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A Comparison Between Graphene and WS2 as Solid Lubricant Additives to Aluminum for Automobile Applications

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A COMPARISON BETWEEN GRAPHENE AND WS$_2$ AS SOLID LUBRICANT ADDITIVES TO ALUMINUM FOR AUTOMOBILE APPLICATIONS

A thesis submitted in partial fulfillment of the requirements for the degree of
MASTER OF SCIENCE
in
MATERIALS SCIENCE AND ENGINEERING

by
Sara Rengifo

2015
To: Dean Amir Mirmiran  
College of Engineering and Computing

This thesis, written by Sara Rengifo, and entitled A Comparison Between Graphene and WS$_2$ as Solid Lubricant Additives to Aluminum for Automobile Applications, having been approved in respect to style and intellectual content, is referred to you for judgment.

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

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Benjamin Boesl, Co-Major Professor

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Arvind Agarwal, Co-Major Professor

Date of Defense: March 20, 2015

The thesis of Sara Rengifo is approved.

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Dean Amir Mirmiran  
College of Engineering and Computing

________________________________________________________________________
Dean Lakshmi N. Reddi  
University Graduate School

Florida International University, 2015
DEDICATION

This is just the beginning of an amazing career. I dedicate this thesis to my beautiful family and my lovely husband. Thank you for supporting me during this path, and being my main motivation to achieve all my professional goals.
ACKNOWLEDGMENTS

I would like to acknowledge and sincerely thank all who contributed during the development of this thesis. Special thanks to Dr. Agarwal for his guidance, trust, understanding and most importantly his friendly advice during my graduate process. His mentorship was the mainstay of an excellent experience in the aim of reaching my goals as a scientist, not only helping me to grow in academia, but also as an independent thinker. Thank you Dr. Agarwal for giving me the opportunity to be part of your team. I thank all the members of the research group, Dr. Benjamin Boesl, Cheng Zhang, Sadegh Behdad and Chris Rudolf. Thank you all for your guidance, assistance and support on giving my graduate career the foundation of science.

I am very grateful with all the AMERI staff for your unconditional help and friendship especially Dr. Alex Franco, Dr. Yusuf Emirov, Ali Hadjikhan, and Neal Ricks. Also, I would like to thank my committee members Dr. Munroe, Dr. Boesl and Dr. Agarwal for providing helpful suggestions.

Finally, and most importantly, I would like to thank God for having my beautiful family and my lovely husband. You are my amazing miracle. You have taught me how to face challenges to find my inner strength and know that every step forward is a step towards achieving something bigger. All your love and support have given me the strength to never give up. I love you all and I owe you my deepest thanks.
The purpose of this thesis was to compare graphene nanoplatelets (GNP) and WS$_2$ as solid lubricant additives to aluminum in order to reduce friction and wear. The central hypothesis of this work relied on lubricating properties of 2D materials, which consist layers that slip under a shear force.

Two aluminum composites were made (Al-2 vol.% GNP and Al-2 vol.% WS$_2$) by spark plasma sintering. Tribological properties were evaluated by ball-on-disk wear tests at room temperature (RT) and 200°C.

WS$_2$ not only presented the lowest COF (0.66) but also improved the wear resistance of aluminum by 54% at RT. Al-2 vol.% GNP composite displayed poor densification (91%) and low hardness resulting in poor wear resistance. The wear rate of Al-2 vol.% GNP composite increased by 233% at RT and 48% at 200°C as compared to pure aluminum. GNP addition also resulted in lower COF (0.79) as compared to pure aluminum (0.87).
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CHAPTER I: INTRODUCTION

Overcoming friction in engine systems can significantly improve the fuel efficiency of automobiles. As a consequence, a significant amount of research has been done to develop automotive materials and techniques that reduce friction in automobile components. Some potential solutions to reduce friction in automotive components include development of (i) novel aluminum alloys and composites, (ii) advanced surface engineering techniques and anti-friction coatings and (iii) solid lubricant additives. The introduction of solid lubricant as additives to coatings, bulk aluminum alloys and composite materials is an emerging technology for reducing friction. 2D materials such as graphene, WS$_2$ and MoS$_2$ have emerged as novel solid lubricants with a greater potential to reduce friction. The goal of this thesis is to study and compare the tribological performance of graphene and WS$_2$ as solid lubricant additives to aluminum.

1. Motivation

1.1. Reducing Friction in Engine Systems

The principal motivation of this thesis focuses on tribological technology as a solution for reducing fuel energy consumption in passenger cars. Since 1966 the term “tribology” has caused apprehension among scientists, who defined tribology as “the science and practices relating to interacting surfaces in relative motion”. It encompasses friction, lubrication and wear [1]. Research on tribology has far-reaching economic implications, having made possible significant savings in the US gross national product. By 1976 a survey established that if tribology program were applied, 11% of total US energy consumption could be saved in the areas of road transportation, power generation,
and industrial machinery; those areas count for about 80% of the total energy consumption in the US [1]. In the 90’s, research on wear of metals brought a deeper understanding and help with progress in the tribology field [2]. However, nowadays in passenger cars, one-third of the fuel energy is used to overcome friction in the engine transmission, tires, and brakes [3]. In order to have a better view, friction loss distribution has been subdivided into four groups: 11.5% to overcome the rolling friction in the tire–road contact, 11.5% to overcome friction in the engine system, 5% in the transmission system, and 5% in the brakes (Fig 1.1).

![Fig 1.1. Breakdown of Passenger Car Energy Consumption [3]](image)

The friction loss by engine systems in a passenger car can be sub-divided into four groups: (i) the piston assembly (45%), (ii) the bearings and seals (30%), (iii) the valve train (15%), and (iv) the pumping and hydraulic (10%) [3]. The piston assembly includes the most critical part (45%) in tribological terms as it includes the ring pack. The ring pack has three main roles on piston’s performance; the first is to maintain an effective gas seal between the combustion chamber and the crankcase, the second is to transfer heat from the
piston into the cylinder wall, and then into the coolant, and the third function is to limit the amount of oil that is transported from the crankcase to the combustion chamber [4].

Fig 1.2. Piston assembly and Piston Ring Function from an Internal Combustion Engine [4].

A schematic representation of a piston assembly from a modern automotive engine is presented in Fig 1.2. The top two piston rings are denoted as the compression rings and the third as oil-control ring. Pressure of combustion pushes the two first rings out in order to seal the combustion chamber and prevent leaking, and the entire surface of the rings engages the cylinder wall. The oil-control ring is in charge of wiping excess oil from the cylinder wall [4]. Chong [5], who studied the piston ring forces without excluding the secondary motion (piston lateral motion) (Fig 1.3), determined the piston ring packs friction from the difference between the total friction force and the piston skirt friction force. Based on this finding, the piston posture is important in determining the piston ring friction force during the existence of piston secondary motion. In contrast, the same author [6] studied the piston forces excluding the secondary motion and concluded that the piston
ring friction force only depends on the lubrication system when the piston secondary motion is excluded.

Fig 1.3. Schematic Representation of Power Piston System [7]

When investigating friction in bearings, the lubricant plays a decisive role (Fig 1.4), hence, the mechanism in charge of causing friction in bearings is very different from pistons. Allmaier [8] described hydrodynamic losses and losses due to the metal–metal contact as two main mechanisms which cause friction in bearings. In describing the two mechanisms, Allmaier underscored the fact that reducing the sliding surface area of the crankshaft or increasing bearing clearance is effective in reducing friction in bearings, however, these methods can increase the wear, seizure or knocking [9]. In addition, the method to reduce the inherent hydrodynamic losses is effective by reducing the viscosity itself. Nevertheless, the consequence of a thin oil increases the likelihood of asperity contact [10].
Fig 1. 4. The Different Regimes of Lubrication According to Strubeck: Hydrodynamic (HD), Elastohydrodynamic (EHD), Mixed and Boundary Lubrication [10]

In a valve train friction analysis, considerable efforts have been made in order to reduce friction. Some investigators have proposed the reduction of weight and size of the design of the valve train. However, the results are not so motivating and have shown that further reduction is very difficult to achieve [9]. Another attempt includes the investigation into new materials such as titanium and different ceramics, these materials have performed well in reducing friction and so in fuel cost saving. However, the cost of these materials is very high, which makes their construction in industry level very complicated [9]. Also, analytical investigations based on elastohydrodynamic lubrication theory has taken place to predict the cyclic variation of important parameters such as minimum film thickness, maximum Hertzian stress and power loss, and the results show that not only does hydrodynamic play a main role, but the chemical reactions in the thickness of the film related to temperature are vital as well [11]. Likewise, pumping and hydraulic losses are
highly related to temperature. For instance, increased viscosity at low temperatures results in higher frictional consequences [12].

1.2. Fuel Saving and Reduction of Environmental Impact by Using Tribology Technology in Engine Parts.

An understanding of the tribology of the piston assembly, valve train, bearings and pumps in automobile engines points out that the most critical source of friction is in pistons and so in piston rings. The piston ring is the most complicated tribological component in the internal combustion engine to analyze because of large variations of load, speed, temperature and lubrication [4], thus challenges for engineers and designers in improving lubrication mechanisms to enhance energy efficiency with reduced environmental impact become a target for the automotive industry. Despite all the efforts and developments in engine design and oil formulation so far to reduce these, friction in engines remains an issue in both fuel saving and environmental impact.

Nowadays, the economy is deeply concerned with energy savings, and for this reason development of new automotive technologies and environmental impacts are being improved at the same time. However, worldwide, 208,000 million liters of fuel were used in 2009 to overcome friction in passenger cars [3]. With such large numbers of liters of fuel consumed, even the smallest improvement in engine efficiency, emission levels and durability of fuel can have a major effect on the worldwide fuel economy and the environment in the long-term (15–25 years) [4]. New technology for friction reduction in passenger cars, can result in a reduction in friction losses by 18% in the short term (5–10 years) and by 61% in the long term. This would represent in world economic savings the
amount of 174,000 million euros in the short term and 576,000 million euros in the long term. Those numbers represent in fuel savings the total of 117,000 million and 385,000 million liters, respectively; and CO₂ emission reduction of 290 million and 960 million tons, respectively [3].

The benefits of improving tribological properties in engines are [4]: reduced fuel consumption, increased engine power output, reduced oil consumption, reduction in harmful exhaust emissions, improved durability, reliability and engine life, reduced maintenance requirements and longer service intervals. However, in spite all these benefits, friction in one passenger car consumes an average of 340 liters of fuel per year Table 1.1 corresponding to a distance of 13,000 km [3].

<p>| TABLE 1.1 Friction Losses in One Global Average Passenger Car According To Type of Triboccontact [3] |</p>
<table>
<thead>
<tr>
<th>Friction loss source type</th>
<th>Energy consumed (MJ/car/a)</th>
<th>Fuel used (liters/car/a)</th>
<th>Percentage (%)</th>
</tr>
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<tr>
<td>Tire–road contact</td>
<td>1452</td>
<td>119</td>
<td>35</td>
</tr>
<tr>
<td>Hydrodynamic lubrication</td>
<td>1993</td>
<td>57</td>
<td>16.8</td>
</tr>
<tr>
<td>Mixed lubrication</td>
<td>899</td>
<td>26</td>
<td>7.6</td>
</tr>
<tr>
<td>EHD lubrication, sliding</td>
<td>747</td>
<td>21</td>
<td>6.3</td>
</tr>
<tr>
<td>EHD, sliding and rolling</td>
<td>979</td>
<td>28</td>
<td>8.2</td>
</tr>
<tr>
<td>EHD lubrication, rolling</td>
<td>356</td>
<td>10</td>
<td>3</td>
</tr>
<tr>
<td>Boundary lubrication</td>
<td>187</td>
<td>5</td>
<td>1.6</td>
</tr>
<tr>
<td>Viscous losses</td>
<td>771</td>
<td>22</td>
<td>6.5</td>
</tr>
<tr>
<td>Braking contact</td>
<td>1779</td>
<td>51</td>
<td>15</td>
</tr>
<tr>
<td>Total</td>
<td>11.863</td>
<td>340</td>
<td>100</td>
</tr>
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Approximately, 10 million automobiles are disassembled each year in the US, and the environmental impact created is shocking. Even, when about 75% of the weight of a
passenger car comes from recycled parts, and knowing that cleaner fuels generally result in lower emissions of toxics, there may be very harmful gases called heat trapping or greenhouse gases (GHG), in which some of their effects are severe weather patterns, ecosystem change, droughts and floods, sea level rise, and increases in incidence of infectious diseases [13].

It is the role of the materials engineers to research into new materials and meet humankind desire for energy and at the same time prevent the adverse environmental effects it produces, especially if it is known that only 21.5% of the fuel energy is used to move the car (Fig 1.1)[3].

1.3 Improving Wear and Coefficient of Friction

Aluminum alloys are widely preferred due to their light weight and excellent corrosion resistance, however the main limitations of these alloys are the lower fatigue strength and poor wear resistance [14]. In the effort of overcoming aluminum limitations in tribological performance, some hard ceramic particles have been used as reinforcement, such as SiC [14], TiB$_2$ [15], TiC [16], also called aluminum matrix composites (AMCs) [17]. However, some investigators have reported that AMCs reduce wear resistance at high temperature in comparison with performance at room temperature [17]. Also, the addition of self-lubricating materials, such as graphene to Al, resulted in an enhancement of strength properties with excellent distribution [14].

Due to solid lubricants’ intrinsic property of easy shear in addition to many advantages over liquid lubricants, GNP and WS$_2$ are added to aluminum to find a better coefficient of friction and wear resistance. Graphene nanoplatelets (GNP), consisting of
approximately 20–30 graphene sheets and with a thickness of 5–10 nm, is a 2D material that [18] exhibits many of graphene's unique properties. The increased thickness of GNP makes them less prone to agglomeration and entanglement as compared to single layer graphene [19]. GNPs have been explored as lubricants, both in liquid [20] and solid forms [19]. Likewise, WS₂ is recognized for high lubricating performance under a high load, high pressure and high temperature. A large amount of W can give excellent anti-wear properties and reacts with the metal to generate a protective tribofilm [21]. The coefficient of friction of the WSx film decreases slightly with an increasing S/W ratio [22].

The biggest concern about WS₂ as a lubricant is its high reactivity in humid environments and formation of WO₃, which affects tribological performance [23]. The biggest drawback in the performance of GNP as a lubricant is the tendency to increase porosity due to agglomeration, affecting the tribological performance [24].

1.4. Novelty of 2D Materials as Reinforcement

Since the separation of graphene into atomic arrangement of carbon sheets, and the demonstration of the novelty properties of graphite, two dimensional materials have been in the sight of researchers. Two widely studied 2D materials are Boron nitride (BN) and Molybdenum disulfide (MoS₂). In comparison with GNP, Boron nitride (BN) exhibits many of graphene’s astonishing properties in electrical applications [25]. Likewise, in comparison with WS₂, Molybdenum disulfide (MoS₂) presents excellent performance in tribological and electrical fields, and works as an excellent solid lubricant, giving a low coefficient of friction and high wear resistance [26].
2. Research Objectives

The overall objective of this study is to compare graphene and WS$_2$-based two dimensional (2D) materials as solid lubricant additives to aluminum in order to reduce the friction and improve energy efficiency of automobile engine components. This study involves the addition of 2D solid lubricant materials (GNP and nano size WS$_2$ powders) to an aluminum matrix to synthesize a composite material with improved tribological properties. The mixed powders were consolidated by spark plasma sintering (SPS) into dense composites for studying their tribological behavior under ball-on-disk sliding wear conditions at room temperature and 200°C. The specific novel scientific features include the following:

i. The effect of chemistry (GNP vs WS$_2$) on wear and friction

ii. The effect of temperature on wear and friction. A higher temperature of 200°C is selected, as it is close to the operating temperature of gasoline engines [11, 27].

In developing this thesis, six chapters are presented. Chapter one presents the introduction and background of the topics related with aluminum alloys, their usage in automotive engine components, and the benefits of developing anti-friction systems that can reduce gas consumption worldwide. The following chapter is Chapter 2, in which the state of art is presented. Some previous investigations on graphene and WS$_2$ as solid lubricants are discussed. Chapter 3 details the experimental procedure, assumptions and evaluations that are made during the study of these novel materials. Chapter 4 provides the results obtained by applying the experimental procedure proposed in Chapter 3. The mechanisms of wear and friction for each material are discussed and compared in this
chapter. Finally, Chapter 5 presents the major conclusions and Chapter 6 sets some suggestions for future studies to expand knowledge about reduction of friction through solid lubricants’ implementation.
CHAPTER II: REVIEW OF THE STATE OF THE ART

1. Aluminum in Automotive Applications

The demand for aluminum composites in automotive applications has increased due to their lightweight, excellent corrosion resistance with good strength, and low density compared with steel [28]. The density of aluminum (2700 kg/m³) is one third the density of steel, which allows 60% saving in weight in the engine and transmission parts, 50% in bumper systems and more than 30% in chassis and suspension parts [29]. Weight reduction of the car is strongly related with the energy consumption, and thus with environmental impact. Some reports assert that, on average, 100 kg (220.5 lbs.) of mass reduction can save 0.35 liters of fuel per every 100 km (62 miles), and 10 grams of CO₂ per km in emission [29, 30]. It translates that 10% in weight reduction improves the economy related with fuel consumption by 5.5% [31].

Currently, varieties of aluminum alloys are used [32] for different automotive components, as shown in Table 2.1 [33]. Al-Si alloys have presented high potential to be used in automotive, aerospace and mineral industries because of their enhanced properties such as strength, wear resistance, and stiffness [34]. Aluminum matrix composites (AMC) reinforced with ceramic particles or fibers have also been widely investigated to obtain higher specific modulus and strength, as compared with conventional alloys. It has been found that on increasing ceramic particles size or volume, AMCs exhibit higher wear resistance in applications such as automotive brake rotors, pistons, cylinder liners, and turbine compressors. However, wear resistance cannot be considered to be a core material property of these kinds of composites [34].
Reducing the friction between two surfaces can be achieved by introducing an intermediate lubricating layer, which effectively separates the two metals [35]. Most metals like aluminum get oxidized in air and create a thin oxide layer with 1 to 10 nm thickness [36]. This oxide layer creates a natural lubricant by allowing shear between metal contacts under low applied load, presenting features of low shear strength and ductility, resulting in low friction between the surfaces [36, 37, 38]. However, the layer becomes damaged under high loads, resulting in higher wear rate and coefficient of friction [36]. Since aluminum is a soft metal, its intrinsic properties do not present good wear resistance, and the high coefficient of friction can be attributed to the high contact area and small elastic recovery [36]. In order to keep the lubricating characteristics of oxide film, pure aluminum must be converted into an alloy or an additive should be used, which could provide improved lubrication while maintaining or improving the wear resistance. Surface engineering and solid lubricant addition to aluminum have been used for the same purpose. The addition of boron nitride (h-BN) to Al demonstrated that h-BN particles with size of 30µm and concentration up to 10%, presents a wear rate decrease of 25% in comparison with pure grease [39]. Likewise, 5 wt% graphite resulted in a reduction in COF in the 7075 Aluminum matrix, while enhancing wear resistance with increments in sliding distance [40]. Some other researchers have demonstrated the formation of the lubricious tribofilm on the sliding surface, preventing surface damage and reducing wear and friction on aluminum alloys such as 6061 [41] and Al-Si [42].
### TABLE 2.1. Aluminum Applications in the Automotive Industry [33].

<table>
<thead>
<tr>
<th>Aluminum Applications in the Automotive Industry (1993 North American Car)</th>
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<tbody>
<tr>
<td><strong>Powertrain (Casting)</strong></td>
</tr>
<tr>
<td>Pistons (100%)</td>
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<tr>
<td>Engine blocks (19%), oil pans, engine cover, water pumps</td>
</tr>
<tr>
<td>Cylinder heads (75%)</td>
</tr>
<tr>
<td>Intake manifolds (85%)</td>
</tr>
<tr>
<td>Transmission: cases (100%), valve bodies, and channel plates (90%); rear axle and differential housings; driveshafts</td>
</tr>
<tr>
<td><strong>Chassis (castings)</strong></td>
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<tr>
<td>Wheels (40%)</td>
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<tr>
<td>Brackets</td>
</tr>
<tr>
<td>Brake components: master brake cylinders</td>
</tr>
<tr>
<td>Suspension: control arms, supports</td>
</tr>
<tr>
<td>Steering components: air-bag supports, steering shafts, wheels knuckles, housings</td>
</tr>
<tr>
<td>Instrument panels</td>
</tr>
<tr>
<td>Electric motor, alternator, and pump housings</td>
</tr>
<tr>
<td><strong>Chassis (wrought)</strong></td>
</tr>
<tr>
<td>Wheels</td>
</tr>
<tr>
<td>Heat shields</td>
</tr>
<tr>
<td>Bumper reinforcements</td>
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<tr>
<td>Body (wrought)</td>
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<tr>
<td>Hoods (9%)</td>
</tr>
<tr>
<td>Hoods, front fenders, and doors</td>
</tr>
<tr>
<td>Body structure</td>
</tr>
<tr>
<td>Air conditioning (wrought)</td>
</tr>
<tr>
<td><strong>Condensers, evaporators/compressors (90%)</strong></td>
</tr>
<tr>
<td><strong>Heater cores, radiators (72%)</strong></td>
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</table>

### 2. 2D materials as Solid Lubricants

Solid lubricants are those materials that present very low friction and low wear, when rubbed against another material in the absence of a liquid lubricant [36]. The most studied solid lubricants are graphite and molybdenum disulfide (MoS$_2$) [36, 43]. Both materials present hexagonal layered structures (lamellar structure), in which the atoms
within each layer are held by strong covalent bonds. The individual layers have significant
distances between one another and are held by weak Van der Waals bonding. The
interplanar bond energy is about 1/10 to 1/100 of that between atoms within the layers [36].
These two materials share many of solid lubricant’s properties, however, in many
applications, graphite performs quite superior to MoS2 [43]. Molybdenum disulfide
belongs to the family of binary compounds, known as metals dichalcogenides. These
metals present excellent self-lubricating properties. Some of these metals are WS2, MoSe2,
NbSe2 [43, 44]. A brief review of synthesis technique and properties of 2D materials used
as solid lubricants is discussed in the following sub-sections.

2.1. Synthesis Methods of 2D Materials

In the 1920s, Langmuir introduced the single molecular layer to investigate the air-
water interface in a hydrophobic field [45]. Then, Blodgett used the same monomolecular
layer method, but this time to study the air-water interface onto solid surfaces [45]. Since
then, research regarding monoatomic surfaces has grown for multiple interests in
engineering applications, such as corrosion protection, the increasing of wear life in tools
and solid lubricants, among others. In accordance with Erdemir [46], the historical
development of very thin solid lubricants is composed of three stages (Fig 2.1): (i) the
research of specific material compositions, (ii) the development of the deposition
processes, and (iii) the development of successive generations of films. The development
of the vacuum deposition techniques, such as plasma vapor deposition (PVD), chemical
vapor deposition (CVD), and ion-beam-assisted deposition (IBAD) has made it possible to
deposit thin layers with a thickness of a few nanometers [46]. In the application of these
techniques, the geometrical shape of the powder is conserved and so is the ability to shear [47, 48].

![Fig 2.1. Historical Development of Thin Solid Lubricant Films [46]](image)

In principle, the layered structure of 2D materials is what gives the lubricating feature. This structure is achieved due to the synthesis of 2D sheets, which are largely formed by micromechanical cleavage, chemical exfoliation, chemical vapor deposition, and surface-assisted epitaxial growth [49]. Micromechanical cleavage was initially used to peel off graphene from graphite. This method requires repeatedly peeling off sheets of 1 to 10 atomic layers from the top layer. Nevertheless, this method is not used for large-scale synthesis due to poor yield. Chemical exfoliation is another synthesis process, which involves sonication of layered bulk materials in polar solvents, resulting in separation into thin layers by centrifugal forces. This method is used for large scale synthesis of 2D materials. The third method is chemical vapor deposition (CVD), which allows the
production of 2D materials by high-temperature chemical reactions of molecular precursors on a surface. This method also has limitations for large scale production. The fourth method, which is surface-assisted epitaxial growth, is a modification of the CVD method. Here, the substrate surface serves as a seed crystal other than a template or a catalyst. This method has achieved the formation of monoatomic thick Si sheets [49].

The focus of the present work is on using Graphene and WS₂ 2D materials as solid lubricant additives to Al alloy. Hence, a brief review of Graphene and WS₂ is provided in the following sections.

3. Graphene

Graphene is a 2D allotrope of carbon consisting of a single sheet of hexagonal arrangements with a pseudo-two-dimensional sp₂ bonding (Fig 2.2 b) [50, 51]. The Bravais lattice of graphene is triangular, where the lattice vector \( \approx 1.42 \text{ Å} \) is the nearest distance between carbon-carbon atoms. The hexagonal lattice contains two atoms A and B per unit cell. It consists of two sublattices, where A is surrounded by three atoms from sublattice B, and B is surrounded by three atoms of the lattice A (Fig 2.2 a) [52]. Graphene is an isolated plane or monolayer of graphite (one atom thickness). Although graphite was been used 6000 years ago to decorate pottery, it was in 2004 that one atom sheet of graphene was obtained by peeling graphite off with an adhesive tape [53]. Since then, the unique electronic and physical properties of graphene have been extensively studied for a myriad of application, such as electrical devices, chemical and biological sensors, solid lubricant additives in liquid and solid state, composites and alloys, preventing materials to be corroded, and photovoltaic cells [50].
Fig 2.2 a) Graphene Hexagonal Lattice with Sublattices A and B [52] b) Representation of Hexagonal Layered Structure of Graphite Showing Three Staggered Layers [36]

The theoretical density of graphene is 1.822 g/cm³ with a very high Young’s modulus (0.5–1 TPa) and tensile strength (130 GPa) in the planar direction [54]. These properties of graphene make it an ideal reinforcement for light-weight and high-strength composite materials. Graphene nanoplatelets (GNPs) are another version of graphene, which consist of approximately 20–30 graphene sheets with a thickness of 5–10 nm [18]. GNPs present hydrogen or covalent bonding capability on the edges of the platelets; allowing them to present enhancement barrier properties and excellent mechanical properties, such as stiffness, strength, and surface hardness [55]. In addition, these nanoplatelets have shown excellent performance in electrical and thermal applications due to their carbon-carbon composition. The multilayer structure of GPNs makes them very useful for tribological applications and ideal as additives with metals [56], polymers [57], and ceramics [58]. GNPs retain several of graphene’s unique properties and provide a low-cost alternative for development of novel materials and composites. Some of the
theoretical properties of GNPs that have been previously measured are: the tensile modulus (1000 GPa), the tensile strength (5 GPa), and the electrical conductivity (10^7 Siemens/m) measured parallel to the surface [55]. These mechanical properties allow GNPs to improve barrier properties and scratch resistance without reducing toughness. Moreover, the increased thickness of GNPs makes them less prone to agglomeration and entanglement, compared to single layer graphene [19]. GNPs have been explored as lubricants, in both liquid [20] and solid forms [19].

3.1. Synthesis of Graphene

The primary methods used to produce graphene in the large-scale are: (i) epitaxial growth of graphene, (ii) chemical vapor deposition (CVD), (iii) chemically derived graphene and (iv) electro-chemical exfoliation.

Epitaxial growth of graphene is used in both semi-conductive and metallic materials that have good lattice matching with the hexagonal structure of graphene [59]. The growth of epitaxial graphene is influenced by thermal decomposition of carbide (SiC), decomposition of hydrocarbon gas on carbide substrates (TiC, TaC, WC, etc.) or metallic substrates (Ni, Pt), and segregation of the atoms of carbon from the bulk [59]. Epitaxial growth on SiC is based on thermal decomposition of the SiC substrate, so, Si atoms sublimate and the remaining carbon atoms become activated, thus the C-face is fully relaxed and adopts the bulk graphite parameter at the high temperature of 1350°C. This fact describes the weak interaction between graphite and the SiC substrate due to Van der Waals epitaxy, where the substrate is in charge to orient the epitaxial layer while the lattice parameters remain intact, and the interfaces between 2D and 3D materials are free from
defects [53, 60]. Graphene has also been grown by decomposition of hydrocarbons, which are compounds made of C and H atoms [61]. Some of these hydrocarbons can be found in the natural state from crude oil and natural gas. Others are obtained by synthesis, for instance: CH₄, C₂H₆, C₂H₄, C₃H₆, and C₃H and some plastics. When these hydrocarbons get oxidized, carbon dioxide and water are produced with a significant release of energy; therefore, they are widely used as fuels [62]. 2D bands of monolayer graphene have been successfully obtained by hydrogen intercalation (sublimation of Si from the substrate) and characterized by a honeycomb structure with lattice constant of 2.5 ± 0.1 Å [63].

Chemical vapor deposition (CVD) has become the most promising method to grow graphene. Since 2006, when the first successful monolayer of graphene was obtained by CVD, this process has been widely explored and it was subdivided into three categories, thermal CVD, plasma enhanced CVD, and thermal decomposition on SiC and other substrates [53]. The first subdivision can be made with high carbon solubility (>0.1 atomic %) and low carbon solubility (<0.001 atomic %). The growth mechanism of graphene with high carbon solubility utilizes substrates, such as Co and Ni. The process works through the diffusion of the carbon into the metal; a thin film of carbon goes out of the bulk metal during the cooling. A typical CVD process is performed using Ni as a substrate. In this case, carbon is dissolved into nickel substrate, and finally the carbon is precipitated on the substrate by cooling. The characteristics of the process include a CVD chamber, where the Ni substrate is placed at a vacuum of 10⁻³ Torr with a temperature <1000 °C, while the hydrocarbon gas is diluted. Similar to the carburization process, CVD starts with the diffusion of a specific number of carbon atoms into the Ni substrate at low temperatures [53]. The second subdivision of CVD is plasma enhanced CVD deposition (PECVD),
which offers another way of acquiring thin layered graphene. The first use of PECVD successfully obtained ultrathin sheet-like carbon nanostructure by synthesizing CH$_4$ diluted in H$_2$ by radio-frequency plasma. These graphene sheets were produced from a gas mixture of 5–100% CH$_4$ in H$_2$ atmosphere while varying the temperature of the substrate from 600-900°C [64]. The last subdivision of CVD, called thermal decomposition on SiC and other substrates, is commonly preferred for the semiconductor industry. The SiC substrate is heated under ultrahigh vacuum (UHV). Silicon atoms are extracted from the substrate. The removal of Si allows the formation of carbon surface into graphene layers. Depending on time and temperature (approximately 1200°C), the thin thickness of graphene layers is obtained. Vapor phase annealing has shown improvement on the homogeneity of the few graphene layers, typically at 400 °C above of ultrahigh vacuum temperature [53].

Graphene is also synthesized by reduction of Graphene oxide (GO), which is a chemically derived compound from carbon molecules and oxidizers [53]. Graphite oxide was first prepared almost 150 years ago, by replacing graphite for a mixture of potassium chlorate and nitric acid [65]. Years later, Hummers and Offemann developed a much safer mixture of sodium nitrate, potassium permanganate and concentrated sulfuric acid to grow GO [50, 65, 66]. Hummers’ method is broadly used in laboratories nowadays. Graphite oxide is similar to graphite from the chemical properties’ point of view (carbonaceous), however, is worth noting that the structures are different [50, 53]. Different methods have been used for reduction of graphene oxide, including hydrazine, and sodium borohydrate. The first mentioned method, hydrate, demonstrated excellent results in the production of very thin sheets. During the reduction process, the hydrazine does not react with water and the reduced sheets are precipitated; the color of the dispersed graphene oxide goes from
brown to black, thus removing oxygen, the reduced graphene oxide becomes less hydrophilic [53].

The mechanical exfoliation process could be the simplest of all processes to synthesize graphene. It arises from the fact that graphene layers are held by weak interlayer Van der Waals forces. The interlayer Van der Waals energy 2 eV/nm, which translates to a force of 300 nN/µm² necessarily exfoliates or isolates a monolayer of graphene [67]. This (300n N/µm²) was the same required force by the adhesive-tape process in 2004, when the first graphene layer was obtained [53]. Mechanical exfoliation techniques have been able to extract ~10 micron graphene flakes [68]. Therefore, this process is commonly preferred for electronic applications and devices as it readily provides high quality of a single layered graphene [67]. However, despite all investigations, even nowadays, the production of high-quality large area monolayer graphene remains very challenging.

3.2. Characterization of Graphene

Characterizing one atom thick sheets represent an enormous challenge. Various techniques are used in the characterization of graphene, such as transmission electron microscopy (TEM), scanning electron microscopy (SEM), Raman spectroscopy, atomic force microscope (AFM) and X-ray diffraction (XRD). In general, TEM is a technique, which works at the nano and micro scale by using a beam of electrons and transmitting it through an ultra-thin sample. The image is formed when the electrons pass through the specimen and interact with the detector. TEM is a very accurate tool when characterizing graphene. The atomic scale of graphene can be depicted (Fig 2.3).
TEM has allowed researchers to appreciate graphene and GNPs dispersion properties with composites [69]. Significant observations of the graphene monolayer of carbon atoms have been reported for TEM. Overlapped GNPs with a thickness of about 20 nm have been observed between two grains by TEM [70]. The nature of interface bonding between GNP/matrix can be obtained by TEM to understand load transfer behavior between GNPs and the matrix [70]. Also, researchers can observe imperfections and abnormal topology on graphene sheets [53] and the number of layers by using TEM [71]. However, the use of TEM in the atomic scale has the restriction of low operating voltage; otherwise, at high voltage, it can be very harmful to the monolayer [69].

Raman spectroscopy is a very important tool to characterize the structure of graphene, by giving information regarding interactions of phonons and electrons in the material while measuring the wavelength and intensity of light scattered of the molecules. Raman spectra of graphite and a few layers of graphene is shown in Fig. 2.6 a. Typically, Carbon allotropes present three distinguished peaks G (1580 cm\(^{-1}\)), D (1350 cm\(^{-1}\)), and 2D
(2700 cm\(^{-1}\)) [53, 73]. The G peak (~1580 cm\(^{-1}\)) [53, 73, 74] is the crystallization peak of graphene and arises due to the in plane vibration of the sp\(^2\) carbon atoms. The G peak is present in all carbon materials but the peak intensities and widths can vary. The D peak (~1360 cm\(^{-1}\)) is related with defects in the layer and can, therefore, be used to observe damage in the graphene structure [73]. The absence of the D peak implies a high quality single layer defect free graphene. In the G’, also called 2D band (~2650 cm\(^{-1}\)), intensity decreases with increasing numbers of graphene layers [75]. This 2D peak is the second order of the D peak. The 2D peak in bulk graphite consists of two components, 2D\(_1\) and 2D\(_2\), measuring 1/4 and 1/2 the height of the G peak, respectively. An increment in layers leads to a decrease of intensity of the lower frequency 2D\(_1\) peaks (Fig 2.4 b) [74].

![Graphene and Graphite few Layered Raman Spectra](image)

Fig. 2.4. a) Graphite and Graphene few Layered Raman Spectra [73] b) 2D Band Changes with Increment on Layers [74]

The ratio of the intensity of the D peak divided by the intensity of the G peak (I(D)/I(G)) gives a measure of the level of defects in a graphene structure[73, 74]. The intensity ratio of peak (I (2D)/I(G)) is almost 4 for single-layer graphene and decreases
when the number of layers increases [53, 73, 74]. Raman spectroscopy can also provide information about the presence of mechanical strain at the molecular level [76]. Tsoukleri [76] reported that in tension; the 2D peak decreases with strain and the peak tends to be shifted to the left. In contrast, the peak is shifted to the right side in compression and presents higher frequency than tension.

3.3 Graphene Nanoplatelets (GNPs)

Graphene nanoplatelets (GNPs) consist of approximately 20–30 graphene sheets held by week Van der Waal forces and with a thickness of 5–10 nm [18]. Due to their unique structure, these carbon-based sheets are an excellent representation of 2D materials. GNP presents very high surface area and higher thickness. These two properties allow GNPs to have higher interfacial energy and make them less prone to agglomeration and entanglement as compared to single layer graphene. [19]. Due to their high surface area, these platelet-like sheets also presents strong interfacial bonding, making GNPs an excellent nano filler in composite with a very tough bonding mechanism. An astonishing property of 2D materials is the high strength connection due to the high surface area. This ability provides higher resistance when propagating cracks in comparison with 1D materials. The increased area of GNPs is effective under applied load, meaning the shear force needed to pull a GNP layer out has to be higher than the amount of force (higher energy) needed for another material (1D for instance). This phenomenon can be understood as an effective and tougher mechanism to prevent crack propagation.

Recent studies have posed fundamental questions like how friction behavior depends on the number of graphene layers. That is why GNPs are being explored as
lubricants, both in liquid [20] and solid states [19, 77]. Wenzheng et al. [77] revealed that the addition of small amounts of GNPs in the Ni₃Al matrix composite helps to radically reduce the friction coefficient and wear rate at both room and high temperatures (400°C). Also, the authors explained that the ability of GNPs sheets to slip easily provides a source of stress dissipation. Moreover, GNPs create a protective layer while sliding. This protective layer leads to the reduction of wear rate as well as the friction coefficient [77].

The 2D nature of GNP also enables an easier dispersion and processing, as compared to their 1D counterpart carbon nanotubes. Some studies have even shown that the easy-dispersion characteristic of GNPs works as an excellent stress dissipator by creating a smooth worn surface and randomly disperses GNPs on this worn surface during tribological tests [77]. The stress concentrates along the edge of the wrinkles and folds during the sliding process, so the initial shape with wrinkles becomes a laminated-like shape. This characteristic combined with enough thickness, allows GNPs to avoid becoming curled up or rolled up into CNT-like tubes, as occurs with single-layer graphene. The 2D nature of graphene also benefits during sintering processes that employ high pressures. During the high-pressure sintering process, these 2D materials get oriented in one direction during axial compression while reorienting the sheets before the fracture with minimum damage. Andy et al [78] found that GNPs endured the spark plasma sintering (SPS) process at a very high temperature of 1850 °C and a pressure of 80 MPa, presenting minimum damage to the structure [78]. GNPs are much cheaper and easier to consolidate than single-layer graphene, but still exhibit many of the amazing properties of graphene.
4. Tungsten Disulfide (WS₂)

Tungsten disulfide (WS₂) is a transition metal made up of WS₂ sheets. These sheets are characterized by an S-W-S sandwich-like structure, in which the elements form a hexagonal crystal structure. The adjacent layers along [0 0 1] direction are held by weak Van der Waals forces, while the bonding between the same elements is characterized to be a strong covalent interaction [79, 80]. WS₂’s lamellar structure is characterized by a silvery-gray-black color, a space group P6₃/mmc and parameters a = 3.154 Å and c = 12.362 Å (Fig 2.5) at room temperature conditions [80]. This anisotropic material has shown high potential in applications such as catalysis, lithium batteries, photoconductors and semiconductors, shock absorbers, solar cell films, wet and solid lubricants [80, 81]. Selvi [80] demonstrated that the WS₂ structure is dependent on the pressure, showing an a-axis decrease by 3.8% and a c-axis decrease by 10.2% when 25.5 GPa pressure was applied at room temperature. It was concluded that c-direction of the hexagonal structure is much more compressible than the a-direction. It can be explained by analyzing the Van der Waals

![Fig 2.5. Crystal Structure of WS₂ [80]](image-url)
bonding at lower pressure. For instance, at 3.3 GPa, Van der Waals bonds are very weak, so it permits larger compression, different from a high pressure, where the force between S-S planes increases due to repulsive forces. This ability has made of WS\textsubscript{2} one of the most promising lubricous material ever known. WS\textsubscript{2} presents a lower coefficient of friction and improvement in oxidation resistance and thermal stability, even for about 100°C above the well-known MoS\textsubscript{2} [82].

4.1 Synthesis of WS\textsubscript{2}

The growth of thin-layered WS\textsubscript{2} has mainly been performed by different types of exfoliation such as liquid exfoliation mechanical exfoliation [83], exfoliation of bulk [84] and chemical vapor condensation (CVC) [85]. WS\textsubscript{2} samples used for this project were grown through liquid exfoliation, where the powder is sonicated in a range of liquid solvents, such as benzyl benzoate, isopropanol, acetone, and methanol. Although metal dichalcogenides can be exfoliated by liquid exfoliation, this is a very time-consuming method, which is extremely sensitive to the environment [86]. Fang [83], obtained single-layer WS\textsubscript{2} (~0.7nm thickness) by mechanical exfoliation from a bulk crystal on Si/SiO\textsubscript{2} substrate. It was noticed that the surface roughness of WS\textsubscript{2} is analogous to Si/SiO\textsubscript{2}, indicating that the layer is uniform with minimum roughness. This method has also proved successful under vacuum and at high temperatures (750–950 °C), to create a monolayer of WS\textsubscript{2} with an area of ~1cm\textsuperscript{2} [84]. Other researchers, have obtained nanosheets of WS\textsubscript{2} through exfoliation of bulk, in which WS\textsubscript{2} powders were sonicated in N-Dimethylformamide (DMF) (Fig 2.6) [87].
Gas phase reactions of the chemical vapor condensation process have been used to successfully synthesize WS$_2$ nanosheets. This method has the ability of producing 2D material with little agglomeration. During this process, the decomposition of tungsten hexacarbonyl W(CO)$_6$ is conducted at temperatures from 420 to 1000°C; the following is the expression of the decomposition: $\text{W(CO)}_6 \rightarrow 6\text{CO+W}$. The process includes two main zones. In the first zone, the vapor of sulfur is mixed with carbonyl vapor and then this mixture is transferred to the second zone, which has been heated. Here, the carbonyl pyrolysis process takes place. Therefore, the formation of nanopowder is achieved in the sulfur [85].

Ultrathin-film of WS$_2$ can be grown and deposited as well by other techniques like sputtering, pulsed laser deposition (PLD) [88, 89], and chemical bath deposition (CBD) [90]. In 1994, the PLD technique was used to grow single layer WS$_2$. The results showed a high-quality monolayer, fully dense, which presented excellent performance in tribological applications with a low coefficient of friction. Since then, PLD has been widely used for growing WS$_2$ thin layers in coatings [89]. Thin films of WS$_2$ growth by PLD demonstrated only 3% structural mismatch occurs with the strontium titanate SrTiO$_3$ (STO)
substrate at temperatures between 200-400°C [88]. Hankare [90] successfully obtained thin film of WS₂ (0.35µm) by chemical bath deposition in an aqueous alkaline environment, with uniform shape and well adherent characteristics. It is worth noting that in growing polycrystalline films, the substrate plays a major role in the crystal formation. WS₂ has a minuscule tolerance for the lattice mismatch. It is not surprising to end up with an amorphous film of WS₂ [88]. In order to characterize the lattice of the film and the defect zones, it is important to know effective characterization tools for WS₂.

4.2 Characterization of WS₂

The techniques used in characterization of WS₂ particles are transmission electron microscopy (TEM), X-ray diffraction (XRD), scanning electron microscopy (SEM), Raman spectroscopy, and atomic force microscope (AFM). The typical observation of WS₂ through TEM is a 2D thin flake or nanosheet. Fig 2.7a shows a WS₂ nanosheet with a lateral size of about 100 to 500 nm. Fig 2.7b displays the hexagonal structure of WS₂ with some defects. Fig 2.7c allows a higher magnification of the structure by using low-pass Butterworth filtering to make the measurement of distance or width of the hexagon which was 3.8 Å [87]. Wang [91] confirmed the formation of the hexagonal structure of WS₂ through HRTEM, with a distance equal to 0.276 nm in (100) plane, and compared with the results obtained from XRD. Another researcher [92] implemented HRTEM to characterize the WS₂ solid lubricant coating (Fig 2.8a) and the transfer film formed by WS₂ (Fig 2.8b). Randomly oriented WS₂ lattice planes were observed in the transfer film, with no evidence of reorientation of the basal planes with the c-axis. It is worth noting that TEM and XRD work conjointly in characterizing WS₂-based materials.
Wang [81] synthesized WS$_2$ nanosheets (10 nm thickness) by the mechanical activation process at different temperatures (500°C, 600°C, 700°C, and 840°C), and implemented XRD to characterize and confirm the presence of hexagonal layers. The typical (002) peak was present in all the samples, which reveals the presence of WS$_2$. However, at synthesis temperature greater than 840 °C, the phase WO$_2$ appears due to the lack of sulfur, which is evaporated at high temperature (Fig 2.9).
Raman spectroscopy is another technique used to characterize the structure of the 2D material, such as WS$_2$. The typical Raman spectra obtained from WS$_2$ is depicted in Fig 2.10. WS$_2$ is represented by two main peaks located at 355 cm$^{-1}$ and 420 cm$^{-1}$. Tungsten trioxide (WO$_3$) shows up with peaks at 715 cm$^{-1}$ and 809 cm$^{-1}$ [93].

**5. Comparison of GNP and WS$_2$ for Tribological Applications**

Graphene and WS$_2$ are 2D materials, which are well known for their intrinsic properties of wear and coefficient of friction reduction [95, 95]. In order to better
understand the behavior of these 2D materials under applied force, it is important to know their tribological mechanisms at both the micro and nano scale.

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<thead>
<tr>
<th>TABLE 2.2</th>
<th>Tribological Mechanisms of GNP And WS₂</th>
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<tr>
<td><strong>FEATURE</strong></td>
<td><strong>GNP</strong></td>
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<tr>
<td>Structure</td>
<td>Monolayer of carbon atoms arranged in a hexagonal structure with the SP₂ bonding between carbon atoms. The sheets are held with Van de Waals forces [96]</td>
</tr>
<tr>
<td>Characteristics</td>
<td>• Less prone to agglomeration in comparison with graphite [97]</td>
</tr>
<tr>
<td></td>
<td>• Excellent corrosion protection [97]</td>
</tr>
<tr>
<td>Pros for Tribological applications</td>
<td>• High flexibility and mechanical stiffness under applied loads, preventing the direct contact between asperities on the steel ball counterpart and the surfaces [98]</td>
</tr>
<tr>
<td></td>
<td>• Absorbs energy generated by compression and shear during friction sliding [98]</td>
</tr>
<tr>
<td>Cons for Tribological applications</td>
<td>• Needs from a humid environment to provide lubrication [20]</td>
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</tbody>
</table>
In the aim of comparing these two solid lubricants, important research has been conducted. As mentioned before, the number of layers is very important for these solid lubricants. In the case of GNP sheets, the ability to easily slip provides a source of stress dissipation and allows GNPs to create a protective layer while sliding [77]. Thus, with fewer layers, the top layer deflects more, and the friction per unit rises. The top surface of the stack becomes less yielding and more slippery as graphene layers are added [94, 98]. The 2D nature of GNP also enables a very easy dispersion that helps GNPs to work as an excellent stress dissipator [77]. Some of the disadvantages of graphene in tribological performance are folding and wrinkling [99]. Another disadvantage, such as free edges, can be reduced in GNPs due to the multilayered structure. Two of the remarkable capabilities of GNPs are the unusual flexibility and the ability of kinking; these two astonishing abilities have never been observed in other atomic-sized carbon materials [99]. Also, the weak interlayer bonds allow slippage when GNPs are kinked, becoming an affective stress releaser during applied load without fracture. Therefore, instead of plastic deformation, GNPs experience layer sliding or layer buckling to reduce the stress. GNPs present two
important behaviors, which are wrinkling and tearing. Wrinkling is the deflection effect of the thin sheets, and it is unable to recover the initial shape due to insufficient elasticity of the material. On the other hand, tearing is an effect of unusual fracture behavior of GNPs. All those characteristics regarding deformation and fracture behavior can help to better understand anti-wear properties of graphene nanoplatelets. GNPs present unique flexibility and mechanical stiffness under the application of loads. *The structure of GNPs works by absorbing the energy generated by compression and shear during frictional performance* [98]. A probable explanation for this mechanism is the *formation of a protective layer* on the sliding contact interfaces. This fact is due to the two dimensional shape of the membrane, which acts a facilitating shear and considerably reduces the wear [20].

**WS₂** has crystal structure formed by the hexagonal of S-W-S sandwich. These three layers are held by weak interlayer VDWs interactions, while the bonding in-between the layers is covalent [79, 80]. **WS₂ has excellent tribological properties due to the ability of the layers to slide back and forth by intercrystalline slip under a shear force.** This mechanism allows a surface to rub against another surface by *creating a smooth transfer film*. This transfer film, working in conjunction with the intercrystalline slip of the layers, provides incredible lubricating properties [101]. Some researchers [93, 102] attribute this fact to the assertion that it is a synergistic effect more than lubricating itself. Some other hypotheses have been made to explain the tribological performance of tungsten disulfide. Prasad [93] hypothesized how the crack propagation effect influences the tribological behavior of WS₂. A crack in WS₂ film will generate a breakdown in the covalent bonds, creating activated surfaces with unsaturated bonds. The formation of these unsaturated bonds accelerates the crack propagation and also leads to the formation of WO₃.
response of WO$_3$ is due to the dangling bonds of WS$_2$ when reacting with oxygen. It has been proved that the presence of WO$_3$ highly affects the friction coefficient during tribological performance at room temperature [23]. Prasad [95] and coworkers explained that hard particles, randomly distributed in the WS$_2$ substrate, are able to deflect or end crack propagation. Thus, an excellent opportunity to improve the performance of the solid lubricant is through the formation of secondary phase protrusions, which provides the growth of microscopic reservoirs of the solid lubricant improving the quality and life of the transfer film.

The growing demand for reducing friction makes graphene and WS$_2$ composite materials a target-research for materials engineers. It is important to understand the tribological mechanism at the nanoscale as well. By exploring the entire range of available tensile orientations in graphene, it is found that this material presents a quasi-isotropic behavior for all tensile angles. This behavior is different from fracture response, where graphene behaves anisotropic [103]. It is worth noting that GNPs tend to align perpendicularly to the pressure, resulting in anisotropic mechanical properties of composites [104]. It has been demonstrated that the structure of graphene is unstable due to the atomic thickness [105], however, the $sp2$ bonding in graphene yields to effective lubricating properties of easy shearing and delamination of the 2D graphene planes. By observing the atomistic thickness of the layer, it is possible to explain the microscopic origin of friction and tribological properties of graphene. Mechanical energy is the beginning of the tribological process, and here the interfacial forces play a main role. Thus, it is demonstrated that the friction behaves differently for one layer than for multilayer-graphene, showing that the higher friction on 1 layer could be caused by higher interfacial
forces as compared with 2 or multi layered graphene [51]. When the counterpart exerts compressional force on the graphene layer, the energy between counterpart and graphene is higher than the energy between graphene layers, this effect leads to the attachment of the top layer with the counterpart, forming a conformal layer around it [106].

At small scale WS₂ tribological behavior is explained by the crystal reorientation of the layers, forming the transfer film on the counterpart and providing long wear life and lowering friction coefficient [100, 107]. When pressure is exerted on a WS₂ layer, the c-direction of the hexagonal structure is much more compressible than the a-direction, where c-direction is the distance reduction between S-S planes. It can be explained by analyzing the Van der Waal bonding at lower pressure. For instance at 3.3 GPa, the Van der Waals bond is very weak. It permits larger compression different from a high pressure where the force between S-S planes increases due to repulsive reaction [80]. It was found that the variation in the flake sizes of WS₂ layers can make a difference in tribological properties. It has been observed, that in different conditions such as high pressure or high temperature, nanoparticles of WS₂ react with the metal substrate generating a thick tribofilm. These nano-particles fill the gaps and disperse themselves through the reacted tribofilm, and reduce the friction at the boundaries [108].

Both materials graphene and WS₂ present intrinsic properties as solid lubricants. However, it is important to notice a big difference between them. WS₂ works as an effective lubricant in air, inert gases, and high vacuum over a wide temperature range (-190°C to over 800°C). This fact, represents and advantage over graphene, which losses its low
friction properties in environments in absence of condensable vapors or water [80]. However, WS$_2$ in humid conditions forms WO$_3$ and degrades its tribological behavior.

The next chapter discusses experimental procedures to synthesize GNP and WS$_2$ reinforced aluminum composites and their tribological behavior.
CHAPTER III: EXPERIMENTAL METHODS

In this chapter, the experimental methods of the preparation and consolidation of powders, tribological testing and characterization of the samples is described. The spark plasma sintering process (SPS) is utilized to consolidate aluminum composites with 2D additives such as GNP and WS₂. The tribological tests are conducted at room and high temperature (200°C). Microstructural characterization of the powder, composite and post wear samples is performed at every stage using different methods such optical microscope (OM), scanning electron microscope (SEM), electron diffraction spectroscopy (EDS), X-ray diffraction (XRD), Raman spectroscopy and transmission electron microscope (TEM).

1. Powders

Spherical atomized aluminum powder (H3, Valimet Inc., Stockton, CA, USA) with an average particle size of 2-10.5 microns and purity greater than 99.7wt.% was used as the matrix material and control sample. Fig 3.1 (a, b) show SEM images of the appearance of the as-received H3 aluminum powder at low and high magnification.

Fig 3.1. SEM Image of As-received Aluminum Powder at a) Low and b) High Magnification.
Graphene nanoplatelets ((xGPN-M-5)) were obtained from XG Sciences, Lansing, MI, USA. The GNPs have a thickness of 6-8 nanometers with an average diameter of about 15 microns and the average platelet consists of 20 graphene sheets (Fig 3.2).

![Fig 3.2. a) SEM Image and b) TEM Image of as Received Graphene Nanoplatelet Powder](image)

Nano-powder of tungsten disulfide (WS₂) was obtained from Graphene Supermarket, USA. Preliminary characterization of nano-WS₂ powder as received is obtained by SEM (Fig 3.3a). The nano-powder used in this study has an average size of 90 nm with hexagonal morphology. The average surface area of nano-WS₂ particles is ~30 m²/g with a true density of 7.5 g/cm³ [18].

![Fig 3.3. SEM Image of as Received WS₂ Powder at a) Low and b) High Magnification](image)
Our objective was to synthesize powders containing 2 vol.% of 2D additives to Al powder. The powders were mixed by the wet chemistry technique. Pure Al powder was added to acetone and ultrasonicated for 90 minutes using Branson 3510 Ultrasonic Cleaner. Similarly, GNP and WS$_2$ powders were added to acetone and sonicated in two different beakers for 90 minutes. The amount of GNP, WS$_2$ and Al powders added were computed to result in 2 vol.% composite. After sonication of GNP and WS$_2$, the slurries were respectively added to the separate beakers containing sonicated Al powder. Al-GNP and Al-WS$_2$ slurries were further sonicated for 60 minutes. Three mixtures pure Al, Al-2 vol.% GNP, and Al-2 vol.% WS$_2$ slurries were put into oven at 75°C for 24 hours for drying.

2. Spark Plasma Sintering

Sintering is a fundamental process in which the powder is compacted. During spark plasma sintering, an electric pulse is conducted through the powder while exerting compression in a graphite die, providing high densification in a very short time [109, 110]. The significant advantage of SPS is the fast heating rate and uniform heating condition with significant saving of time during densification, which allows retention of the nano-sized grains reducing coarsening.

During this study, the mixed powders were consolidated at Oklahoma State University (OK, USA), using a Spark Plasma Sintering 10-3 model. The consolidation was carried out in a 20 mm diameter graphite die using a holding pressure of 50 MPa. The maximum temperature used during sintering was of 500 °C with a holding time of 10 minutes. A heating rate of 50°C/min in vacuum was used. The consolidated samples were cooled in the furnace.
3. Characterization

The characterization process has been made by three steps which are (i) density measurements, (ii) microstructural and chemical composition and (iii) phase identification.

*Density Measurement:* The density of a powder and sintered composite was measured using AccuPyc II 1340 Series Pycnometer utilizing Helium gas. Helium atoms fill the vacancies and pores up to 2Å providing more accurate results [111, 112]. The density obtained via helium pycnometer was qualitatively compared using image analysis technique from the polished cross-section of the sintered compacts.

*Microstructure and Chemical Composition:* The cross-sections of the sintered samples were grinded and metallographically polished up to 0.5µm alumina powder. A Buehler Versamet 3 Optical Microscope was used to observe the cross-sectional microstructure and the sub-surface formed after wear test. Scanning electron microscopy (JEOL JSM-633OF field emission scanning electron microscope (FE-SEM)) with an operating voltage of 15 kV was used to characterize the powders, microstructure of the sintered pellet and the fracture surfaces. A combination of low and high magnification SEM images provided information about dispersion of the additives into the matrix and morphology of the 2D GNPs and WS₂ additives. Also, these images allow to identify how these additives interact with Al matrix, giving information about grain size and trapped porosity of the sintered samples. Energy-dispersive Spectroscopy (EDS) was carried out using a JEOL JIB 4500 SEM to study the chemical composition and elemental analysis in the sintered composites, as well as analysis of phases formed and oxygen content after wear.
**Phase Identification:** X-ray diffraction is a completely non-destructive technique for identification of phases. A D5000 Siemens diffractometer was used for 2θ scan between 10 and 90 degrees. Scan speed was set at 2 deg/min. The operating voltage and current are 40 kV and 35 mA respectively. The radiation used is Cu Kα which has a wavelength of 1.542 Å. This technique is used to obtain information about phases present in the powder and pellets and if any phase transformation occurred during SPS.

Raman spectroscopy was used to characterize the structure of graphene and WS₂. Raman spectroscopy is a technique where the incident light produces inelastic scattering in which the interaction of phonons and electrons forms the Raman spectra. Micro-Raman spectroscopy analysis was conducted using a Spectra Physics (Model 3900S, California, USA) with Ti-sapphire crystal as the target (514 nm), a laser power of 18mW and a detector.

**4. Hardness Measurement**

Hardness of a material provides useful information which can be correlated to its tensile strength, wear resistance, ductility, and other physical characteristics. Hardness of the sintered pellets was measured using Vickers Microhardness tester and a nanoindenter at micro and nano-scale respectively. For the microhardness tests, 100 gram load was used in 15 different attempts and the average was taken. Vickers pyramid hardness number (HV) is calculated as shown in Eqn. (3.1), where \( L \) is the average length of the diagonal of the indent produced in square shape, and \( P \) is the load.

\[
HV = \frac{1.854 \ p}{L^2}
\] (3.1)
Nanoindentation experiments were carried out using Hysitron TI 900 Triboindenter with a 100 nm Diamond Berkovich tip. The tip-area calibration was done using a standard fused quartz substrate of known modulus (69.6 GPa). The triboindenter was used in quasi-static indentation mode in order to measure the elastic modulus and hardness of the sintered compacts. Nanoindentations were performed on top surface, a total of 16 indentations were made with separation of 5μm from one another. The indentation process consisted of a 10 second period to reach a load of 1500 μN, with a holding time of 3 seconds, followed by a 10 second unloading to 0N. The elastic modulus was calculated from the load-displacement unloading curves using Oliver-Pharr method.

5. Tribological tests

Tribological test were conducted in rotative Mode (ASTM G99). The wear tests were conducted using a ball-on-disk tribometer (Nanovea, Irvine, CA), using a static 3 mm diameter alumina oxide ball as the counter material. Sintered samples rotated at 100 rpm making 4mm diameter wear track during a total period of 30 minutes. A normal load of 1N was applied. A real-time coefficient of friction was also recorded using LVDT method. Wear tests were conducted at room temperature and 200°C. High temperature tests were conducted at 200°C to better simulate gasoline operated engines [11, 27]. For high temperature conditions a furnace surrounding the sample was preheated until it reaches 200°C, and subsequently wear tests were started. The initial surface roughness determines the rate at which the asperities from the softer surface are either fractured or deformed during sliding [113]. All samples were polished to obtain very similar surface roughness (Ra = 0.62-0.69 um) prior to wear tests.
6. Post Wear Characterization

After wear, all the tracks were scanned by a noncontact three-dimensional (3-D) optical profilometer PS50 (Nanovea). Wear depth was obtained using a line profile with SPIP software (Image Metrology A/S, Horsholm, Denmark). Five different curves of the cross-section of the wear track are measured. Then, using Origin 6.0 and taking the integral of these curves the average cross-section area of the track and its standard deviation are computed. After getting the average this new cross-sectional area is multiplied by $2\pi r^2$, where $r$ represents the half of the diameter of the wear track. Thus, the volume loss during wear test is found. Fig 3.4a, b depicts the cross section of the wear scar. Likewise, wear volume was computed and subsequently wear rates.

Fig 3.4. a) Wear Track of Al-2vol.%GNP scanned at RT by Nanovea PS50 Optical Profilometer, b) Cross-sectional area of wear track (width and height are specified)

Each sample is weighed before and after conducting the wear tests, and the weight loss is obtained as well as the sliding distance. All these parameters define the wear rate as shown in Eqn. (3.2), where $V$ is volume loss (mm$^3$), $S$ represents sliding distance (mm)
and $P$ is the applied load (N) during wear test. In addition, some other calculations are made to better understand the wear mechanisms of the different samples i.e effective surface area ($m^2$) (Eqn. (3.3)), where $r_i$ is the radius of the counterface ($Al_2O_3$ ball) and $w$ is the width (See Fig 3.5), and lateral stress (Pa), where $\mu$ is the average coefficient of friction (Eqn. (3.4)).

\[
\text{Wear rate } [mm^3/(Nmm)] = \frac{V}{*P} \tag{3.2}
\]

\[
\text{Effective surface area} (E_{sa}) [mm^2] = \frac{r_i * r_i * w^2}{2} \tag{3.3}
\]

\[
\text{Lateral stress } [pa] = \frac{P}{E_{sa}} \tag{3.4}
\]

JEOL-JIB 4500 Multi Beam Focused Ion Beam (FIB) is a technique used to characterize nanoparticles and observe subsurface. This tool helps to observe the subsurface characteristics, such as crack distribution, crack propagation, formation of the tribofilm and wear mechanisms after tribological test is performed. The use of this tool helps to understand how stress contributes to the formations of the tribofilm and how the grains are deformed and the cracks are distributed due to stress. The way to use this technique is making a trench in the middle of the wear track (Fig 3.5) with ions using tungsten gas nozzles. After protective layer deposition a rough cut was performed by introducing a high dosage of ions ~3000Pa and then it was followed by a smooth polish using a lower dosage of ions ~500Pa. The dimensions of the trenches were ~40x40 microns. The whole procedure takes about 5 hours and then the cross-section of the wear
track is exposed and can be characterized using SEM. Fig 3.5 depicts the use of the focused ion beam tool, showing the location of the trench (by the red circle) and also a higher magnification image of it (observe the red arrow). The trench observed in this figure belongs Al-2 vol.%WS₂ sample.

![Fig 3.5: FIB Performed on the Surface of the Wear Track](image)

XRD analysis is performed between 10 and 90 degrees in the 2θ menu. Scan type is set as locked-coupled and the scan speed is set at 2 deg/min. The operating voltage and current are 40 kV and 35 mA respectively. The radiation used is Cu Kα which has a wavelength of 1.542 Å. The sample has to fit within the holder and the top of the sample is positioned flat against the pin located in the sample mount. This technique is used to obtain information about phases present in the pellets and any phase transformation occurred during SPS, also this tool helps to visualized if any phase is formed after wear.

Energy-dispersive Spectroscopy (EDS) was carried out using a JEOL JIB 4500. This tool works by bombarding a specimen with an incident electron beam inside of a scanning electron microscope. Qualitative analysis of the results involves identification of wear morphology and prediction of wear mechanism as well as formation of tribolayers.
Scanning electron microscopy (SEM) is used to characterize the powders, the composites or pellets, the fracture surface of each pellet and the wear track. The equipment implemented was JEOL JSM-633OF field emission scanning electron microscope (FE-SEM) with an operating voltage of 15 kV. After wear, this tool allows the observation of mechanism in charge of wear, formation of tribolayers, formation of cracks and porosity of wear surface.
CHAPTER IV: RESULTS AND ANALYSIS

This chapter presents experimental results and discussions on the effect of adding GNP and WS\textsubscript{2} as solid lubricant additives to aluminum matrix. The overall microstructure of Al, Al-2 vol.% GNP and Al-2 vol.% WS\textsubscript{2} is described in terms of densification, and dispersion of GNP and WS\textsubscript{2} in aluminum matrix. The phases present in the composites are characterized by x-ray diffraction and Raman spectroscopy. Subsequently, tribological behavior of Al-2 vol.% GNP and Al-2vol.% WS\textsubscript{2} is presented in terms of wear rate, and coefficient of friction at room temperature and 200°C. Wear mechanisms are described in terms of tribolayer formation. Pure aluminum pellet is used as a point of reference for the sake of comparison.

4.1. Microstructure of Sintered Al, Al-2 vol.% GNP and Al-2 vol.% WS\textsubscript{2}

The overall cross-sectional microstructure of Al, Al-2 vol.% GNP and Al-2 vol.% WS\textsubscript{2} is shown in low magnification SEM micrographs in Fig 4.1. Pure Al (Fig 4.1a) and Al-2 vol.% GNP (Fig 4.1b) show sintered spherical aluminum particles with high degree of porosity. However, Al-2 vol.% WS\textsubscript{2} shows a very dense structure with fine WS\textsubscript{2} particles homogeneously dispersed in the matrix (Fig 4.1c). The poor density in Al-2 vol.% GNP is attributed to agglomeration of GNPs in the starting powder which caused a barrier in the sintering. Agglomerated GNPs are observed as long stringers in Fig. 4.1b. The high density of Al-2 vol.% WS\textsubscript{2} sample is attributed to nanosize WS\textsubscript{2} powder which tends to fill in the pores between spherical Al powders resulting in a better packing density due to bimodal powder size distribution. Pure Al displays poor sintering due to large spherical particles which forms small necks during sintering but leaves large trapped pores.
Fig. 4.1. SEM of Cross-section for a) Pure Al, b) Al-2 vol. % GNP, c) Al-2 vol.% WS₂

A similar phenomenon is observed in fracture surface of three samples as shown in Fig 4.2. Fig 4.2 a, shows large aluminum spherical particles which are poorly sintered particles. Fig 4.2b shows an agglomerated GNP stringer surrounded by poorly sintered aluminum particles. The GNPs are wrinkled and agglomerated. Although the wrinkled structure indicates that GNPs may have experienced damage, it also has the ability to transfer the load to the matrix, leading to stronger interfaces in the sintered sample. However, due to poor sintering of surrounding aluminum particles will results in weak interface. Fig 4.2c is a low magnification SEM image of Al-2 vol.% WS₂ showing highly
dense structure as compared to pure Al and Al-2 vol.% GNP samples. The high magnification image of Al-2 vol.% WS$_2$ clearly shows WS$_2$ nanoparticles in the void region between Al particles which leads to denser packing.

Table 4.1 shows the density of sintered pellets and their hardness measured using microhardness tester and nanoindenter. The densification measured by helium pycnometer is a reflection of microstructure described earlier. The lowest hardness is observed for Al-2 vol.% GNP, which is attributed to the low percent of densification caused by agglomeration of GNP. It has been known that GNP tends to agglomerate and restack allowing the formation of porosity [56]. The highest hardness and percent of densification
are observed for Al-2 vol.% WS$_2$. Although absolute hardness values measured by nanoinserter and microhardness tester are different due to measurement at different length scales, they show a similar trend.

4.2. Phase Identification

<table>
<thead>
<tr>
<th>Composition</th>
<th>Nanohardness (GPa)</th>
<th>Microhardness (HV)</th>
<th>Percent of densification (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pure Al</td>
<td>1.1±0.3</td>
<td>59.2±3.9</td>
<td>96.9</td>
</tr>
<tr>
<td>Al- 2 vol.% GNP</td>
<td>0.8±0.2</td>
<td>55.7±4.9</td>
<td>91.0</td>
</tr>
<tr>
<td>Al-2 vol.% WS$_2$</td>
<td>1.2±0.1</td>
<td>59.4±5.1</td>
<td>99.9</td>
</tr>
</tbody>
</table>

X-Ray diffraction of sintered pellets was carried out to observe if any new phase(s) were formed during sintering. All major peaks representing pure Al are located at 2θ equals to 38.74°, 44.96°, 65.29°, 78.29° and 82.81 in three samples. Pure Al also shows trace amount of Al$_2$O$_3$ formation as observed from the lower portion of Fig 4.3. Al-2 vol.% GNP sample resulted in the formation of trace amount of Al$_2$O$_3$ and Al$_4$C$_3$ phases as shown in Table 4.2. In addition, the main peak representing the retention of carbon-base material can be also observed at (006) plane. WS$_2$ peaks at 2θ equals to 14.17° (002), 34.79° (100), and 39.87° (103) confirm the retention of the hexagonal structure in sintered Al-2 vol.% WS$_2$ sample. Trace amount of WO$_2$ and WO$_3$ oxides are also formed due to oxidation of WS$_2$ as shown in Table 4.2. It must be recalled oxide amounts are in trace due to spark plasma sintering processing done under vacuum condition.
TABLE 4.2
Reactants Products and $\Delta G$ During Sintering Process

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Delta Gibbs Free Energy At Sintering Temperature (500°C) (J)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$2 \text{Al} + \frac{3}{2} \text{O}_2 \rightarrow \text{Al}_2 \text{O}_3$</td>
<td>-1433270.1</td>
</tr>
<tr>
<td>$\text{WS}_2 + \text{O}_2 \rightarrow \text{WO}_2 + 2\text{S}$</td>
<td>-221244.3</td>
</tr>
<tr>
<td>$\text{WS}_2 + \frac{3}{2} \text{O}_2 \rightarrow \text{WO}_3 + 2\text{S}$</td>
<td>-415865.5</td>
</tr>
<tr>
<td>$4 \text{Al} + 3 \text{C} \rightarrow \text{Al}_4 \text{C}_3$</td>
<td>-183260.1</td>
</tr>
<tr>
<td>$\text{H}_2 + \text{S} \rightarrow \text{H}_2\text{S}$</td>
<td>-45340.5</td>
</tr>
</tbody>
</table>
In order to have a better understanding of the retention of the 2D structure of the solid lubricants (GNP and WS₂) after SPS, the Raman spectroscopy was performed on the starting powders and sintered pellets. Fig 4.4 shows Raman spectra from the starting GNP powder and sintered Al-2 vol.% GNP pellet. Three representative peaks of carbon are known as D, G, and 2D located around 1336 cm⁻¹, 1583 cm⁻¹ and 2702 cm⁻¹ respectively.

![Raman Spectroscopy](image)

**Fig 4.4. Raman Spectroscopy for GNP Powder and Al-2 vol.% GNP**

The D peak represents the presence of edges of graphene, ripples, and atomic disorder. The G peak is associated to the doubly degenerated E₂g phonon mode at the Brillouin zone center and represented the in-plane vibration of the sp² carbon atoms, and the third peak (2D) presents more intensity in comparison with peak D and represents the second order...
Raman scattering process giving information about the number of graphene layers present in the structure [46]. The number of graphene layers is determined by the observation of the band’s width (2D peak), the sharper the peak the fewer layers the structure has [46, 114]. The intensity ratio between 2D and G bands (I_{2D}/I_{G}) is a measure of graphitic vs. graphene nature. An I_{2D}/I_{G} ratio of less than 1 indicates graphite. I_{2D}/I_{G} ratio for the starting GNP powder is ~5.5 indicating few layered structure (Fig. 4.4). The I_{2D}/I_{G} ratio for the sintered Al-2 vol.% GNP composite is ~2.85 which shows that more layers are present in the GNP’s structure in the composite. This could be due to stacking of several GNPs under high pressure of SPS. From Fig 4.4 the ratio I_{D}/I_{G} for starting GNP is ~0.079, which increases to 1.19 in the composite showing that defects has increases due to sintering process. It can be concluded from Raman Spectroscopy that GNP structure is retained with some damage after SPS consolidation. A typical Raman spectra of WS2 shows two main peaks located at 355 cm^{-1} and 420 cm^{-1}, which correspond to the E_{2g} and A_{1g} modes [115].

![Raman Spectroscopy](image)

Fig 4.5 Raman Spectroscopy for WS2 Powder and Al-2 vol.%WS2
The $A_{1g}$ peak represents the $S$–$S$ vibration between two different WS$_2$ layers. The relative intensity of the $A_{1g}$ peak suggests that the vibration energy is small and that the particles, therefore, contain only a few stacked layers. It has been demonstrated that the ratio $I_{2LA}/I_{A_{1g}}$ is always greater than 2 for monolayer films [116]. In the present study, the ratio $I_{2LA}/I_{A_{1g}}$ for the powder is $\sim$0.86 and for the composite is $\sim$1.68 indicating the presence of few layered structure in both WS$_2$ powder and Al-2 vol.% WS$_2$ composite (Fig 4.5).

Based on all results obtained from microstructural characterization and phase identification, it was confirmed that 2D solid lubricants are retained in the composite. The following section will discuss wear behavior of these composites with solid lubricant additives.

### 4.3. Tribological Behavior

#### 4.3.1 Wear Rate

Wear resistance of pure Al, Al-2 vol.% GNP and Al-2vol.% WS$_2$ is characterized in terms of wear rate ($\frac{V}{s}$), which is expressed as volume loss (V) divided by the sliding distance (S). The results are presented in Table 4.3.

<table>
<thead>
<tr>
<th>Sample Composition</th>
<th>Wear Rate-RT (mm$^3$/N mm)</th>
<th>Wear Rate (200 °C) (mm$^3$/N mm)</th>
<th>COF-RT</th>
<th>COF (200°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pure Al</td>
<td>$5.7 \pm 1.6 \times 10^{-6}$</td>
<td>$7.5 \pm 1.9 \times 10^{-6}$</td>
<td>0.87±0.04</td>
<td>0.82±0.03</td>
</tr>
<tr>
<td>Al-2 vol.% GNP</td>
<td>$1.9 \pm 0.28 \times 10^{-5}$</td>
<td>$1.1 \pm 0.31 \times 10^{-5}$</td>
<td>0.79±0.09</td>
<td>0.69±0.10</td>
</tr>
<tr>
<td>Al-2 vol.% WS$_2$</td>
<td>$2.6 \pm 0.17 \times 10^{-6}$</td>
<td>$5.3 \pm 0.56 \times 10^{-6}$</td>
<td>0.66±0.09</td>
<td>0.55±0.10</td>
</tr>
</tbody>
</table>

TABLE 4.3
Wear Rate and Coefficient of friction at Room Temperature and 200°C
distance (S) and applied normal load (P). Wear track is scanned using optical profilometer which provides wear depth and cross-sectional area of the track to enable estimation of wear volume. The wear rate for Al, Al-2 vol.% GNP and Al-2 vol.% WS₂ at room

Fig 4.6. Wear rate for Ball-on-disk sliding Experiments at Room Temperature.

Fig 4.7. Wear rate for Ball-on-disk Sliding Experiments at 200°C
temperature and 200°C are presented in Fig 4.6 and 4.7 respectively. The data for wear rates is also presented in Table 4.3.

It was observed that Al-2 vol.% GNP shows the highest wear rate among three samples whereas Al-2 vol.% WS₂ shows the lowest wear rate. Al-2 vol.% WS₂ shows 54% improvement in the wear rate at room temperature as compared to pure aluminum. At 200°C, Al-2 vol.% WS₂ shows 29% improvement in the wear rate as compared to pure aluminum. This can be attributed to densification and hardness of three samples as listed in Table 4.1. Al-2 vol.% WS₂ shows highest densification (99.8%) and hardness resulting in the lowest wear rate. Al-2 vol.% GNP presents the lowest densification (91.0%), and is the softest sample among three. Archard’s equation illustrates that: the harder the material, better the wear resistance will be [118]:

$$dV = \frac{kP}{H} \cdot dx \quad (1)$$

In Eqn 1, dV is the wear volume loss, P and H are the load and the hardness respectively, k is called the wear resistance factor, and dx is the wear distance. The overall wear behavior and trend of three samples is similar at room temperature and 200°C. Table 4.3 shows that wear rate increases at higher temperature except with the Al-2 vol.% GNP sample. It is well known that materials become softer at higher temperature resulting in more wear. However, the GNP-containing sample displayed the contradictory behavior at high temperature which is explained by two phenomena: (i) localized sintering and densification of 91% dense Al-GNP sample at high temperature, in which the plastic deformation and the soft characteristic due to high temperature contribute to compact the
sample due to stress from the counterball and (ii) the ability of GNP s to act as sealers against oxygen which contributes to improve wear [119].

4.3.2 Coefficient of Friction

In addition to the wear rate, coefficient of friction (COF) is the critical parameter to obtain an indirect measurement of energy efficiency in automotive materials. Fig 4.8 shows the coefficient of friction variation during the room temperature wear test along with the average value for Al, Al-2 vol.% GNP and Al-2 vol.% WS$_2$. COF at 200°C is shown in Fig 4.9. The average values of COF are listed in Table 4.3.

![Coefficient of Friction vs Time](image)

**Fig 4.8.** Average COF and COF Variation During RT Tests

It was observed that aluminum has the highest COF (0.87) whereas Al-2 vol.% WS$_2$ has the lowest COF (0.66) at room temperature. A similar trend is observed for average COF at 200°C; however COF values are lower than those at room temperature (Table 4.3).
It is also observed that COF variation is more stable at 200°C as compared to RT tests. However, Al-2 vol.% WS$_2$ sample displayed most stable COF without significant oscillations both at RT and 200°C. It is worth noting that Al-2 vol.% GNP presents an improvement in COF in comparison with pure Al in spite of having the highest wear rate. This is largely due to the presence of GNP sheets which act as solid lubricant and contribute to reduce friction.

![Coefficient of Friction](image)

**Fig 4.9. Average COF and COF Variation at 200°C Tests**

The worn surfaces of the samples were studied using SEM and x-ray diffraction. The SEM images of the worn surfaces provide important information to understand wear mechanisms of the different samples. The sub-surface of the wear was studied using FIB and optical microscopy to understand the formation of tribolayer and dynamic wear features such as cracks progression. Fig 4.10 is that x-ray diffraction of the wear surface of Al, Al-2 vol.% GNP and Al-2 vol.% WS$_2$ before and after wear. It provides information about the phase(s) formed during wear at RT and 200°C.
4.4. Wear Mechanisms at Room Temperature

4.4.1 Wear mechanism of pure Al at room temperature

A common type of wear in Al is adhesive wear [113, 120], which occurs when two surfaces are initially rubbed against each other. However, due to localized heating and oxygen diffusion, the surface forms Al₂O₃ layer which is brittle in nature and undergoes abrasive wear. It is expected that hard and brittle Al₂O₃ phase will generate large particles
of wear debris, thus increasing wear rate and COF. Fig. 4.11 shows a schematic of wear mechanisms of pure Al at room temperature.

![Schematic Representation of Behavior of Pure Al at RT](image)

**Fig 4.11. Schematic Representation of Behavior of Pure Al at RT**

Figure 4.12a shows the wear track of pure Al which is very rough and presents formation of cracks. The surface of the wear track, reveals crack formation, and the EDS (Appendix 1) shows high amount of oxygen. The brittle Al$_2$O$_3$ surface layer generates large amount of wear debris (Fig. 4.12b). These wear debris present average bimodal size distribution of 20 µm and 60 µm. These big detachments reveal adhesive wear for pure Al, which is a very common type of wear in this material [113, 117], whereas finer debris is indicative of mixed nature due to presence of brittle Al$_2$O$_3$. The wear particles are lifted up and allow the observation of some cracks propagating through the grain boundaries, which can be attributed to surface crack propagation [120]. Fig 4.12c exposes surface crack propagation of the wear track. The thickness of the tribofilm in a range of 5-10 µm as indicated by the red arrow in Fig. 4.12c and 4.12d. The formation of subsurface zone is the result of severe contact stresses during the wear [121]. Aluminum forms a brittle Al$_2$O$_3$ tribofilm which is rough due to cracks resulting in a higher COF as observed earlier.
4.4.2 Wear mechanism of Al-2 vol.% GNP at room temperature

The addition of GNP to Al matrix is expected to promote the formation of lubricious tribofilm that will reduce COF and wear rate of Al composites [14, 21]. The stated hypothesis of GNP in Al matrix expects to find lower wear rate and COF than for pure Al. However, it was seen in section 4.1 that Al-2 vol.% GNP displayed lowest wear resistance and intermediate COF. Fig 4.12 shows a schematic of the proposed wear mechanism in Al-2 vol.% GNP at RT. Figure 4.14 a shows the wear track of Al-2 vol.%
GNP. The smooth region indicates formation of a tribofilm. However this tribofilm is not homogeneous and contains rough region indicating large wear. The red arrow marks the thickness of the tribofilm which is ~5 µm (Fig 4.14 c and d). Fig 4.14c also shows the crack-end and the beginning of the new tribofilm formed by deflection as pressure from the counterball is exerted during wear. This tribofilm consists of Al₂O₃ and Al₄C₃ in addition to smooth region of graphene as indicated in x-ray diffraction pattern shown in (Fig 4.10). In addition, the EDS of the worn surface (Appendix 1) shows higher oxygen, suggesting that oxide formation occurs due to faster oxygen diffusion in low density (91%) composite. Homogeneous detachments are found as wear debris with average size of 60 µm (Fig 4.14b). No plastic deformation is observed in debris particle, suggesting that the wear debris is lifted up before plastic deformation. Thus, it is clear that GNPs are detached from the surface by delamination and crack propagation (Fig 4.14c, d). These GNPs undergo high contact stress under the wear counter face and forms a tribofilm which is shown as smooth region in Fig 4.14a. Weak interlayer forces between graphene layers
enable sliding or layer buckling to reduce the stress [99]. Hence, COF for Al-2 vol.% GNP is lower than pure Al.

![SEM images of the a) Wear Track and b) Wear Debris, c) Optical Microscope Image of the Sub-surface of the Wear Track, d) FIB Image of the Sub-surface of the Wear Track for Al-2 vol.% GNP after RT Tribological Test](image)

**4.4.3 Wear mechanism of Al-2 vol.% WS$_2$ at room temperature**

The addition of nanosize WS$_2$ particles resulted in homogeneous dispersion and a dense microstructure as observed in section 4.1. It is expected that uniform dispersion of nanosize WS$_2$ would facilitate the formation of a uniform tribofilm, which will promote the reduction of COF and improved wear resistance. Fig 4.15 presents the schematic of the wear mechanism of Al-2 vol.% WS$_2$ at RT. During the wear test, a chemical reaction occurs
between freshly exposed WS\textsubscript{2} nano-particles and surrounding humid air forming H\textsubscript{2}S gas with a foul odor.

![Diagram of Al-2 vol.% WS\textsubscript{2} at RT](image)

Fig 4.15. Schematic Representation of Behavior of Al-2 vol.% WS\textsubscript{2} at RT

Figure 4.16a shows wear track of Al-2 vol.% WS\textsubscript{2} which is rough in appearance and looks similar to that obtained of pure Al at room temperature. This is due to the nano-size of WS\textsubscript{2} particles, which allows these nano-particles to act as solid lubricants without altering the matrix’s properties. Two different debris sizes are found, one in which the average size of the wear particles is \( \sim 1 \mu \text{m} \) and for bigger particles the average size is around 20 \( \mu \text{m} \) (Fig 4.16b). All the wear debris is adhered to the counterball and in order to be collected these debris were peeled off from the Al\textsubscript{2}O\textsubscript{3} counterball. The tribofilm is formed by small clusters of WS\textsubscript{2} particles homogeneously distributed as shown in Fig 4.16d. A crack is seen in sub-surface which shows the delamination under shear contact stresses. The thickness of the tribofilm is \( \sim 7 \mu \text{m} \) and it is pointed out by the red arrow in Fig 4.16c. The same can be seen in Fig 4.16d, in addition this figure shows the cracks propagating all along the tribofilm similar to that behavior observed in pure Al at room temperature conditions.
Fig 4.16. SEM images of the a) Wear Track and b) Wear Debris, c) Optical Microscope Image of the Sub-surface of the Wear Track, d) FIB Image of the Sub-surface of the Wear track for Al-2 vol.% WS$_2$ after RT Tribological Test

It is summarized that at RT the addition of both GNP and WS$_2$ solid lubricants leads to reduction in COF during sliding wear. However, the distribution of GNPs particles is not as homogeneous as WS$_2$ nano-particles resulting in poor density and lowest wear resistance. The composite with WS$_2$ additives presents few wear debris with small size mostly located around the counter ball as mentioned before. This demonstrated that WS$_2$ nano-particles can effectively provide lubricious and anti-wear properties in aluminum creating a tribofilm and a transferfilm when a tribological test is performed at room temperature conditions.
4.5 Wear Mechanisms at 200°C

4.5.1. Wear mechanism of Al at 200°C

It is expected that high temperature would result in aggressive wear which is also reflected by wear rates shown in Table 4.1. Fig 4.17 presents schematic of wear mechanism for pure Al at high temperature. Due to higher temperature, oxygen diffusion would be faster resulting in larger amount of Al₂O₃ layer formation on the surface. The Al₂O₃ phase in tribofilm is confirmed by x-ray diffraction spectrum of the wear track shown in Fig 4.10.

Fig 4.17. Schematic Representation of Behavior of Pure Al at 200°C

Fig 4.18a shows a smooth wear surface which is due to slight softening of aluminum at 200°C. The wear debris presents some small particles but predominantly is composed by big particles with average size of ~100 µm, what refers to high surface stress with heavy deformation on the surface. The thickness of tribofilm is ~15 µm as it is pointed out in Fig 4.18c by the red arrow. The large thickness of the deformed film demonstrates high plastic deformation. No signal of grains or cracks are observed in higher magnification image of the sub-surface obtained from FIB (Fig 4.18d).
4.5.2. Wear mechanism of Al-2 vol.% GNP at 200°C

The tribological behavior for Al-2 vol.% GNP at 200°C is largely similar to that described at RT with an exception of more severe oxidation. Fig 4.19 presents the schematic representation of the behavior of GNP in the Al matrix at 200°C. It has been reported GNPs can act as sealers preventing oxygen diffusion through the grain boundaries [119]. At high temperature more oxidation is present on the surface which is also proven by EDS on the wear track (Appendix 1). Figure 4.20a shows the wear track of Al-2 vol.%
Fig 4.19. Schematic Representation of Behavior of Al-2vol.%GNP at HT.

GNP. A smooth surface with mixed layered morphology is observed. The wear debris presents a sheet-like form that works as a mechanism of stress dissipation, allowing

Fig 4.20. SEM Images of the a) Wear Track and b) Wear Debris, c) Optical Microscope Image of the Sub-surface of the Wear Track, d) FIB Image of the Sub-surface of the Wear Track for Al- 2vol.%GNP after HT Tribological Test.
themselves to slip upon the movement of the shear stress (Fig 4.20). The tribofilm is formed by Al₂O₃ and Al₄C₃ phases (Fig 4.10b) that are wrapped by GNP sheets. The thickness of the tribofilm is ~20 µm (Fig 4.20c) which is the largest observed in this study and it is pointed out by the red arrow. This figure reflects the ability of GNPs (which are the grey lines pointed out by the blue arrow) to act as sealers being located parallel to the sliding motion. This ability of acting as inhibitors of oxygen help to reduce wear as well as the fact of being compressed by the counterball reducing porosity, thus improving wear rate as compared with room temperature. Different from at RT, at HT no cracks are observed on the sub-surface (Fig 4.20d), and this can be attributed to the ability of GNPs to wrap that have reflected to contribute on reducing crack propagation [78].

4.5.3. Wear mechanism of Al-2 vol.% WS₂ at 200°C

At high temperature it is expected that Al-2 vol.% WS₂ would have higher wear rate than at RT due to accelerated oxidation and formation of WO₃. The results point out that this sample presented the lowest COF at 200°C. However, the wear rate is higher at 200°C than at RT, as expected. Appendix 1 shows low amount of oxygen on the wear

Fig 4.21. Schematic Representation of Behavior of Al-2vol.%WS₂ at HT.
surface and the XRD (Fig 4.10c) reflects the formation of WO₃. Fig 4.21 represents the mechanism developed during tribological test at HT. The worn surface obtained after HT test is observed in Fig 4.22a. This figure confirms the formation of the tribofilm to protect the surface. This tribofilm is smooth, however it is not homogeneously distributed on the surface. The wear debris got stuck on the countersurface (Al₂O₃ ball) which suggest that sliding occurred between the tribolayer and the transfer film on the counterface, resulting in a very stable and low COF.

From Fig 4.22b, it can be inferred a very dense wear debris, with a smooth surface and well distributed WS₂ particles is formed. The smooth and lamellar morphology of wear debris confirm that the detachment is due to delamination. Some amount of oxygen is found on the wear track. The tribofilm is mainly formed of WO₃ and WO₂ (Fig 4.10c). This tribofilm is the dense part and has been pointed out by the red arrows, it has a thickness of 5-10 μm as can be observed in Fig 4.22c and d. A FIB image in Fig 4.22d, shows the subsurface of the tribofilm, WS₂ nanoparticles can be seen as white spots with intact shape as it is observed as well on the wear debris images, reflecting the ability of WS₂ nanoparticles to transfer and cleaves to the counterface due to the weak bonding forces between sandwiches of Sulfur-Tungsten-Sulfur (S-W-S). The wear mechanism is schematically summarized in Fig. 4.23. Fig 4.10 depicts the phases formed for each sample after SPS, room temperature test and high temperature test.

The schematic representation of the wear mechanisms observed during this study for GNP and WS₂ are depicted in Fig 4.23 a and b respectively. When a GNP sheet experiences shear the weak van der Waals forces break apart and few layers of GNP tend
to wrap the deformed material forming sheet-like structures that facilitate shear sliding with one another. Also, as the counterball has contact with the surface, the GNP provides the surface with the ability to deflect. It must be noted that this is possible only in thin GNP and not thick graphite as ability to wrap and deflect is more difficult to be achieved (Fig 4.23a) when large number of layers are present. Another mechanism present in this sample, is the ability of the GNPs to act as inhibitor at 200°C [119], preventing oxygen to go through the sample as observed from Appendix 3. On the other hand, Fig 4.23 b describes the wear mechanism for WS₂. A triofilm is formed on the surface due to breaking of the weak van der Walls forces that hold the S-W-S structure. Due to shear forces, a portion of the triofilm is transferred on the counter surface. The wear occurs between transfer film and tribolayer providing lubrication properties and helping to reduce COF.
Fig 4.23. Schematic Representation of a) GNP and b) WS₂ Wear Mechanism.
CHAPTER V: CONCLUSIONS

This thesis presents a detailed investigation on using GNP and WS$_2$ based 2D materials to aluminum as solid lubricant additives for improved tribological properties. 2 vol.% 2D material was added to aluminum and consolidated as pellets by spark plasma sintering (SPS). Ball-on-disk wear tests in dry sliding conditions were carried out at room temperature and 200°C. The overall finding of this thesis suggests that WS$_2$ is an excellent 2D solid lubricant additive which not only reduces coefficient of friction (COF) but also improved the wear resistance of aluminum. GNP also provide a good alternative for reducing coefficient of friction, but show poor wear resistance. The overall finding of this thesis is suggested by following specific conclusions:

- Al-2 vol.% GNP composite displayed poor densification (91% dense) and low hardness due to agglomeration of GNP which resulted in insufficient sintering of spherical aluminum powder that lead to poor wear resistance.
- Al-2 vol.% WS$_2$ composite displayed excellent densification (99% dense) and high hardness as nano-size WS$_2$ particles filled the void between spherical aluminum particles resulting in full densification.
- At room temperature (RT), the wear rate of Al-2 vol.% WS$_2$ composite reduced by 54% as compared to pure aluminum. Al-2 vol.% WS$_2$ has the lowest COF (0.66) at room temperature whereas pure Aluminum has a highest COF of 0.87.
- At room temperature (RT), the wear rate of Al-2 vol.% GNP composite increased by 233% as compared to pure aluminum. The COF for Al-2 vol.% GNP at room temperature was 0.79.
At 200°C, the wear rate of Al-2 vol.% WS₂ composite *reduced* by 29% as compared to pure aluminum. Al-2 vol.% WS₂ has the lowest COF (0.55) at 200°C whereas pure Aluminum has a highest COF of 0.82.

At 200°C, the wear rate of Al-2 vol.% GNP composite *increased* by 48% as compared to pure aluminum. The COF for Al-2 vol. % GNP at room temperature was 0.69.

The wear mechanism in pure Al is adhesive in initial stages but becomes brittle due to formation of Al₂O₃ tribolayer at the surface.

The wear mechanism in Al-2 vol.% GNP involves formation of excessive wear debris due to poor densification. However, GPNs from the debris form a tribofilm on the surface resulting in lower COF as compared to pure Al. This is possible due to slippage of interlayers of GNP which makes the surface more slippery, and thus, lower friction. At 200°C, *GNPs act as inhibitors to oxygen diffusion* resulting in larger oxide content at the surface.

The wear mechanism of Al-2 vol.% WS₂ involves the formation of the protective transfer film. Thus the sliding occurs between the tribolayer and the transfer film located on the counterface, resulting in a very stable and low COF.
CHAPTER VI: FUTURE WORK

Fig 4.22. SEM Images of the a) Wear Track and b) Wear Debris, c) Optical Microscope Image of the Sub-surface of the Wear Track, d) FIB Image of the Sub-surface of the Wear Track for Al- 2vol.% WS₂ after HT Tribological Test.

The aim of the current research was to explore the potential of GNP and WS₂ as 2D solid lubricant additives to Aluminum alloys for reducing coefficient of friction (COF) and improving energy efficiency. The findings of this study establish WS₂ as an excellent option as a solid lubricant additive. However, some of the topics need further investigation to progress towards use of GNP and WS₂ as solid lubricant additives. Following is the list of recommendation for advancing the research in this thesis towards the final goal.
**Processing:** The density of Al-GNP composite needs to be improved to match that of Al-WS₂ composite. This can possibly be achieved by using smaller GNPs which will fill the voids of Al spherical particles similar to WS₂. However, delamination of graphene layers from a smaller GNP could be a challenge and may not provide sufficient reduction in COF. The packing density of starting powder blend of Al-GNP could also be improved by using irregular shaped aluminum powder mixed with both larger and smaller GNPs. This thesis employed 2 vol.% additive to aluminum matrix. Composition should be varied to evaluate the effect of concentration.

**Characterization:** TEM analysis of Al/GNP and Al/WS₂ should be carried out to better understand the nature of interfacial bonding as it will influence the wear behavior.

**Wear Behavior:** The wear tests in the present thesis were conducted at constant load and sliding distance at RT and 200°C. It is recommended that wear tests should be conducted at multiple loads and sliding distances to create a larger envelope of operating parameters to evaluate efficacy of GNP and WS₂ as solid lubricant additives. It will also enable a deeper understanding of wear mechanism such as tribofilm formation.
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APPENDICES

Appendix 1- EDS of Worn Surface at RT and HT

APPENDIX 1

AMOUNT OF OXGEN ON THE WEAR TRACK

<table>
<thead>
<tr>
<th>Sample Composition</th>
<th>RT (wt%)</th>
<th>HT (wt%)</th>
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</thead>
<tbody>
<tr>
<td>Pure Al</td>
<td>31.43</td>
<td>19.56</td>
</tr>
<tr>
<td>Al- 2 vol.% GNP</td>
<td>14.72</td>
<td>31.24</td>
</tr>
<tr>
<td>Al-2 vol.% WS₂</td>
<td>35.30</td>
<td>14.67</td>
</tr>
</tbody>
</table>

### EDS of the worn surface at RT

**Pure Al**

- **EDS of Pure Al**

**Al- 2 vol.% GNP**

- **EDS of Al-2 vol.% GNP**

### EDS of the worn surface at HT

- **EDS of Pure Al**

- **EDS of Al-2 vol.% GNP**
Al-2 vol.% WS₂
Appendix 2- TEM and SAED of Wear Debris

TEM and SAED of wear debris of pure Al after RT test.

TEM and SAED of wear debris of Al-2vol.%GNP after RT
TEM and SAED of wear debris of Al-2vol.%WS₂ after RT test.

TEM and SAED of wear debris of pure Al after HT test.
TEM and SAED of wear debris of Al-2vol.%GNP after HT test.

TEM and SAED of wear debris of Al-2vol.%WS₂ after HT test.
Appendix 3- EDS to Study Oxidation Behavior of GNP on the Tribofilm

Cross-section of the Al-2vol.% GNP wear track at room temperature:
Cross-section of the Al-2vol.% GNP wear track at high temperature:
Oxygen content through the sample (cross section of the tribofilm)

<table>
<thead>
<tr>
<th>From 0µm (surface) to 25 µm depth</th>
<th>Room Temperature (wt%)</th>
<th>High Temperature (200°C) (wt%)</th>
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</thead>
<tbody>
<tr>
<td>0</td>
<td>12.86</td>
<td>12.20</td>
</tr>
<tr>
<td>1-2</td>
<td>16.01</td>
<td>7.13</td>
</tr>
<tr>
<td>3</td>
<td>27.95</td>
<td>13.85</td>
</tr>
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</tr>
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