface now leads to more nucleating sites (and increased γ-Al\textsubscript{2}O\textsubscript{3} content). In addition to reduced cooling rate (via increased heat absorption by core CNTs), and enhanced chances of merging the precipitated γ-Al\textsubscript{2}O\textsubscript{3} crystallites by amplified CNT content shows increased γ-Al\textsubscript{2}O\textsubscript{3} size.

Generation of metastable phases is inherent in the plasma sprayed structures. Considering the high energy input of plasma process, the free energy change associated with α- Al\textsubscript{2}O\textsubscript{3} to γ- Al\textsubscript{2}O\textsubscript{3} phase transformation can be described as [[214]]:
\[ \Delta G_r = \Delta G_{vol} + \Delta \Gamma_{surface} = \frac{4}{3} \pi r^3 (G_\gamma - G_\alpha) + 4\pi r^2 (\Gamma_\gamma - \Gamma_\alpha) \]  \hspace{1cm} \text{Equation 4.2} \\

where \( G_\alpha \) and \( G_\gamma \) are free energies, and \( \Gamma_\alpha \) and \( \Gamma_\gamma \) are surface energies of \( \alpha-\text{Al}_2\text{O}_3 \) and \( \gamma-\text{Al}_2\text{O}_3 \) respectively of radius \( r \).

Since \( \gamma-\text{Al}_2\text{O}_3 \) is a metastable phase, it has critical radius of transformation much smaller than existing \( \alpha-\text{Al}_2\text{O}_3 \) phase. Considering the crystallite size of \( \alpha-\text{Al}_2\text{O}_3 \to \gamma-\text{Al}_2\text{O}_3 \) phase transformation, the critical radius \( r_c \) is obtained by equating the differential, with respect to \( r \), of eq. 4.2 to zero.

\[ r_c = \frac{-2(\Gamma_\gamma - \Gamma_\alpha)}{(G_\gamma - G_\alpha)} \]  \hspace{1cm} \text{Equation 4.3} \\

Though the \( \gamma-\text{Al}_2\text{O}_3 \) crystallites will grow as coarsening progresses, they will serve as second phase particles and help improving the dispersion strengthening of the composite. Hence, CNTs become critical in manipulating the bimodal phase distribution in the plasma sprayed coatings and take advantage of secondary strengthening mechanisms.

High magnification TEM image of CNT surface, Fig. 4.29a, shows precipitation of \( \gamma-\text{Al}_2\text{O}_3 \) crystallites. High resolution TEM imaging, Figs. 4.29b and c, show lattice fringes of the generated precipitate. Lattice spacing of fine crystallite corresponds to 2.19 Å, which matches with the interplanar crystal spacing of (023) plane for \( \gamma-\text{Al}_2\text{O}_3 \) phase. Presence of fine \( \gamma-\text{Al}_2\text{O}_3 \) precipitates on CNT surface confirms the proposed mechanism of \( \gamma-\text{Al}_2\text{O}_3 \) phase generation.
Fig. 4.29: TEM image showing a) γ-Al₂O₃ precipitation on CNT surface, and High resolution TEM image showing γ-Al₂O₃ lattice fringes corresponding to b) (311) plane (lattice spacing of 2.38 Å, and c) (023) plane (lattice spacing of 2.19Å).
4.6. Role of CNT Dispersion in the Fracture Toughening of Nanocomposite Ceramic Coatings

Role of CNT dispersion is measured by an increase in the fracture toughness of the plasma sprayed coatings. Measure of toughening by CNT pull out, crack bridging, and crack deflection is reflected in the high magnification SEM microscopy of the coatings. Vicker indentation is utilized for measuring indentation toughness with a load of 200g with a dwell time of 15 s.

Table 4.9 details the hardness and fracture toughness of the plasma sprayed coatings. It can be noted that A4C-B there is no significant variation in the hardness value of the different coatings. Though it might be expected that higher density of coatings result higher hardness, but it is counter balanced by the reduction of hardness due to increasing PM content. Hence, overall hardness of the plasma sprayed coatings show similar hardness values in Vicker indentation.

Table 4.9: Hardness and fracture toughness of plasma sprayed coatings

<table>
<thead>
<tr>
<th>Coating</th>
<th>A-SD</th>
<th>A4C-B</th>
<th>A4C-SD</th>
<th>A8C-SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Density g/cc (% theoretical)</td>
<td>3.47 (87%)</td>
<td>3.40 (87.2%)</td>
<td>3.52 (90.2 %)</td>
<td>3.53 (94%)</td>
</tr>
<tr>
<td>Hardness (V_H)</td>
<td>714.2 ± 14.9</td>
<td>709.6 ± 8.4</td>
<td>738.7 ± 12.9</td>
<td>712.7 ± 2.3</td>
</tr>
<tr>
<td>Fracture Toughness (MPa m^{1/2})</td>
<td>3.22 ± 0.22</td>
<td>3.86 ± 0.16</td>
<td>4.60 ± 0.27</td>
<td>5.04 ± 0.58</td>
</tr>
</tbody>
</table>

Fracture toughness \( (K_c) \) of the ceramic coatings, Fig. 4.29, is calculated from Antis Equation as [215]:

130
\[ K_c = \chi \left( \frac{E}{H} \right)^{1/2} \frac{P}{c^{3/2}} \]  

Equation 4.4

where \( P \) is the applied load, \( E \) is the Young's modulus (taken as 390 GPa), \( H \) is the Vickers hardness, \( c \) is the radial crack length (measured from center of indent), and \( \chi \) is the calibration constant (taken as 0.016).

Vicker indentation fracture toughness of coatings is calculated as 3.22 MPa m\(^{1/2}\) for A-SD coating, whereas it increases to 3.86 MPa m\(^{1/2}\) for A4C-B coating, an improvement of 20 %, Fig. 4.30. Enhancement of fracture toughness is solely arising from the introduction of 4 wt.% CNTs in the matrix. But, A4C-SD coating shows fracture toughness of 4.60 MPa m\(^{1/2}\) (Fig. 4.30), an improvement of 43 % over A-SD coating. Though CNT contents are same in A4C-B and A4C-SD coating, tremendous improvement in the fracture toughness is attributed to uniform CNT dispersion in the A4C-SD coating when compared to A4C-B coating. Further increase in fracture toughness (5.04 MPa m\(^{1/2}\)) to 56% is achieved for A8C-SD coating, which is attributed to increase in the CNT content. Student t-test was performed to confirm significant statistical difference between the mean fracture toughness values with more than 95% confidence level.

It must be noted that porosity content, porosity size and porosity distribution can also alter the fracture toughness of the nanocomposites [10, 45, 56]. The dispersion of CNTs (in A4C-B, A4C-SD and A8C-SD) has shown contrasting change in the thermal exposure to inflight powders (Table 4.6), which consequently results in the variation in porosity in the plasma sprayed coatings, as seen in Fig. 4.26. Table 4.5 shows that higher plasma power of 30.8 kW for A4C-SD and 32.3 kW for A8C-SD was used when
compared to A4C-B powders (power of 25.7 kW). Higher plasma power results in enhanced degree of sintering for composite spray dried powders. Hence, higher degree of consolidation, both between (i) CNTs and Al₂O₃ interface and ii) Al₂O₃- Al₂O₃ interface leads to reduced porosity content in A4C-SD and A8C-SD coatings (when compared to that of A4C-B coating. Thereby, synergetic densification of coatings (in addition to CNT addition and dispersion) assisted in enhanced fracture toughening of plasma sprayed nanocomposite coatings.

![Fracture Toughness of Plasma Sprayed Coatings](image)

Fig. 4.30: Fracture toughness of plasma sprayed coatings.

To solely estimate the role of porosity in evaluating the fracture toughness of ceramic nanocomposite coatings, precise control on porosity should be required without disturbing (i) porosity size, and (ii) porosity distribution. In order to estimate effect of porosity, processing should prevent CNT damage, retain the grain size, and should not disturb CNT distribution even after consolidation. This in itself is a big challenge. Post
spray treatment such as hot-isostatic processing (HIPping) is recommended for future endeavors to separate the effect of porosity and CNT content on fracture toughness. Microstructural observation of CNT reinforcement in coatings is explained in the following section.

4.6.1: Enhancement of Fracture Toughness by CNT Distribution in A4C-B Coating

Long fibrous-CNT rope structures form anchors between the splats of the aluminum oxide matrix as seen in Fig. 4.31. CNTs are also fused with the molten and resolidified Al₂O₃ matrix (Fig 4.31). In addition, CNT form hooks and loops with other CNTs as shown in Fig. 4.32. Looping of CNTs, Fig. 4.31, initiate entangling effect upon tensile loading and aids in restraining the separation of fused regions. This behavior abets superior fracture toughness of plasma sprayed nanocomposite coating. Entangling CNT network has also been observed by other researchers [166].

Tensile stretching of the anchored hook structure, Fig. 4.33, is restrained by high bending stiffness of CNTs. Bending stiffness \( B \) of CNTs has been largely accepted as [216]:

\[
B = \frac{NEt^3}{12(1-v^2)} \tag{4.5}
\]

where \( t \) is the thickness of CNT wall and \( N \) is the number of walls in CNT. This clearly implies the direct proportionality of increase in bending stiffness with increasing number of CNT walls in multiwalled carbon nanotubes. Hence, it is expected that loop and hook observed in Fig 4.32 will improve the fracture resistance of A4C-B coating.
Fig. 4.31: Fibrous CNT fused onto Al$_2$O$_3$ surface anchoring the splats.

Fig. 4.32: Hook and loop formation of CNTs.
Bending Stiffness

Hook Formation

Fig. 4.33: Hook formation by entanglement of CNTs in A4C-B coating.

CNT alignment along the splat interface is also observed in A4C-B coating (Fig. 4.34). Since the A4C-B powder has CNTs dispersed only on the surface of the powder, surface melting during plasma spraying gathers most of the CNTs upon the impact onto the substrate. Surface melting of the powder agglomerate eases the CNT movement in the semi-molten/molten pool of Al₂O₃. And upon impact onto substrate, CNTs align parallel to the substrate in order to absorb the high collision shock. Consequent impact and deposition of splats entraps CNTs along the intersplat region. Preferential alignment of CNTs brings about the concept of directional strengthening towards enhancing the fracture toughness. Energy absorption upon shock via interfacial sliding also assists in improving the fracture toughness of the coating. Hence, *trimodal microstructure* (CNTs, partially-melted (PM), and fully-melted (FM) regions) observed in the cross-sectional images of the plasma sprayed nanocomposite coating, Fig. 4.34, rivet fracture toughness enhancement owing to differential phase properties [94].

Interfacial strengthening of the nanocomposite ceramic can be realized in terms of load transfer between the CNTs and matrix as shown in schematic Fig. 4.35. Based on the
Fig. 4.34: Impact alignment of CNTs along the intersplat region in the plasma sprayed A4C-B Coating.
assumptions of elastic interfacial shear stress between reinforcement and ceramic matrix, without any sliding [217], modified axial stress (\(\sigma\)) and critical shear stress (\(\tau\)) equations for the parallel aligned MWNT at the interface can be expressed as.

\[
\sigma = \frac{L^2}{L^2 - r^2} \left[ \sigma_{CNT} - \frac{r^2}{L^2} \left( \frac{E_{CNT}}{E_{splat}} \sigma_{splat} \right) \right] \quad \text{Equation 4.6}
\]

\[
\tau = \frac{r E_{CNT} (w_{interface} - w_{CNT})}{L^2 (1 + v_{CNT})} \quad \text{Equation 4.7}
\]

Fig. 4.35: Increased axial shear and absorption of energy by impact alignment of CNTs along splat interface.

where \(r\) is the radial distance, and \(L\) is the length of CNT, \(w\) is the displacement along axial direction, and \(v\) is the Poisson’s ratio, and \(E\) is the Young’s modulus identified with subscripts. Equations 4.6 and 4.7 suggest that parallel alignment of CNTs increase interfacial area and thereby results improved load transfer at the interface and reduction in axial stress (\(\sigma\)). Splat interface therefore experiences smooth transitioning of stresses without causing stress concentration. In addition, increased surface area of CNTs
requires higher critical shear stress ($\tau$) to debond and pull out from $\text{Al}_2\text{O}_3$ matrix resulting higher dissipation of fracture energy [217].

Fig. 4.36 shows restraining of the crack damage and consequent crack termination by CNTs. Hence, on one hand, interfacial alignments of CNTs stick the splats, and on the other hand, transverse alignment of CNTs prevents the crack progress. Eventually CNT alignments both along longitudinal and transverse directions contribute to the improved fracture toughness of the plasma sprayed A4C-B nanocomposite coating.

![Fig. 4.36: Restraining of crack-damage by presence of CNTs transverse to crack propagation path.](image)
High magnification SEM image of the A4C-B coating, Fig. 4.37, shows fine (150-250 nm) Al₂O₃ particles attached onto CNT surface. Also, CNTs form a rope structure by aligning with each other. Each CNT in the rope gets coated with molten Al₂O₃. Apart from the melt coating of Al₂O₃ on CNT surface, increase in the CNT diameter (Fig. 4.37) to form CNT ropes can be attributed to CNT fusion. Ceramic-coated CNTs easily fuse with the surrounding matrix and show good anchoring with the splats.

Fig. 4.37: SEM image of nano Al₂O₃ particles sticking to CNT rope surface in AC-B coating.
Fig. 4.38 is a bright field TEM image and shows features similar to Fig 4.37. Fig. 4.38a shows entanglement of CNTs and Al₂O₃ particles hanging on the CNT network. CNT fusion is also observed in Fig. 4.38b indicating superheating of CNTs to fuse with other CNTs. However, CNT structure is intact and not damaged due to fusion. It only forms a CNT network, which will further assist in improving the fracture toughness. Embedded SAD pattern (in Fig. 4.38b) shows diffraction spots corresponding to α-Al₂O₃ phase, as indexed in Fig. 4.39. This is indicative of signal arising primarily from large size (~95 nm) α-Al₂O₃ crystallites (~68.4%), (also see Table 4.8).

Fig. 4.38: a) TEM micrograph depicting non-preferred dispersion of Al₂O₃ particles onto CNT surface, and b) incomplete sintering of Al₂O₃ particles embedded in CNT network and CNT fusion is observed. SAD pattern is embedded along with.
The role of CNT dispersion in enhancing fracture toughness can be understood by comparing A4C-B coating with A4C-SD coating as content of CNT is same in both coatings. CNT distribution in A4C-SD is explained in the next section.

4.6.2: Enhancement of Fracture Toughness by CNT Dispersion in A4C-SD Coating

Figure 4.40 shows CNT dispersion in all the regions of fully melted, partially melted and retained nano Al₂O₃ regions of A4C-SD coating. Presence of CNTs throughout the matrix shall provide homogeneous and consistent properties of the coatings. A4C-SD coating shows additional toughening mechanisms such as CNT pinning of splats, CNT bridge formation and CNT mesh formation.
Fig. 4.40: CNT retention and dispersion is observed in the PM, FM and nano-Al$_2$O$_3$ regions.

CNTs are present on the surface, in between the splat regions, and embedded in the melted region of plasma sprayed A4C-SD coating, Fig. 4.41. CNTs entrapped in the fully melted region indicate their survival in the plasma plume and consequent retention in the deposited coating. CNTs in the fully molten region can directly add onto the excellent mechanical properties. CNTs present on the surface assist shock absorption during impact and serve as energy absorption regions to result enhanced fracture toughness. CNTs embedded between the splat regions serve as glue to hold the splats together and at the same time act as strengthening agents.
Fig. 4.41: Dispersion of CNTs in FM, PM and splat regions is observed in A4C-SD coating.

Fig. 4.42: Non-agglomeration of CNTs in the nano-\(\text{Al}_2\text{O}_3\) particle matrix.
Fig 4.42 shows non-agglomerated CNTs with the nano $\text{Al}_2\text{O}_3$ particles. Retention of nanoparticles in the semi-porous region acts as energy absorption sink in terminating the crack-propagation.

Fig. 4.43 shows sintered $\text{n-Al}_2\text{O}_3$ particles attached to extended CNT emanating from the matrix. $\text{Al}_2\text{O}_3$ particles are entrapped by curved CNTs indicating strong interaction of the surfaces resulting in enhanced wettability. Wetting of surfaces, therefore, becomes a dominant factor in terms of reinforcing the ceramic nanocomposite.

Fig. 4.43: Engulfing of $\text{Al}_2\text{O}_3$ particles by CNTs. $\text{Al}_2\text{O}_3$ sintering is also observed in the SEM image.
Fig. 4.44 shows a CNT bridge between matrix splats. Thickening of CNT bridge is caused by the flow of molten Al₂O₃ over CNT surface. Curvature effect at the ends of CNT is due to the capillary action responsible for covering the CNT along its surface. Excellent wetting of CNT by molten Al₂O₃ is evident from Fig. 4.44. Though increased wetting of CNT with molten Al₂O₃ is observed, currently there is no data on the determination of wetting angle [218] and interaction of Al₂O₃ and CNTs. The existing literature on Al₂O₃-CNT and ceramic-CNT nanocomposite does not refers to wetting phenomenon as these materials are processed via solid state processes such as sintering, spark plasma sintering, and HIP [16, 172, 176, 186].

Fig. 4.44: CNT bridge formation between two splats in A4C-SD coating.
Fig. 4.45 shows an interesting phenomenon of CNT mesh formation in A4C-SD coating which can be explained in terms of wetting kinetics. Surface tension holds the molten Al₂O₃ over CNT surface as a coating during plasma spraying. Since increased contact area leads to reduced surface energy, Al₂O₃ coated CNTs come closer to each other. As the volume of molten Al₂O₃ over CNTs increases, surface tension is not large enough to hold the molten liquid in place. Now capillarity has enough time to allow seeping down of molten surfaces to leave behind a meshed network of CNTs, Fig. 4.45 [197, 219]. These phenomena have been validated by modeling the CNT-Al₂O₃ interface, which is explained in the later section.

![CNT Mesh](image)

**Fig. 4.45:** CNT mesh formation observed due to capillarity in the plasma sprayed A4C-SD coating.
TEM image of plasma sprayed A4C-SD coating, Fig. 4.46a, shows a single dispersed CNT anchored at Al₂O₃ surface. This is in contrast with A4C-B (Fig. 4.38) coating that shows agglomerated CNTs. Enhanced fracture toughness observed for A4C-SD coating is attributed to improved dispersion of CNTs in the matrix, along with enhanced wetting of CNT surface by molten Al₂O₃ during plasma spraying. Embedded SAD pattern depicted presence of two phases, viz. α-Al₂O₃ and γ-Al₂O₃. An enlarged image of the SAD pattern is shown in Fig. 4.46b.

Diffraction spots from γ-Al₂O₃ are observed as secondary diffraction away from the primary transmitted beam, Fig. 4.46b. One of the diffraction spots of α-Al₂O₃ is acting as primary beam for diffraction of γ-Al₂O₃ crystallites. Presence of dual phases is indicative of precipitation of fine γ-Al₂O₃ crystallites (~15 nm) in the α-Al₂O₃ matrix (~73nm).

Vicker indentation radial crack as observed in Fig. 4.47a is restrained by CNT bridges across the propagating crack in A4C-SD coating [220]. Bridge structure reinforces the splat separation during crack propagation leading to topsy-turvy crack path. Hence, higher energy absorption regulates the reduced crack length to reinforce the integrity of the structure. Herein role of dispersed CNTs in A4C-SD coating (Fig. 4.40 and Fig. 4.41) provides multiple sites of anchoring and bridging the splats (Fig. 4.44).

Solid-state sintered region, Fig. 4.47b, (also Fig. 4.42) fulfils the objective of terminating the propagating crack by providing the necessary sink for energy relaxation. Hence, combination of powder treatment to uniformly distribute CNTs, and controlling the plasma parameters to obtain bimodal microstructure can greatly influence improvement in the fracture toughness of the plasma sprayed nanocomposite coating.
Fig. 4.46: a) TEM image of dispersed CNT in Al$_2$O$_3$ matrix showing CNT fusion with Al$_2$O$_3$. Al$_2$O$_3$ particle sintering is also observed. Embedded SAD pattern is enlarged, and b) dual crystalline phases are seen in the diffraction image.
Fig. 4.47: Vicker indentation crack in A4C-SD coating depicting: a) radial crack generation, and b) crack deflection and restraining by CNTs and crack termination at solid state sintered region.
4.6.3: Enhancement of Fracture Toughness by CNT Dispersion in A8C-SD Coating

Plasma sprayed A8C-SD coating shows enhanced CNT content throughout the matrix, i.e., entrapped in the fully melted region, partially melted region and present with nano Al₂O₃ particles (Fig. 4.48). In additions, some agglomerated and undispersed CNT are also observed indicating that CNT-agglomeration effect is not counteracted because of increased CNT content. The role of CNT as reinforcing agent is beneficial in enhancing the toughening of ceramic nanocomposite only in dispersed state. Fig. 4.49 shows clumps of agglomerated CNTs. Agglomeration tendency in the A8C-SD coating is attributed to excessive CNT content (8 wt%) indicating that the optimized CNT content in the composite spray dried agglomerate should be between 4 wt.% CNT and 8 wt. % CNT. Presence of CNTs in the solid state sintered region and presence with nano Al₂O₃ grains, Fig. 4.50, still assists in energy absorption upon impact. But it is possible that increase of fracture toughness by addition of CNTs is getting neutralized by CNT agglomeration. This is indicative of further optimization (of uniform CNT dispersion with maximum CNT content) required for excavating a maxima of fracture toughness in the provided composition range.

CNT fusion is also observed in the plasma sprayed A8C-SD coating, Fig. 4.51, resulting in a CNT rope. Though nanotubes are known for offering tensile- and bend-toughness [221], advantage of nanorope CNTs to serve as strong fortification can be extracted efficiently in the nanocomposite coating. In addition to the intra-nanotube sliding potential of graphitic planes in CNTs, superior toughness of CNT-rope in comparison to carbon fiber of equivalent diameter must be realized owing to higher torsion stiffness and inter-nanotube sliding-capability of CNT-rope structure.
Fig. 4.48: CNT distribution in the plasma sprayed A8C-SD coating.

Fig. 4.49: SEM image showing CNT retention and agglomeration in A8C-SD coating.
Fig. 4.50: Fracture toughening enhancement via presence of CNTs in solid state sintered region and its presence with nano $\text{Al}_2\text{O}_3$ particles. CNT agglomeration as CNT network is also observed.

Fig. 4.51: CNT fusion occurring in A8C-SD coating to result a rope structure.
Schematic of torsional stiffening by rope structure is presented in Fig. 4.52. Shear modulus of nanoropes may be contrasting to that of individual nanotubes since shear in between the nanotubes will dominate over graphitic shear modulus [222]. Intertwining of CNTs induce torsion stiffening upon loading, which improvises the overall toughness of the nanocomposite, Fig. 4.52. Torsion stiffness (K) of CNT is given as [223]:

\[
K = \frac{1}{L} \frac{d^2 E}{d^2 \theta} = \frac{r^3}{L} G(2\pi h)
\]

Equation 4.8

where \( \theta \) is torsion angle and \( G \) is shear modulus of CNT. Strong dependence of torsion stiffness on CNT-diameter imparts enhanced reinforcing effect to the Al\(_2\)O\(_3\) matrix making torsion stiffness an essential toughening parameter.

![Fig. 4.52: Schematic of torsional stiffening by CNTs.](image)

TEM of plasma sprayed A8C-SD coating depicts sintering of Al\(_2\)O\(_3\) particles in Fig. 4.53a. Exaggerated fusion of CNTs is observed in Figs. 4.53a and 4.53b. Entrapment of Al\(_2\)O\(_3\) particles by CNTs confirms the CNT dispersion by high surface energy of nano-
Fig. 4.53: TEM micrograph of A8C-SD coating showing, a) Al₂O₃ sintering with embedded SAD α-Al₂O₃ pattern, and b) CNT fusion and agglomeration with embedded SAD pattern of γ-Al₂O₃.
Al₂O₃ particles. But, excessive agglomeration of CNTs is attributed to the enhanced CNT content in the plasma sprayed A8C-SD coating. Light rings pattern in the embedded SAD image (as two light rings in embedded Figs. 4.53 a and b) correspond to [10\overline{1}0] and [30\overline{3}0] family of planes of CNT. Embedded SAD diffraction pattern in Fig. 4.53a corresponds to α-Al₂O₃ (indexed earlier), whereas SAD diffraction pattern in Fig. 4.53b corresponds to γ-Al₂O₃. Diffraction pattern embedded in Fig. 4.53b is indexed in Fig. 4.54, and zone axis of [1\overline{1}2] is measured.

Fracture toughening of the nanocomposite depends on the load transferability from Al₂O₃ matrix onto CNTs. CNT/ Al₂O₃ interface is critical for stress transfer.
Toughening mechanisms provided by excellent-bending stiffness (eq. 4.5), axial and shear strength (eqs. 4.6 and 4.7), and torsional stiffness (eq. 4.8) can play a part in the presence of a strong CNT/Al₂O₃ interface. Interface is affected by wettability between the two surfaces. Hence wettability between CNT and Al₂O₃ is an underlying phenomenon towards understanding of the interface properties.

### 4.7. Wettability of CNT-Al₂O₃ Interface

Surface tension is defined as the surface force acting along a line per unit length. Interatomic distances are stretched to balance the extra energy associated with the unsatisfied surface atoms, Fig. 4.55a. Contact of the surface with another body often reduces the surface tension and makes the system more stable. Since the nature has tendency to minimize potential energy, adhesion and cohesion interplay to adjust contact area/angle and equilibrate at dynamically stable state, Fig. 4.55b. It is clearly evident that when cohesive force between solid and liquid is stronger than that of liquid and vapor

![Fig. 4.55: a) High energy at surface results from unbalanced atoms, and b) Wetting characteristic of droplet on a substrate.](image)

θ > 90° indicating poor wettability of droplet onto substrate  
θ < 90° indicating good wettability of droplet onto substrate
wetting occurs. Wetting is required in realizing good bonding between the substrate and freezing droplet.

It is important to emphasize that high melting points of Al₂O₃ (~ 2333 K) and CNT (~ 3773 K) makes it very difficult to partake liquid droplet angle studies in such a system. Since no theoretical literature is available on such a study, an effort is made to calculate the freezing dynamics involved in the plasma spraying. Surface tension and capillarity are taken as reflective measures of understanding wettability between CNT and Al₂O₃. Pressure jump across a curved surface can be defined by Young-Laplace Equation:

\[ \Delta p = \gamma \cdot \frac{dA}{dV} \quad \text{Equation 4.9} \]

where \( \gamma \) is the surface energy at constant pressure and temperature, given as:

\[ \gamma = \left( \frac{dG}{dA} \right)_{p,T} \quad \text{Equation 4.10} \]

which can be further simplified as

\[ \Delta p = \frac{2\gamma}{r} \quad \text{Equation 4.11} \]

Equations 4.9 through 4.11 clearly indicate the enhanced free energy of nano particles due to enhanced surface area (at reduced particle radius). CNTs tend to agglomerate in order to reduce their surface energy. But, this aspect is used to our advantage by dispersing CNTs in the nano-Al₂O₃ matrix, fig. 4.4 and fig. 4.5. Owing to increased surface energy of nano Al₂O₃ particles [123], strong seeking and adherence to other surface becomes a requirement to reach stability. Instead of CNT agglomeration,
now selective agglomeration is negated and surface forces dominate toward uniform distribution of particles (in the nanometer range).

CNTs and nano Al₂O₃ particles are seen to distribute without biased agglomeration, Fig. 4.4 and 4.5. The importance of attaching one surface in order to reduce the surface energy is seen via CNT-rope formation in Fig. 4.51. Non-preferential attachment of Al₂O₃ and CNTs is observed by scattered particles in TEM micrograph, Fig. 4.38 and 4.53.

Interfacial studies on alumina-CNT are almost non-existent in the literature owing to increased complexity of ultra high temperatures and solid state processing by other researchers [193]. Since surface is the sole direct contact with surrounding environment, capillarity and surface tension are the only direct measures in defining wettability. A system of Al₂O₃ wetting on CNT is defined through Fig. 4.56. In considering theoretical/computational modeling, it is assumed that molten Al₂O₃ layers onto CNT surface by capillary action, where it freezes instantaneously mimicking the dynamic equilibrium depending on surface tension, 4.57. Rapid kinetics inherent to plasma spraying (dwell time of ~ 4 x 10⁻⁴ s) is fast enough to freeze frozen Al₂O₃ droplet before it dropped from CNT surface, Fig. 4.57. Effects of gravity are neglected and resulting microstructures are taken as representative model for surface forces towards defining wettability. Since constant equilibrium contact angles are not experimentally observed owing to changing ‘true’ contact angles, it is a valid assumption to assume dynamic freezing as a representative model [195].
Fig. 4.56: Theoretical representation of the CNT wetting by molten Al₂O₃.

Fig. 4.57: Rapid kinetics of plasma spraying demonstrating freezing of Al₂O₃ droplet before surface tension is overcome by gravity.
Capillarity force $f_c$ is defined as [219]:

$$f_c = \frac{\gamma(\cos \theta_1 + \cos \theta_2) \Omega}{h_m} \quad \text{Equation 4.12}$$

where $\gamma$ is the surface energy (taken as $\sim 1.59$ J/m$^2$ for Al$_2$O$_3$), and $\Omega$ is surface projection area and $h_m$ is the meniscus height. Higher meniscus infers reduced capillarity, but increased surface tension leading to enhanced wetting of CNTs by Al$_2$O$_3$.

Surface tension $T_s$ is defined as [197]:

$$T_s = l \gamma \cos \alpha \quad \text{Equation 4.13}$$

where $l$ is the perimeter contact, and $\gamma \cos \alpha$ is the vertical component of the surface tension, Fig. 4.56. Surface tension is caused by difference in the magnitude of surface forces where difference in forces of adhesion and cohesion results wetting/dewetting of liquid droplets onto substrates.

Theoretical calculations performed on the system are presented in Table 4.10. CNT diameter is assumed to be 70 nm, approximate angles of contact are calculated from the experimental results (SEM micrograph, Fig. 4.43), and various meniscus heights are assumed to theoretically calculate the perimeter contact, and the surface projection area. Meniscus height, perimeter contact and surface projection area are dominating factors, which define the capillary force and surface tension occurring at Al$_2$O$_3$ coated CNT interface.

Figure 4.58 describes various interactions in assimilating interfacial capillarity and surface tension with respect to meniscus height of Al$_2$O$_3$ over CNT in differential microstructures. This depends on time and temperature experienced during plasma spraying. As the meniscus height increases, the surface tension of molten Al$_2$O$_3$ should
Table 4.10: Theoretical calculations of capillary force and surface tension on Al₂O₃-CNT interface (CNT diameter assumed to be 70 nm): (*) represents calculated value from observed micrograph, Fig. 4.44

<table>
<thead>
<tr>
<th>Meniscus Height (h_m) nm</th>
<th>θ₁ degrees</th>
<th>θ₂ degrees</th>
<th>α degrees</th>
<th>Perimeter contact, (l) nm</th>
<th>Surface Projection area (Ω) nm²</th>
<th>Surface Energy, (γ=1.59 J/m²)</th>
<th>Capillary force (f_c) nN</th>
<th>Surface Tension (T_s) nN nm/nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>16.39</td>
<td>13.34</td>
<td>64.29</td>
<td>30</td>
<td>40</td>
<td>1.59</td>
<td>12.61</td>
<td>40.4</td>
</tr>
<tr>
<td>25</td>
<td>15.36</td>
<td>18.85</td>
<td>25.72</td>
<td>75</td>
<td>90</td>
<td>1.59</td>
<td>11.32</td>
<td>116.3</td>
</tr>
<tr>
<td>30</td>
<td>24.20</td>
<td>17.86</td>
<td>0.00</td>
<td>105</td>
<td>110</td>
<td>1.59</td>
<td>9.83</td>
<td>167.0</td>
</tr>
<tr>
<td>50</td>
<td>14.04</td>
<td>15.64</td>
<td>-38.57</td>
<td>150</td>
<td>160</td>
<td>1.59</td>
<td>10.09</td>
<td>225.1</td>
</tr>
<tr>
<td>70*</td>
<td>14.87</td>
<td>16.32</td>
<td>-90.00</td>
<td>210</td>
<td>195</td>
<td>1.59</td>
<td>8.78</td>
<td>236.2</td>
</tr>
</tbody>
</table>

(* represents calculated value from observed micrograph, Fig. 4.44)
suffice its holding against capillarity to seep down. Similarly low meniscus height insinuates ease of holding down molten Al₂O₃ rather than allowing it to rise as a coating over CNT. It must be clarified that seeping down of Al₂O₃ can occur even at a later stage when enough mass of molten Al₂O₃ dragged over CNT cannot be supported by surface tension. These phenomena can be further split into three stages namely:

i) Stage 1: Surface tension dominated region

ii) Stage 2: Intermixed mode

iii) Stage 3: Capillarity dominated region
Stage 1 is characterized by occurrence of high surface tension (~ 236.2 nN nm/nm), wherein molten Al₂O₃ freezes onto CNT to form a thick coating, Fig. 4.59. The process has just enough time to allow quick flow of molten Al₂O₃ over CNTs via capillary action, and cause consequent freezing of Al₂O₃ as a coating. Though CNTs assist reduction force seeping down of molten Al₂O₃ by capillary flow, surface tension force dominates and holds molten Al₂O₃ as a coating (upto 40-50 nm) over CNT. This bridge-like structure improves the fracture toughness of the nanocomposite by serving as anchors to the adjoining splats.

Fig. 4.59: Stage 1 showing surface tension dominated bridge structure.
Intermediate mode (stage 2) is described by entrapment of CNTs along the solid state sintered Al₂O₃ regions, where wetting by Al₂O₃ is just occurring, and there is just sufficient time to entrap a few CNTs before they can merge as a mesh. Both capillarity and surface tension of Al₂O₃ play a key role in generating the trimodal microstructure, Fig. 4.60. Trimodal microstructure is highly beneficial in enhancing the fracture toughness of nanocomposite, i.e., second phase strengthening by nanoparticles and CNTs, structural strengthening by fully molten regions, energy absorption at partially molten regions, and ductility enhancement by grain sliding of nano particles.

Fig. 4.60: Stage 2 showing CNTs entrapped in Al₂O₃.
Once the molten Al$_2$O$_3$ flows onto CNT surface, the surface tension of liquid attracts the surrounding Al$_2$O$_3$-coated CNT and merges them together. After enough Al$_2$O$_3$-surfaced CNTs are accumulated altogether, it becomes difficult to hold molten alumina by surface tension alone. Owing to the increased volume of surrounding liquid (molten Al$_2$O$_3$), the capillarity reduces the surface tension by its seeping out molten Al$_2$O$_3$ and leaving CNT structure as a mesh, Fig. 4.61 (stage 3). This mesh has a very thin layer (few nanometers) of molten Al$_2$O$_3$, which in true sense is reinforcement by flow of Al$_2$O$_3$ ceramic into intricate surface of CNT. Thickening and smoothening of CNTs is visible in Fig. 4.61.

![CNT Mesh](image)

Fig. 4.61: Stage 3 showing capillarity dominated CNT-mesh.
Hence, excellent wettability of Al₂O₃ on CNT surface is observed via formation of Al₂O₃ coated CNT, solid state sintering and CNT mesh formation (Fig. 4.56-4.61). Rapid solidification kinetics, inherent to plasma spraying, might be reasoned to attribute enhanced wettability observed in the current work. Nucleation frequency (Iᵣ) of Al₂O₃ on CNT surface at a temperature (T) can be expressed by eq. 4.14 as [224]:

\[ Iᵣ = Kᵣ \exp \left( - \frac{ΔG^* f(θ)}{kT} \right) \]

Equation 4.14

where \( Kᵣ \) is kinetic parameter, \( θ \) is the wetting angle, and \( ΔG^* \) is the excess free energy of the critical nucleus, and \( k \) is the Boltzmann constant. Since rapid solidification offers enhanced excess free energy for nucleation, it must be balanced by reduction in the wetting factor \( f(θ) \). The wetting factor expression [225] \( f(θ) = (2 + \cos θ)(1 - \cos θ)^2/4 \) further clarifies reduction of wetting angle (and enhanced wettability). Moreover, heterogeneous nucleation reduces the excess energy barrier of nucleation [225], allowing rapid solidification of Al₂O₃ by wetting CNT surface.

Wetting, therefore immediately raises the question of secondary phase generation at the CNT-Al₂O₃ interface. In order to understand the fundamental thermodynamic predictions, FactSage thermochemistry software is used to construct phase diagram of Al-O-C system (Fig. 4.62). Al₂O₃ and C emerge as the only stable phases present in the current systems (A4C-B, A4C-SD, and A8C-SD). Thermodynamics calculations convey very low activity (~6.8889 x 10⁻¹⁹ at 2200 K) of Al₄C₃ even near the melting point of Al₂O₃, and on the other hand, CO and CO₂ product gases will tend to destabilize the
interface. Consequently, there is no stable reaction product of $\text{Al}_2\text{O}_3$ reaction with $\text{C}$, eq. 4.15.

$$2\text{Al}_2\text{O}_3 + 6\text{C} \rightarrow \text{Al}_4\text{C}_3 + 3\text{CO}_2$$  \hspace{1cm} \text{Equation 4.15}

$\Delta G = +700.2$ KJ (at 2200 K)

$\Delta G = +558.4$ KJ (at 2500 K)

Fig. 4.62: Phase stability diagram calculated from FactSage software [226]. Activity of $\text{Al}_2\text{O}_3 = 1$, activity of carbon =1, and activity of $\text{Al}_4\text{C}_3$= $1.7429 \times 10^{-18}$. 

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Hence, formation of secondary phases such as carbides is not preferred in the Al-O-C system. In addition, absence of prism planes in CNT also restrict free C bonds being available for bonding with Al and O. Surface wetting of CNTs by Al₂O₃ is therefore not influenced by presence of any secondary phases, and can be extracted as direct interacting interface. After the interface is defined, *ab-initio* computational modeling can provide atomic interaction at molecular level. Visualization of interfacial atoms can evolve understanding of interfacial reactions enhancing the fracture toughness of ceramic nanocomposite.

4.8. *Ab-initio* Computational Modeling of CNT-Al₂O₃ Interface

Since capillarity and surface tension are defined by the interfacial reaction of Al₂O₃–CNT, it becomes critical to visualize the molecular phenomenon at such level. Surface energy of oxygen terminated Al₂O₃ is in the range 4.45 – 10.83 J/m², whereas Al terminated Al₂O₃ possesses surface energy of 1.59 J/m² [196]. Though CNT surface is much stable with surface energy of 0.2 J/m², nano surfaces often are understudied and depict non-intuitive behavior. Al-graphite can drain the surface energy down in the 0.02-0.4 J/m² range, making a strong possibility of a stable system [194]. Ooi’s molecular modeling of Al-graphite depicted no bonding at the interface [194]. In addition to the absence of oxygen in the system, representation with specific cross-sectional plane in demonstrating Al-C interface, overall bonding at the interface was absent.

In the current Al₂O₃-CNT system, stable 1x1x1 α-Al₂O₃ (a=4.76 Å x b=4.76 Å x c=12.99 Å, α=90⁰, β=90⁰, γ=120⁰) crystal is interfaced with 2x2x2 CNT surface (a=2.46 Å, b=2.46 Å, c=6.71 Å, α=90⁰, β=90⁰, γ=120⁰), Fig. 4.63. Since Al-terminated Al₂O₃
shows a more stable interface, Al-terminated Al₂O₃ crystal is considered for modeling the interface. Schematic of atomic positions of Al₂O₃-CNT system is presented in Fig. 4.63, which define the system considered for atomistic simulation. Ab-initio SIESTA 1.3 computational modeling details are provided in section 3.6. Conjugate gradient optimization of atomic positions provides equilibrium positions of the resulting structure. This consequently provides electronic density surfaces of various layers that can be plotted to understand the interactions occurring near/at CNT-Al₂O₃ interface. Planes A, B, C and D are considered to describe various interactions occurring at various levels of influence from the interface.

Fig. 4.63: Al₂O₃-CNT system defined in ab-initio computational modeling.
The iso-surface contours for unaffected graphite (0001) layer (plane A) are presented in Fig. 4.64a. This shows the periodic and regular energy contours between the carbon atoms. As the interface approaches, the interference from the Al₂O₃ surface on the graphite layer is depicted by the distortion of the periodic energy contours, Fig. 4.64b. Influence from the surface atoms of Al₂O₃ crystal depicts clustering of electron cloud between aluminum and carbon along the interface. Oxygen terminated Al₂O₃ surface might further destabilize the interface by formation of CO or CO₂. Thermodynamics of the reaction presented as eq. 4.15 clearly obviates the necessity of considering this reaction.

![Periodic and regular iso-surface contours of carbon layer](image1)

![Interference by alumina on the iso-surface contours of carbon](image2)

Fig. 4.64: Isosurface contours of graphite layers at a) uninfluenced planes, and b) near interface plane.

Aluminum terminated Al₂O₃ crystal illustrates presence of three aluminum atoms at the interface (Plane C), Fig. 4.65a, which further insinuates the contribution of
aluminum atoms in distorting the energy contours at the interface (plane D), presented in Fig. 4.65b. Pseudo bonding over the aluminum atoms (in the Al₂O₃ crystal) is implying pseudo bonding with carbon. Since high polarity indicates strong metallic bonding in aluminum [194], distorted iso-surface contours confirm the pseudo metallic bond at Al₂O₃-CNT interface. On the Al-terminated surface, weak binding energy with silver

![image](image_url)

Fig. 4.65: Isosurface of a) aluminum surface-terminated alumina crystal, b) aluminum-carbon pseudo metallic bond interference at alumina-CNT interface.

(~0.5 eV) and low activation barrier (~0.25 eV) might allow rapid diffusion and bonding on this surface [227]. Direct evidence from molecular modeling result clearly shows the possibility of enhanced interfacial bonding between Al and C. Enhanced interfacial strength of Al₂O₃-CNT system will provide enhanced fracture toughening. It makes sense because crystal with high surface energy will try to adhere to a new surface in order to minimize its overall energy.
Some experimental data already demonstrates the stability of Al-C interface (energy ~ 0.02-0.4 J/m²) in comparison to self-existing Al₂O₃ or graphite crystals [194]. It is quite viable that partial bonding at interface interconnects the CNTs by introducing humps and encouraging strong wettability. Moreover, the unconnected region acts as energy sink during impact to further enhance the interfacial strength. Though these molecular simulations consider Al₂O₃-graphite interface, this model closely mimics the surface properties of Al₂O₃-CNT interface. This combination of interfacial linking abridges the gap that had existed in describing the wettability of the CNTs with Al₂O₃. Consideration of bulk structure, gravity effects, and rapid solidification can further alter the interpretations from the molecular modeling studies. An exhaustive modeling of this work is recommended for computational researchers in describing the surface interaction more effectively.

Interfacial atomic interaction can be scaled through interpretation of load transfer at interfaces via formulating i) contact points between Al₂O₃ surface coating the CNT surfaces and ii) Al₂O₃-Al₂O₃ splat interface, which completely describes stress transfer between Al₂O₃ coated CNTs and Al₂O₃ matrix. Since cohesion is stronger in similar adjoining materials (Al₂O₃- Al₂O₃), the weaker interface (Al₂O₃ – CNT interface) is elucidated herewith. Stress distribution along Al₂O₃-CNT interface can be modeled by considering longitudinal stress of σ_f occurring within a distance of dx, balanced by interfacial shearing, τ_i, as shown in Fig. 4.66 using eq. 4.16 as:

\[ 2\pi rt(\sigma_f + d\sigma_f) - 2\pi r2\sigma_f - 2\pi r\tau_i dx = 0 \]

Equation 4.16

which simplifies to
\[
\frac{d\sigma_f}{dx} = \frac{\tau}{t}
\]

Equation 4.17

where \( t \) is the wall thickness of the CNT.

Wettability influences the transition from interfacial to longitudinal stresses. Thereby shearing between \( \text{Al}_2\text{O}_3\)-CNT interface reinforces energy absorption resulting in enhanced toughening.

\[
\sigma_f + d\sigma_f
\]

Fig. 4.66: Interfacial Shear at \( \text{Al}_2\text{O}_3\)-CNT interface.

Dominance of surface tension in conjunction with rapid freezing (section 4.7) can evolve fascinating toughening junctions, as seen in Fig. 4.67. To reduce its surface energy, molten \( \text{Al}_2\text{O}_3 \) accumulates coated CNTs as ropes. Incomplete termination of process can result formation of Y-junction ropes, which act as toughening anchors in the matrix. Multi directional load bearing capability (shear and axial strength, eqs. 4.6 and 4.17) and flexibility (bending stiffness, eq. 4.5) at other end of the junction-ropes can improve the fracture toughness of the ceramic nanocomposite.
Though various mechanisms interplay in deciding the fracture toughness, representative mechanical properties of generated structure can also be mapped from its elastic modulus. Next section deals with the use of nanoindentation technique in evaluating modulus of the plasma sprayed coatings.

![CNT Rope Formation by Al₂O₃ Freezing](image)

**Fig. 4.67:** Dominance of surface tension in forming CNT Y-junction ropes.

4.9. Nanomechanical Modulus Mapping of Plasma Sprayed Coatings

Nanomechanical dynamic analyzer provides unprecedented characterization of materials, such as modulus mapping of the plasma sprayed coatings over an area (to render the stiffness associated with the content and distribution of CNTs). Nanoindenter tip oscillates with small forces and rasters along the surface to monitors the displacement and phase lag arising from material’s response. Though surface asperities also influence the modulus of the coating, uniformity of modulus can insinuate the dispersion of CNTs in the coatings. Surface profile and corresponding modulus mapping is correlated with
the distribution of modulus values to associate the overall range of modulus pertaining to
the coating. Instead of complex modulus, which involves the loss modulus, storage
modulus is taken as direct representative of the Young’s modulus considering the non-
plastic fracturing of ceramics. Area of 4μm x 4 μm is selected for modulus mapping in
order to scan wide enough section in commenting the role of CNT content and dispersion
in contributing enhanced modulus of the plasma sprayed coatings.

4.9.1. Modulus Mapping of A-SD Coating

Surface profile of A-SD coating, Fig. 4.68a, is showing fine surface undulations
and nano scratches. Corresponding modulus mapping, Fig. 4.68b, show uniform structure
with modulus average of approximately around 210 GPa as observed in the color spread.
Blue streaks correspond to the scanning limit of the SPM (Scanning Probe Microscopy)
and edge effects arising from sharp corners. Neglecting those features, distribution of the
modulus along distance, Fig. 4.68c, and histogram of storage modulus, Fig. 4.68d,
provide average storage modulus around 210 GPa. Lower modulus values are typical to
the plasma sprayed coating, which are attributed to layered structure and porosity, which
is inherent to plasma sprayed structure. These modulus values can be taken as base values
for comparison with modulus mapping in presence of CNTs.

4.9.2. Modulus Mapping of A4C-B Coating

Topography of A4C-B coating is observed in Fig. 4.69a, which shows uniform
structure. Modulus mapping of A4C-B coating, Fig. 4.69b, shows storage modulus in the
range of 200-300 GPa, with some regions reaching values upto 650 GPa, Fig. 4.69c. Localized high value of modulus is attributed to the presence of CNTs at such locations. Modulus mapping histogram shows average modulus of ~ 250 GPa, Fig. 4.69d. Shift of average modulus in A4C-B to higher vale (when compared to A-SD coating) is attributed to the presence of CNTs in the structure.

4.9.3. Modulus Mapping of A4C-SD Coating

In the uniform area of the topographical image shown in Fig. 4.70a, average modulus value is ~ 300 GPa in the mapped storage modulus (Fig. 4.70b). Distribution of modulus along the distance is presented in Fig. 4.70c. Reduced modulus values (<250 GPa) is arising from the valleys present on the surface. Dominating regions of modulus appear bimodally at 250 GPa and 350 GPa, as presented in the histogram, Fig. 4.70d. Modulus values average at ~ 300 GPa, which is higher than modulus of A4C-B coatings (~250 GPa). Increase in the modulus is attributed to uniform CNT dispersion in the matrix. Bimodal modulus distribution can be attributed to the presence of FM and PM regions.

Reduced modulus of ~ 250 GPa can be reasoned to presence of high solid state sintered region where particles are consolidated by surface sintering. Increased modulus of ~ 350 GPa can be reasoned to FM region which is densified by surface melting and resolidification (along with entrapped CNTs). Hence the role of CNT dispersion (in A4C-SD coating) is lucidly detailed by modulus mapping in comparison to that of A4C-B coating.
Fig. 4.68: Plasma sprayed A-SD coating showing, a) Surface topography, b) Modulus map, c) Line scan distribution of storage modulus with distance, and d) Histogram of modulus distribution.
Fig. 4.69: Plasma sprayed A4C-B coating showing, a) Surface topography, b) Modulus mapping, c) Line scan distribution of storage modulus with distance, and d) Histogram of modulus distribution.
Fig. 4.70: Plasma sprayed A4C-SD coating showing, a) Surface topography, b) Modulus mapping, c) Distribution of storage modulus with distance, and d) Histogram of modulus distribution.
4.9.4. Modulus Mapping of A8C-SD Coating

Topography of plasma sprayed A8C-SD coating appears rough, Fig. 4.71a, but modulus mapping shows enhanced overall modulus, Fig. 4.71b. Dispersion of modulus line scan, Fig. 4.70c shows modulus in the range of 200 – 500 GPa. Here again, bimodal modulus peaks are observed at ~325 GPa and ~ 420 GPa in Fig. 4.71d. Increase in the modulus is attributed to enhanced CNT content in the coating. Dispersion of CNTs in the solid state sintered region is resulting modulus ~ 325 GPa, whereas CNTs reinforcing the fully molten region further increase the modulus to ~ 420 GPa. Hence, modulus mapping vividly differentiates the role of CNT content in enhancing the elastic modulus and fracture toughness of the A8C-SD coating when compared to A4C-SD coating.

Since storage modulus falls in phase with the materials response, it is the direct measure in relating the resilience of a material. Stiffness associated with the content and distribution of CNTs is captured in the nanomechanical dynamic characterization. Corresponding fracture toughness of the plasma sprayed coatings directly correlate with the modulus maps. Local area evaluation of modulus in the coatings, therefore, is a direct measure of increase in fracture toughness. Nanomechanical analysis becomes an unmatched tool in mapping mechanical properties of the plasma sprayed nanocomposite coatings. Localized effects of dispersion strengthening by CNT content and CNT dispersion are clearly visualized through nanomechanical data. Modulus mapping, therefore, completes the picture of associated fracture toughening via CNT reinforcement.
Fig. 4.71: Plasma sprayed A8C-SD coating showing, a) Surface topography, b) Modulus mapping, c) Line scan distribution of storage modulus with distance, and d) Histogram of modulus distribution.
5. CONCLUSIONS

CNT reinforced aluminum oxide nanocomposite coatings have been successfully synthesized using plasma spray technique. Fracture toughness enhancement of upto 57% have been attained in the engineered coatings. Role of CNT retention and dispersion is elucidated towards understanding various toughening mechanisms. A detailed list of conclusions regarding the powder treatment, microstructural and phase characterization, Al₂O₃/CNT interface, and mechanical properties are presented herewith.

1. Blending of A-SD powder with 4 wt. % CNTs (A4C-B powder) has resulted dispersion of CNTs on the surface of powder agglomerate, and accumulation of CNTs in the dome cavity of the powder agglomerate. Composite spray drying of nano Al₂O₃ and 4 wt. % / 8 wt. % CNTs (A4C-SD and A8C-SD respectively) have resulted dispersion of CNTs uniformly throughout the powder agglomerate. CNTs are also dispersed in the core of the powder agglomerate.

2. Plasma spray parameters were effectively controlled using inflight particle sensor. In combination with pretreatment of the powder and controlled process parameters, a bimodal microstructure (fully melted and resolidified structure, and solid state sintered structure) was engineered in Al₂O₃ matrix.
3. CNTs are successfully retained in the plasma sprayed A4C-B, A4C-SD, and A8C-SD coatings as confirmed via microstructural characterization and Raman analysis. Retention of CNTs is attributed to the rapid kinetics (dwell time of $\sim 4 \times 10^{-4}$ s) involved in the plasma spraying.

4. Distribution and content of CNTs in the powder agglomerate have strong effect on the thermal exposure observed by powder particles in the plasma plume (measured via inflight particle monitor). A4C-B coating experienced temperatures of 2898 K (with velocity of 271 m/s), which is higher than that of A-SD coating (2512 K, 289 m/s). A4C-SD and A8C-SD coatings experienced reduced thermal exposure (2332 K (244 m/s), and 2241 K (232 m/s) respectively) owing to core-CNTs serving as ‘heat-sinks’.

5. Generation of metastable $\gamma$-Al$_2$O$_3$ phase is dependent on the thermal exposure experienced by CNTs available on the surface of powder agglomerate. Bimodal crystallite size ($\sim 20$ nm for $\gamma$-Al$_2$O$_3$ and $\geq 40$ nm for $\alpha$-Al$_2$O$_3$) is obtained.

6. Addition of 4 wt.% CNT (in A4C-B coating) showed an improvement of 20 % in fracture toughness (from 3.22 to 3.86 MPa m$^{1/2}$) when compared to AC-SD coating. But, addition of dispersed 4 wt.% CNTs (A4C-SD) coating showed an improvement of 43% (to 4.60 MPa m$^{1/2}$). This evinces enhancement of fracture toughness just because of CNT dispersion owing to powder treatment. Further, increasing the CNT content to 8 wt.% (A8C-SD coating), fracture toughness
increased to 5.04 MPa m$^{1/2}$ which is indicative of fracture toughness enhancement (by 57%) because of increase in CNT content.

7. Toughening features such as impact alignment of CNTs along interface, CNT hooks, CNT bridge formation, CNT fusion on Al$_2$O$_3$ surface, and CNT chain loop formation are observed. Enhancement of toughness is attributed to mechanisms such as interfacial shear strengthening, high bending stiffness, enhanced energy absorption, and good torsional stiffness associated with excellent mechanical properties of CNTs.

8. Improved wetting is observed between molten Al$_2$O$_3$ and CNT surface. Phenomenon like CNT bridging, CNT entrapment in the solid-state sintered region, and CNT mesh formation are dominated by interplay of surface tension and capillarity. As estimated from theoretical computation modeling, high surface tension (~236 nN) was pinned for promoting high meniscus height, whereas increased capillarity (~12.6 nN) was reasoned for CNT meshing.

9. Computational ab-initio modeling was performed to evince electronic charge density perturbations observed along the Al$_2$O$_3$-CNT interface. Overlapping of electron cloud at the Al$_2$O$_3$-CNT interface in forming pseudo metallic bonding (charge density > 0.3), which is indicative of adhesion and good wettability at the interface. Effect of rapid kinetics inherent to the plasma spraying is attributed to enhanced-wettability associated in the Al$_2$O$_3$-CNT system.
Effect of CNT dispersion in enhancing the elastic modulus is identified using nanoindentation dynamic modulus mapping. A-SD (∼210 GPa), A4C-B (∼250 GPa), A4C-SD (∼280 GPa) and A8C-SD (∼325 GPa and 420 GPa), clearly indicate increase in the modulus with increasing CNT content and distribution.
6. RECOMMENDATIONS FOR FUTURE

Every dissertation is a connecting link between new paradigm of science and engineering tying its free end with the advancement of scientific knowledge. Plasma spraying of CNT reinforced Al₂O₃ was performed to produce enhanced toughening of the nanocomposite coatings using pretreatment of powders. Detailed optimization, microstructural characterization, phase analysis and mechanical property evaluation of the coatings have elicited toughness improvement of upto 57% when compared to that of unreinforced coating. During the completion of the dissertation or a major research project, it is always the desire to hover on a few more dimensions of research. Perspectives from different directions and endless discussions on some of the undiscovered aspects germinate the seeds of further development and are recommended for future dwelling as follows:

6.1. Optimization of CNT Content in Aluminum Oxide Matrix

Fracture toughness enhancement of 43% (4.60 MPa m¹/²) was observed in A4C-SD coating when compared to that of A-SD coating. Agglomeration of CNT in A8C-SD coating could not be fully avoided even though it displayed fracture toughness enhancement of 57% (5.04 MPa m¹/²). This is indicative of intermediate composition of CNT (between 4 and 8 wt.% CNTs) in Al₂O₃ matrix where CNTs can be full dispersed without agglomeration to provide highest toughness. A systematic parametric study could be performed to optimize CNT content and its uniform dispersion in providing maximum fracture toughness.
6.2. Plasma Spraying of Insitu grown CNTs On Al₂O₃ Powder Particles

As observed in the current research that CNT dispersion is critical in imparting enhanced fracture toughness. Instead of adding CNTs separately in the powder feedstock via blending and spray drying, CNTs can be grown insitu on Al₂O₃ powders. Advantage of insitu grown CNTs can provide the required anchoring and dispersion throughout the matrix in each powder particle. However, a critical control is required on insitu grown CNT length and their density on each Al₂O₃ powder particle. CNT length should be short enough to avoid entanglement and subsequent clogging of powders during plasma spray. On the other hand, CNTs should be long enough to allow inter-splat anchoring.

6.3. Molecular Dynamics Simulation of Al₂O₃/CNT Interface

Energy of formation could be compared for different structures such as Al₂O₃, α-Al₂O₃/CNT interface and γ-Al₂O₃/CNT interface in deriving stable atomic configuration. Molecular dynamics approach can be applied to calculate force of separating CNTs from Al₂O₃ surface. Measuring the force of separation points out the work of adhesion required that would be absorbed by interface before fracture. This can directly indicate the dependence of interface on the fracture toughness related to the modeled system.

6.4. Hot-Isostatic Pressing of Plasma Sprayed Nanocomposites

Role of secondary densification processing in enhancing the fracture toughness of the plasma sprayed nanocomposites can throw light on the role of porosity in imparting toughening behavior. Retention and distribution of CNTs in the hot-isostatically pressed (HIPped) coatings will elucidate the damage tolerance of CNTs under high temperature
and pressure. Consequently, balance between the retention of CNTs (and its damage) and the porosity closure will decide the overall toughening of the nanocomposite. HIP study will differentiate the role of porosity versus CNT on the fracture toughness of Al₂O₃-CNT nanocomposite coating.

6.5. Estimating Flexural Strength of Plasma Sprayed Al₂O₃-CNT Nanocomposite

To estimate the flexural strength of the bulk structure, three-point bend test can be utilized on freestanding samples. But, one of the major challenges is to machine such a thick and large sample for 3-point test from these brittle materials. Flexural strength should be evaluated in plasma sprayed and hipped conditions to differentiate the role of porosity in toughening (and/or strengthening) the ceramic nanocomposite.

6.6. Sliding Wear Evaluation of Al₂O₃ with CNT Addition

One of the major applications of Al₂O₃ is as wear resistant coatings. Effect of CNT content in improving the wear resistance by anchoring the splats should be studied for its tribological performance. Hence sliding wear resistance of the novel coating with different CNT reinforcement should be evaluated. In addition, investigation of Al₂O₃-CNT frictional coefficient and abrasion resistance can explain its usage in thermal liners and insulating systems.
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APPENDICES

Appendix 1: As-Received CNT Characterization

SEM image of CNTs show an average diameter ranging between 40-70 nm, Fig. A1. Though CNTs show XRD diffraction peak of 2-theta around 26 degrees, Fig. A2, high intensity peaks arising from Al₂O₃ suppress CNT peaks in plasma sprayed nanocomposite coatings.

Fig. A1: SEM image showing CNT diameter in range of 40-70 nm.

Fig. A2: XRD spectrum of CNTs showing intense peak at 26 degrees.
Appendix 2: Modulus Enhancement with CNT Addition (Micromechanics Approach)

Micromechanics illuminate the interaction of matrix and reinforcement to determine the overall properties of composite depending on the individual properties constituents [41]. Hence the intention of adding CNT reinforcement in the Al₂O₃ matrix can be insinuated from micromechanics approach. Anisotropy, aspect ratio and morphology of the reinforcement are also accounted in mechanics model to describe the resulting properties of resulting combination. When overall response of the composite can be predicted via constituent relationships, it becomes easier to design the homogeneous (and generally isotropic) material [42].

Tensile and shear modulus of randomly distributed CNTs in a matrix can be described by eq. A1 and A2 as [43]:

\[
E = \frac{3}{8} E_{11} + \frac{5}{8} E_{22} \quad \text{Equation A1}
\]
\[
G = \frac{1}{8} E_{11} + \frac{1}{4} E_{22} \quad \text{Equation A2}
\]

where longitudinal modulus \((E_{11})\) is given by eq. A3 as:

\[
E_{11} = \frac{1 + 2(l_{\text{CNT}} / d_{\text{CNT}}) \eta_L v_{\text{CNT}}}{1 - \eta_L v_{\text{CNT}}} E_m \quad \text{Equation A3}
\]

and transverse modulus \((E_{22})\) is given by eq. A4 as:

\[
E_{22} = \frac{1 + 2 \eta_T v_{\text{CNT}}}{1 - \eta_T v_{\text{CNT}}} E_m \quad \text{Equation A4}
\]

and the other parameters are expressed as:

\[
\eta_L = \frac{(E_{\text{CNT}} / E_m)^{-1}}{(E_{\text{CNT}} / E_m) + 2(l_{\text{CNT}} / d_{\text{CNT}})} \quad \text{Equation A5}
\]
\[
\eta_T = \frac{(E_{\text{CNT}} / E_m)^{-1}}{(E_{\text{CNT}} / E_m) + 2} \quad \text{Equation A6}
\]
Properties of such a composite are same in all the directions. By substituting the tensile moduli of $\text{Al}_2\text{O}_3$ (taken as 390 GPa, and Poisson’s ratio of 0.22), one can calculate the overall effect of volume fraction of CNTs, and effect of $l/d$ ratio of CNTs on the composite stiffness. In addition, upper and lower range limits of Young’s modulus of CNTs, a working range of response can be obtained for composite material. Fig. A3 considered various $l/d$ ratios of the CNTs and $E_r/E_m (=E_r)$ ratios for various combinations to state that there was not much effect of $l/d$ ratio of CNT on the effective stiffness. This can be attributed to very small lengths of CNTs considered (0.5 to 15 microns) and randomly aligned CNTs display average constant-volume-behavior. Young’s modulus of CNT is considered between 300 GPa (lower curve in Fig. A3) to 1 TPa (Upper curves in Fig. A3).

![Variation of E with fiber vol. fraction](image)

**Fig. A3**: Effective stiffness of randomly dispersed CNTs in $\text{Al}_2\text{O}_3$ matrix.
Young’s modulus of CNT (ECNT) is of critical importance since no significant effect of l/d ratio plays a part in deciding E11 of composite. But, at the same time, volume percent of CNT has stronger effect on tensile and shear stiffness of the composite.

Eshelby and Hashin Shtrikman approach do not consider L/D ratio or particulate geometry. As observed for randomly oriented fibers, that l/d ratio do not impart much effect on determining the overall stiffness of the composite, this is an acceptable assumption. In addition linear elastic approach considered in this model is true in our case owing to brittle nature of the ceramic composites. Tremendous improvement in the properties of the CNT reinforced composite justifies the strengthening via CNT addition.

Rule of mixture (ROM), eq. A7, considers one-dimensional model to determine elastic modulus (Ec) [44].

\[ E_c = V_{CNT} E_{CNT} + V_m E_m \]  

Equation A7

where the \( V_{CNT} \) and \( V_m \) are the volume fraction of CNT, and matrix respectively, and \( E_{CNT} \) and \( E_m \) are the Young’s modulus of CNTs and matrix respectively.

Eshelby approach considers effective Young’s modulus of heterogeneous material with linearly elastic material response. It assumes randomly distributed spherical particles in the matrix leading to isotropic nature of properties. Standard bulk, shear and elastic modulus equations can be used in converting moduli dependencies, i.e, K=E/3(1-2ν); G=E/2(1+ν); E=9KG/(3K+G). Bulk and shear modulus are given by eqs. A8 and A9 respectively:

\[ \bar{K} = K^M - \frac{fK^M}{K^M - \frac{K^M - K^{CNT}}{1 + \nu^M}} \]  

Equation A8
\[
\bar{G} = G^M - \frac{fG^M}{G^M - G^{CNT}} \left[ \frac{2(4 - 5v^M)}{15(1 - v^M)} \right]
\]

Equation A9

where \( K^M, G^M \) are bulk and shear moduli of matrix, \( v^M \) is matrix’s Poisson’s ratio, \( K^{CNT}, G^{CNT} \) are bulk and shear moduli of CNTs, and \( f \) is volume fraction of CNTs.

Hashin-Shtrikman shows narrowed bounds of isotropically heterogeneous material. Equations describing the bounds are expressed as:

\[
\frac{f}{1 + (1 - f)(K^{CNT} - K^M)} \leq \frac{\bar{K} - K^M}{K^{CNT} - K^M} \leq \frac{f}{1 + (1 - f)(K^{CNT} - K^M)}
\]

Equation A10

\[
\frac{f}{1 + (1 - f)(G^{CNT} - G^M)} \leq \frac{\bar{G} - G^M}{G^{CNT} - G^M} \leq \frac{f}{1 + (1 - f)(G^{CNT} - G^M)}
\]

Equation A11

Condition I: If \((G^{CNT} - G^M)(K^{CNT} - K^M) \geq 0\)

then \( K^* = 4G^M/3; K^+ = 4G^{CNT}/3 \)

\[
G^- = \frac{3}{2 \left( \frac{1}{G^M} + \frac{10}{9K^M + 8G^M} \right)}
\]

Equation A12

\[
G^+ = \frac{3}{2 \left( \frac{1}{G^{CNT}} + \frac{10}{9K^{CNT} + 8G^{CNT}} \right)}
\]

Equation A13

Condition II: If \((G^{CNT} - G^M)(K^{CNT} - K^M) \leq 0\)

then, \( K^- = 4G^{CNT}/3; K^+ = 4G^M/3 \)

\[
G^- = \frac{3}{2 \left( \frac{1}{G^M} + \frac{10}{9K^{CNT} + 8G^M} \right)}
\]

Equation A14
Independency of particle shape in Hashin Shtrikman’s approach, provides enough leverage in calculating the bounds without worrying about the aspect ratio, and the isotropy of the second phase. Fig. A4 compares the theoretical Young’s modulus calculations by various approaches. Experimental modulus values of Al₂O₃-CNT coatings are compared herewith.

Fig. A4: Effective Young's modulus calculated by various micromechanics models.

Often plasma sprayed coatings depict reduced modulus owing to presence of lamellar microstructure and presence of defects (such as porosity, inclusions, etc) inherent to plasma spraying, as described in section 4.9.1.
Appendix 3: This section includes list of directly related publications (peer-reviewed journal articles) during the current Ph.D. research work. First page of the published papers directly related to the current research is provided here.

Peer-Reviewed Journal and Proceedings


Role of Powder Treatment and CNT Dispersion in the Fracture Toughening of Plasma-Sprayed Aluminum Oxide – Carbon Nanotube Ceramic Nanocomposite

Kantesh Balani, Srinivasa Rao Bakshi, Yao Chen, Tapas Laha, Arvind Agarwal*
Mechanical and Materials Engineering Department, EAS 3464, 10555 W. Flagler St., Miami, FL-33174, USA

ABSTRACT:

Al₂O₃ ceramic reinforced with 4-wt.% multiwalled carbon nanotube (CNT) is plasma sprayed for improving the fracture toughness of the nanocomposite coating. Two different methodologies of CNT addition have been adopted in the powder feedstock to assist CNT dispersion in the nano-Al₂O₃ matrix. First, spray-dried nano-Al₂O₃ agglomerates are blended with 4 wt.% CNT as powder-feedstock, which is subsequently plasma sprayed resulting in the fracture toughness improvement of 19.9%. Secondly, spray dried composite nano-Al₂O₃ and 4 wt.% CNT powder was used as feedstock for attaining improved dispersion of CNTs. Plasma sprayed coating of composite spray dried powder resulted in increase of 42.9% in the fracture toughness. Coating synthesized from the blended powder displayed impact alignment of CNTs along splat interface, and CNTs chain loop structure anchoring the fused Al₂O₃ melt whereas coating synthesized from composite spray dried powder evinced anchoring of CNTs in the solid state sintered region and CNT mesh formation. Enhanced fracture toughness is attributed to significance of CNT dispersion.

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*Corresponding Author: Dr. Arvind Agarwal, Ph: 305-348-6533, Fax: 305-348-1932, Email: agarwala@fiu.edu
Challenges and advances in nanocomposite processing techniques

V. Viswanathan a, T. Laha b, K. Balani b, A. Agarwal b, S. Seal a,*

*Advanced Materials Processing and Analysis Center, Department of Mechanical, Materials and Aerospace Engineering, Nanoscience and Technology Center, University of Central Florida, Orlando, FL 32826, USA
"Mechanical and Materials Engineering, Florida International University, Miami, FL 33174, USA

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Abstract

Of late, nanotechnology seems to be rapidly thrusting its applications in all aspects of life including engineering and medicine. Materials science and engineering has experienced a tremendous growth in the field of nanocomposite development with enhanced chemical, mechanical, and physical properties. A wide array of research has been conducted in the processing of nanocomposites. Consolidation of these systems from loose particles to bulk free form entities has always been a challenge. To name a few, traditional consolidation techniques such as cold pressing and sintering at high temperatures, hot pressing, and hot isostatic pressing have strong limitations of not being able to retain the nanoscale grain size due to the excessive grain growth during processing. This article reviews in detail the results from numerous studies on various methods for manufacturing nanocomposites with improved properties and retained nanostructures. Both challenges and recent advances are discussed in detail in this review. © 2006 Elsevier B.V. All rights reserved.

Keywords: Nanostructures; Processing techniques; Mechanical properties; Processing challenges

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* Corresponding author. Tel.: +1 407 882 1119; fax: +1 407 882 1462.
E-mail address: seal@mail.ucf.edu (S. Seal).

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Plasma-sprayed carbon nanotube reinforced hydroxyapatite coatings and their interaction with human osteoblasts in vitro

Kantesh Balani, Rebecca Anderson, Tapas Laha, Melanie Andara, Jorge Tercero, Eric Crumpler, Arvind Agarwala*

*Department of Mechanical and Materials Engineering, Florida International University, EC 2464, 10555 West Flagler Street, Miami, FL 33174, USA

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Abstract

Carbon nanotubes (CNT) possess excellent mechanical properties to play the role as reinforcement for imparting strength and toughness to brittle hydroxyapatite (HA) bioceramic coating. However, lack of processing technique to uniformly distribute multiwalled CNTs in HA coating and limited studies and sparse knowledge evincing toxicity of CNTs has kept researchers in dispute for long. In the current work, we have addressed these issues by (i) successfully distributing multiwalled CNT reinforcement in HA coating using plasma spraying to improve the fracture toughness (by 56%) and enhance crystallinity (by 27%), and (ii) culturing human osteoblast hFOB 1.19 cells onto CNT reinforced HA coating to elicit its biocompatibility with living cells. Unrestricted growth of human osteoblast hFOB 1.19 cells has been observed near CNT regions claiming assistance by CNT surfaces to promote cell growth and proliferation.

Keywords: Hydroxyapatite coating; Carbon nanotube (CNT); Plasma spraying; Titanium alloy; Crystallinity; Human osteoblast

1. Introduction

Hydroxyapatite (HA), Ca₁₀(Po₄)₆(OH)₂, is an attractive biomaterial owing to its close chemical resemblance (Ca/P = 1.67) with bone and teeth [1-3]. Osteoblasts proliferate onto HA owing to its bioactivity and biocompatibility [3,4] and therefore HA coatings have long been applied to dental implants, bone repair scaffolds, skeletal implants, and body/biointerface material [5]. Microstructure, crystallinity, and phase composition of HA coating is critical in deciding its cell response and mechanical performance. Plasma sprayed HA coating often result in the generation of secondary phases such as tricalcium phosphate (TCP), tetracalcium phosphate (TTCP), calcium oxide (CaO), and amorphous calcium phosphates (ACPs) [6]. Though HA is very stable in the body environment, presence of secondary phases causes dissolution leading to degradation of the implant in vivo. Hence, higher crystallinity content is required for the increased implant life. In addition, researchers have used carbon nanotubes (CNT), Ti-alloys, yttria stabilized zirconia (YSZ), Ni₃Al, and alumina (Al₂O₃) reinforcements to HA coating [7-9] for improving its fracture toughness and wear resistance [1,5,10-13].

Chen et al. investigated mechanical properties of laser processed HA-multiwalled CNT coating showing strong improvement in the fracture toughness and marginal improvement in the elastic modulus [1,14]. However, laser synthesized coating result in the formation of undesired TiC phase [1]. The purpose of addition of CNT was to enhance the mechanical performance of the coating without deteriorating the biocompatible properties of HA [15,17]. On the contrary, CNTs have also been debated as toxic [18] under organic environment [1,19]. Though in a recent study, osteosarcoma cell growth has been observed on functionalized CNT [20], researchers have also depicted micro patterning of CNTs to result directed growth of osteoblasts [21], or surface modification with DNA- and HA-nanostructured films for enhanced detection and biosensitivity [22]. Haddon et al. has concluded that CNT
Effect of carrier gas on mechanical properties and fracture behaviour of cold sprayed aluminium coatings

S. R. Bakshi1, T. Laha1, K. Balani1, A. Agarwal*1 and J. Karthikeyan2

Two different coatings of 1100 aluminium were cold sprayed onto similar substrates, using He and He–20N2 (vol.-%) mixture as carrier gases. Three point bend testing was carried out. The elastic moduli of the coatings were found to be close to each other and the substrate. The He processed coating showed higher fracture strength which was attributed to the higher degree of strain hardening. The He–20N2 processed coating failed at lower stress owing to its strain relaxed structure. The mode I fracture of the coating substrate system was found to be higher for the helium processed coating. The toughness was correlated to the microstructure. The delaminated coating showed a higher degree of brittle failure of the interface for the He processed coating.

Keywords: Cold spraying, 1100 aluminium, Fracture surface, Fracture toughness, Three point bend test, Elastic modulus, Brittle fracture, Delamination, Notch, Porosity, Carrier gas, Splats, Dislocation pile up, Subgrain formation

Introduction

Cold spraying is a relatively new coating technique where metallic powder particles are accelerated to very high velocities (600–1500 m s−1) by a carrier gas (He, N2) flowing at very high pressure (up to 3.5 MPa) and impact the substrate through a converging diverging de laval nozzle. The carrier gas is preheated to a temperature between 300 and 800°C, partly in order to compensate for the cooling produced by expansion of the gas and to reduce the density which in turn increases the sonic velocity, but below the melting point of the particles. The loss of kinetic energy on impact causes plastic deformation of the particles. This process is also known as cold kinetic spraying. The low temperature solid state coating process eliminates problems owing to oxidation and defects owing to solidification. The disadvantage is that a large amount of carrier gas is lost, unless recycled, and that only plastically deformable materials can be deposited. Many materials have been deposited till date by cold spraying including pure metals,7,8 alloys9,10 and composite materials.11,12 The exact mechanism of the bonding is not fully understood. It is believed that impact of the particles results in rupture of oxide layers which provides clean surfaces for bonding. Bonding has been attributed to adiabatic shear instabilities at particle/particle and particle/substrate interfaces owing to impact and has been modelled using finite element analysis.10,11 The parameters affecting the process and spraying efficiency are particle size, density of particles, temperature of gas and density of gas and spraying angle, and various models have been proposed for the effect of various parameters.12–14

In the authors’ earlier work,6,16 cold spraying of 1100 Al on 1100 Al substrate was reported. Two different carrier gases, namely He and He 20N2 (vol-%), were used to study the effect of carrier gas on the properties of the coating. It was observed that the 100% He processed coating had higher hardness than the He-20N2 processed coating. TEM analysis revealed dislocation pile-ups and oxide layers in the He processed coating. The He-20N2 processed coating showed subgrain formation and a relaxed structure.16 There was more strain hardening in the He processed coating as compared with He 20N2 processed coatings. Potentiodynamic polarisation experiments in 0.9 pH H2SO4 revealed that the corrosion current density was higher for the He processed coating than for the He-20N2 processed coating indicating superior corrosion resistance of the He 20N2 processed coating.6

The purpose of the present work is to study the effect of the carrier gas on the fracture strength and interfacial properties of the coating/substrate and to establish correlation with the microstructure. Three point bend testing of the samples was carried out. The elastic modulus and fracture properties were studied.

Experimental

The schematic of the three point bend test is shown in Fig. 1. A notch was introduced at the midpoint of the specimen using a diamond saw. The initial crack length was measured by optical microscopy of the cross-section. Table 1 tabulates the dimensions of the samples used.

For the three point bend test geometry the elastic modulus of the coating/substrate system is given by the
Synthesis, Microstructural Characterization, and Mechanical Property Evaluation of Vacuum Plasma Sprayed Tantalum Carbide

Kantesh Balani, Gabriela Gonzalez, and Arvind Agarwal

Mechanical and Materials Engineering, Florida International University, Miami, Florida 33174

Robert Hickman and J. Scott O’Dell

Plasma Processes Inc., Huntsville, Alabama 35811

Sudipta Seal

Advance Material Process and Analysis Center (AMPAC), Department of Mechanical, Materials and Aerospace Engineering (MDMAE), University of Central Florida, Orlando, Florida 32816

Tantalum carbide (TaC) is an ultra-high-temperature ceramic for potential applications as protective coating, furnace components, propulsion liners for space shuttles and aircrafts, etc. Microstructural and mechanical behavior of vacuum plasma-sprayed (VPS) TaC has been investigated in the present study. Apart from major TaC phase, microstructural definitions elucidated TaC, non-stoichiometric TaC, phases (0.82 ≤ x ≤ 0.94), partial grain formation, polymerization of grains, and inhomogeneous C:Ta ratios in the sprayed material. Near-isotropy in the fracture toughness ratio (K/Ia/K/Ic = 1.01) is attributed to compact coating, fine-closed porosity, and distribution of non-stoichiometric phases.

1. Introduction

THERMAL protection systems, nozzle throat inserts, and high erosion resistant thrusters require ultra-high-temperature capability of material for withstanding exit gases under propulsion environment. Candidate materials involve refractory metals, ceramic matrix composites, carbon-carbon composites, cermet and intermetallics, but no single material has the desired combination of physical, mechanical, and structural properties. Ceramic oxides are chemically inert, but they are more brittle and susceptible to thermal shock than non-oxide ceramics. On the one hand refractory metals excel in high-temperature strength, wear, and erosion resistance, but on other hand possess low oxidation resistance, high density, poor machinability, and high cost. Ceramic borides possess high melting temperature, high hardness, and high thermal and electrical conductivity, but oxidation of boride leads to formation of particles prone to thermal shock. Plasmas sprayed TaC is an alternative to overcome fabrication hurdle of high melting point materials and their brittleness. Plasma powder reaches temperatures in excess of 10,000 K and impacts the powders in molten or semi-molten states, generating typical mechanically bonded layered structure. It is important to emphasize that much of the literature research material for the applications of TaC, because of military applications, is sparingly available. In the current work, microstructural characterization is correlated with the observed mechanical property of the vacuum plasma-sprayed (VPS) TaC.

II. Experimental Procedure

As-received TaC powder (composition of 93.4 wt% C, 0.1034 wt% O, and rest Ta), shown in Fig. 1, has faceted morphology and is 1–2µm in size. Spray drying of these fine powder particles resulted in spherical agglomerates of porous cakes. (Fig. 2) between 10 and 45 µm, which is adequate for plasma spraying. This spherical morphology and ideal size help reduce interparticle friction, enhance flowability of powder during plasma spray and thereby improve the density of the sprayed structure. Vacuum plasma spray formed tantalum carbide (TaC) provides an alternative to overcome fabrication hurdle of high melting point materials and their brittleness. Plasma reaches temperatures in excess of 10,000 K and impacts the powders in molten or semi-molten states, generating typical mechanically bonded layered structure. It is important to emphasize that much of the literature research material for the applications of TaC, because of military applications, is sparingly available. In the current work, microstructural characterization is correlated with the observed mechanical property of the vacuum plasma-sprayed (VPS) TaC.
NEAR NET SHAPE FABRICATION VIA VACUUM PLASMA SPRAY FORMING

K. Balani*, A. Agarwal* and T. McKechnie*

*Mechanical and Materials Engineering, Florida International University,
10555 W. Flagler Street, CEAS 3464, Miami, FL-33174, USA.
**Plasma Processes Inc., 4914 Moores Mill Road, Huntsville, AL 35811, USA

E-mail: kbalani@yahoo.com

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ABSTRACT

Near net shape fabrication substantiates the manifestation of final structure near to the desired shape, therefore can easily cut down the price of finished product contributed by machining. Though conventional powder metallurgy techniques are prominent in fabricating near net shape structures with controlled porosity, it still lags in producing thin walled complex shapes with functionally graded/ differential structures. Hence, fabrication of complex near net shapes using Vacuum Plasma Spray (VPS) forming has emerged as innovative rapid prototyping technique for a variety of applications. In the present work, the nuances of near net shape fabrication by VPS technique have been explained. Few examples of spray formed structures such as hypereutectic aluminum-silicon, nanostructured aluminum oxide, and intermetallic iron aluminide thin sheet have been discussed. A detailed case study of VPS formed thin TaC structure for ultrahigh temperature application has been elucidated via near net shape fabrication.

1. NEAR NET SHAPE FABRICATION USING VPS FORMING

Owing to extreme brittle nature, conventional fabrication techniques find difficult to shape refractory materials and intermetallics into desired shapes. Moreover machining techniques incur extreme wastage of material and require dead time for the finishing of the structural component 1-2. Though few processes satisfy the near-net-shape fabrication criteria, a serious limitation is posed by the extreme melting points of refractory materials. Ultrahigh temperature material application range between 2500-3300 K and therefore the selection of crucibles to hold the melt and help to form the finished product becomes quite difficult. Plasma spraying has been used as a versatile technique for depositing metals, alloys, polymers and ceramics as coatings 3. Temperature in excess of 10,000 K are easily reached in the plasma flame, hence can melt any known material with great ease. With an impact velocity of 1-3 Mach, plasma sprayed structure depict a typical mechanically bonded layered structure. VPS utilizes HF (high-frequency) started, DC plasma to heat and accelerate powder feedstock for deposition onto negative shaped substrate/ mandrel for fabricating positive shape structure, Fig. 1. Generally Ar is used as the primary plasma gas with secondary He or H2 gas for increased heat transfer. Plasma gun can be manipulated by computer control in six axes of motion. With the cooling rates usually observed in the range of 103-108K/s, generation of ultrafine grain microstructure and non-equilibrium phases is not so surprising in the Vacuum Plasma Spray (VPS) formed structured 2-3. Both metals and ceramics have been processed with densities exceeding 97% of theoretical densities 2-6. Plasma gun and the mandrel are computer-controlled in VPS forming, hence fabrication of complex structures can be designed accordingly. Deposition rates can go as high as 20 lb/h (9 kg/h). Near net shapes, spray deposited onto preformed mandrel, help reduce the finish machining of the deposited structure. Removal of mandrel from the deposited structure hold a critical importance in the near-net fabrication of VPS formed structures. A comparison between VPS forming and conventional P/M processing route is schematically
Molecular modeling of metastable FeB$_{49}$ phase evolution in laser surface engineered coating

K. Balani and A. Agarwal

Department of Mechanical and Materials Engineering, Florida International University, EAS 3400, Miami, Florida 33174

Narendra B. Dahotre

Department of Materials Science and Engineering, The University of Tennessee, Knoxville, Tennessee 37996 and Materials Processing Group, Metals and Ceramic Division, Oak Ridge National Laboratory, Oak Ridge, Tennessee 37831

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Interstitial iron-boride phases have been a subject of research interest for a long time owing to their useful properties. Metastable FeB$_{49}$ phase evolved during nonequilibrium laser surface engineering was investigated along with FeB, Fe$_2$B, and Fe$_3$B phases. Theoretical x-ray diffraction spectrum derived from numerically constructed FeB$_{49}$ crystal matched with the experimental diffraction pattern from laser surface engineered boride coating on the steel substrate. Furthermore, employment of ab initio SIESTA 1.3 molecular modeling for computation of total crystal energy elucidated instability of the FeB$_{49}$ phase. The generation of thermodynamically nonequilibrium phase FeB$_{49}$ along with Fe$_2$B phase was further confirmed by selected area diffraction and high-resolution transmission electron microscopy analyses. © 2006 American Institute of Physics. [DOI: 10.1063/1.2172704]

1. INTRODUCTION

Lasers have been used for surface modification for improvement in material properties such as resistance to wear, friction, erosion, fatigue, corrosion, and high-temperature oxidation. Rapid laser material processing involves extremely high cooling rate ($10^9 \text{ K/s}$) that results in the generation of metastable and nonequilibrium phases due to extended solid solubility. It is often observed that these phases are unknown to equilibrium phase diagram.

In our earlier work, we have synthesized TiB$_2$ coating on AISI 1010 steel substrate using a 2.5 kW continuous wave neodymium-doped yttrium aluminum garnet (Nd:YAG) laser. Rapid cooling and solidification during laser processing generated "composite" coating consisting of borides in iron-rich matrix. TiB$_2$ was the major phase along with Ti and iron borides such as FeB, Fe$_2$B, and Fe$_3$B.

In continuation of the above mentioned efforts, in the present work, phases of iron-boride system (FeB, Fe$_2$B, Fe$_3$B, and FeB$_{49}$ phases) were considered for ab initio total energy simulation using SIESTA 1.3 (Spanish initiative for electronic simulations with thousands of atoms) molecular modeling package. Evolution of rarely reported metastable FeB$_{49}$ phase was further validated employing high-resolution transmission electron microscopy (HRTEM). An effort was made to compare these phases with theoretically generated Fe-B phases and correlate the total energy to the relative stability of the structure.

Apart from major TiB$_2$ peaks, experimental x-ray diffraction (XRD) spectrum depicted some extra peaks corresponding to metastable phases. These extra peaks were identified for FeB$_2$, FeB$_{49}$, and Fe$_2$B$_{12}$B$_{10}$X phases. The experimentally obtained XRD pattern was compared with theoretically computed XRD pattern (Fig. 1) for confirmation of the presence of FeB$_{49}$ and Fe$_2$B$_{12}$B$_{10}$X phases. FeB$_{49}$ crystal construction [Fig. 2(a)] and its computed XRD pattern were obtained using CARINE 3.1 (Divergents A, Compiegne, France) crystallographic software. The strongest peak corresponding to FeB$_{49}$ appeared to be overlapped by (001) peak of TiB$_2$. Even though Table I presented lattice parameters of various phases in laser synthesized composite coating, the primary interest of the present study resides with the characterization of metastable FeB$_{49}$ phase, which so far is not reported in the open literature.

Thermodynamic calculations have shown that the free energy of TiB$_2$ is the least and therefore is a most stable
Transmission electron microscopy of cold sprayed 1100 aluminum coating

K. Balani a, A. Agarwal a,*, S. Seal b, J. Karthikeyan c

* Department of Mechanical and Materials Engineering, Florida International University, EC 3464, 10555 West Flagler Street, Miami, FL 33174, United States
b Advanced Material Process and Analysis Center (AMPAC), Department of Mechanical, Materials and Aerospace Engineering (MMAE), University of Central Florida, Eng 381, 4000 Central Florida Blvd, Orlando, FL 32826, United States
c ASB Industries Inc., Barberton, OH 44203, United States

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Abstract

Cold-spraying of 1100 Al powder particles onto 1100 Al substrate was accomplished using carrier gases 100 vol.% He and a mixture of He-20 vol.% N₂. TEM characterization correlates very well the occurrence of dislocations, presence of oxide, and surface instability phenomenon with observed mechanical and corrosion properties of the cold-sprayed coating.

Keywords: Cold spraying; 1100 Aluminum coating; TEM

1. Introduction

In the 1990s, the cold spraying process emerged in North America, having been developed in the mid-1980s by the Siberian Division of the Russia Academy at the Institute of Theoretical and Applied Mechanics [1-3]. A high-pressure (~3.5 x 10⁶ N/m²) supersonic gas jet is utilized for accelerating fine powder particles to above a critical velocity (~500-1200 m/s), to deposit them as coatings [1-8]. The kinetic energy subsequently dispensed during the impact of the powder particle with the substrate ruptures the surface oxide, plastically deforms the particle and it approaches the clean surface of the substrate, thereby bonding particle as a deposition coating [9].

In our earlier work, 100 vol.% helium (He) and a mixture of helium and 20 vol.% nitrogen (He-20 vol.% N₂) were used as carrier gases for depositing 1100 Al onto 1100 Al substrate via the cold spray technique [3]. The higher sonic velocity and higher degree of tamping in 100 vol.% He when compared to that of He-20 vol.% N₂ carrier gas resulted in denser and harder coatings [3,10]. The addition of N₂, being a diatomic gas, into He increased the enthalpy of the carrier gas for better heat-transfer with spray particles [11]. However, addition of N₂ in He reduced the velocity of sprayed particles due to the heavier atomic mass resulting in coatings with reduced density and hardness [3]. Also, 100 vol.% He processed coating displayed inferior corrosion resistance when compared to that of He-20 vol.% N₂ at 0.9 pH using sulfuric acid (H₂SO₄) as an electrolyte [3]. Both the cold-sprayed coatings were more corrosion resistant compared to the 1100 Al substrate.

In the present study, high magnification scanning electron microscopy (SEM) and transmission electron microscopy (TEM) were utilized to elucidate and substantiate results and findings of our earlier work [3]. TEM imaging elucidated dislocation piling, surface...
Effect of carrier gases on microstructural and electrochemical behavior of cold-sprayed 1100 aluminum coating

K. Balani*, T. Laha*, A. Agarwal*, J. Karthikeyanb, N. Munroc*

*Department of Mechanical and Materials Engineering, Florida International University, 10555 W Flagler Street, Miami, FL 33174, United States

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Abstract

1100 Aluminum has been cold sprayed onto similar substrate using 100 vol.% He and mixture of He–20 vol.% N₂ as carrier gases. Analysis and characterization of sprayed deposits under different processing conditions was performed through optical micrography, stereological analysis, scanning electron microscopy, and microhardness depth profiles. Potentiodynamic polarization curves and Tafel extrapolation experiments were carried at 0.9 pH value using sulfuric acid (H₂SO₄) as an electrolyte. Helium processing condition displayed more compact structure and higher hardness for the coating when compared to He–20 vol.% N₂ processing. Electrochemical studies depicted better corrosion resistance of the He–20 vol.% N₂-processed coatings when compared to coatings processed with helium alone.

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Keywords: X (C) Cold Spray

1. Introduction

Cold spray is a relatively new coating process developed in the mid-1980s at the Institute of Theoretical and Applied Mechanics of the Siberian Division of the Russian Academy of Science in Novosibirsk [1,2]. High-pressure carrier gas (-3.5 MPa or 500 psi) is required to impart high kinetic energy to the powder particles and achieve supersonic velocities (~500–1200 m/s; [2,7]). After the powder particles are accelerated to these high velocities, they impact the substrate causing the thin surface oxide layers to rupture, plastically deform at surface, come in close proximity of clean surfaces, thereby reconsolidating the particles bonding them together. A high degree of plastic deformation occurs at the surface regions due to the impact of the accelerated particles onto the substrate. The source of bonding is the kinetic energy achieved by the particles under high-pressure accelerating gas, to impinge onto substrate at temperatures ranging much below the melting point of the powder particles.

No microstructural evidence has been reported indicating the melting of the spray particles [6]. The bonding in particles in this process is attributed to the adiabatic shear instabilities at the particle substrate or particle particle interfaces, caused due to the heavy particle impact [8].

The high velocity is developed by compressed gas flow (usually helium, nitrogen, or their mixture) through high inlet pressure in a converging diverging de Laval type of nozzle [4,6]. With increasing velocities, the particles undergo process of erosion and abrasion to deposition onto substrate. Hence, critical velocity is a definite requirement for generating sufficient kinetic energy to cause plastic deformation of the surfaces, thereby leading to deposition of spray material onto substrate with a strong bonding [6,7]. Therefore, velocity of particles, attained prior to impact, holds a very important place in cold spraying. Brittle spray materials, such as ceramics, do not undergo the required plastic deformation to form a good bond with the substrate, thereby the solution to the above problem is using ductile matrix or filler materials [9]. Hence, the cold-spray process is ideally suited for synthesizing pure metallic coatings. This process resembles explosive welding in a manner that the
Synthesis of Nanostructured Spherical Aluminum Oxide Powders by Plasma Engineering

T. LAHA, K. BALANI, A. AGARWAL, S. PATIL, and S. SEAL

Irregularly shaped aluminum oxide particles were plasma atomized resulting in narrow size range distribution of spherical nanostructured powders. Cooling rates, on the order of 10^6 to 10^8 K/s, were obtained from the different quenching medias, viz. air, water, and liquid nitrogen. Plasma-engineered powder particles developed nanosize crystallites, while solidification provided insight into the morphological feasibility in refinement of grain size. X-ray diffraction (XRD) methods have been used to quantify the crystallite size obtained with different quenching media. Raman peak shift validated the X-ray analysis in anticipating the grain refinement with increasing cooling rates. Salient structural morphology characteristics and a detailed understanding of spheroidized plasma-sprayed alumina powders were analyzed through scanning electron microscopy (SEM) studies. Formation of nanograins, novel metastable phases, and amorphous structure were endorsed by transmission electron microscopy (TEM) investigations.

I. INTRODUCTION

Nanostructured materials exhibit superior properties such as excellent strength, toughness, and hardness, as obvious from the presence of refined grains. The nanostructured coatings/structures experience superior and novel properties; thus, the importance of achieving such tremendous improvement in performance cannot be neglected by researchers. Plasma spraying is one of the methods used to form nano composite structure, but it must overcome obstructing and difficult processing challenges. The first challenge is that the low mass and poor flow characteristics restrict the smooth flow of individual nanoparticles in the spray nozzle during thermal spraying. In addition, nano-sized powders tend to agglomerate and form clogs in the plasma gun nozzle, hindering the smoothness of the flow and thereby imparting nonuniform coating. The other challenge is to retain the nanograins in the powder because grains coarsen, experiencing the high temperatures in the plasma flame. Several techniques are being developed to get nanocoating/composites by plasma such as employment of spray dried agglomerates of nanopowders, blending and spraying of bimodal powders with controlled plasma parameters, explosion of loose nano agglomerates in plasma flame, etc. In this study, we are trying to modify the feedstock powder by two ways simultaneously, viz. spheroidization and nanocrystallization/grain refinement of spray particles. It is anticipated that synthesis of nanostructured ceramic spheres by plasma spray technique would result in development of an ideal free flowing powder feedstock to synthesize nanostructured coatings, and small complex parts with improved physical properties. Spheroidization by plasma spraying has been studied earlier, but the novelty of this article lies in the idea of using different quenching media (viz. air, water, and liquid nitrogen) to achieve nanostructured grains in individual spherical powder particles through the plasma spraying process. Al2O3 (alumina) powder has been selected for the present study because spherical nanostructured alumina powder is an excellent starting material for near-net shape forming, spray powders for coatings and soft abrasives, surface adsorbents in chemical industries, biocompatible coatings, and as carriers for catalyst. This powder is also being used potentially in thermal insulation as well as in inertial confinement fusion of nuclear materials.

Thermal plasma spraying has been used extensively within many industries for a variety of applications because of its versatility to synthesize fine metal, alloy, ceramic, polymer, and composite powders. Plasma spheroidization is a modified form of atomization technique, where the temperature reaches more than 15,000 K in plasma flame with gas velocities reaching speeds of 1 to 3 Mach. Because of the short residence time of the particle in the plasma, impurity limits in the product are generally low. The added beauty of the process is the use of inert carrier gases minimizing the chemical reaction of the processed powder, which possesses higher surface energy than the starting powder, due to its refined size. Processing of ceramics such as alumina powders becomes much easier since the temperatures achieved in plasma are much higher than the melting point of any known material. The surface tension on the molten particles, upon disintegration, fragments it into spheroids of fine particles added with quenching from water, and liquid nitrogen helps to nucleate and retain the nanocrystalline/grain structure of the particle. Plasma-spheroidized powder experiences rapid cooling, resulting in a high nucleation rate for solidified particles, without any significant time for grain growth.

II. EXPERIMENTAL PROCEDURE

Commercially available aluminum oxide powder (99.8 pct purity) of irregular shape in the 15- to 45-μm particle size
Tribological Behavior of Plasma Sprayed Carbon Nanotube Reinforced Hydroxyapatite-Coating in Physiological Solution

Kantesh Balani¹, Yao Chen¹, Sandip P. Harimkar², Narendra B. Dahotre², Arvind Agarwal¹*

¹Department of Mechanical and Materials Engineering, Florida International University, Miami, FL 33174, USA
²Department of Materials Science and Engineering, University of Tennessee, Knoxville, TN 37996, USA

Abstract: Wear behavior of plasma sprayed carbon nanotube (CNT) reinforced hydroxyapatite (HA) coating is evaluated in the simulated body fluid environment. Apart from enhancing the fracture toughness and providing biocompatibility, CNT reinforced HA coating demonstrated superior wear resistance (> 1.5 times) compared to that of hydroxyapatite coating without CNT. Initiation and propagation of microcracks during abrasive wear of plasma sprayed hydroxyapatite coatings was suppressed by CNT reinforcement. Surface characterization and wear studies have shown that in addition to acting as underprop-lubricant, CNTs provide reinforcement via stretching and splat-bridging for enhanced abrasion resistance in vitro.

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* Corresponding author. Tel:305-348-1701, Fax:305-348-1932
Email address: agarwala@fiu.edu (Dr. Arvind Agarwal)
VITA

KANTESH BALANI

1999  B. E., Metallurgical Engineering, 
PSG College of Technology 
Coimbatore, India  
Best Outgoing Student

2001  M.Tech., Metallurgical and Materials Engineering, 
Indian Institute of Technology (IIT) Madras 
Chennai, India  
Sudharshan Bhat and S. Ananthramakrishnan Memorial Prize  
Deutscher Akademischer Austausch Dienst (DAAD) Scholarship

2002  M.S., Materials Science and Engineering 
University of Kentucky 
Lexington, KY  
RCTF (Research Challenge Trust Fund) Fellowship

2003-  Doctoral Candidate, Materials Science and Engineering 
Florida International University 
Miami, FL  
Best PhD Student  
Dissertation Year Fellowship  
2006-07 Dean’s Award of Highly Productive Doctoral Students

Publications in Peer-Reviewed Journals:


