MELT INFILTRATION OF CERAMIC PREFORMS FOR FUNCTIONALLY GRADED MATERIALS

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I hereby declare that all information in this document has been obtained and presented in accordance with academic rules and ethical conduct. I also declare that, as required by these rules and conduct, I have fully cited and referenced all material and results that are not original to this work.

Name, Surname: Caner Erdamar

Signature:
ABSTRACT

MELT INFILTRATION OF CERAMIC PREFORMS FOR FUNCTIONALLY GRADED MATERIALS

Erdamar, Caner
Master of Science, Metallurgical and Materials Engineering
Supervisor: Prof. Dr. Ali Kalkanlı

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The aim of this work was to produce ceramic-metal composite which shows combination of high hardness and high flexural strength by a combination of hard front layer and tough back layer by Functionally Graded Materials (FGM) process. Composition is the most important variable to obtain gradual changing green/bulk density and/or porosity which are the requirements of an FGM sample. Three different layers of FGM with gradual changing green density and/or porosity were pressed following by ball milling of different weight % of boron carbide and silicon carbide particles with different particle sizes. Each green body was sintered, and then, pressure melt infiltration process was applied to fill porosities with aluminum alloy in order to reach the near theoretical density. Finally, the material was characterized by phase analysis, particle size and distribution analysis, green/bulk density and porosity measurements, optical microscope and SEM analyses, three point bending test, hardness measurement and XRD analysis. Flexural strength measured for FGM B₄C-SiC-Al composites having dimensions of 12 cm length, 4 cm width, 2.3 cm thickness is nearly 348 ± 40 MPa.

Keywords: Functionally Graded Materials, Ceramic-Metal Composite, B₄C-SiC-Al Composite, Melt Infiltration
ÖZ

FONKSİYONEL DERECELİ MALZEMELER İÇİN ÖN ŞEKİLLENDİRİLMİŞ SERAMİKLERİN İNFİLTRASYONU

Erdamar, Caner
Yüksek Lisans, Metalurji ve Malzeme Mühendisliği
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Bu çalışmanın amacı; ön sert tabaka ve arka sünek tabakanın birleşimiyle yüksek sertlik ve yüksek eğme dayanımı özelliği gösteren seramik-metal kompozitlerin Fonksiyonel Dereceli Malzeme (FDM) süreci ile üretilmesidir. FDM numunenin gereksinimlerinden olan dereceli değişen yaş/yığınsal yoğunluk ve/veya porozite elde edilebilmesi için kompozisyon en önemli değişkenleridir. Farklı tane boyutlarına sahip bor karbür ve silisyum karbür tozlarının farklı ağırlık yüzdelerinde bilyalı deformede öğütülmesi ve daha sonra preslenmesi ile dereceli değişen yaş yoğunluklara ve poroziteye sahip olan FDM’nin üç farklı tabakası üretildi. Üretilen her bir numuneye ısıl işlem uygulandı ve azami teorik yoğunluğa ulaşabilmek için basınçlı infiltrasyon süreciyle poroziteler alüminyum alaşımı ile dolduruldu. Son olarak; faz analizi, tane boyut ve dağıtım analizi, yaş/yığınsal yoğunluk ve porozite ölçümü, optik mikroskop ve SEM analizi, üç nokta eğme testi, sertlik ölçümü ve XRD analizi ile kompozit malzemelerin karakterizasyonu gerçekleştirildi. 12 cm uzunluk, 4 cm genişlik ve 2,3 cm kalınlığa sahip FDM BaC-SiC-Al kompozitlerinin eğme dayanımı yaklaşık olarak 348 ± 40 MPa ölçüldü.
Anahtar Kelimeler: Dereceli Değişen Malzemeler, Seramik-Metal Kompozit, Bor Karbür-Silisyum Karbür–Alüminyum Kompozit, İnfiltrasyon
To My Dear Family
ACKNOWLEDGEMENTS

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CHAPTER 1

INTRODUCTION

In defense applications, the development of armor materials with low density, high hardness and high flexural strength is very important to obtain weight materials with improved performance properties such as wear resistance and energy absorption. Three main material classes which are ceramics, metals and composites are preferred for ballistic threats. Engineering ceramics have superior properties such as high elastic modulus, high wear and corrosion resistance, low density and low thermal expansion coefficient. However, they have poor fracture toughness and high expense which limits their industrial applications. Metals are strong, ductile, good thermal and electrical conductor. However, they are not well suited for high temperature applications, shrink or expand with temperature changes and are prone to chemical and corrosion attacks. Composite materials are advance materials consist of one or more materials with distinct properties. They exhibit the best of properties of combinations which are different from individual materials. Achieving the best properties with eliminating undesirable ones led to come up with a new class of material called FGM which is the shortening of “Functionally Graded Materials”. FGM can be considered as an effective armor material because it serves different actions at different layers [1, 2, 3].

In this thesis, B$_4$C – SiC – Al composites that have low density and a combination of high hardness and flexural strength were produced by cost effective melt infiltration process to obtain FGM which may be used against ballistic threats. Also, the production of high ceramic fraction composites with minimized porosity at lower temperatures (<1400 °C) was aimed. The specimens were characterized by phase analysis, green/bulk density and porosity measurements, particle size and distribution
analysis, X-ray diffraction (XRD) analysis, light microscope and scanning electron microscope (SEM) analyses, hardness measurement and three point bending test.
CHAPTER 2

LITERATURE REVIEW

2.1. THE CONCEPT OF FGM

Achieving the best properties with eliminating undesirable ones led to come up with a new class of material called FGM which is the shortening of “Functionally Graded Materials”. Due to their unique properties, FGM have attracted great attention from researchers in many fields such as future space programs, biomaterials, armor materials. FGM have a continuous variation of material properties like thermal, mechanical and electrical properties with respect to position due to non-homogeneous microstructure, atomic order or chemical composition. With the FGM concept, specific materials for specific applications can be produced. Some applications of FGM are as follows: smart structures, MEMs and sensors, electronics and optoelectronics, medicine, biomaterials, power plant, manufacturing, aerospace and aeronautics, energy, defense [4].

Powder metallurgy process with two or more powders may be used to obtain FGM with a stepwise structure. The schematic view of the FGM structure is illustrated in Figure 2.1. There are phase changes from the upper part to the lower part as the volume fraction of B phase increases. Both A and B phases may serve as a matrix. The upper part consists of phase-B particles embedded in Phase-A matrix called A dominant part. Whereas, the lower part consists of phase-A particles embedded in phase-B matrix called B dominant zone. There is a transition zone in the middle part because the dominant phase can’t be determined easily.
Gradation provides the material to have desirable properties of each component at a specific layer. For instance, ceramic-metal FGM have properties of metal such as high fracture toughness and high electrical and thermal conductivity without eliminating required properties of ceramic such as high hardness and high thermal resistance. Gradation can be supplied by gradual changing of microstructure, atomic order or chemical composition from one end to the other [5]. Ceramic-metal FGM may be obtained by building-up a ceramic preform with graded porosity (green density) and
its melt infiltration with molten metal. A graded preform is built up layer by layer with each layer containing an appropriate amount of constituent powders. Layers are compacted and/or sintered into a bulk form that exhibits variation of porosity. According to porosity content, the required amount of metal to fill all pores is calculated then infiltrated in appropriate conditions. The resulting material exhibits a differing ceramic-metal ratio and so properties [6].

The differences in structure and properties between FGM and non-FGM (conventional composite materials) are illustrated in Figure 2.2. For non-FGM, ceramic and metal phases are distributed homogeneously so thermal conductivity and mechanical strength don’t change with position. While for FGM, ceramic and metal phases are distributed non-homogeneously so thermal conductivity and mechanical strength change with position. Because the left part mostly consists of ceramic phases, mechanical strength is higher and thermal conductivity is lower. Because the right part mostly consists of metal phases, thermal conductivity is higher and mechanical strength is lower.

![Figure 2.2. The differences in structure and properties between FGM and conventional composite materials [7]](image)
The joining of dissimilar materials is so difficult and problematic. Because different materials have different thermal expansion coefficient, the combination of them results in residual stress due to ceramics do not undergo plastic deformation to accommodate volume changes. Also, different volumes of different materials cause undesirable reactions at the interface. To cope with these problems, FGM as interlayers has been established. FGM annihilates the sharp interfaces in composite materials in which failure is initiated. It transforms this sharp interface to a gradient interface and a smooth transition from one material to others is obtained. Some problems due to the joining of ceramic and metal and the role of gradient for interlayer are illustrated in Figure 2.3. Inclusions, unjoined islands or pores, reaction layer with cracks, surface damage, unjoined edge, interface flaw and crack into ceramic may form between ceramic and metal without gradient. When the interlayer has no gradient (top), the pieces do not join. However, when a graded interlayer is used (bottom), the two pieces can be successfully joined to each other.

**Figure 2.3.** (a) Some problems due to joining of ceramic and metal, (b) The role of gradient for interlayer [2]
2.2. THE FIRST APPLICATION OF FGM

FGM concept has arisen in 1984 by material scientists in Japan for the spacecraft industry. The outer surface of the spacecraft should be durable against high temperature and high temperature changes. At that time there was no suitable material that could resist this high thermomechanical loading. Thus, Dr. Niino et al. prepared a thermal barrier material that was a new composite by using heat resistant ceramic at high temperature zone and high thermal conductive metal at low temperature zone with gradual changing of ceramic/metal composition. By this way, the outer surface of the spacecraft was protected by gradual distribution of temperature [7].

2.3. DEFENCE APPLICATION OF FGM

The development of armor materials that have low density and high protection ability is very crucial for defense applications. There are a lot of parameters that should be considered for choosing materials in armor system. In ballistic attacks, the most widely used threat type is the projectile with various calibers, core hardness and velocity. Therefore, the ballistic resistance of armor against these kinds of projectiles should be considered. Personnel armor can be classified into soft armor (textile based systems with fibers) and hard armor (ceramic, metal or ceramic-metal composite). Soft armors are used against low level threats (NIJ level III or less), whereas hard armors are used against higher level threats (NIJ level IV) Hence, there is no single best armor material against all threats which are given in Table 2.1 [8].
Some important properties that affect ballistic properties are required for armor systems as shown in Table 2.2. Firstly, the weight of the armor system is based on density so materials that have low density should be used to enhance maneuver capacity. Only materials that have higher hardness value of that of the projectile can break up the projectile when it strikes on the armor system. The ballistic performance of the armor system depends on also compressive and tensile strength. Compressive strength is related with the thickness of the armor system for protection. Materials that have low tensile strength fail at an early stage of the perforation process without exhibiting high compressive strength. Hence, high strength provides multi-hit resistance.

Table 2.1. NIJ Standard 0101.04 [9]

<table>
<thead>
<tr>
<th>Armor Type</th>
<th>Test Round</th>
<th>Test Bullet</th>
<th>Bullet Weight</th>
<th>Reference Velocity (ft/s)</th>
<th>His Per Armor Part at 0° Angle of Incidence</th>
<th>His Per Armor Part at 30° Angle of Incidence</th>
<th>Shots Per Panel</th>
<th>Shots Per Sample</th>
<th>Shots Per Threat</th>
<th>Total Shots Req’d</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>1</td>
<td>.22 caliber LR LRN 2.6 g</td>
<td>2.6 g</td>
<td>329 m/s (1080 ft/s)</td>
<td>4</td>
<td>44 mm (1.73 in)</td>
<td>2</td>
<td>6</td>
<td>12</td>
<td>24</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>.380 ACP FMJ RN 6.2 g</td>
<td>6.2 g</td>
<td>322 m/s (1055 ft/s)</td>
<td>4</td>
<td>44 mm (1.73 in)</td>
<td>2</td>
<td>6</td>
<td>12</td>
<td>24</td>
</tr>
<tr>
<td>II A</td>
<td>1</td>
<td>9 mm FMJ RN 8.0 g</td>
<td>8.0 g</td>
<td>341 m/s (1120 ft/s)</td>
<td>4</td>
<td>44 mm (1.73 in)</td>
<td>2</td>
<td>6</td>
<td>12</td>
<td>24</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>40 S&amp;W FMJ 11.7 g</td>
<td>11.7 g</td>
<td>322 m/s (1055 ft/s)</td>
<td>4</td>
<td>44 mm (1.73 in)</td>
<td>2</td>
<td>6</td>
<td>12</td>
<td>24</td>
</tr>
<tr>
<td>II B</td>
<td>1</td>
<td>9 mm FMJ RN 8.0 g</td>
<td>8.0 g</td>
<td>367 m/s (1205 ft/s)</td>
<td>4</td>
<td>44 mm (1.73 in)</td>
<td>2</td>
<td>6</td>
<td>12</td>
<td>24</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>.357 Mag JSP 10.2 g</td>
<td>10.2 g</td>
<td>436 m/s (1430 ft/s)</td>
<td>4</td>
<td>44 mm (1.73 in)</td>
<td>2</td>
<td>6</td>
<td>12</td>
<td>24</td>
</tr>
<tr>
<td>III A</td>
<td>1</td>
<td>9 mm FMJ RN 8.2 g</td>
<td>8.2 g</td>
<td>436 m/s (1430 ft/s)</td>
<td>4</td>
<td>44 mm (1.73 in)</td>
<td>2</td>
<td>6</td>
<td>12</td>
<td>24</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>.44 Mag JHP 15.6 g</td>
<td>15.6 g</td>
<td>436 m/s (1430 ft/s)</td>
<td>4</td>
<td>44 mm (1.73 in)</td>
<td>2</td>
<td>6</td>
<td>12</td>
<td>24</td>
</tr>
<tr>
<td>III B</td>
<td>1</td>
<td>7.62 mm NATO FMJ 9.8 g</td>
<td>9.8 g</td>
<td>838 m/s (2700 ft/s)</td>
<td>6</td>
<td>44 mm (1.73 in)</td>
<td>0</td>
<td>6</td>
<td>12</td>
<td>12</td>
</tr>
<tr>
<td>IV</td>
<td>1</td>
<td>.30 caliber M2 AP 10.8 g</td>
<td>10.8 g</td>
<td>569 m/s (1880 ft/s)</td>
<td>1</td>
<td>44 mm (1.73 in)</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>2</td>
</tr>
</tbody>
</table>

Panel = Front or back component of typical armor sample.
Sample = Full armor garment, including all component panels (F & B).
Threat = Test ammunition round by caliber.
Table 2.2. Material properties and their roles in ballistic performance [3,10]

<table>
<thead>
<tr>
<th>Property</th>
<th>Role / Effect in ballistic performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microstructure</td>
<td>Affects all properties listed in the left-hand column below</td>
</tr>
<tr>
<td>Grain Size</td>
<td></td>
</tr>
<tr>
<td>Minor phases</td>
<td></td>
</tr>
<tr>
<td>Phase transformation or Amorphization</td>
<td></td>
</tr>
<tr>
<td>Porosity</td>
<td></td>
</tr>
<tr>
<td>Density</td>
<td>Weight of the armor system</td>
</tr>
<tr>
<td>Hardness</td>
<td>Damage to the projectile</td>
</tr>
<tr>
<td>Strength</td>
<td>Multi hit resistance</td>
</tr>
</tbody>
</table>

In ballistic attacks, projectile should be stopped without harming threat. The time which projectile is kept at the ceramic surface prior to penetration is described as dwell. A cone that spreads out below the projectile is formed at damaged material. This causes spreading of load of the impact over larger areas of the backing material, thus stress reduces. Backing material should exhibit high flexural strength to not allow penetration so it stops the projectile. Backface deformation should be limited to not to harm threat. Hence, the ratio of dwell, erosion and backing deformation is determined by the armor system and the threat [30].

Functionally Gradient Material (FGM) can be considered as an effective armor material because it serves different actions at different layers. A combination of the hard outer layer consists of mostly ceramic phases and the tough inner layer consists of mostly metal phases provides effective protection against ballistic threats. Projectile – armor interaction in FGM is illustrated in Figure 2.4. The outer hard layer of FGM shatters the projectile whereas the inner tough layer of FGM stops the projectile and shrapnels via absorbing its kinetic energy.
To provide maximum protection against high level ballistic threats, a new type of material consisting of different layers that serve different actions must be developed. The use of high volume fraction ceramic materials for front plate and high volume fraction metal materials for backing plate in the form of FGM has been considered to replace with non-FGM in armor technologies [11].

Increasing the thickness of steel and Al armor materials to provide maximum protection against high ballistic threats is not acceptable because it increases the weight of the armor system. Thus, ceramic based armors that have low density and extreme ballistic performance may be considered as a good solution [10]. The most common ceramics used for armor applications are aluminum oxide (alumina), silicon carbide and boron carbide. Alumina is the most economical choice, but it has higher density and less ballistic efficiency than boron carbide and silicon carbide. Boron carbide is the hardest ceramic and has very low density among them, but it is not effective alone against armor piercing projectiles with high impact velocity because of the weakening of ceramic by the amorphization process and it is very expensive. Boron carbide is a good option for one-hit protection. When multi-hit protection is needed, boron carbide can be used with silicon carbide which has close hardness that of B₄C and doesn’t undergo amorphization process. The density of silicon carbide is
a bit higher than that of boron carbide, but the hardness of silicon carbide is very close to that of boron carbide and the price of silicon carbide is lower than that of boron carbide. Also, the thermal expansion coefficient value of silicon carbide is very close to that of boron carbide that minimizes residual stress formed because of different thermal expansion of different materials. Thus, it can be considered that silicon carbide is a very effective material for armor technologies. Pressure transmitted to the backside of the ceramic causes high tensile stresses that result in cracks propagating radially outwards on the backside of the ceramic tile. Al$_2$O$_3$ ceramic is broken up into larger fragments which provide blocking of the penetrator. Hence it gives higher penetration and abrasion resistance to composite. The choice between these types of ceramic materials depends on which threat level, weight restrictions and price are dominant. Figure 2.5 and Figure 2.6 show some of the important parameters for armor materials [3].

![Figure 2.5. Image of CES Edupack program that shows density and hardness of ceramic materials](image-url)

Figure 2.5. Image of CES Edupack program that shows density and hardness of ceramic materials
Boron carbide is the hardest third material that can exhibit 30 GPa followed by diamond and cubic-BN. At temperatures higher than 1200 °C, the hardness of B₄C may be higher than the hardness of diamond [12]. Since B₄C has high hardness and low density, it is best as armor material. Nevertheless, industrial use of B₄C is restricted due to its high cost, sintering difficulties, low toughness and machinability [13]. Also, B₄C is not well suited against high speed armor piercing projectiles because of the amorphization process in B₄C when it is exposed to high pressure in which protection capability of B₄C decreases since shear strength of B₄C decreases. Also, because B₄C is very brittle, it does not provide efficient protection against shot more than one [3].

The use of silicon carbide in the armor system is very popular since price/performance ratio is getting better [14]. The availability of SiC is easier than that of B₄C since the use of SiC is higher than the use of B₄C in other sectors. SiC is cheaper than B₄C and it behaves like B₄C in armor system. Thus, the use of B₄C is tried to keep low for this system by scientists work for this sector [15]. Even monolithic SiC can be used in the armor system, it can also be mixed with B₄C. The density of SiC is 3.21 g/cm³ which is a bit higher than the density of B₄C and lower than the density of Al₂O₃. Also, the
hardness of SiC is so close to the hardness of B₄C. Hence, even SiC has higher density than B₄C, SiC can be considered as a good choice against high level ballistic threats such as NIJ IV [3]. Figure 2.7 illustrates the phase diagram of the B₄C-SiC system. The presence of silicon carbide decreases the melting temperature of boron carbide to below 2250 °C. This result is supported by the observed eutectic reaction between B₄C and SiC. Researchers have found different eutectic temperatures with different SiC addition.

![Phase Diagram of B₄C-SiC System](image)

**Figure 2.7.** The phase diagram of the B₄C–SiC system [17]

Solid state sintering of SiC can be achieved via sintering aids such as B₄C as sources of B and C elements and glucose as source of C elements. B elements enable diffusion in SiC by increasing the energy of the structure. It causes to formation of stacking faults, so stress field to balance these. Also, B elements cause more grain growth during sintering. C elements enable direct contact between SiC particles by removing SiO₂ coating on SiC particles. Also, C elements cause lowering the temperature of the formation of Si-B-C liquid [30].
However ceramic armors are mostly used nowadays, the use of ceramic armors is restricted because of their poor fracture toughness and high cost. They do not exhibit enough protection against shots more than one in a short time. After first shot strikes to armor, armor is fractured and it allows to the penetration of following projectiles [18]. Armor system which includes a combination of ceramic and metal can exhibit better protection. Thus, the formation of the ceramic preform with porosities and infiltration of this ceramic preform with metal can solve this problem. Aluminum and its alloys are used for infiltration by many researchers due to their low density, low melting point and high ductility. They are also cheaper and available. Melted aluminum and its alloys can be easily infiltrated, thus materials with almost 0 % porosity can be developed. Produced ceramic-metal composites display both high hardness and high flexural strength with low density [19].

Table 2.3 shows some properties which are important for the armor system of B₄C, SiC, Al₂O₃ and Al.

<table>
<thead>
<tr>
<th>Materials</th>
<th>Density (g/cm³)</th>
<th>Melting Point (°C)</th>
<th>Hardness (Knoop)</th>
<th>Fracture Toughness (MPa.m½)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B₄C</td>
<td>2.52</td>
<td>2445</td>
<td>2750</td>
<td>2.9 - 3.7</td>
</tr>
<tr>
<td>SiC</td>
<td>3.21</td>
<td>2730</td>
<td>2480</td>
<td>4.3</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>4.00</td>
<td>2070</td>
<td>2100</td>
<td>3.3 – 5.0</td>
</tr>
<tr>
<td>Al</td>
<td>2.70</td>
<td>660</td>
<td>120</td>
<td>29</td>
</tr>
</tbody>
</table>

Table 2.3. Some properties of B₄C, SiC, Al₂O₃ and Al [16]
2.5. MELT INFILTRATION

Production of highly dense boron carbide – silicon carbide - alumina composites is so difficult because of hard covalent bonds, high endurance to grain boundary sliding, lack of plastic behavior which restrict their diffusion in sintering. In addition to the high price of powders, both high temperature and high pressure applied in sintering make this process very expensive [19]. In contrast to traditional sintering processes, the melt infiltration process is widely used to develop composite from ceramic preforms with porosities. Some advantages of the melt infiltration process are as follows. Near or near-net shape composites which include high vol. % ceramic can be produced. Also, the microstructure of composites may show a uniform distribution of ceramic phases [20]. The formation of residual stress because of different thermal expansion coefficient of different material may be prohibited, hence residual porosities may be eliminated. With the use of suitable materials, physical and mechanical properties of the composite may be adjusted [21]. By this process, the reaction rate between constituent materials and reaction products may be managed.

Wetting conditions between solid ceramic and liquid metal is very important for the melt infiltration process. Wettability is the capability of a liquid to cover the solid surface when liquid and solid get in touch [15]. The reduction of free energy of the system is the driving force for wetting. When there is no wetting, ceramic and metal do not react with each other. Thus, the wetting of the solid ceramic by liquid metal should be provided to achieve the infiltration process [22]. To do that, the suitable temperature and atmospheric situation should be selected. By this way, liquid metal is sucked into porosities in ceramic preform by capillarity thermodynamic criteria [21]. Figure 2.8 illustrates the wetting and non-wetting conditions between solid ceramic and reactive liquid metal. When the contact angle is lower than 90° wetting condition is obtained, whereas when the contact angle is higher than 90° non-wetting condition is obtained.
Wettability of ceramics with liquid Al metals is generally poor due to the existence of contamination, humidity, oxide and gas layer which covers the solid ceramic powders surface and aluminum oxide layer that covers the liquid Al. Because of these obstacles, the melted metal may not get in touch with the solid ceramic powders surface. In order to enhance the wetting of solid ceramic powders by liquid metal, researchers have found some procedures: (i) increasing temperature of liquid metal, (ii) addition of alloying elements like Ca, Mg, Ti, Ca, Zr, Li, (iii) oxidizing or coating of ceramic particles, (iv) cleaning the particles. By this way surface energy of the ceramic particles increases and surface tension of the liquid metal decreases at the ceramic particle – liquid metal interface [15].

Melt infiltration process is very economical because exposure of the material to high temperature and high pressure is eliminated. If wettability is ensured, even exposure of the material to pressure is not necessary. Melt infiltration may be achieved via two ways: pressureless melt infiltration and pressure melt infiltration.
2.5.1. Pressureless Melt Infiltration

In pressureless melt infiltration, a solid metal block is placed on top of a ceramic preform and compacted in a furnace in an inert atmosphere. Figure 9 shows the pressureless melt infiltration process.

![Diagram of pressureless melt infiltration process]

**Figure 2.9. Illustration of the pressureless melt infiltration [29]**

In this thesis, B₄C-SiC ceramic preform was infiltrated with Al alloy.

B₄C has more reactivity with Al in the excess of 500 °C. At temperature range between 900 – 1125 °C, the main reaction products are Al₃BC and AlB₂. At higher temperatures, Al₃BC may be replaced by Al₄C₃ which is a detrimental reaction product. Thus, the reaction rate between B₄C and Al should be controlled. Otherwise, the metallic phase may be consumed before complete infiltration that results in considerable porosity in the composite and the reaction layer may form which may inhibit further infiltration of liquid metal [24]. This situation may cause the failure of the infiltration process due to the earlier formation of reaction products which may lock microstructure. Pyzik et al have worked to decrease the reactivity of B₄C with Al by passivating B₄C particles prior to infiltration process. Passivation provides controlling of the kinetics of molten metal infiltration and rate of formation of reaction products. Reaction products like AlB₂, Al₄BC, AlB₁₂C₂, AlB₁₂ and Al₄C₃ affect
negatively some mechanical properties of the composite. For example, $\text{Al}_4\text{BC}$ is a detrimental phase because it is more brittle than $\text{B}_4\text{C}$ and Al. Lower reactivity provided by the passivation process decreases the formation of undesirable ceramic phases. There are two different forms of surface boron ($\text{B}_3$ and $\text{B}_3'$) in ceramic preforms that are passivated in the temperature range of 1250 – 1400 °C. $\text{B}_3'$ is more reactive than $\text{B}_3$. $\text{B}_3'$ content is zero or near zero when the passivation process is applied at temperatures higher than 1400 °C. All the surface borons are in the form of $\text{B}_3$. Infiltration of preform takes place much faster at lower temperatures in the presence of the passivation process prior to sintering. For example, passivation at 1400 °C with 2 hours provides infiltration at lower than 1000 °C [25].

Arslan and Kalemtaş have worked to develop $\text{B}_4\text{C}$ - $\text{SiC}$ - Al composites which contain low $\text{B}_4\text{C}$ contents by using cost effective pressureless melt infiltration method for impact applications. $\text{B}_4\text{C}$ addition improves the wettability of $\text{SiC}$ by Al. However, its high cost limits its industrial applications. Thus, $\text{B}_4\text{C}$ content should be optimized to the minimum value as much as possible. Table 4 shows the required temperatures and times for different $\text{B}_4\text{C}$ and $\text{SiC}$ compositions for this process. When there is no $\text{B}_4\text{C}$ in the composite, even at temperatures in the excess of 1400 °C with 60 minutes, only partial infiltration is achieved. When there is 5 wt. % $\text{B}_4\text{C}$ in the composite, required temperature for full infiltration drops to 1130 °C. Nevertheless, at temperatures in the excess of 1050 °C, detrimental reaction product $\text{AlC}_3$ is formed in significant amounts. When there is 10 wt. % $\text{B}_4\text{C}$ in the composite, the required temperature for full infiltration decreases towards 985 °C and required time decreases to 30 minutes. Also, the amount of porosity is so low for composites that contain minimum 10 wt. % $\text{B}_4\text{C}$. 
Table 2.4. Limit conditions of infiltration [15]

<table>
<thead>
<tr>
<th>Composition</th>
<th>SiC (wt.%)</th>
<th>B₄C (wt.%)</th>
<th>Temperature (°C)</th>
<th>Time (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>50S50B</td>
<td>50</td>
<td>50</td>
<td>985</td>
<td>60</td>
</tr>
<tr>
<td>60S40B</td>
<td>60</td>
<td>40</td>
<td>1035</td>
<td>10</td>
</tr>
<tr>
<td>70S30B</td>
<td>70</td>
<td>30</td>
<td>1035</td>
<td>10</td>
</tr>
<tr>
<td>80S20B</td>
<td>80</td>
<td>20</td>
<td>935</td>
<td>60</td>
</tr>
<tr>
<td>90S10B</td>
<td>90</td>
<td>10</td>
<td>985</td>
<td>30</td>
</tr>
<tr>
<td>95S5B</td>
<td>95</td>
<td>5</td>
<td>1130</td>
<td>60</td>
</tr>
<tr>
<td>100S⁴</td>
<td>100</td>
<td>0</td>
<td>1420⁴</td>
<td>60⁴</td>
</tr>
</tbody>
</table>

* Partial infiltration (~2.4 mm).

Figure 2.10 shows green density variation with respect to B₄C content. The lowest green density is obtained with 0 wt. % B₄C, the highest green density is obtained with 30 wt. % B₄C. Green density decreases gradually from 30 wt. % to 10 wt. % B₄C. But, further addition of B₄C causes decreasing of green density. Hence, FGM design with 3 different layers containing 30 wt. %, 20 wt. % and 10 wt. % of B₄C can be considered to provide different amounts of molten metal due to gradual changing of green density and by this way, gradual changing mechanical properties can be obtained.
Figure 2.11 shows the infiltration mechanism of B₄C - SiC - Al composite. During the infiltration process, Al₃BC formation takes place due to interaction between B₄C and Al. Coarse SiC particles are confined with Al₃BC particles and suppress a significant amount of Al₄C₃ formation. It is reported that 10 wt. % fine B₄C is enough to cover the surfaces of coarse SiC particles with Al₃BC.

Zahedi et al. have worked to produce SiC – Al composites based on the SiC porous preforms by pressureless melt infiltration. Due to the non-wetting system of SiC – Al, pressure melt infiltration may be applied to assist the infiltration process. If the wetting condition is enhanced by adding preferably more than 3 wt. % magnesium to the aluminum melt and changing the internal atmosphere of the furnace to 100% nitrogen, pressureless melt infiltration may also be applied. Magnesium is an effective surfactant that clean oxygen from the melt surface. The formation of MgAl₂O₄ provides the consumption of oxygen present in the atmosphere and decreases the thickness of the oxide layer. Hence, magnesium affects positively infiltration process by enhancing wetting behavior. Also, the used nitrogen atmosphere avoids the formation of the oxide layer and enhances wetting behavior [23].
2.5.2. Pressure Melt Infiltration

In pressure melt infiltration, firstly metal block is melted and sintered ceramic preform is preheated. The preform is placed inside the preheated die. Molten metal is poured and infiltrated under pressure. Then ejection and cooling are applied. Figure 2.12 shows the pressure melt infiltration process.

![Pressure Melt Infiltration Process Diagram](image)

**Figure 2.12. Illustration of the pressure melt infiltration [26]**

SiC – Al composites are very difficult to produce by pressureless melt infiltration because of poor wetting system. It results in the presence of residual porosity because of poor wetting of silicon carbide by melted aluminum and the formation of undesirable reaction products Al₄C₃, Al₅SiC₄ due to the dissolution of the SiC by molten Al which decreases mechanical properties of the composite. Because these phases are developed at high temperatures (>660°C) and atmospheric pressure, SiC
becomes thermodynamically unstable and interfacial reactions take place. Also, these phases hydrolyze with the atmospheric humidity and resulted in the formation of Al(OH)₃ that causes crack propagation in composite via corrosion mechanism. B₄C addition provides enhancing wettability of SiC by Al [15]. Thus, B₄C – SiC – Al composites can be produced by pressureless melt infiltration process in which application of external pressure for filling of voids of ceramic preforms by liquid metal is not necessary. To produce SiC – Al composites, an external pressure should be applied to exceed threshold pressure which is the pressure required to handle capillary back pressure by pressure melt infiltration process [23].

Chedru et al. have worked to produce AlN-Al composites based on the AlN porous preforms by pressure melt infiltration. The whole infiltration process was ended in just 120 seconds. As a result of this study, the composite was considerably dense and microstructure showed homogeneous distribution of the aluminum and small amount of porosity. During this study different kind of aluminum alloys and pure aluminum were used as the infiltrated metal. Only pure aluminum failed that partial infiltration occurred. This study shows also that some infiltrating agents are required to achieve full infiltration [26].

H. S. Lee and S. H. Hong have worked to produce SiC – Al composites that contain high volume fraction SiC particles up to 70 vol. % by pressure melt infiltration. Two different kinds of SiC particles which particle sizes of 8 µ and 48 µ, 0 - 4 wt. % colloidal silica as inorganic binder and 1 wt. % cationic starch as organic binder were ball milled and then cold pressed at 1 – 2 MPa to have porous ceramic preforms. Preforms heated at 1100 °C for 4 hours. Al was melted and the temperature of the Al melt was kept between 700 °C and 800 °C. The preform was reheated to 450-550 °C. Molten Al was infiltrated by applying 10 -50 MPa. Importance of some parameters such as the temperature of molten Al, the temperature of preform before infiltration, infiltration pressure and infiltration time was emphasized. These parameters should
be controlled to avoid early solidification since it may cause the formation of pores and blockage of microstructure that means partial infiltration. The volume fraction of SiC could be controlled by controlling the particle sizes of SiC particles and their mixing ratio. The colloidal silica provided higher bond strength between contacting SiC particles since it transforms the cristobalite phase. As a result, H. S. Lee and S. H. Hong concluded that full infiltration may be achieved via process parameters such as the temperature of molten al 800 °C, the temperature of preform before infiltration 550 °C, infiltration time 20 -70 seconds, infiltration pressure 30-50 MPa [27].
CHAPTER 3

AIM OF THE STUDY

3.1. MOTIVATION OF THE STUDY

In this thesis, the development of ceramic-metal FGM for defense applications was aimed. In defense applications, development of material with low density, high hardness at the impact point of the projectile and high flexural strength at the backing point are very important. B₄C, SiC and Al which have low density were used to obtain lightweight armor material. FGM process that provides different mechanical properties at different sections was used to obtain both high hardness and high flexural strength in the same body. Figure 3.1 illustrates the expected structure and mechanical properties. The black part of the sample includes the highest number of ceramic particles, so hardness is the highest at that part. Gray part of the sample includes the highest number of metal particles, so flexural strength is the highest at that part. These mechanical properties increase or decrease gradually with gradually changing compositions (ceramic -metal ratio).

![Figure 3.1. Expected structure and mechanical properties](image-url)
Pressure melt infiltration process was applied to produce near or near-net shape ceramic-metal FGM. High ceramic fraction composites were produced at lower temperature, 1400 °C without Al4C3 phase formation which is detrimental. Since the cost of B4C is higher than the cost of SiC, B4C-SiC-Al composites which contain low B4C wt. % were produced. Also, it was aimed that composite exhibits low density and high hardness despite decreasing wt. % of B4C.

3.2. MAIN CONTRIBUTION TO THE LITERATURE

FGM can be considered as a potential candidate for armor applications since it exhibits different mechanical properties at different parts of the material. Thus, a combination of two contrasting properties which are high hardness and high flexural strength in the same body can be obtained. With this thesis, a new kind of material that can be used for defense applications was produced.

In conventional ballistic plates, ceramic plates are joined to metal or polymer layers. Because differences between thermal expansion coefficients of ceramics and metals/polymers are high, a combination of them without gradient may result in residual stress, unjoined edges, unjoined pores, reaction layer with cracks which are detrimental for armor applications. By the FGM process, difficulties of the joining of dissimilar materials were overcome by gradually changing composition so mechanical properties.

B4C does not provide effective protection against armor piercing projectiles because of the amorphization process. It is a good candidate for one-hit protection. When multi-hit protection is required SiC whose hardness is close to that of B4C can be used. Also, adding metal to composite improves the flexural strength of the plate, so increases penetration resistance.
In this chapter, the production of B₄C – SiC – Al composites by pressure melt infiltration of ceramic preforms with aluminum alloy are explained. Also, applied characterization methods are indicated. Figure 4.1 shows the experimental flow chart.

**Figure 4.1.** Experimental flow chart
4.1. PRODUCTION OF FGM LAYERS

In this thesis, firstly each potential layer of FGM was produced separately to investigate the effects of each layer to the composite in terms of green/bulk density, porosity, microstructure and some mechanical properties. In the light of the literature review, at least 10 wt. % of B₄C is required as sintering aids. 10 wt. % of Al₂O₃ was used at all layers to provide higher penetration and abrasion resistance. The composition is the most important variable to obtain gradual changing green/bulk density and/or porosity which are the requirements of an FGM sample. So, it was planned to prepare compositions to create each layer of FGM which is indicated in Table 4.1.

<table>
<thead>
<tr>
<th>Layers</th>
<th>Composition</th>
<th>B₄C (weight %)</th>
<th>SiC (weight %)</th>
<th>Al₂O₃ (weight %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>80S10B10A</td>
<td>10</td>
<td>80</td>
<td>10</td>
</tr>
<tr>
<td>2</td>
<td>70S20B10A</td>
<td>20</td>
<td>70</td>
<td>10</td>
</tr>
<tr>
<td>3</td>
<td>60S30B10A</td>
<td>30</td>
<td>60</td>
<td>10</td>
</tr>
</tbody>
</table>

B₄C powders which have 1 µm (d50) particle size, SiC powders which have 35 µm (d50) particle size and Al₂O₃ powders which have 6 µm (d50) particle size were used. To obtain a green body, all the ceramic particles, 15 wt. % PVA or PEG as binders and 1 wt. % Cu-Zn stearate as lubricant were mixed by ball milling in alcohol media. Three different compositions in Table 4.1. which indicated wt. % of B₄C, SiC and Al₂O₃ ceramic particles were produced separately. PVA or PEG solution was prepared by mixing 50 ml of boiled distilled water with 5 grams of PVA or PEG powder until completely dissolved. The amount of zirconia ball and the amount of alcohol was equal to the amount of total ceramic powders. Ball milling operation was applied for 90 minutes. Figure 4.2 illustrates ball milling machine and jar filled with ingredients.
When ball milling operation finished, the milled solution was poured into a beaker and then beaker was placed in a furnace and operated to remove alcohol and water in milled solution. Thus, the preparation of the compositions section was completed.

![Ball milling machine and jar filled with ingredients](image)

**Figure 4.2.** Ball milling machine and jar filled with ingredients

Ceramic preforms were produced by pressing operation which involves the compaction of powder into a rigid die by applying pressure in a single axial direction through a rigid punch or piston. Before pressing operation, 15 ml glucose - distilled water solution which has 50 wt. % glucose was added to the composition to provide moisture and binding. The manual press was applied under 70 tons pressure and it means that green ceramic bodies were produced under 100 MPa pressure according to dimensions of the mold. Width and length of the sample depend on dimensions of the mold which is cylindrical with 9 mm diameter; whereas the height of the sample depends on the weight of composition pressed. Pressed samples were kept at room temperature at least one day to release their humidity before sintering operation. Thus, the production of green ceramic bodies section was completed. Figure 4.3 illustrates pressing operation and green ceramic body.
Due to compacts with high amounts of binder decreases mechanical properties of the ceramic preform and may affect the sintering process negatively, the binder should be removed prior to sintering. Debinding process at 550 °C for 2 hours followed by the sintering process at 1400 °C for 2 hours were done in a nitrogen atmosphere by using Protherm furnace. The heating rate was applied 3 °C / min until 550 °C, 5 °C / min until 900 °C and 10 °C / min until sintering temperature which is 1400 °C. Thus, the sintering of green ceramic bodies section was completed. Figure 4.4 shows furnace and temperature vs time graph of the sintering process.
Due to the problem encountered at the sintering process because of $\text{Al}_2\text{O}_3$ ceramic powder, it was planned to prepare new compositions to create each layer of FGM which is indicated in Table 4.2. All the experimental procedure was followed as before except for rectangular mold which has 120 mm X 40 mm dimensions used for pressing and melt infiltration process with an applied pressure of 200 MPa. More pressure could be achieved by double action press. 100 tons of pressure could be applied and it means that ceramic green bodies were produced under 200 MPa pressure.

<table>
<thead>
<tr>
<th>Layers</th>
<th>Composition</th>
<th>$\text{B}_4\text{C}$ (weight %)</th>
<th>SiC (weight %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>90S10B</td>
<td>10</td>
<td>90</td>
</tr>
<tr>
<td>2</td>
<td>80S20B</td>
<td>20</td>
<td>80</td>
</tr>
<tr>
<td>3</td>
<td>70S30B</td>
<td>30</td>
<td>70</td>
</tr>
</tbody>
</table>

The porosity of the sintered ceramic bodies should be filled with metal to reach near theoretical density. It was achieved by the pressure melt infiltration process which is illustrated in Figure 4.5. After Al alloy was melted, the sintered ceramic preform was
preheated to 900 °C. Preform placed inside the preheated die on alumina thermal paper to decrease temperature differences between metal mold and liquid metal. Molten aluminum alloy was poured at 700 °C and infiltrated under 200 MPa pressure. Then ejection and cooling were applied.

![Pressure melt infiltration process diagram](image)

**Figure 4.5.** Pressure melt infiltration process

7075 Al alloy is widely used as infiltrated metal in literature due to its low melting point, low density and high ductility. The new composition was developed by adjusting 7075 Al alloy with 11.2 wt. % Si and 6.9 wt. % Mg to increase the infiltration rate in the light of the literature review as illustrated in Figure 4.6. Silicon addition increases Al alloy’s fluidity and decreases its melting temperature. Silicon prevents interfacial reactions between SiC and Al. While magnesium reduces the surface tension and the wetting angle between Al melt and B₄C – SiC ceramics. Magnesium vaporizes from Al melt as a result of high gas pressure and it may destroy the Al₂O₃ layer on the surface of Al alloy and helps direct contact between the Al melt and B₄C – SiC ceramics.
Thus, Al melt infiltration into the sintered ceramic section was completed. Figure 4.7 shows preheated die and 90S10B composite after aluminum pressure melt infiltration process.
4.2. PRODUCTION OF FGM

After investigating the effects of each layer to the composite in terms of green/bulk density, porosity, microstructure and some mechanical properties, FGM with different compositions which are indicated in Table 4.2 was produced. All the experimental procedure was followed as before. Green ceramic bodies were produced by respectively addition and pressing of each composition as layers of FGM. Before pressing operation, 5 ml glucose - distilled water solution which has 50 wt. % glucose was added to each composition to provide moisture and binding.

4.3. CHARACTERIZATION OF COMPOSITES

The composites were characterized by phase analysis, particle size and distribution analysis, green and bulk density/porosity measurement, optic emission spectrometer analysis, XRD analysis, optical microscopy and SEM-EDX analyses, hardness measurement and three point bend test.

For sample preparation, composites were cut with silicon carbide cutting disc into required dimensions. Mounting was done by bakelite powder. After that, samples were ground by abrasive papers with different grids and polished by broadcloths with diamond solutions with 6 and 1 μ particle size respectively.

Particle size and distribution analysis is important in terms of creating compositions which result in different green/bulk density and so porosity. It was done to each composition of each layer after ball milling operations with the Malvern Mastersizer 2000 device. This device also provides specific surface areas of each composition.
Green density measurement was done to evaluate how tightly packed the powder particles. Green densities of each layer were calculated, so whether gradual changing green density for FGM was obtained or not could be understood.

For green ceramic bodies produced by manual press, the calculation is as indicated in Equation 4.1 according to cylindrical mold where \( r = 4.5 \text{ cm} \), \( d_{B4C} = 2.52 \text{ g/cm}^3 \), \( d_{SiC} = 3.21 \text{ g/cm}^3 \):

\[
V_{\text{obtained}} = \pi \cdot r^2 \cdot h
\]

\[
V_{\text{theoretical}} = \left( \frac{m_{B4C}}{d_{B4C}} \right) + \left( \frac{m_{SiC}}{d_{SiC}} \right)
\]

\[
\% d_{\text{green}} = \left( \frac{V_{\text{theoretical}}}{V_{\text{obtained}}} \right) \cdot 100
\]

**Equation 4.1.** Green density calculation for manual pressed green ceramic bodies

For green ceramic bodies produced by double action press, the calculation is as indicated in Equation 4.2 according to rectangular mold where \( a = 12 \text{ cm} \), \( b = 4 \text{ cm} \), \( d_{B4C} = 2.52 \text{ g/cm}^3 \), \( d_{SiC} = 3.21 \text{ g/cm}^3 \):

\[
V_{\text{obtained}} = a \cdot b \cdot c
\]

\[
V_{\text{theoretical}} = \left( \frac{m_{B4C}}{d_{B4C}} \right) + \left( \frac{m_{SiC}}{d_{SiC}} \right)
\]

\[
\% d_{\text{green}} = \left( \frac{V_{\text{theoretical}}}{V_{\text{obtained}}} \right) \cdot 100
\]

**Equation 4.2.** Green density calculation for double action pressed green ceramic bodies
Bulk density and porosity measurements were done by Archimedes principle to evaluate the required amount of Al alloy for each composition to reach near theoretical density and amount of porosity after the infiltration process. The theoretical density of each layer was calculated according to the rule of mixture to use them in porosity (%) formula given in Figure 4.11, so whether gradual changing porosity percentage for FGM was obtained or not could be understood.

Archimedes principle was applied with xylene solution to determine the bulk density and porosity % of sintered layers of FGM. Firstly, samples were weighed as dry which is termed as \( m_{\text{dry}} \). Then, after samples were kept in xylene solution for 24 hours, samples were weighed in xylene solution which is termed as \( m_{\text{submerged}} \). Finally, samples were extracted from the solution, the surfaces of the samples were wiped out and the samples were weighed in the air which is termed as \( m_{\text{wet}} \). Density of xylene solution is 0.861 g/cm\(^3\). Calculations of bulk density (\( d_{\text{bulk}} \)) and porosity (P) % are illustrated in Equation 4.3.

\[
V = \frac{m_{\text{wet}} - m_{\text{submerged}}}{0.861}
\]
\[
d_{\text{bulk}} = \frac{m_{\text{dry}}}{V}
\]
\[
P (\%) = \frac{\left(\text{theoretical density} - \text{bulk density}\right)}{\text{theoretical density}} \times 100
\]

**Equation 4.3.** Calculation of bulk density and porosity % by using xylene solution

Archimedes principle was applied with distilled water to determine the bulk density and porosity % of Al alloy infiltrated FGM. Firstly, samples were weighed as dry which is termed as \( m_{\text{dry}} \). Then, after samples were boiled in distilled water for 1 hour and cooled to room temperature, samples were weighed in distilled water which is termed as \( m_{\text{submerged}} \). Finally, samples were extracted from distilled water, the surfaces
of the samples were wiped out and the samples were weighed in the air which is termed as $m_{\text{wet}}$. Calculations of bulk density ($d_{\text{bulk}}$) and porosity (P) % are illustrated in Equation 4.4.

$$d_{\text{bulk}} = \frac{m_{\text{dry}}}{(m_{\text{wet}} - m_{\text{submerged}})}$$

$$P \% = \frac{[(m_{\text{wet}} - m_{\text{dry}}) \times 100]}{(m_{\text{wet}} - m_{\text{submerged}})}$$

Equation 4.4. Calculation of bulk density and porosity % by using distilled water

The composition of Al alloy used for pressure melt infiltration process was measured with optic emission spectrometer analysis. The results are given as weight percentage. The composition analysis was carried out by WAS Foundry Master after alloy preparation in the induction furnace.

Processes used in this thesis study may yield a mixture of several ceramic and metal phases that differ from the starting materials. XRD measures reaction products (phases) formed. With this technique formation of Al$_4$C$_3$ which is a detrimental phase and other phases can be investigated. Reference pdf cards were used to detect which peaks belong to which phases. XRD analysis was applied by using the Bruker D8 Advance device after breaking up composites into small particles. Samples were scanned with 1°/min speed between 20° and 80°.

Optical microscopy was used to examine microstructure and to do phase analysis. During phase analysis, different colors created for different phases thanks to contrast differences. SEM analysis was done for microstructural examination in high magnifications and resolutions. Microscope analysis provides to see the gradual
changing of microstructure between each layer of FGM. Quantitative phase analysis was conducted using EDS.

Three point bend tests and hardness measurements were applied for each layer separately to determine the effect of each layer to the flexural strength and hardness. Hardness and flexural strength of each layer were detected, so whether gradual changing hardness and flexural strength for FGM was obtained or not could be understood. Hardness values were calculated in the HRC scale by using 150 kg load. Three point bend tests were done to evaluate the bending and fracture behavior of the Al alloy and composites by using Instron 5582 tensile test machine. Calculations of flexural strength are given in Equation 4.5. The letters M, c, I, F, L, w, h represent bending moment, area moment of inertia, distance from neutral plane, load at span center, distance between supports (80 mm), width (40 cm) and height of the specimen respectively. Specimen subjected to three point bend test and stress/strain variations are illustrated in Figure 4.8. The material remains unstressed at the transition from tensile to compressive stress. This transition plane is called as neutral plane. Neutral plane for ceramic metal composite is the interlayer between ceramic and metal. h_b is measured from the neutral plane to the bottom of the specimen (metal part) and h_t is measured from the neutral plane to the top of the specimen (ceramic part). When h_b = h_t, c=h/2.

\[
\sigma = \frac{(M \cdot c)}{I} \text{ where; } M = \frac{(F \cdot L)}{4}, I = \frac{(w \cdot h^3)}{12} \text{ and } c = \text{distance from neutral plane} \\
\sigma = 1.5 \cdot \frac{(F \cdot L)}{(w \cdot h^2)}
\]

**Equation 4.5.** (a) Calculation of flexural strength, (b) Calculation of flexural strength when c = h/2
Figure 4.8. (a) Specimen geometry, (b) Normal stress variation, (c) Shear stress variation, (d) Normal strain variation [31]
CHAPTER 5

RESULTS AND DISCUSSION

5.1. PHASE ANALYSES

Phase analyses were done by Thermo-Calc software. B$_4$C-SiC composites at 1400 °C temperature were investigated by drawing ternary phase diagrams by selecting B, C and Si elements. Figure 5.1 shows that there are B$_4$C and SiC phases in solid state at each composition which are 90S10B, 80S20B and 70S30B.

The mass percent of elements were calculated for each composition. For example, Figure 5.2 shows the calculation of mass percent of elements in 90S10B composition which includes 10 wt. % B$_4$C and 90 wt. % SiC. According to that calculation, 90S10B composition includes 7.8 weight % B, 63.1 weight % Si and 29.1 weight % C elements. The interception point of two elements (C and Si) in this figure represents the phase region.
Figure 5.1. Ternary phase diagram of B, C and Si elements at 1400 °C

\[ \text{B}_4\text{C} \rightarrow 4 \times M_B + M_C = 43.2 + 12 = 55.2 \]

\[ \text{SiC} \rightarrow M\text{Si} + M_C = 28.1 + 12 = 40.1 \]

Mass percent B = \( \frac{10 \times 43.2}{55.2} = 7.8 \)

Mass percent Si = \( \frac{90 \times 28.1}{40.1} = 63.1 \)

Mass percent C = 100 – (7.8 + 63.1) = 29.1

Figure 5.2. Calculation of mass percent of elements in 90S10B composition
Al alloy at 700 °C temperature was investigated by drawing ternary phase diagrams by selecting Al, Mg and Si elements. The Al alloy includes 74.2 wt. % Al, 11.2 wt. % Si and 6.9 wt. % Mg elements. The interception point of two elements (Al and Si) in Figure 5.3 shows the phase region which is the liquid phase. Thus, 700 °C was chosen as molten metal temperature for the infiltration process.

![Ternary phase diagram of Al, Mg and Si elements at 700 °C](image)

**Figure 5.3.** Ternary phase diagram of Al, Mg and Si elements at 700 °C

### 5.2. PARTICLE SIZE AND DISTRIBUTION ANALYSIS

Composite with high ceramic fraction is required for armor applications. To reach high green densities, high coordination number which is defined as the number of particles that a particle holds as its nearest neighbors in a coordination compound is required. In this thesis, B₄C powders which have 1 µm particle size and SiC powders
which have 35 µm particle size were ready to use. To get high coordination number Al₂O₃ powders which have 6 µm particle size were procured.

Particle size and distribution analyses are illustrated in Figure 5.4, 5.5 and 5.6 for 90S10B, 80S20B, 70S30B respectively. There are two peaks and they are called bimodal because there are both fine grain size B₄C and coarse grain size SiC. Also, particle size distribution is wide.

We can see from these figures that as B₄C content increases from 10 wt. % to 20 wt. %, the intensity of first peaks on the left side increases because B₄C has smaller particle size. 80S20B composition has the highest first peak among them. Also, we can see from these figures that as SiC content increases from 80 wt. % to 90 wt. %, the intensity of second peaks on the right side increases because SiC has larger particle size. 90S10B composition has the highest second peak among them. It was concluded that these results may be wrong because B₄C has smaller particle size than SiC, the intensity of first peak should increase and the intensity of second peak should decrease as B₄C content increases from 20 wt. % to 30 wt. These may result from agglomeration of 70S30B composition and some measuring errors of device.

The d(0.5) is the diameter of the particle that 50% of a sample's mass is smaller than and 50% of a sample's mass is larger than. The d(0.5) is also known as the "mass median diameter" as it divides the sample equally by mass. We can see from these figures that as B₄C content increases from 10 wt. % to 20 wt. %, d(0.5) value dramatically decreases from 27.23 µm to 3.30 µm, because B₄C has lower particle size. 80S20B composition has the lowest d(0.5) value among them. However, as B₄C content increases from 20 wt. % to 30 wt. %, d(0.5) value slightly increases from 3.30 µm to 5.02 µm.
Figure 5.4. Particle size and distribution analysis of 90S10B

Figure 5.5. Particle size and distribution analysis of 80S20B

Figure 5.6. Particle size and distribution analysis of 70S30B
Particle size and distribution analysis also provides specific surface area. Table 5.1 shows the specific surface area of each composition. Specific surface area increases from 1.3 m²/g to 2.78 m²/g as B₄C content increases from 10 wt. % to 20 wt. %. But the specific surface area decreases as B₄C content increases from 20 wt. % to 30 wt. % in spite of increasing wt. % of B₄C which has smaller particle size. That result was supported by d(0.5) values of these compositions. Figure 5.7 shows the variation of d(0.5) values of each composition with respect to B₄C content. Figure 5.8 shows the variation of the specific surface area of each composition with respect to B₄C content. It was concluded that these results may be wrong because B₄C has smaller particle size than SiC, d(0.5) value should decrease and specific surface area should increase as B₄C content increases from 20 wt. % to 30 wt. These may result from agglomeration of 70S30B composition and some measuring errors of particle size and distribution analysis device.

Table 5.1. Specific surface area of compositions

<table>
<thead>
<tr>
<th>Composition</th>
<th>Specific Surface Area (m²/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>90S10B</td>
<td>1.3 (± 0.02)</td>
</tr>
<tr>
<td>80S20B</td>
<td>2.78 (± 0.01)</td>
</tr>
<tr>
<td>70S30B</td>
<td>2.17 (± 0.04)</td>
</tr>
</tbody>
</table>

Figure 5.7. Variation of d(0.5) values with respect to B₄C content
Green densities of each composition were calculated as mentioned in Section 4.3. To obtain nearly accurate green density value, four ceramic green bodies of each composition were prepared and green density results averaged as shown in Table 5.2. Green density varies according to composition because of the changing weight percentage of fine B₄C particles and coarse SiC particles. Figure 5.9 shows green density variation with B₄C content. As B₄C wt. % increases from 10 to 20, green density increases dramatically while further addition up to 30 wt. % leads to a slight increment in green density. That result was supported with d(0.5) values and specific surface area of each composition. In conclusion, it was understood that green density values increase from 90S10B composition to 70S30B composition to obtain FGM. However, the increment is not gradual.
Table 5.2. Green densities of each composition

<table>
<thead>
<tr>
<th>Composition</th>
<th>Green Density (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>90S10B</td>
<td>62.78 (± 0.63)</td>
</tr>
<tr>
<td>80S20B</td>
<td>68.03 (± 0.55)</td>
</tr>
<tr>
<td>70S30B</td>
<td>68.08 (± 0.63)</td>
</tr>
</tbody>
</table>

Figure 5.9. Green density variation with B₄C content

Bulk density and porosity % of each sintered layer of FGM were calculated as mentioned in Section 4.3. To obtain nearly accurate bulk density and porosity % value, three pieces of samples were prepared. The theoretical density of each layer was calculated according to the rule of mixture, then porosity % of each layer was determined with formula by using theoretical and bulk density values as shown in Table 5.3. Figure 5.10 shows porosity % variation with B₄C content. As B₄C wt. % increases from 10 to 20 and 20 to 30, porosity % decreases gradually from 34.2 to 29.6 and 29.6 to 23.2 respectively. However gradual changing of green density from 80S20B layer to 70S30B layer could not be achieved, gradual changing of bulk density/porosity % from 80S20B layer to 70S30B layer could be achieved due to
increment in wt. % of B₄C that is densification aid. In conclusion, the gradual changing of porosity % from 90S10B layer to 70S30B layer to obtain FGM was achieved.

### Table 5.3. Porosity % values of each composition

<table>
<thead>
<tr>
<th>Composition</th>
<th>Theoretical Density (g/cm³)</th>
<th>Bulk Density (g/cm³)</th>
<th>Porosity %</th>
</tr>
</thead>
<tbody>
<tr>
<td>90S10B</td>
<td>3.13</td>
<td>2.06 (± 0.03)</td>
<td>34.2 (± 0.91)</td>
</tr>
<tr>
<td>80S20B</td>
<td>3.04</td>
<td>2.14 (± 0.02)</td>
<td>29.6 (± 1.21)</td>
</tr>
<tr>
<td>70S30B</td>
<td>2.97</td>
<td>2.28 (± 0.03)</td>
<td>23.2 (± 0.88)</td>
</tr>
</tbody>
</table>

![Figure 5.10. Porosity % variation with B₄C content](image)

In the light of these analyses, it was noted that FGM in Table 4.3 can be considered as a potential candidate for this study because as bulk density increases/porosity % decreases from 90S10B layer to 70S30B layer, amount of Al alloy to achieve full infiltration decreases from 90S10B layer to 70S30B layer.
Bulk density and porosity % of Al alloy infiltrated FGM were calculated as mentioned in Section 4.3. To obtain nearly accurate bulk density and porosity % value, three pieces of samples were prepared. Bulk density of FGM was calculated as 2.70 (± 0.02) g/cm³. FGM was successfully infiltrated with porosity % lower than 0.88 (± 0.3).

5.4. OPTIC EMISSION SPECTROMETER ANALYSIS

During this study, the new Al alloy was produced from 7075 Al alloy. In the light of the literature review, it was aimed to use Al alloy with 4-8 wt. % Mg and up to 12 wt. % Si for pressure melt infiltration process to increase the infiltration rate. 7075 Al alloy was melted and required Mg and Si elements were added to the melt. Table 5.5 shows optic emission spectrometer analysis results of 7075 Al alloy and new composition which was modified from 7075 Al alloy for this study. The chemical composition of the new composition which was modified from 7075 Al alloy could be arranged as aimed.

Table 5.4. Optic emission spectrometer analysis results of 7075 Al alloy and new composition which was modified from 7075 Al alloy for this study

<table>
<thead>
<tr>
<th>Elements</th>
<th>Weight Percentage of Elements</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>7075 Al Alloy</td>
</tr>
<tr>
<td>Al</td>
<td>89.42 (±0.32)</td>
</tr>
<tr>
<td>Mg</td>
<td>2.12 (±0.05)</td>
</tr>
<tr>
<td>Si</td>
<td>0.27 (±0.03)</td>
</tr>
</tbody>
</table>

5.5. XRD ANALYSIS

During this thesis, some problems occurred due to Al₄C₃ phase formations which are illustrated in Figure 5.11 with peaks represented by triangle shape as a result of XRD
analysis. Al₂O₃ reacts with carbide-based ceramics such as B₄C and SiC at prolonged temperatures higher than 1000 °C and Al₄C₃ phase formation occurs. Al₄C₃ is very detrimental since it hydrolyzes with atmospheric moisture and Al(OH)₃ forms according to reactions illustrated in Equation 5.1. 80S10B10A sample was sintered, then was put into water. Some bubbles were observed and then the sample crashed into pieces as seen in Figure 5.12. Thus, it was decided to not to add Al₂O₃ powder.

Figure 5.11. XRD analysis of sintered 80S10B10A

\[
\begin{align*}
3\text{SiC (s) + 4Al (l)} & \rightarrow \text{Al}_4\text{C}_3 \text{ (s) + 3Si (in Al)} \\
\text{Al}_4\text{C}_3 \text{ (s) + 12H}_2\text{O (g)} & \rightarrow \text{4Al(OH)}_3 \text{ (s) + 3CH}_4 \text{ (g)} \\
\Delta G^{\circ}_{1100} & = -293.9 \text{ kJ/mol} \\
\text{Al}_4\text{C}_3 \text{ (s) + 18H}_2\text{O (g)} & \rightarrow \text{4Al(OH)}_3 \text{ (s) + 3CO}_2 \text{ (g) + 12H}_2 \\
\Delta G^{\circ}_{1100} & = -634.6 \text{ kJ/mol}
\end{align*}
\]

Equation 5.1. Al₄C₃ and Al(OH)₃ formation reactions
XRD analyses were applied to Al alloy and Al alloy infiltrated 70S30B, 80S20B, 90S10B, FGM composites which are illustrated in Figure 5.13, 5.14, 5.15, 5.16 and 5.17 respectively. For Al alloy used in this thesis study, Mg\(_2\)Si peaks were detected in addition to 7075 Al alloys peaks because the weight percentage of Mg and Si of new composition was higher than that of 7075 Al alloy and cause to formation of Mg\(_2\)Si phase. For Al alloy infiltrated 70S30B, 80S20B, 90S10B, FGM composites, no phases other than B\(_4\)C, SiC, Al, Mg\(_2\)Si and Si were formed. In this thesis, it was aimed that B\(_4\)C and SiC ceramic particles do not react with Al alloy in order to avoid the formation of intermetallic phases which decrease hardness of composite and formation of Al\(_4\)C\(_3\) which hydrolyzes with atmospheric moisture. These figures confirm that this aim was achieved.

**Figure 5.12.** 80S10B10A sample was put into water and crashed
Figure 5.13. XRD analysis of Al alloy

Figure 5.14. XRD analysis of Al alloy infiltrated 70S30B composite
Figure 5.15. XRD analysis of Al alloy infiltrated 80S20B composite

Figure 5.16. XRD analysis of Al alloy infiltrated 90S10B composite
5.6. OPTICAL MICROSCOPY ANALYSIS

Microstructural analysis and phase analysis were done with optical microscopy. But the images taken from optical microscopy were not clear due to lower magnification and resolution.

Figure 5.18 shows the microstructure of Al alloy which was infiltrated into ceramic preforms. The image was taken with 1000X magnification. Angular shape Mg$_2$Si phases distributed in the alloy matrix can be seen in Figure 5.18. These phases are brittle and cause decrease in percent elongation of the metal matrix alloy.
Al alloy infiltrated FGM was investigated with optical microscopy to detect gradually changing of microstructure. The image was taken with 50X magnification to obtain lower magnification so investigate all layers of FGM as illustrated in Figure 5.19. White and bright particles represent Al alloy. Since the highest porosity % is obtained for 90S10B, the vol. % of infiltrated Al alloy is the highest. Since the lowest porosity % is obtained for 70S30B, the vol. % of infiltrated Al alloy is the lowest. Thus, the vol. % of Al alloy decreases from 90S10B to 70S30B gradually with decreasing volume percent porosity. Microstructure shows inhomogeneous distribution of Al alloy. This may result from low ball milling time and hand-mixing of glucose-distilled water with powder compositions.

Phase analysis in terms of volume percent of infiltrated Al alloy was done to reveal variation in Al alloy infiltrated through section of FGM’s layer as illustrated in Figure 5.20. The green areas represent Al alloy infiltrated. Phase analysis of FGM’s layers which are (a) 90S10B, (b) 80S20B, (c) 70S30B as vol. % are given in Table 5.5. Vol. % of Al alloy decreases gradually from 90S10B layer to 70S30B layer.
Figure 5.19. Microstructure of FGM

Figure 5.20. Phase analysis of FGM’s layers which are (a) 90S10B, (b) 80S20B, (c) 70S30B
Table 5.5. Phase analysis of FGM’s layers which are (a) 90S10B, (b) 80S20B, (c) 70S30B as vol. %

<table>
<thead>
<tr>
<th>FGM’s Layers</th>
<th>Black Area (vol. %)</th>
<th>Green Area (Al Alloy) (vol. %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>90S10B</td>
<td>66.6</td>
<td>33.4</td>
</tr>
<tr>
<td>80S20B</td>
<td>70.5</td>
<td>29.5</td>
</tr>
<tr>
<td>70S30B</td>
<td>77.4</td>
<td>22.6</td>
</tr>
</tbody>
</table>

Figure 5.21 shows each layer of Al alloy infiltrated FGM which are 90S10B, 80S20B and 70S30B and their vol. % of Al alloy. 5 images were taken with 100X magnification to do phase analysis. The white areas represent Al alloy infiltrated. The results were very close to the results represented in Table 5.5.

a  Vol. % of Al alloy: 33.89 (±0.46)

b  Vol. % of Al alloy: 29.92 (±0.55)

c  Vol. % of Al alloy: 22.93 (±0.28)

Figure 5.21. Phase analyses of Al alloy infiltrated (a) 90S10B, (b) 80S20B and (c) 70S30B layers
5.7. SEM-EDX ANALYSIS

Microstructural analysis was done with scanning electron microscopy (SEM) and phase analysis was done with energy dispersive X-ray spectrometry (EDX). The images taken from SEM were clear due to higher magnification and resolution. The phases detected with XRD analysis can be confirmed by microstructure and peaks detected by SEM – EDX analysis.

Firstly, microstructure and particle size of B₄C and SiC powders were examined by SEM analysis. B₄C and SiC powders were embedded in epoxy resin separately, ground and polished prior to SEM analysis. Particle size and distribution results of B₄C and SiC particles were confirmed that B₄C has 1 µm particle size and SiC has 35 µm particle size with SEM analyses which are illustrated respectively in Figure 5.22 and Figure 5.23. Aspect ratios of B₄C and SiC powders were also measured by dividing the long edge of the particle to short edge of the particle as shown in Table 5.6. The aspect ratio of B₄C and SiC are very close to 1 which indicates the cubical or spherical shape.
Figure 5.22. SEM analysis of B₄C powder

Figure 5.23. SEM analysis of SiC powder
Table 5.6. Aspect Ratio of B₄C and SiC powders

<table>
<thead>
<tr>
<th>Powders</th>
<th>Aspect Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>B₄C</td>
<td>1.38 (±0.22)</td>
</tr>
<tr>
<td>SiC</td>
<td>1.41 (±0.14)</td>
</tr>
</tbody>
</table>

SEM-EDX analyses show that each layer of FGM and FGM were successfully infiltrated with Al alloy. It can be concluded that there is no difficulty in wetting of B₄C – SiC ceramic by liquid aluminum alloy containing Mg and Si. Almost all the pores between ceramic particles were filled with Al alloy.

Bigger particles illustrated in Figure 5.24 with red circle represent SiC particles. Since the particle size of SiC is very large (35 µm) and the wt. % of SiC is more, SiC particles can be easily seen in the microstructure. This can be confirmed by EDX analysis as illustrated in Figure 5.24. EDX analysis was applied to this particle and it revealed only Si and C peaks that confirmed this particle is SiC. This image was taken from Al alloy infiltrated 80S20B composite. The particle size of SiC was also confirmed by the magnification bar at right below the image.
Figure 5.24. SEM-EDX analysis of SiC region of Al infiltrated 80S20B composite

Figure 5.25 shows the SEM image of the Al alloy region of infiltrated 80S20B composite. This can be confirmed by EDX analysis as illustrated in Figure 5.25. EDX analysis was applied to this region and it revealed only Al, Mg, Zn, Cu and Si peaks that are ingredients of infiltrated Al alloy. Figure 5.25 also shows the SEM image of the B₄C region of Al alloy infiltrated 80S20B composite. Small black particles in the Al alloy represent B₄C particles as illustrated in Figure 5.25 and it can be seen at higher magnification such as 7416X. Since the particle size of B₄C is very small and the wt.
% of B₄C is low, B₄C particles can be hardly seen in the microstructure. This could not be confirmed by EDX analysis because boron and carbon elements could not be detected correctly with this analysis due to, they are light elements.

Figure 5.25. SEM-EDX analysis of Al alloy / B₄C region of Al alloy infiltrated 80S20B composite
Figure 5.2 shows \( \text{Mg}_2\text{Si} \) phase as the angular shape which was drawn with red lines detected at Al alloy infiltrated 80S20B composite. This phase is embedded in Al alloy and it can be seen at higher magnification such as 13000X. By EDX analysis Mg and Si elements in \( \text{Mg}_2\text{Si} \) and Al element around \( \text{Mg}_2\text{Si} \) were detected. Although these elements dissolved at the time of infiltration, \( \text{Mg}_2\text{Si} \) formed again at the time of solidification. \( \text{Mg}_2\text{Si} \) phase was also detected by XRD analyses and confirmed by SEM-EDX analysis.

Figure 5.26. SEM-EDX analysis of \( \text{Mg}_2\text{Si} \) region of Al alloy infiltrated 80S20B composite
5.8. HARDNESS MEASUREMENTS

Hardness were measured on Al alloy infiltrated 90S10B, 80S20B and 70S30B composites whose results are given in Table 5.7. Hardness measurements were done 5 times in the HRC scale by using 150 kg load. Then results were averaged. Since they are ceramic-metal composite, hardness of aluminum and ceramic rich regions revealed low and high values. Therefore, standard deviations are high among each layer and datas were scattered. But the average hardness values of each layer is close to each other. However, 70S30B layer has the highest hardness value and this layer can be considered as a front layer that breaks up to projectile into small pieces due to its highest hardness value if this composite is to be used for ballistic application. 70S30B layer contains the highest wt. % B₄C having higher hardness value than the hardness of SiC and the lowest vol. % Al phase which increases the hardness value due to increase in volume fraction of ceramic phases.

<table>
<thead>
<tr>
<th>Layer</th>
<th>Hardness (HRC)</th>
</tr>
</thead>
<tbody>
<tr>
<td>90S10B</td>
<td>36.96 (± 3.53)</td>
</tr>
<tr>
<td>80S20B</td>
<td>33.14 (± 2.05)</td>
</tr>
<tr>
<td>70S30B</td>
<td>38.82 (± 2.24)</td>
</tr>
</tbody>
</table>

5.9. THREE POINT BEND TEST

Three point bend tests were done to Al alloy and Al alloy infiltrated 90S10B, 80S20B and 70S30B composites whose results are given in Table 5.8. Al alloy exhibits the highest flexural strength. In terms of Al alloy infiltrated 90S10B, 80S20B and 70S30B composites, because 90S10B layer includes less wt. % B₄C and more vol. % Al alloy
due to its highest porosity %, flexural strength is the highest for 90S10B layer. Since 70S30B layer includes more wt. % B₄C and less vol. % Al alloy due to its lowest porosity %, flexural strength is the lowest for 70S30B layer. Flexural strength decreases gradually as illustrated in Figure 5.27. 90S10B layer may be considered as a backing layer that absorbs the remaining kinetic energy of propellants due to its highest flexural strength if this composite is to be used for ballistic application.

Flexure load versus flexure extension graphs are illustrated in Appendix.

<table>
<thead>
<tr>
<th>Al Alloy</th>
<th>280 (± 17)</th>
</tr>
</thead>
<tbody>
<tr>
<td>90S10B</td>
<td>193 (± 12)</td>
</tr>
<tr>
<td>80S20B</td>
<td>131 (± 1)</td>
</tr>
<tr>
<td>70S30B</td>
<td>111 (± 3)</td>
</tr>
</tbody>
</table>

Table 5.8. Three point bend test results

Figure 5.27. Flexural strength with respect to B₄C content
Three point bend tests were done FGM. 1 cm height ceramic-metal composites which consist of 3 different layers of FGM (90S10B, 80S20B and 70S30B) with 1,3 cm height Al alloy base were tested. Compression was applied from the ceramic-metal composite part (70S30B layer) which was developed for the front layer that projectile strikes. Flexure load versus flexure extension graphs are illustrated in Appendix and flexural strength was found 348 (±40) MPa. M. V. Silva et. al. [28] was applied three point bend test to 1 cm height sintered alumina plates and the results were 210 to 300 MPa. These alumina based ballistic plates (0,3 cm polyurethane + 1,2 cm sintered alumina + 0,4 cm SAE 1045 Steel) were also exposed to ballistic tests according to NIJ Level IV threat type and none of the plates were penetrated. This result shows that FGM can be considered as a potential candidate for armor material because the flexural strength of FGM is higher than the values reported in the literature.

Figure 5.28 shows FGM specimen subjected to three point bend test.

![Figure 5.28. FGM with Al alloy base specimen subjected to three point bend test](image-url)
CHAPTER 6

CONCLUSIONS

6.1. CONCLUSIONS

The major conclusions drawn from the results of experiments as follows:

- **B₄C – SiC – Al** composite which has low density \((2,7 \pm 0,02 \text{ g/cm}^3)\) and a combination of high hardness \((\text{up to } 38,82 \pm 2,24 \text{ HRC})\) and high flexural strength \((348 \pm 40 \text{ MPa})\) was produced by pressure melt infiltration process to obtain FGM.

- Composition is the most important variable to obtain gradual changing bulk density and porosity which are the requirements of an FGM sample. To develop FGM, three different layers \((90S10B - 80S20B - 70S30B)\) with different bulk densities \((2,06 \pm 0,03 - 2,14 \pm 0,02 - 2,28 \pm 0,03 \text{ g/cm}^3)\) and different porosity % \((34,2 \pm 0,91 - 29,6 \pm 1,21 - 23,2 \pm 0,88)\) was produced by altering composition respectively.

- Since the highest porosity % is obtained for 90S10B, the vol. % of infiltrated Al alloy is the highest. Since the lowest porosity % is obtained for 70S30B, the vol. % of infiltrated Al alloy is the lowest. Thus, the vol. % of Al alloy decreases from 90S10B to 70S30B gradually with respect to porosity % results.
FGM produced exhibits gradually increment of flexural strength (111 ± 3 – 131 ± 1 - 193 ± 12 – 280 ± 17 MPa) for 70S30B, 80S20B, 90S10B and Al alloy respectively which make it a potential candidate for armor applications.

Flexural strength for FGM B₄C-SiC-Al with dimensions of 12 cm length, 4 cm width and 2,3 cm thickness (1 cm height ceramic-metal composite with 90S10B, 80S20B and 70S30B layers and 1,3 cm height Al alloy base) was found 348 ± 40 MPa. This result shows that FGM can be considered as a potential candidate for armor material because this value is higher than the values (210 – 300 MPa) reported in the literature.

The new composition of Al alloy which had 11,2 wt. % Si and 6,9 wt. % Mg for pressure melt infiltration process was modified from 7075 Al alloy. Microstructure investigation showed that this composition was suitable to achieve process and to produce an almost pore-free composite.

Near or near-net shape products were produced as mass production with this pressure melt infiltration process.

With pressure melt infiltration, the sintered body was exposed to Al metal only a few seconds. Thus, there was no time to react Al element with carbide to yield detrimental Al₄C₃ phase together with B₄C and SiC particles. Elimination of this detrimental phase increased mechanical performance of composites developed.
6.2. RECOMMENDATIONS FOR FUTURE STUDY

- For future study, FGM may be exposed to the ballistic tests and modified according to results. To do that a bigger specimen dimensions suitable for ballistic tests should be studied. Figure 6.1 illustrates the armor design for future study. Fabric cover to protect ballistic plate from weathering conditions and to provide integrity of the materials inside the ballistic plate can be used. And some polymer backing such as aramid, UHMWPE, glass fiber, carbon fiber may be considered in addition to ceramic-metal composite.

![Armor design for future study](image)

**Figure 6.1. Armor design for future study**

- Since sintered ceramic exhibits high hardness, there is a machining problem. When the same mold is used for both pressing of powders and pressure melt infiltration, sintered ceramic does not fit mold due to thermal expansion. Thus, it is recommended that a mold used for pressure melt infiltration should be larger than a mold used for the pressing of powders to obtain green body.
REFERENCES


APPENDIX

\[ c = \frac{h}{2} \]
\[ h = 18 \text{ mm} \]

<table>
<thead>
<tr>
<th></th>
<th>Flexure load at Maximum Flexure stress (N)</th>
<th>Maximum Flexure stress (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>31577.74414</td>
<td>292.13495</td>
</tr>
</tbody>
</table>

Figure A. 1. Flexure load-Flexure extension graph of Al alloy specimen 1
\[ c = \frac{h}{2} \]
\[ h = 15.7 \text{ mm} \]

<table>
<thead>
<tr>
<th></th>
<th>Flexure load at Maximum Flexure stress (N)</th>
<th>Maximum Flexure stress (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>21971.73633</td>
<td>266.98361</td>
</tr>
</tbody>
</table>

**Figure A. 2.** Flexure load-Flexure extension graph of Al alloy specimen 2
\[ c = 0.565h \]
\[ h = 23 \text{ mm} \]

<table>
<thead>
<tr>
<th>Flexure load at Maximum Flexure stress</th>
</tr>
</thead>
<tbody>
<tr>
<td>50001.99609 (N)</td>
</tr>
</tbody>
</table>

**Figure A. 3.** Flexure load-Flexure extension graph of FGM specimen 1
Figure A. 4. Flexure load-Flexure extension graph of FGM specimen 2

$c = 0.565h$

$h = 23 \text{ mm}$

<table>
<thead>
<tr>
<th></th>
<th>Flexure load at Maximum Flexure stress (N)</th>
<th>Maximum Flexure stress (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>58738.65625</td>
<td>338.40494</td>
</tr>
</tbody>
</table>
\[ c = \frac{h}{2} \]
\[ h = 11 \text{ mm} \]

<table>
<thead>
<tr>
<th>Flexure load at Maximum Flexure stress (N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
</tr>
<tr>
<td>4412.75146</td>
</tr>
</tbody>
</table>

Figure A. 5. Flexure load-Flexure extension graph of Al alloy infiltrated 70S30B specimen 1
c = h/2
h = 20.1 mm

Figure A. 6. Flexure load-Flexure extension graph of Al alloy infiltrated 70S30B specimen 2
\( c = \frac{h}{2} \)
\( h = 10 \text{ mm} \)

<table>
<thead>
<tr>
<th>Flexure load at Maximum Flexure stress (N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
</tr>
</tbody>
</table>

**Figure A. 7.** Flexure load-Flexure extension graph of Al alloy infiltrated 80S20B specimen 1
\[ c = \frac{h}{2} \]
\[ h = 20.4 \text{ mm} \]

<table>
<thead>
<tr>
<th></th>
<th>Flexure load at Maximum Flexure stress (N)</th>
<th>Maximum Flexure stress (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>18158.55078</td>
<td>127.14342</td>
</tr>
</tbody>
</table>

Figure A. 8. Flexure load-Flexure extension graph of Al alloy infiltrated 80S20B specimen 2
$c = \frac{h}{2}$

$h = 12.5\text{ mm}$

<table>
<thead>
<tr>
<th>Flexure load at Maximum Flexure stress (N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
</tr>
</tbody>
</table>

Figure A. 9. Flexure load-Flexure extension graph of Al alloy infiltrated 90S10B specimen 1
$c = \frac{h}{2}$
$h = 20.5 \text{ mm}$

<table>
<thead>
<tr>
<th></th>
<th>Flexure load at Maximum Flexure stress (N)</th>
<th>Maximum Flexure stress (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>28120.45508</td>
<td>187.23705</td>
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</table>

**Figure A. 10.** Flexure load-Flexure extension graph of Al alloy infiltrated 90S10B specimen 2