Review Article

Sara I. Ahmad*, Hicham Hamoudi, Ahmed Abdala, Zafar K. Ghouri, and Khaled M. Youssef

Graphene-Reinforced Bulk Metal Matrix Composites: Synthesis, Microstructure, and Properties

https://doi.org/10.1515/rams-2020-0007
Received Aug 12, 2019; accepted Dec 12, 2019

Abstract: This paper provides a critical review on the current status of graphene-reinforced metal matrix composites (GRMMCs) in an effort to guide future work on this topic. Metal matrix composites are preferred over other types of composites for their ability to meet engineering and structural demands. Graphene is considered an ideal reinforcement material for composites due to its unique structure and extraordinary physical, thermal, and electrical properties. Incorporating graphene as a reinforcement in metals is a way of harnessing its extraordinary properties, resulting in an enhanced metallic behavior for a wide variety of applications. Combining graphene with bulk metal matrices is a recent endeavor that has proven to have merit. A systematic study is needed to critically examine the efforts applied in this field, the successes achieved, and the challenges faced. This review highlights the three main pillars of GRMMCs: synthesis, structure, and properties. First, it discusses the synthesis techniques utilized for the fabrication of GRMMCs. Then, it highlights the resulting microstructures of the composites, including graphene dispersion and interfacial interactions. Finally, it summarizes the enhancements in the mechanical, electrical, thermal, and tribological properties of GRMMCs, while highlighting the effects of graphene type and content on those enhancements.

Keywords: graphene, metals, composites, interface, properties

Table of Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| 2D           | two dimensional |
| 3D           | three dimensional |
| ARB          | accumulative roll bonding process |
| at%          | atomic % |
| BM           | ball milling |
| CMC          | ceramic matrix composites |
| CNT          | carbon nanotube |
| COF          | coefficient of friction |
| CP           | cold pressing |
| CS           | compressive strength |
| CTAB         | cetrimonium bromide |
| CVD          | chemical vapor deposition |
| CYS          | compressive yield strength |
| CTE          | coefficient of thermal expansion |
| DMD          | disintegrated melt deposition method |
| DI water     | deionized water |
| EC           | ethyl cellulose |
| FLG          | few layer graphene |
| FS           | flexural strength |
| FSP          | friction stir processing |
| GNF          | graphene nanoflake |
| GNP          | graphene nanoplate |
| GNR          | graphene nanoribbon |
| GNS          | graphene nanosheet |
| GO           | graphene oxide |
| GQD          | graphene quantum dot |
| Gr           | graphene |
| GRMMC        | graphene-reinforced metal matrix composite |
| HP           | hot pressing |
| HPS          | hot press sintering |
| HRDSR        | high-ratio differential speed rolling |
| HRTEM        | High-resolution transmission electron microscopy |
| HV           | Vickers hardness |
| IACS         | international annealed copper standard |
| IPA          | isopropyl alcohol |
| LS           | laser sintering |
| MLG          | multilayer graphene |

*Corresponding Author: Sara I. Ahmad: Department of Sustainable Development, Hamad Bin Khalifa University, Doha 5825, Qatar; Email: sahmed@mail.hbku.edu.qa; Tel.: +974 66633557
Hicham Hamoudi: Qatar Environment and Energy Research Institute, Doha 34110, Qatar
Ahmed Abdala, Zafar K. Ghouri: Department of Chemical Engineering, Texas A&M University, Doha 23874, Qatar
Khaled M. Youssef: Department of Materials Science and Technology, Qatar University, Doha 2713, Qatar

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MLM molecular level mixing  
MM mechanical mixing  
MMC metal matrix composite  
MO metal oxide  
MS magnetic stirring  
PASE particle-assisted shear exfoliation  
PCA process control agent  
PG pristine graphene  
PM planetary milling  
PMC polymer matrix composite  
PMMA Poly (methyl methacrylate)  
PVA polyvinyl alcohol  
PVP polyvinyl pyrrolidone  
RGO reduced graphene oxide  
SAED selected area electron diffraction  
SEM scanning electron microscopy  
SLM selective area electron diffraction  
SPS spark plasma sintering  
TC thermal conductivity  
TEM transmission electron microscopy  
TYS tensile yield strength  
UTS ultimate tensile strength  
UHP uniaxial hot pressing  
VHP vacuum hot pressing  
vol% volume %  
wt% weight %  
XRD x-ray diffraction  
YM Young’s modulus

1 Introduction

Graphene, known as “the wonder material of today’s scientific world,” is not only the strongest material to be discovered to date but also is the thinnest material [1, 2]. The two-dimensional (2D) structure of graphene is defined by its one-atom thick, 2D sp² hybridized carbon sheet consisting of hexagonal lattices with a height of 0.33 nm [3, 4]. For many decades, graphene has been identified by the scientific world as the basic structural unit of bulk graphite. Theoretically, it had been regarded as unstable to exist physically; hence, it attained no specific interest as an individual entity. However, the pioneering work of Geim and coworkers [5] in 2004, which led to the separation of single layers of graphite, led to rapidly growing attention to graphene, which would later be known as the wonder material. It has been proven to have exceptional properties as an isolated entity, such as excellent thermal and electrical conductivities of 5×10³ W/m·K [6] and 6×10³ S/m [7], respectively. It is harder than a diamond and 100 times stronger than steel, with a tensile strength and elastic modulus of 130 GPa and 1.2 TPa, respectively [8]. Yet it is flexible and can take any form or shape [9]. Moreover, it is capable of absorbing 2.3% of light [10], making it the first and only one-atom-thick material visible to the naked eye. Those exceptional properties elevated graphene from being the building unit of graphite to an independent material with countless possibilities to be used in a wide range of applied technologies. Applications of graphene include electronics, energy storage, sensors, biomedical devices, and as the newest member in bulk composites for structural applications [11–16].

Harnessing the extraordinary properties of graphene can be accomplished through the incorporation and dispersion of graphene as a reinforcement in bulk composites such as polymers, ceramics, or metal composites [17, 18]. Composites are advanced materials and are considered the most promising class of materials in the modern applied science world. Combining different materials to provide unique properties is highly attractive for all sorts of applications [19, 20]. Composite materials can be classified into metal matrix composites (MMCs), ceramic matrix composites (CMCs), and polymer matrix composites (PMCs). More research has been conducted on CMCs and PMCs than MMCs. This is mainly due to the simpler fabrication routes of such composites, which does not involve high temperature and pressure requirements. Yet MMCs have higher abilities to meet engineering demands and are the most widely used in manufacturing industries such as aerospace, automotive, and construction [21]. Even in small concentrations, the interaction of graphene as a reinforcement within the matrix is facilitated by its nanometric nature [18] and its large surface area of 2626 m²/g [22]. The 2D structure of graphene provides a higher surface area than other carbon-based materials, thereby offering a larger area to interact with the matrix for enhanced mechanical stress and electrons or phonons transfer [1, 23]. In addition, its large specific area eliminates the need to use large fractions of the reinforcements, reducing the tendency for agglomeration within the matrix [24, 25], and it can also act as a barrier to dislocation motion, resulting in enhanced strengthening of materials [26]. Another feature that makes graphene suitable as a reinforcement is its flat morphology, which is more susceptible to withstand the high-pressure processing routes involved in the fabrication of composites [1]. Its lower production cost than carbon nanotubes (CNTs) and its availability in high-quality and bulk quantities adds to its potential as a reinforcement material [1, 27].

The alteration in the microstructure of metallic materials after adding nanoscale reinforcements allows for tai-
loring of the properties of those materials, which makes it highly suitable for structural and functional applications [20]. However, compared with graphene-based PMCs [22, 28–30] and CMCs [31–33], the field of graphene-based MMCs has been little researched. In addition, the majority of the existing literature in this field has focused on the deposition of metallic nanoparticles, metallic oxides, and metallic carbides on the surface of graphene and its derivatives to explore its applications in energy storage, sensory applications, and photocatalytic systems [34–43]. Increasing interest has been reported in using graphene and its derivatives as a reinforcement in bulk MMCs, yet the field is still at its early stages [44]. This can be attributed to many challenges in fabricating MMCs reinforced with graphene, such as the dispersion of the reinforcement in the matrix and the undesired interfacial reaction due to the metals’ high reactivity [45]. Even with limited number of research studies, graphene has been proven to fulfill the requirements of an excellent nanofiller in MMCs, leading to the production of superior materials that are lightweight [24], demonstrate high strength [46, 47], have high ductility [48, 49], and have enhanced thermal [50, 51] and electrical conductivity [52, 53].

Graphene can be used to describe a graphene-based family of materials that have been produced with varying structures. In this paper, those materials will be referred to as graphene “Gr” unless specified otherwise and include graphene nanoplates (GNPs), graphene oxide (GO), reduced graphene oxide (rGO), graphene quantum dots (GQDs), and graphene nanoribbons (GNRs). It is important to note that unless single-layered graphene is mentioned, the abbreviation “Gr” is used often in this paper to represent all the above-mentioned graphene derivatives for simplicity. It is important to note that per the author’s knowledge, no studies so far have reported the reinforcement of GQDs in MMCs. In addition, only one paper reported the fabrication of GNRs in MMCs [52]. The main focus of this field thus far had been on developing GNPs [18, 23–27, 54–57] and GO-reinforced MMCs [49, 58–73].

Figure 1 shows the increased attention toward utilizing Gr as a reinforcement material in bulk MMCs. According to Web of Science and Science Direct databases, the number of publications investigating Gr-reinforced MMCs per year has increased from 2010 to 2018. This increasing trend shows the increased scientific research interest in studying the effect of Gr as a reinforcement material in MMCs. Among the reported studies of MMCs, Gr-reinforced Al, Cu, and Mg matrix composites have gained the most interest.

Research on GRMMCs has developed in a short period of time and is still at its beginning. A systematic study on the efforts applied in this field and the successes and challenges is required. The aim of this review paper is to provide a deeper insight into the status of the GRMMC research field and to serve as a comprehensive reference to guide future work on GRMMCs. It provides a full review on the work that has been performed in this field while focusing on the most studied metallic systems (Al, Cu, Mg). The review starts with a brief background introducing Gr and its derivatives. Then, it sheds light on the various processing techniques for GRMMCs, the microstructures of GRMMC and how it is affected by the dispersion and distribution of Gr in metal matrices, as well as the interfacial interactions between Gr and metals. Finally, the review discusses the effect of Gr addition on the mechanical, thermal, electrical, and surface properties of GRMMC.

2 Graphene Properties, Derivatives, and Synthesis

The research boom on Gr is directly related to its birth in 2004 when Geim et al. [5] successfully isolated it from graphite. It was regarded for the first time as a thermally stable entity regardless of its atomic thin structure [22, 45]. However, the history of Gr dates back to as early as the 19th century. A timeline of the history of Gr is presented in Figure 2. A more detailed historical background on Gr research can be found in Geim’s 2012 review paper [74].

The fabrication of monolayer Gr requires overcoming the Van der Waals forces connecting the Gr layers in graphite. Exfoliating high-quality and large quantities of monolayers of Gr is still a challenging process. In contrast, attaining graphene nanoplatelets (GNPs), which consist of a few (10–40) Gr layers [1], is easier and faster
Figure 2: Timeline of the most important milestones in graphene history.

than exfoliating monolayer Gr [55, 75]. The thin structure of GNPs can retain the structural integrity and outstanding properties of monolayer Gr, such as its high surface area and excellent mechanical, electrical, and thermal properties [1]. In the literature, a variety of names are used to describe few layers Gr (FLGs) or GNPs such as graphene nanosheets (GNSs), graphene platelets, and graphene nanoflakes (GNFs).

Just like graphite, which consists of stacked layers of Gr connected via Van der Waals forces, graphite oxide consists of stacked layers of GO. In graphite oxide, the stacked GO sheets are functionalized with O, OH, or COOH groups. The rich presence of functional groups increases the interlayer spacing between the sheets to 6–10 Å, compared with 3 Å for unfunctionalized graphite [22, 70]. Graphite oxide was first prepared in the 19th century by Brodie [76] by treating graphite with strong materials such as potassium chlorate and nitric acid. However, graphite oxide possesses different properties than Gr. Today, a safer synthesis technique of graphite oxide is favored in laboratories, as reported by Hummers and Offeman [77]. It involves the mixing of graphite with sodium nitrate, sulfuric acid, and potassium permanganate. After the discovery of Gr in 2004, GO has been used in research as the starting material for large scale production of GNP through exfoliation of graphite oxide into GO and the reduction of GO into rGO [22]. The functional groups in GO allow for its dispersibility in water and other solvents. This produces monodispersed GNPs via chemical or thermal reduction of GO in colloidal suspensions [78, 79]. Unfortunately, an inversely proportional relationship exists between the physical properties of Gr and dispersibility [80]. The reduction of GO is an important process in order for the material to possess the outstanding properties of Gr, especially electrical properties, as GO is electrically nonconductive [22, 78].

GNRs are thin, elongated Gr sheets with nanometric width around 10–20 nm and a high aspect ratio [81]. GNR materials have attracted the attention of many research studies due to their capability of generating a band gap, which normal Gr sheets lack. This unique characteristic of GNRs is attractive for optoelectronic applications. However, controlling the quality and the type of the edge is the main challenge in utilizing Gr in electronic devices [81, 82]. GNRs have been produced by a variety of methods, such as lithography [83], chemical vapor deposition (CVD) [84], and unzipping CNTs by the plasma etching technique [85]. Figure 3 presents a collection of transmission electron microscopy (TEM) images of monolayer Gr, GNPs, GO, and GNRs.

The first successful fabrication method of Gr was performed in 2004 [5] using the micromechanical cleavage method. In this method, a scotch tape was used to exfoliate a single layer Gr from graphite. Several Gr synthesis processes have been developed, any of which can be categorized as a top-down or a bottom-up technique. Top-down techniques involve the isolation of Gr layers from graphite or graphite derivatives to produce single or multilayer Gr sheets. Graphene samples produced using top-down methods are usually of small sizes and large quantities. Bottom-up techniques involve building the Gr sheet atom by atom using specific substrates and carbon-containing compounds. Graphene samples provided by
such techniques are usually of small quantity, high quality, and large size [3, 28, 86].

A detailed explanation of all techniques is out of the scope of this work. Therefore, related references are included within the text. Top-down techniques have been used mainly in the fabrication of Gr for bulk GRMMCs [64, 67]. Only a few studies have used in-situ bottom-up Gr growing techniques [88–90]. Recent reviews on the synthesis of Gr and its derivatives can be found in References [91, 92]. Tables 1 and 2 summarize the information on commercially available Gr used in certain GRMMC studies and the main techniques adopted for the synthesis of Gr and its derivatives in other GRMMC studies, respectively.

3 Graphene-Reinforced Metal Matrix Composites (GRMMCs)

3.1 Synthesis Techniques

The synthesis of GRMMCs has faced several obstacles that challenge the successful fabrication of such composites. The first challenge is the poor affinity of Gr toward metals, suggesting poor resultant interfacial bonding. The second challenge is the agglomeration of Gr sheets within the metal due to the Van der Waals forces between Gr sheets [97] and its high surface energy [1], which restricts the homogenous distribution of Gr in the metal matrices. Thus, attaining high-quality GRMMCs is directly related to the synthesis technique of choice. Several authors have argued that the usual powder methodology techniques do not fully solve the dispersion problem of Gr in metal matrices, thus not capturing Gr’s full potential and resulting

Figure 3: TEM images of (a) a graphene sheet, (b) GO, (c) GNR, and (d) GNP. Reproduced from References [52, 86–88], respectively.
Table 1: Summary of Commercial Gr and its Derivatives Used in GRMMC Studies.

| Product                  | Company                                               | Size (µm)/Surface Area (m²/g) | Thickness (nm) | Reference(s)     |
|--------------------------|-------------------------------------------------------|------------------------------|----------------|------------------|
| GNP, GNS, FLG, and multilayer graphene (MLG) | Tanfeng Tech Co. Ltd                                 | 5–30                         | 5–10           | [93]             |
|                          | Skyspring Nanomaterials                                | —                            | 5–10           | [47]             |
|                          | Qingdao Huagao Graphene Technology Corp. Ltd. (China) | —                            | 10 layers      | [94]             |
|                          | Sixth Element Materials Technology Co. Ltd. (China)   | 14.17 SA                     | 20 layers      | [87]             |
|                          | Ningbo Morsh Technology Co., Ltd., China.             | 5–15                         | 2–10           | [2, 71, 95–97]   |
|                          | US Nano Company, (USA)                                | 2                            | 20             | [98]             |
|                          | XG Sciences, (USA)                                    | 1–15                         | 1–20           | [23, 24, 54, 99–102] |
|                          | Angstron Materials                                    | <14                          | 10–100         | [63, 103]        |
|                          | Nanjing Xian Feng Nano Material Technology Co. Ltd (Jiangsu, China) | 5–25                         | 5–20           | [11, 17, 55, 104–107] |
|                          | Nanografi Co. Ltd.                                    | 750 SA                       | 5–8            | [108]            |
|                          | American Elements, USA                                | 5–25                         | 5–25           | [109]            |
|                          | XG Science, South Korea                               | —                            | 6–8            | [46]             |
|                          | Chengdu Organic Chemistry Co. Ltd.                    | 5–10                         | 4–20           | [48, 110, 111]   |
|                          | Cheap Tubes, Inc.                                     | <2                           | —              | [112]            |
| GO                       | Graphenea (Gipuzkoa, Spain)                           | —                            | >95% monolayer |                  |
|                          | Institute of Coal Chemistry, Chinese Academy of Sciences | 0.3–0.8                     | 0.7–1.2        | [60]             |
|                          | Nanjing Xian Feng Nano Material Technology Co. Ltd (Jiangsu, China) | —                            | —              | [61]             |
|                          | Nanomaterials LS                                      | <0.1                         | 2–4            | [63]             |

Table 2: Summary of Synthesized Gr and its Derivatives in GRMMC Studies.

| Product                  | Synthesis                                               | Reference(s)     |
|--------------------------|---------------------------------------------------------|------------------|
| GO                       | Modified Hummers' method                               | [49, 64–72, 113] |
| GNP, GNF, MLG, FLG, and GNS | Reduction of GO                                        | [49, 57, 113–118] |
|                          | Exfoliation of graphite                                | [18, 56, 119–121] |
|                          | In-situ growing Gr in bulk metal matrix                | [89]             |
|                          | In-situ growing 3D Gr by CVD using NaCl template       | [90]             |
|                          | Ex-situ growing of Gr sheets by CVD                    | [122]            |
| GNR                      | Facile chemical unzipping of multiwalled CNTs          | [52]             |

in a direct negative impact on the fabricated Gr-metal composite properties [123]. Researchers are striving to discover and develop more advanced techniques for the fabrication of GRMMCs, aiming to produce higher-quality composites by achieving better Gr distribution within metal matrices. Table 3 summarizes the synthesis techniques used for the fabrication of GRMMCs. In this review, the synthesis section is divided into three main sections according to the steps involved in the synthesis technique: pretreatment, synthesis, and postsynthesis and consolidation.
Table 3: Summary of Synthesis Techniques Used for the Fabrication of GRMMC.

| Synthesis Step         | Techniques                          | Reference(s)                                                                 |
|------------------------|-------------------------------------|-------------------------------------------------------------------------------|
| Pretreatment           | Gr pretreatment                     | Mechanical mixing / sonication in a solvent [11, 17, 24–27, 44, 49, 51, 52, 54, 58, 59, 64, 65, 68, 69, 71, 73, 93, 96, 99, 104, 107, 108, 115, 118, 120, 123–126, 128–131] |
|                        | Surface modification                 | [69, 71, 90, 96, 113, 127]                                                   |
| Metal pretreatment     | Mechanical mixing / sonication in a solvent [11, 17, 52, 58, 59, 65, 104, 107, 116, 129, 130] |
|                        | Surface modification                 | [51, 64, 66, 68, 71, 73]                                                     |
|                        | Ball milling                         | [88, 128, 129]                                                               |
| Mixing Techniques      | Ball milling                         | [18, 20, 23, 25, 27, 46, 48–50, 53–57, 63, 65, 70, 95, 99, 101, 109, 114–116, 118, 119, 131–133] |
|                        | Solution-assisted mixing             | Sonication in a solvent [24, 44, 50, 108, 115, 131]                          |
|                        | Mechanical mixing                    | [11, 17, 57, 58, 66, 68, 71, 73, 89, 93, 104–108, 117, 120, 125, 127, 130] |
|                        | Magnetic stirring                    | [51, 52, 58, 59, 69, 124, 126]                                               |
|                        | Molecular-level mixing               | [67, 72, 96, 97, 134, 135]                                                   |
|                        | High shear mixing                    | [121, 135, 136]                                                             |
|                        | Pestle and mortar                    | [137]                                                                       |
|                        | Vortex mixing                        | [93, 128]                                                                    |
|                        | Molten/liquid metal processes        | [87, 94, 98, 103, 110, 111, 120, 138–141]                                    |
|                        | Other techniques                     | In-situ synthesis [88–90, 122, 140, 142–144]                                |
|                        | Rolling processes                    | [100, 123, 145]                                                             |
| Postsynthesis/consolidation | Cold pressing followed by hot treatment | [11, 17, 18, 20, 26, 27, 44, 50, 58, 59, 65, 66, 73, 95, 104, 105, 107, 124, 125, 129, 130, 132, 137] |
|                        | Hot pressing                         | [23, 49, 51, 54, 55, 63, 64, 68, 70, 88, 99, 101, 108, 118–120, 126, 128]    |
|                        | Hot extrusion                        | [11, 17, 48, 50, 55, 57, 73, 95, 104, 105, 107, 110, 111, 118, 130]         |
|                        | Spark plasma sintering (SPS)         | [24, 52, 69, 71, 72, 93, 96, 97, 109, 127, 134, 135]                         |
|                        | Hot rolling                          | [25, 52, 56]                                                                |
|                        | 3D printing                          | [60, 112, 133]                                                              |

3.1.1 Pretreatment of Graphene Reinforcement and Metal Matrix

Before mixing the composite components to fabricate the GRMMCs, several researchers treated the Gr and the metals separately for several reasons, such as to further exfoliate the Gr sheets, to prevent Gr agglomeration and enhance its dispersion in the metal matrix, or to enhance the attraction and attachment between the Gr and the metal matrix. Thus far, the main pretreatment processes that have been reported are either solvent dispersion of the Gr reinforcement [11, 17, 24–27, 44, 49, 51, 52, 54, 58, 68, 69, 71, 73, 99, 118, 123–126] and/or the metal matrix [11, 17, 52, 58], and the surface modification of the Gr reinforcement [69, 71, 90, 96, 127] and/or the metal matrix [51, 66, 68, 71, 73]. Certain mixing techniques have
been associated with certain pretreatment steps. For example, mechanical mixing or sonication of metals or Gr in solvents is a step mainly used before solution-assisted mixing techniques or ball milling. Pretreatment steps involving surface modifications for metals or Gr have been reported mainly in studies utilizing solution-assisted mixing. Even ball milling has been used as a pretreatment step in some studies [88, 128, 129]. One study reported ball milling of Cu and poly (methyl methacrylate) (PMMA) as a carbon source before in-situ fabricating Cu-Gr composite via CVD [88]. Others used milling to modify the morphology of their metal, such as changing it from spherical to flake-like morphology [128, 129]. However, other studies did not report a pretreatment step because the mixing technique used did not require one or because the authors simply did not opt for one.

3.1.1.1 Graphene Pretreatment
Maintaining the structural integrity of Gr (that is, its high surface area and nanometric structure) is important to extrapolate its unique properties in metal matrices. Several researchers have reported the dispersion of Gr in different solvents in an aim to further exfoliate Gr sheets [68] and to enhance their dispersion and prevent their agglomeration in the metal matrix during the mixing step [58]. Different solvents have been used, such as water [51, 58, 68, 69, 71, 73, 125], ethyl alcohol [25, 118, 124], ethanol [11, 27, 49, 52, 54, 99], and acetone [17, 24, 26, 44, 123, 126], in order to disperse Gr using either ultrasonication, mechanical, or magnetic stirring. Bisht et al. [24] ultrasonicated GNPs in acetone before mixing it with Al matrix and consolidating it by SPS. Figures 4a and 4b show the scanning electron microscopy (SEM) and TEM images of the GNP before and after ultrasonication, respectively. The transparency of the GNPs shown in the TEM image suggested good reduction in the number of GNP layers and exfoliation during ultrasonication. In addition, ultrasonication the GNPs has led to the formation of folds and wrinkles in the GNP sheets, which resulted in limited plastic deformation of the composite due to the mechanical interlocking of GNPs in the Al matrix provided by those features.

The nature of the Gr starting material directly affects the synthesis technique of choice and is critical to their interactions with the matrix material. Several researchers have preferred to start with GO as a precursor [70, 125] in order to avoid the agglomeration of Gr reinforcement. This is attributed to the hydrophilic functional groups in GO, which can successfully suspend GO in aqueous solutions and form chemical bonds with surface-modified metal particles [51, 66, 68, 73]. However, the reduction of GO is hard to control and is often incomplete, which introduces structural defects into the graphitic structure [127]. To avoid the need to reduce GO and eliminate any related structural damages while attaining good affinity of Gr toward metals, pretreating Gr by surface modification has been reported [69, 71, 90, 96, 113, 127]. Zhang et al. [127] performed electroless plating of GNP with Cu and Ni before mechanically stirring the modified GNPs with Cu powders and finally sintering the mixed powder composite via SPS. Electroless plating includes sensitizing and activating GNP by ultrasonication in SnCl$_2$ and PdCl$_2$ solutions followed by washing and filtration before the electroless plating process takes place. At the end of the electroless plating, Ni and Cu were, respectively, adsorbed on the surface of GNPs.
Several studies [69, 90, 113] have reported the surface modification and in-situ synthesis of GNS. Tang et al. [113] produced Ni-decorated GNSs using an in-situ chemical reduction method. In their method, negatively charged GO is first mixed with Ni salts where Ni ions aggregate and nucleate at the functional group’s sites of GO. The mixture then is treated with a reducing agent where nickel ions are reduced in situ to Ni nanoparticles, while the GO are simultaneously reduced to GNS. Finally, this Ni-decorated GNS was sonicated with Cu powders and consolidated via SPS. A schematic diagram of the in-situ chemical reduction steps is shown in Figure 5. A similar procedure was adopted by Ju et al. [69]. They used Mg ions as an anchor binding the GNS to Al. The GO sheets were mixed with the Mg salts and then chemically reduced to GNSs using ascorbic acid. The resultant Mg-decorated GNSs were mixed with Al and consolidated via SPS.

More recently, Fu et al. [90] synthesized 3D Ni-decorated GNSs by an in-situ high-temperature CVD technique using NaCl crystals, C₆H₁₂O₆, and Ni(NO₃)₂ as a template, a carbon source, and Ni precursor, respectively. This was followed by an impregnation-reduction process to produce Ni-decorated GNS-Ni composite powders. A schematic diagram of the process is shown in Figure 6. The final composite exhibited a 188% and a 26% increase in the tensile yield strength (TYS) and the ultimate tensile strength (UTS), respectively.
strength (UTS), respectively. This result was attributed to the homogeneous distribution of GNS in the Ni matrix and the strong interfacial bonding.

3.1.1.2 Metal Pretreatment
Aside from dispersing the Gr-based reinforcement, several researchers reported the dispersion of the metal-based matrix powders in a similar manner. This step was then followed by mixing both reinforcement suspension and the metal slurry for enhancing the uniform dispersion of Gr reinforcement in the metal matrix. The solvents used for metal dispersion include acetone [17, 58] and ethanol [11, 52]. Yang et al. [52] used ultrasonication and magnetic stirring to disperse GNRs and Cu separately before mixing both solutions. Liu et al. [58] mechanically stirred Al in acetone before adding the ultrasonicated GO water dispersion. Rashad et al. [11] ultrasonicated and mechanically mixed GNPs and Al in ethanol, respectively, before mixing the ultrasonicated GNPs and the Al slurry via mechanical mixing.

Several researchers reported the surface modification of the metal matrices ahead of mixing the Gr reinforcement with the metal matrix to enhance the interaction affinity and binding capacity between Gr and metals [51, 64, 66, 68, 71, 73]. Gao et al. modified the surface charge of Cu and Al in separate studies ([51, 64] and [68], respectively) using the same technique to enhance the Gr-metal affinity via charge attraction. They mechanically stirred the negatively charged GO with the positively charged cationic surface agent (CTAB)-coated Cu and Al. In general, GO adheres to a negative surface charge due to its attached functional groups such as epoxide and hydroxyl groups on the basal plane and carboxyl groups at the edges. In contrast, CTAB is cationic in nature, which gives Cu and Al powders a positive charge after they are coated with it. As a result of charge attraction, the GO-Cu and GO-Al composite powders achieved uniform adsorption, and the process is referred to as electrostatic self-assembly. Figure 7 is a schematic diagram of the synthesis process of GO-reinforced CTAB-coated Cu. This process was followed by cold compaction of composite powders and sintering at high temperatures to thermally reduce GO. The authors suggested that this novel process eliminates the agglomeration of the GO sheets and preserves its large size as they uniformly adsorb on the surface of the metal.

Some researchers reported the surface modification of metal matrices by the use of polyvinyl alcohol (PVA) [66, 71, 73]. It is a water-soluble polymer used to introduce a hydrophilic membrane to metals’ surfaces. Wang et al. [73] synthesized the GNS-Al composite by treating Al flakes by PVA aqueous solution before mechanically mixing it with GO aqueous dispersion dropwise, filtering and rinsing the GO-Al composite powders, and heating the GO-Al composite powders to decompose the PVA and reduce the GO nanosheets to GNSs. The same procedure was followed 5 years later by Ponraj et al. [66] using a Cu matrix instead of Al. Similarly, Jiang et al. [71] modified Cu powders with
PVA to produce Gr-Cu composites, but they used Gr as precursor instead of GO. Because Gr lacks the hydrophilicity of GO and hence has no affinity toward the OH functional groups in PVA, Gr was treated with polyvinyl pyrrolidone (PVP) before mixing it with PVA-modified Cu. The two polymers were used to enhance the interactions between Gr reinforcement and the Cu matrix. Jiang et al. [71] explained that the PVP would interact with Gr via π-π interactions, while the carbonyl group in PVP would interact with the hydroxyl groups in PVA. Because Gr has lower defects and higher structural integrity than GO, it is expected to perform better as a reinforcement.

3.1.2 Mixing Techniques

Mixing the composite components is done using a variety of techniques and is considered the main step in the synthesis of GRMMCs. No systematic studies have compared mixing techniques and their effects on the structure and behavior of GRMMCs. Thus, it is impossible to categorize the mixing techniques based on effectiveness.

3.1.2.1 Mechanical Milling

Mechanical milling, which is also called ball milling or mechanical alloying, is a top-down synthesis technique used to produce materials with nanostructures. This is done by disintegrating the coarse-grained structure of those materials’ elemental powders [146, 147]. In mechanical milling, which is represented in Figure 8, selected powders are sealed in a steel vial along with a number of steel balls of certain sizes under an inert atmosphere, typically argon. The high energy rotation of the balls inside the vial causes constant collision of the balls, which leads to crushing the powders trapped in between with a high impact force. The repeated mechanical deformation results in the formation of cell/subgrain structures with high dislocation density. These cellular structures transfer into high-angle grain boundaries with increasing milling time, forming nanocrystals [146, 147].

Mechanical milling has become a widely chosen technique for the mixing of Gr-reinforced MMCs due to its simplicity and scalability in Gr-reinforced Al [18, 20, 23, 25, 27, 50, 54–57, 70, 99, 118, 119, 132], Cu [49, 63], or Mg [48] matrix composites. In most cases where milling was used for the mixing of Gr and metals, a process control agent (PCA) is used to adsorb onto the surface of powders, minimize cold welding, and prevent particle agglomeration. Nonetheless, the structural integrity of Gr in MMCs prepared via milling is a point of debate. Some researchers have suggested that the structural integrity of Gr is damaged during mechanical milling due to its large surface area, which cannot withstand the high impact force from milling [51]. In addition, it was suggested that milling can cause interfacial reactions due to heat generation during milling [68, 104]. In most studies, the structural integrity of Gr in GRMMCs had been characterized using Raman spectra [18, 48, 49, 53, 55, 56, 70]. Ranges between 1333–1359, 1583–1603, and 2665–2727 cm⁻¹ have been reported for typical D, G, and 2D bands, respectively, for GRMMCs synthesized via milling. Generally, the G and D bands are the most important for the evaluation of Gr. The \( I_D/I_G \) ratio represents the integrity of the Gr structure. It was reported that this ratio increases after milling, suggesting an increase in the defect-presenting band [55, 56, 70]. Li et al. [53]...
used milling to disperse 0.2 wt% GNP with Al to form what they called the “master alloy” as the GNP carrier, before adding it to Al melts to form the composite through casting and rolling. The $I_D/I_G$ ratio from the Raman spectrum for GNP before and after milling of the master alloy and in the Al-0.2 wt% GNP composites was 0.13, 1.10, and 1.14, respectively. The ratio increases significantly after milling of GNP, yet it remains almost the same with a slight increase after high-temperature melting, casting, and rolling. The result suggests that most of the structural damage occurred in the milling process.

The intrinsic structure damage of Gr and its derivatives may be unavoidable during milling, yet several researchers have reported a reduction in the damage degree by controlling the milling time [18, 49, 95] and the milling media size [55]. The ball-milling time is a main factor that greatly affects the milling synthesis and, hence, the structural integrity and dispersibility of Gr [49, 95]. Pérez-Bustamante et al. [18] mixed GNSs and Al powders via milling in methanol as a PCA to avoid agglomeration. They reported a reduction in the peak intensity of Raman spectra as the milling time increased. This finding was justified by the introduction of structural disorders and increase in amorphous fraction of the GNP with increased milling time. However, crystallinity of GNP was still preserved even after 5 h of high-energy milling, which was attributed to the protection provided by the ductile Al matrix against constant collision on the well-dispersed GNP. Similar results were reported by Yue et al. [49], where the Raman spectra ratio of $I_D/I_G$ increased from 0.84 to 1.42 with increasing the ball-milling time. The authors attributed this behavior to the large accumulated milling energy that results in breaking the weaker chemical energy of the C-C bonds.

Jiang et al. [95] reported that implementing an optimum milling time can eliminate the need for a tradeoff between achieving a uniform dispersion of Gr in metal matrices and the structural damage of Gr nanosheets, as well as the dilemma of achieving high strength at the expense of poor ductility. A shift-speed ball milling process, which consisted of various hours of low-speed milling and 0.5 h high-speed milling, was used to prepare 0.5 vol% Gr-reinforced Al composite. The dispersion of Gr in Al was studied via SEM images in Figure 9. It was reported that after 6 h of low-speed milling, Gr was well dispersed with no apparent clustering, compared with the obvious clustering of Gr after 4 h milling and reaggregation after 8–10 h of milling. In addition, a well-balanced combination of strength and ductility, of 295 MPa UTS, and 13.5% tensile elongation, were reported after 6 h low-speed milling and 0.5 h high-speed milling.

Zhang et al. [55] suggested that the use of smaller diameter milling balls or slower rotation speed can reduce the milling-induced degree of damage in the Gr intrinsic structure. It was explained that the reduction in milling collision energy is determined by these two factors, thus the 3-mm balls used in their study, compared with the 5- to 10-mm balls used in other studies, helped to minimize
the reduction in their $I_G/I_D$ ratio to 0.9 only after 24 h of milling. Nevertheless, milling is the most widely used technique in the synthesis of GRMMCs due to its simplicity, low cost, and effectiveness in dispersing Gr into metal matrices [49, 95], as studies utilizing milling in the synthesis of GRMMCs have reported uniform and homogenous dispersion of Gr nanosheets in metals [18, 23, 25, 27, 54, 56, 57, 70, 95, 99, 119].

3.1.2.2 Solution-Assisted Mixing

The techniques in this category involve the mechanical mixing of Gr and metal powders in a solvent. The selection of solvents is based on certain criteria such as nonreactivity with Gr or metals, volatility, and ease of evaporation [1]. There have been several reports on wet powder mixing of Gr and metals using techniques such as ultrasonic dispersion [24, 44, 50], mechanical mixing [11, 17, 57, 58, 66, 68, 71, 73, 89, 93, 104–108, 117, 120, 125, 127, 130], and magnetic stirring [51, 58, 69, 124, 126]. Solution-assisted methods have several drawbacks to be considered. First, the liquid media can be a source of contamination, which if not treated properly can prevent the full densification of the powder composites. Second, these methods are not suitable for large-scale composite fabrication. This is because a large amount of liquid media is required to ensure the adequate mixing of a few grams of composite powders. In addition, a uniform dispersion of Gr and metals might not be possible to achieve in liquid media mixing with all types of metals. This is because of the much higher density of metals than Gr, which can cause metal powders to settle instead of mixing uniformly with Gr [1].

Molecular-level mixing (MLM) is a synthesis technique that involves the mechanical mixing of the Gr reinforcement and the metal matrix in a solvent, yet with an aim of enhancing the interfacial bonding between the composite’s reinforcement and matrix through chemical bonding [72]. MLM has been used in the synthesis of Gr-reinforced Cu [51, 72, 96, 97, 134, 135] and Ni [67] matrix composites. Usually, the MLM synthesis technique involves several chemical steps. First, GO is mechanically mixed with the metal ions in a solvent, where the metal ions would attach to the GO sheets, which prevents its agglomeration. Second, metal ions are oxidized to form a chemical bond between the GO and the metal ions, thereby producing GO-reinforced metal oxide (MO) composites. This enhances the adsorption and dispersion of Gr in metals due to the decreased density difference between Gr and metal ions compared with Gr and metals. Finally, thermal reduction takes place to reduce GO-MO to GRMMCs [72, 134]. A schematic diagram of the MLM process adopted by Hwang et al. [72] for the synthesis of Gr-reinforced Cu matrix composites is shown in Figure 10. After preparing the GO from graphite using the Hummers method, an aqueous solution of Cu acetate was added to the GO suspension. NaOH was added after heating the
mixed solution to 80°C and the reaction reduced Cu ions to CuO, forming the GO-CuO mixture. Performing this step should enhance the dispersion between Cu and Gr through chemical bonding. Finally, the composite powders were thermally reduced at 450°C to Gr-Cu and consolidated by SPS. A Raman spectrometer was used to examine the effect of the synthesis process on the rGO structural integrity. The \( \frac{I_D}{I_G} \) ratio increased from 0.78 to 0.81 for GO and GO-Cu ions, respectively, then decreased to 0.40 for rGO-Cu composite. The authors related the increase in the ratio to the damage of the sp\(^2\) network of Gr as a result of the interaction of the Cu ions with the GO surface. The reduction in the ratio was, however, related to the partially recovered structure of Gr in the composite after the reduction process.

Chen et al. [97], Si et al. [96], and Zhang et al. [134] used MLM to fabricate GNP-reinforced Cu composite by starting with GNPs instead of GO. Zhang et al. [134] reported a modified MLM process. They suggested that the NaOH used in Reference [72] can have an adverse effect on chemical bonding by rapidly reducing CuO and GO during heating. In their study, they activated GNPs by using a hydrochloric acid solution before dispersing it in deionized water via ultrasonication. Then, they added the aqueous solution of Cu acetate to the GNP suspension with continuous sonication to facilitate the adsorption of Cu\(^2+\) ions. Heating and magnetic stirring followed to evaporate the solvent and dry the composite powders. The Cu acetate was then decomposed and thermally reduced in a tube furnace. Finally, composite powders were sintered and densified via SPS. The \( \frac{I_D}{I_G} \) intensity ratio from Raman spectra before and after the reduction were found to be 0.25 and 0.20, respectively. The drop in the ratio after the reduction suggests the partial restoration of the original structure.

Another promising solution-assisted technique is the particle-assisted shear exfoliation (PASE) technique [121, 135, 136]. The operating mechanism in PASE requires a high-shear mixer to induce a high-shear force on materials in liquids. In the synthesis of GRMMCs, PASE had been reported to effectively exfoliate graphite into fewer Gr sheets and to positively influence a homogenous distribution of Gr sheets in the metal matrix. It is claimed that the resultant high-shear force is sufficient to successfully exfoliate Gr sheets and eliminates the need for Gr pretreatment steps such as intercalation to weaken the interlayer binding strength. Wang et al. [121] used PASE to synthesize a Gr-Cu bulk composite using graphite and Cu particles as the starting materials without any additional pretreatment steps. The starting materials were dispersed in 1,3-butandiol and mixed using a rotor-stator mixer. The mixed slurry was then centrifuged for 1 h, heated at 200°C, and consolidated by SPS to obtain a bulk composite. The successful exfoliation of Gr into Cu powders was verified using x-ray diffraction (XRD), SEM, and TEM. Most importantly, high resolution TEM (HRTEM) investigations revealed that the sheets were multilayered but with a thickness less than 20 nm. SEM images of the powder composite revealed that the Gr sheets were embedded in the copper powder, suggesting good distribution; the copper particles prevented the Gr sheets from restacking. In another study, Wang et al. [136] combined PASE with MLM to produce a rGO-reinforced Cu matrix composite. The authors compared the properties and strengthening efficiency of the rGO-Cu composite produced by combining PASE and MLM with that produced using MLM alone. The microstructural and mechanical results verified the positive effect the PASE method imposed on the final composite. The composite produced using MLM alone suffered from agglomeration, a nonhomogenous structure, thicker GO sheets, holes and cracks, and lower yield strength values when compared with the composite fabricated using PASE and MLM. The effective synthesis technique included a typical MLM process where a GO colloid was added to cuprammonia and sonicated for 30 min. The PASE step was introduced during the solution evaporation step where, instead of the usual magnetic stirring, a high-shear rotor-stator mixer was used to offer high-shear force in the solution, aiding in the exfoliation and distribution of the GO.

### 3.1.2.3 Rolling Processes

Kim et al. [100] and Liu et al. [123] prepared the Gr-metal composites using different rolling procedures that yielded uniform microstructures and enhanced the composite properties. Kim et al. [100] used a combination of ball milling and high-ratio differential speed rolling (HRDSR) processing for the synthesis of MLG-reinforced Cu matrix composites. It was reported that ball milling resulted in significant fragmentation of MLG followed by a large shear strain provided by HRDSR, which accelerated the breakage of MLG into nanosizes, resulting in a dense uniform dispersion of MLG in the Cu matrix. The yield strength of the 1 vol% MLG-Cu composite was reported to be 361 MPa and the strengthening mechanism was concluded to be Orowan strengthening due to the homogeneous dispersion of high-density nanosized MLG particles. Liu et al. [123] reported the effect of changing the number of cycles in the accumulative roll bonding (ARB) process on the dispersion of GNSs in the Cu matrix. As the number of rolling cycles increases, the tensile strength reaches a maximum of 496 MPa after 6 ARB cycles, which is higher than that of unrolled GNS-Cu by 275 MPa. However, elon-
Figure 11: SEM images of tensile fracture of (a) annealed Cu showing ductile features, (b) accumulative roll bonded Cu, and (c) accumulative roll bonded GNS-Cu composite after 6 h rolling, where (b) and (c) show features of shallow dimples and shear zones. Reproduced from Reference [123].

gation drops significantly after the first cycle by 38% compared with annealed Cu and remains almost constant (5%) with increasing the cycle number. The sharp drop in elongation was attributed to the increment in the number of interfaces that act as sources for crack initiation after the first cycle. However, cycling was reported to enhance the uniformity of the GNS distribution within the Cu matrix, as well as the Cu-GNS and Cu-Cu bonding, leading to constant elongation with further cycling. This finding agrees with the SEM images in Figure 11, where the fracture surface of annealed Cu is represented by a typical ductile fracture with dimple-like morphology, while that of accumulative roll bonded Cu and accumulative roll bonded Cu-GNPs combines the features of shallow dimples and shear zones. Therefore, the authors [100, 123] concluded that rolling processes are effective methods for the fabrication of GR-MMCS.

3.1.2.4 In-Situ Synthesis
In-situ synthesizing of Gr in metal matrices has been reported to be effective for enhancing the strengthening efficiency of GRMMCs. Recently in-situ synthesis of Gr has been reported in Cu [88, 142], Al [140, 143, 144], and Ni [89, 90] bulk metal matrices. Chen et al. [88] reported the fabrication of a bulk Gr-Cu composite by CVD. They suggested that the in-situ growing of Gr results in better dispersion of Gr and a stronger adhesion between Gr and the metal matrix. In addition, their in-situ method resulted in the fabrication of a 3D Gr structure, which has a better strengthening effect than the usual 2D Gr produced by other methods. Initially, PMMA was used as the carbon source and was milled with Cu powders. The PMMA distributes within the Cu powders during milling and prevents Cu welding by forming interspaces between the agglomerated Cu powders in which the Gr will be grown into during CVD. The milled powders went through a CVD pro-
cess in a quartz tube during which carbon atoms decomposed from PMMA and diffused into the porous structure of the Cu powder agglomerates. This is attributed to the high temperature environment and low solubility of C in Cu. As a result, Gr copies the porous structure of the agglomerated Cu powders, and a 3D Gr structure is formed following the shape of the porous Cu framework. The Cu-Gr composite powders were then hot pressed under a vacuum. TEM images of in-situ grown Gr in Cu are shown in Figure 12(a–c). In addition, the 3D structure of Gr was demonstrated and captured by TEM, as shown in Figure 12(d–f). This was done by removing the Cu matrix via CuCl$_2$ solution acidized with hydrochloric acid, leaving the interconnected 3D framework of Gr with a porous structure corresponding to the empty spaces where the Cu matrix was placed.

Jiang et al. [89] reported the in-situ fabrication of Gr-reinforced bulk Ni matrix composites. They used simple powder metallurgy steps to fabricate in-situ grown Gr into Ni matrix. A schematic diagram of the in-situ fabrication process is shown in Figure 13. They used sucrose as Gr precursor and mechanically stirred it with Ni in water until the water evaporated. Later, the composite powders were green compacted and sintered. The composite achieved a 69.8% increase in hardness and 311% increase in UTS. This was attributed to the uniform distribution of Gr in Ni and the foreign impurity-free interface achieved with this in-situ synthesis leading to efficient load transfer.
Producing in-situ Gr grown in an Al matrix via CVD holds several challenges, such as the noncatalytic nature of Al, its pore wettability with carbon, and the $\text{Al}_4\text{C}_3$ formation reaction that can take place due to high-temperature processing. These challenges can be overcome by adding an intermediate material into the interface of Gr and Al to act as a catalyst, enhance interfacial bonding, and prevent carbide formation. Chen et al. [143] and Liu et al. [144] produced Cu-coated GNS-Al and Ni-decorated GNP-6061Al matrix composites, respectively. Both studies utilized CVD to incorporate the intermediate elements. Chen et al. [143] used Cu as the catalyst and methane as the carbon source. First, they used magnetic stirring and ultrasonication to mix Al and CuCl$_2$ and obtain a homogeneous Cu$^{2+}$-Al solution. The solution was then heated to dry the powders before loading it into a tube furnace for the CVD process to obtain Cu-Al powders followed by heating with methane-Ar atmosphere to obtain the Cu-coated GNS-Al powders. This was followed by vacuum hot pressing and sintering. The composite achieved a 200% increase in UTS. A schematic diagram of the process is shown in Figure 14.

3.1.2.5 Molten or Liquid Metal Processes

Fabricating bulk GRMMCs with steps involving metal melts processing, such as Mg [103, 110, 111, 138] and Al [87, 94, 98, 120, 139–141], has been reported. Rashad et al. [110, 111] used the disintegrated melt deposition (DMD) technique to fabricate 1.5 wt% GNP-reinforced Mg-6Zn alloy and 3 wt% GNP-reinforced AZ61 Mg alloy, respectively. The process involves the melting of a 1.3 Kg Mg ingot in a graphite crucible and a protective atmosphere, followed by the addition of the alloying elements and the GNP powders while maintaining the Mg melting temperature of 20°C. The molten mixture was allowed to cool and solidify in a steel mold preheated at 300°C. The solidified mixture was homogenized and hot extruded to 16-mm diameter rods of uniformly dispersed GNP-Al alloy powders.

Du et al. [138] mechanically mixed GNP-Al alloy powders in ethanol under ultrasonication followed by centrifugation and vacuum drying. Mixed composite powders were extruded into rods before melting in an electromagnetic induction furnace with stirring to achieve homogeneity. The molten composite mixture was allowed to solidify under atmospheric conditions followed by hot extrusion. The composite achieved a 62% increase in TYS. Seventy-two percent of the strength enhancement was attributed to load transfer due to the efficient interfacial
bonding and homogenous distribution of GNPs in the Mg alloy matrix.

In 2012, Chen et al. [103] combined liquid-state processing of metal melts with solid-state stirring to develop a novel GNP-reinforced Mg matrix composite technique. The process consists of three main steps. In the first step, GNPs were dispersed into Mg by adding GNPs into the ultrasonicated Mg melt at 700°C using an automatic feeding system and a high-powered ultrasonic probe. The second step was to solidify the ultrasonicated GNP-Mg melt by casting it onto a plate mold to obtain a plate-shaped composite. The presence of large clusters within the composite was captured by SEM images, suggesting poor dispersion of GNPs into the Mg matrix. This was attributed to the large aspect ratio of the GNP, which makes it difficult to achieve homogenous dispersion of GNP into the Mg melt using ultrasonication only. Thus, the third step in the process was needed: solid-state processing. Friction-stir processing (FSP) was the solid-state technique applied to further distribute the GNPs into the Mg matrix. FSP is an emerging solid-state metal working technique, which involves mixing the sample in the solid state, imposing severe plastic deformation on the processed zone. In addition, friction is created between the rotating tool and the processed sample, which leads to localized heating [148, 149]. FSP is a proven technique for homogenous dispersion of reinforcements in metal matrices and homogenizing structures of cast alloys [150, 151]. During FSP, the composite plate prepared by step 2 was treated with a special rotating tool that has a pen-shaped end, which interacts with the plate by heating and mixing it, thus processing it in the solid state. Recently, FSP was employed in several studies for the fabrication of Gr-reinforced Al matrix composites [98, 120, 139]. Zhang et al. [120] reported the fabrication of 1 wt% GNP-reinforced 2009Al alloy matrix composite via a combination of powder metallurgy and the FSP technique. The effect of increasing the number of FSP passes on the distribution of GNPs in the metal composite was studied. It was reported that two passes were enough to achieve homogenous distribution of GNPs and to decrease the number and size of GNP clusters in the matrix. The above investigations indicate that the molten metal-based synthesis techniques are feasible and effective for the fabrication of bulk GRMMCs.

### 3.1.3 Postsynthesis/Consolidation

The consolidation step is important to fabricate GRMMCs. It provides the final product to be studied and controls the extent of the sample’s densification. This step is mostly needed when the composite obtained is in the powder form after the mixing step. This applies to mixing techniques including ball milling, solution-assisted mixing, a
few in-situ synthesis techniques, and after some molten metal mixing procedures. The consolidation step can be divided into three categories: pressing and sintering methods, full densification methods, and the most recently developed, 3D printing.

Pressing/sintering consolidation techniques involve two steps: the pressing of the mixed composite powders (which can be done at room temperature and is known as green compaction, or it can be done simultaneously with heating) followed by sintering or annealing [11, 17, 18, 20, 26, 27, 44, 50, 58, 65, 66, 73, 124, 125, 132, 137]. Composite powders are placed into a steel die, and a vertical pressing motion is applied onto the mixed powders for pressing. Limitations exist such as the size and shape of the final sample as it is controlled by the size and shape of pressing tools, the pressing capacity, and the powder compressibility. Although this technique is a simple approach for composite compaction, full densification is hardly achieved [152].

Achieving approximately full density in samples can be achieved through the second category of consolidation techniques: full densification methods. Several techniques have been developed and applied for the synthesis of GRMMCs, such as hot pressing [23, 49, 51–55, 63, 68, 70, 88, 99, 108, 118, 119, 126], hot rolling [25, 52, 56], extrusion [11, 17, 50, 55, 57, 73, 118], and SPS [24, 52, 69, 71, 72, 97, 127, 134]. Relative densities between 98% and 99.8% have been achieved for GRMMCs via full-densification consolidation techniques [11, 17, 24, 26, 88, 113, 125–127].

3D laser printing techniques have been used for the synthesis of Gr-reinforced Ni [112], Ti [60, 153], and Al [133] matrix composites. Most recently, Hu et al. [133] fabricated the first time a Gr-reinforced Al matrix composite by 3D printing using the selective laser melting (SLM) technology. First, ball milling was used to disperse Gr into Al without the use of process control agents or protective gases. Then, SLM was used to consolidate the ball-milled powder composite and fabricate a 3D bulk composite. A schematic diagram of the 3D printing process is presented in Figure 16. Raman spectra of the laser-melted composite showed an increase in the $I_D/I_G$ ratio from 0.053 to 0.48 for as-received Gr and Gr in the laser-sintered Gr-Al composite. The increase in structural defects was attributed to the thermal damage from the high temperature of the SLM process and to the reaction between Gr and aluminum, leading to the formation of Al$_4$C$_3$.

Ayyappadas et al. [137] compared the effects of two sintering techniques on the densification of their Gr-Cu composite prepared by powder metallurgy and cold pressing. Microwave sintering resulted in higher relative density than conventional tube furnace sintering for their 0.9 vol% Gr-Cu composite. Sintering in a microwave field allowed for higher heating rates, which led to shorter processing times and gave more refined structure of the composite. Similarly, Ghasali et al. [102] studied the effect of different sintering techniques, such as conventional, microwave, and SPS, on the densification and performance of 1 wt% GNP-Al composite. Each sintering technique had a different effect on the formation of Al$_4$C$_3$ and thus resulted in a different structure and mechanical performance. The highest relative density was reported after SPS thanks to its vacuum environment.

A few observations have been made regarding density after consolidation in GRMMCs. One observation is the density of the GRMMCs compared with the pure metal prepared in the same way. In several reports, the density of the GRMMCs was found to be less than that of the pure metal due to the contribution from Gr, which has a lower density than that of metals [11, 17, 24, 27, 58, 108, 129, 140]. However, several authors reported a higher density for the GRMMCs compared with the pure metal [26, 137]. Ayyappadas et al. [137] suggested that Gr acting as a lubricant during cold pressing enhanced the densification and increased the density of the composite compared with pure
Cu processed the same way. Alam et al. [26] reported a poor relative density of 77% for pure Al compared with 98% for 3 wt% GNP-Al composite. They justified the poor Al density to the poor powder compressibility of Al as a result of the Al$_2$O$_3$ layer on the Al surface, which hindered the direct contact between Al powders. In contrast, the addition of GNP particles helped in filling those pores and resulted in a much higher composite densification. Because the density of Al$_2$O$_3$ is higher than the density of Al and Gr, Rashad et al. [17] reported higher experimental densities than theoretical densities and attributed it to Al oxidation and formation of Al$_2$O$_3$ during sintering.

The density of GRMMCs has been reported to vary with Gr content [11, 17, 24, 26, 27, 47, 58, 65, 97, 99, 104, 108, 126, 127, 140]. A decrease in experimental density was reported at higher Gr additions. Some authors attributed this behavior to the lower density of Gr than the metal matrix, thus lowering the density of the composite via Gr addition [11, 17]. Ayyappadas et al. [137] suggested that this behavior is a result of Gr acting as a barrier to Cu diffusion, resulting in lower densification. Li et al. [27] attributed the drop in the relative density of their GNS-Al composite when adding GNSs to the increased air gaps that form in the composite as a result of the absorption of GNSs to gas elements such as O, N, and CO. Alam et al. [26] reported an enhancement in the density of the Gr-Al composite up to 3 wt% GNP addition, after which a sharp decrease in density is observed. The enhancement was attributed to the filling of the voids between Al particles by GNPs. However, beyond 3 wt% GNP addition, authors suggested that the agglomeration of the GNPs in the composite and the formation of Al$_4$C$_3$ phase resulted in poor densification and sinterability [26]. The agglomeration of GNPs in GRMMCs at higher GNP addition and its effect on poor densification and porosity has been reported in other GRMMC studies as well [24, 65, 127].

### 3.2 Structure and Morphology of GRMMCs

During the processing of GRMMCs, the Gr sheets tend to agglomerate. This is caused by the high surface energy of the nanometer-thick sheet that stems from the weak Van der Waals forces that hold the Gr layers. This results in inhomogeneous dispersion of Gr in metal matrices [69]. Enhancing the properties of metals depends on the integrity of the interface between the Gr reinforcement and the metal matrix. Thus, uniform distribution and homogenous dispersion of the reinforcement within the metal matrix must be achieved without sacrificing the structural integrity of the Gr [73]. This is basically controlled by the type and amount of reinforcement material used and the synthesis technique of the composite. The morphology of GRMMC is usually investigated through the atomic-scale resolution images of SEM and TEM, and other analytical tools such XRD and Raman spectroscopy, as they offer detailed morphological and structural information [56].

#### 3.2.1 Dispersion of Graphene in MMCs

One of the main structural features investigated in all prepared GRMMCs is the degree of dispersion of Gr in the metal matrix. A homogenous and good dispersion of Gr and its derivatives has been reported in Al [18, 23–25, 27, 44, 53–57, 69, 70, 94, 139, 141], Cu [51, 52, 66, 71, 88, 93, 97, 123–127, 134, 137], and Mg [65, 103–105, 108, 110, 138], where good dispersion is usually attributed to the effectiveness of the composite's fabrication technique [48, 56, 71, 103, 127]. In some cases, it was difficult to observe Gr by electron microscopic means, suggesting that Gr was well dispersed throughout the metal matrix [18, 48, 56, 70]. Shin et al. [56] produced FLG with an Al matrix using planetary milling followed by attrition milling. Their analysis concluded that FLG can be observed after planetary milling, as the sheets were thin enough to transmit the electron beam and show the Al particles below the Gr sheets. However, those thin sheets were no longer visible on the Al powder surface after attrition milling, which suggests its effectiveness in embedding and dispersing the sheets inside the Al matrix. Similar results were reported by Rashad et al. [48]; they used ball milling to fabricate GNP-reinforced AZ31 Mg alloy. GNP were not traced via TEM in the milled composite powders, suggesting that they are well embedded and uniformly distributed within the metal matrix.

Surface modification of Gr and/or the metal before composite fabrication is suggested to enhance Gr dispersion in metal matrices [66, 71, 73, 127]. Zhang et al. [127] attributed the homogenous distribution of GNPs in the Cu matrix composite to the electroless plating of GNPs by Cu and Ni before the ultrasonication of the treated GNPs with Cu powders. Compared with the treated GNP-Cu composite, the untreated GNP-Cu composite showed heavy agglomeration and clustering. Jiang et al. [71] used PVP and PVA in the surface modification of pristine graphene (PG) and Cu, respectively. The affinity of the carbonyl groups in PVP to the hydroxyl groups in PVA compensates for the lack of affinity between PG and Cu, realizing a homogenous dispersion of PG in Cu. Compared with their modified PG-Cu composite, PG sheets agglomerated and separated from Cu particles in the unmodified sample.
Many reports have indicated that Gr was distributed around or along the grain boundaries of metal matrices such as Al [47, 53–55, 57, 59, 68, 69, 98, 140] and Cu [49, 51, 63, 66, 100, 124, 126, 134, 137, 142]; see Figure 17. This feature suggests good dispersion of reinforcement in the matrix. Ju et al. [69] confirmed the distribution of Gr along the grain boundaries of Al particles for their 0.3 wt% GO-Al composite through HRTEM, where Gr sheets with three to five layers were observed to surround the Al particle. Gao et al. [51] reported the homogenous distribution of rGO along the grain boundaries of Cu matrix, which was attributed to their successful electrostatic self-assembly technique.

### 3.2.2 Interfacial Interactions and Carbide Formation

Another challenge in synthesizing Gr-reinforced MMCs is the high reactivity of metals and the possibility of interfacial interactions at the Gr-metal interface [44]. Good interfacial bonding has been reported in GRMMCs prepared by several synthesis techniques [49, 51, 52, 63, 64, 66, 71, 72, 88, 93, 97, 105, 108, 110, 111, 123–127, 134]. However, metals have different interactions with Gr, and many challenges still persist regarding the interfacial interactions.

In Gr-reinforced Al matrix composites, the formation of Al\(_4\)C\(_3\) phase has been highly investigated. XRD has been used to determine the phase compositions of Gr-Al composite and to investigate the presence of interfacial reac-
The presence of Al₄C₃ has been identified by XRD in some cases [18, 24, 27, 44, 55, 57, 69, 70, 102, 133], while in other cases, XRD was not able to detect any interfacial reactions, such as forming the Al₄C₃ [24, 44, 54, 69, 70, 94]. The undetectability of Al₄C₃ by XRD was attributed to the low content of carbides formed, making it beyond the detection limit of XRD; the absence of any residual carbon in the starting reinforcement material; and the passivation of Al due to the formation of thin stable aluminum oxide preventing any chemical contact between Al and carbon. Furthermore, investigating the presence of second phases and interfacial reactions between Gr and Al was based on TEM images, SAED analysis, and EDX in some studies [23, 24, 27, 55, 57, 87, 94, 95, 120, 133, 154, 155]. Figure 18 shows TEM images that have reported the presence of Al₄C₃ phase in Gr-Al composites. The Al₄C₃ phase was described to have a needle-like or rod-like appearance [23, 27, 55, 87, 95, 120, 133, 154, 155].

Li et al. [27] reported the absence of Al₄C₃ in XRD patterns of Al-0.25 wt% GNS composite prepared by ball milling and hot pressing (HP). However, rod-like Al₄C₃ phases were identified by TEM. Hence, the absence of
Al₄C₃ peaks was attributed to the XRD detection limit. In the same study, Al₄C₃ peaks were detected by XRD for high Gr wt%, and it was reported that the intensity of those peaks increased with increasing the Gr content from 0.5 wt% to 1.0 wt%. This suggests that the Gr content plays a significant role in the intensity of the Al₄C₃ phase.

There seems to be a debate surrounding the formation of Al₄C₃ in Gr-Al composites regarding whether it is caused by the exposure of the composite to high temperatures during synthesis, or it is initiated due to disorder of the graphitic structure induced by synthesis regardless of the temperature. It has been reported that the formation of carbides in Al-C systems can be controlled by thermodynamic and kinetic approaches. Thermodynamically, the reactivity of Al-C systems can be lowered by using carbonaceous materials with fewer defect sites or by adding alloying elements to change the Gibbs free energy of the reactions. Kinetically, controlling the rate of reaction by controlling synthesis temperature or time of exposure to high temperature can inhibit Al₄C₃ formation [94, 102]. The phase diagram of the Al-C system [26] suggests that a facial interaction in Gr-Al composites can occur at a temperature of 550°C with a stochiometric ratio of 4:3. When it comes to Al-Gr in the literature, some studies have used synthesis routes such as sintering, HP compaction, or SPS, at temperatures in the range of 300–600°C, yet no formation of Al₄C₃ was detected by XRD [44, 54, 57, 69, 70], TEM [94], or EDX [53, 59].

It has been reported that defect-free Gr platelets are known to be chemically inert to the Al reaction even at temperatures above the Al melting temperature [87, 94, 154, 156]. However, structural defects on Gr can serve as initiation sites for the interfacial reaction with the Al matrix even at temperatures below the Al melting point [26, 55, 94]. The formation of Al₄C₃ has been attributed to the quality and structural integrity of the Gr reinforcement in the Al-C matrix [26, 44, 55, 57, 69, 70, 156]. Li et al. [53] used both ball milling and high-temperature melting, in addition to rolling and casting, to fabricate their 0.2 wt% GNP-Al composite. No traces of Al₄C₃ were detected at the interface by EDX [53]. Yang et al. [94] fabricated Al₄C₃-free 0.54 wt% GNP-Al composite using the pressure infiltration method; the composite was subjected to high pressures and temperatures. The author suggested that the use of low-defect Gr can hinder the Gr-Al reaction. Jiang et al. [95] reported an increase in structural damage of Gr with milling time as well as higher content of Al₄C₃ at higher milling times. This result showed the relationship between the structural integrity of Gr and carbide formation in Gr-Al composites synthesized by ball milling. Al₄C₃ peaks were detected at all milling times in Raman spectroscopy with significantly higher relative peak intensities at higher milling times. In addition, rod-like phases corresponding to Al₄C₃ were only identified by TEM at higher milling times.

Not only is there a lack of understanding regarding the formation mechanism of Al₄C₃ in Gr-Al composites, but there is also a debate on the role the phase plays on the properties of the composites. Several studies have reported that the carbide phase exhibits a detrimental effect on the mechanical behavior of Gr-Al composites due to the structural damage involved with the formation of Al₄C₃. Other studies attributed the enhancement in mechanical properties (such as strength and hardness) to the Al₄C₃ interfacial product. Ghasali et al. [102] suggested that the amount of carbide formed at the Gr-Al interface determines the effect this phase has, whether positive or negative, on the mechanical behavior of the Gr-Al composites. A more systematic approach toward producing and characterizing these composites is required in order to achieve a better and more comprehensive understanding of the issue.

In studies involving Gr-reinforced Cu composites, second phase formation between Cu and Gr has not been reported [51, 63, 64, 71, 97, 123–127, 134, 137]. This is attributed to the lack of a solid-solution phase in the Cu-C phase diagram [71, 123, 137], resulting in physical bonding only and no chemical reactions between Gr and Cu. However, several researchers have reported the formation of Cu-O-C bonding at the interface between Gr and copper. Several advantages have been related to such bond formation including the enhanced load transfer between the matrix and reinforcement, and the enhanced distribution of the Gr reinforcement in the Cu matrix [49, 52, 72, 97, 127, 134, 137]. Other researchers have suggested decorating Gr sheets with metal or carbide coatings that have better affinity with Cu to achieve a strong interface between Gr and Cu. Several coatings, such as Ni [2, 127], TiC [96], and VC [96], have been reported to have a positive effect on the interfacial bonding and mechanical properties of Gr-Cu composites [128].

While carbides such as MgC₂ and Mg₂C₃ are known in Mg-C systems, the formation of such metal carbides in Gr-Mg studies has not been reported [109]. Similar to Cu, a big challenge in the field of fabricating Gr-reinforced Mg matrix composites is the poor wettability between Mg and carbonaceous material such as Gr [105]. As a result, an alloying element, which possesses better wettability with Gr, is usually added to the Mg matrix before mixing it with the Gr reinforcement in an attempt to increase the wettability between Gr and Mg [48, 104, 105, 110, 111]. Al and Zn have been reported as efficient alloying elements in Mg matrices that enhance the wettability of Mg and Gr and result in...
excellent bonding and uniform dispersion of Gr within the metallic matrix [105, 110].

3.3 Properties of GRMMCs

3.3.1 Mechanical Properties

The main purpose behind incorporating Gr or its derivatives as a reinforcement in metals is to enhance their mechanical behavior [1]. The main challenge in the development of GRMMCs is to improve strength without sacrificing good ductility. It is important to note that enhancement in the mechanical properties of GRMMCs vary from one study to another because of the difference in the starting materials used, Gr concentrations, fabrication methods, or synthesis parameters used; hence, different structures and morphologies are obtained [1, 129]. This section outlines the latest GRMMC research and the reported mechanical behavior. Tables 4, 5, 6, and 7 summarize the reported studies in the field of Gr-reinforced Al, Cu, Mg, and other metals and their resultant mechanical properties.

3.3.1.1 Strength and Hardness

The mechanical properties of GRMMCs vary from one study to another. The two most reported factors to affect the mechanical properties in the same study are the Gr content and Gr type used as a reinforcement.

Effect of Graphene Content

Relatively small amounts of Gr have been used in the majority of reports, with weight percentages ranging from 0.05% to 5% [18, 20, 24, 26, 27, 44, 49–51, 54, 55, 57–59, 63–66, 69, 71, 88, 94, 105, 108, 110, 124, 125, 133, 138, 141], and volume percentages ranging from 0.1% to 10% [23, 25, 52, 56, 70, 72, 97, 98, 109, 126, 127, 134, 137, 154]. Improved tensile and compressive strengths and hardness have been reported in the majority of fabricated and studied Gr-Al [18, 20, 23–25, 27, 44, 50, 54–58, 68–70, 94, 133, 140, 141], Gr-Cu [49, 51, 52, 63, 64, 66, 71, 72, 88, 93, 96, 97, 123–128, 134, 137] and Gr-Mg [48, 65, 103–105, 108–111, 138] composites. In these studies, the concentrations of Gr were optimized; above this level, the mechanical properties gradually decreased with observed deterioration in the behavior. Such behavior was attributed to the difficulty in achieving a homogenous distribution of reinforcement in the matrix at a high reinforcement content. In addition, agglomeration of Gr or its derivatives at a high Gr content can form microvoids and, hence, poor contact between the reinforcement and the matrix. In the case of Al, the deterioration in mechanical strength was related to the formation of interfacial reactions such as Al₄C₃, which had been reported to increase with increasing the Gr content. The enhancement in the strength and hardness of GRMMCs occurred for several reasons, such as the homogenous dispersion of the reinforcement in the matrix, the successful load transfer between the reinforcement and the matrix, and the resultant strong interfacial bonding [18, 23, 49, 51–53, 57, 65, 66, 72, 88, 94, 96, 97, 101, 108, 120, 123, 125–130, 134, 137, 138], dislocation pinning by Gr and grain refinement [20, 27, 49, 54, 56, 57, 65, 70, 71, 88, 93, 96, 97, 101, 108, 123–129, 138, 139], Orowan strengthening [24, 52, 129, 130, 134], the increase in dislocation density due to mismatch in coefficient of thermal expansion (CTE) [18, 27, 48, 49, 52, 97, 103–105, 110, 111, 124, 126, 127, 129, 130, 134, 138], the formation and presence of carbides [102] as obstacles to dislocation motion [44], and the formation of twin boundaries that act as barriers resulting in the discontinuity of the slip systems [137].

Kumar et al. [44] studied the effect of Gr concentrations in GNP-reinforced Al6061 matrix composite. The increase in hardness was attributed to the homogenous dispersion of Gr in the Al alloy matrix at 0.5 wt% concentration, whereas the reduction in hardness was attributed to agglomeration and initiation of microvoids at higher Gr weight concentrations. Similar observations have been reported in References [24, 49, 59, 133]. Hu et al. [133] reported that the hardness increased after adding Gr to the Gr-Al composite. Compared with pure aluminum, Al-2.5 wt% Gr had the highest hardness value, with a 75% increase in hardness. The increase was attributed to the significant intrinsic mechanical properties of Gr. The larger error bars reported for the average hardness values at a higher Gr content suggest that a higher graphene content leads to Gr agglomeration and results in scattering of the hardness values. Dasari et al. [59] reported a higher porosity content in the Al-GO composite at higher GO concentrations. Similar observations have been reported in References [24, 49, 59, 133]. Hu et al. [133] reported that the hardness increased after adding Gr to the Gr-Al composite. Compared with pure aluminum, Al-2.5 wt% Gr had the highest hardness value, with a 75% increase in hardness. The increase was attributed to the significant intrinsic mechanical properties of Gr. The larger error bars reported for the average hardness values at a higher Gr content suggest that a higher graphene content leads to Gr agglomeration and results in scattering of the hardness values. Dasari et al. [59] reported a higher porosity content in the Al-GO composite at higher GO concentrations and attributed that to the agglomeration of GO sheets. Similarly, Ayyappadas et al. [137] reported a decrease in the density of the GNF-Cu composite with increasing Gr content, which was correlated with the increase in porosity when increasing the GNF content. The authors reported that the GNFs acted as barriers to Cu diffusion throughout the sintering process. Hence, the higher the GNF content, the larger the barrier to Cu diffusion, and the higher the porosity in the composite, which resulted in lower density [137].
| Matrix | Reinforcement | Pretreatment | Mixing | Compacting | % Increase | Strength (YS/TS) | Hardness (MPa/HV) | Others | Elongation (%) | Ref. |
|--------|--------------|--------------|--------|------------|------------|----------------|------------------|--------|----------------|------|
| Al6061 | GNP 0.70 wt% | In-situ CVD synthesis of Ni-decorated GNP | Ball milling (BM) | HP | 30% | 75% | 33.7% | – | 22.1% | [144] |
| Al    | GNP 0.2 wt%  | –            | BM     | Cold pressing (CP) / electromagnetic induction heating of CP powders and Al melt / casting and rolling | 36.8% | (156 MPa) | – | 16.5% | – | 4% | [53] |
| Al    | GNP 1.00 wt% | BM in ethanol of GNP | Ultrasonication of GNP with SDS in ethanol | BM | SPS | – | – | 109 HV | – | – | [102] |
| Al    | GNP 0.54 wt% | –            | BM     | HP / Infiltration of molten Al and pressed powder Extrusion / annealing | 93% | 228% | – | – | – | [94] |
| AI5083| GNP 1.8 wt%  | –            | BM     | HP / Infiltration of melted alloy and pressed powder Extrusion / annealing | 11.07% | (331 MPa) | 19.12% | (186.9 MPa) | – | – | 6.3% | (21.3%) | [87] |
| AI5083| GO 1.8 wt%   | –            | BM     | HP / Infiltration of melted alloy and pressed powder Extrusion / annealing | – | 17.6% | (186.5 MPa) | – | – | 9% | [87] |
| Al    | MLG 2.50 wt% | –            | BM     | 3D printing by SLM processing | – | – | 75.3% | (66.6 HV) | – | – | [133] |
| AA2124| GNP 5.00 wt% | –            | Mechanical mixing (MM) BM | CP / sintering / hot extrusion | 21.6% | (425.6 MPa) | 127.3% | (200 MPa) | – | 69% | (120 GPa) | Young’s modulus (YM) | [47] |
| AlMg5 | GO 1.00 vol% | –            | BM     | Uniaxial HP (UHP) | 113.8% | (556 MPa) | 53.8% | (200 MPa) | 137% | (166 HV) | 287.14% | (813 MPa) | Flexural strength (FS) | 10% | (25%) | [70] |
| Al    | GNS 0.50 vol% | –           | BM     | CP / sintering / hot extrusion | 20% | (295 MPa) | 18.6% | (230 MPa) | – | 5.3% YM | 13.5% | (19.3%) | [95] |
| Al    | GNS 1.00 wt% | GNP sonication in ethanol | BM | HP | 56.19% | (164 MPa) | 38.27% | (112 MPa) | 16.13% | (72HV) | – | 15% | (16%) | [27] |
| Matrix  | Reinforcement | Pretreatment | Mixing  | Compacting       | % Increase | Hardness | Others | % Elongation (original) | Ref. |
|---------|---------------|--------------|---------|------------------|------------|----------|--------|------------------------|------|
| Al2024  | FLG 0.50 vol% | –            | BM      | HP / sintering    | 100% (700 MPa) | –  | –      | 4%                      | [25] |
| Al2024  | GNP 0.70 vol% | GNP sonication in anhydrous ethanol | BM | CP / HP / annealing | – – | – – | 26.3% (120 MPa) | – – | [119] |
| Al      | GNP 0.50 wt%  | GNP sonication in ethanol | BM | CP / HP | – – | 8.81% | – – | – – | [54] |
| Al      | GNP 1.00 wt%  | GNP sonication in ethanol | BM | CP / HP | – – | 12.6% drop (97 HV) | – – | – – | [99] |
| Al      | FLG 0.70 vol% | BM of graphite in isopropyl alcohol | BM | Hot rolling | –  | 71.8% (440 MPa) | – – | – – | [56] |
| Al2024  | GNP 0.70 vol% | GNP sonication in ethyl alcohol | Planetary milling (PM) BM | Hot rolling | 100% (700 MPa) | – | –  | – – | [25] |
| Al-1.5 wt% Mg 3.9 wt% Cu | GNF 0.50 wt% | Sonication of GNFs in ethyl alcohol | BM MM | HIP / hot extrusion | 25.5% (467 MPa) | 50% (319 MPa) | – – | – | [118] |
| Al      | GNP 1.00 wt%  | –            | BM      | CP / sintering    | 130% (92 HV) | – – | 8.3% (17.3%) | – – | [18] |
| Al      | GNF 1.00 wt%  | –            | MM      | Hot extrusion / annealing | 68.7% (248 MPa) | 55.2% (19 MPa) | – – | 47% (760 MPa) FS | – | [119] |
| Al      | rGO 0.5 wt%   | –            | Sonication in ethanol BM | CP / sintering | – – | 85.7% (316 HV) | – – | – | [50] |
| Al      | FLGO 1.00 wt% | MM Sonication | – | CP / sintering | 26.3% (120 MPa) | 43% (≈60 MPa) | – – | – | [129] |
| Al      | GO 0.30 wt%   | –            | MM      | Filtration / thermal reduction CP / sintering | – – | 32% (34.5 HV) | – – | – | [58] |
Table 4: ...continued

| Matrix | Reinforcement | Pretreatment | Mixing | Compacting | % Increase Strength TS | % Increase YS | Hardness | Others | % Elongation (original) | Ref. |
|--------|---------------|--------------|--------|------------|------------------------|---------------|----------|--------|------------------------|------|
| Al     | GO 0.30 wt%   | Sonication and mixing of Al in CTAB | MM     | CP Sintering | 30% (110 MPa)          | –             | –        | –      | –                      | [68] |
| Al     | GNP 1.00 wt%  | Sonication in ethanol Al MM in ethanol | MM     | CP sintering / hot extrusion | 9.14% (203 MPa)        | 29.5% (145 MPa) | 18.4% (90 HV) | –      | –                      | [11] |
| Al     | GNP 0.3 wt%   | Sonication of GNP in water MM of Al in acetone | MM in ethanol | CP sintering / hot extrusion | 11.3% (280 MPa)        | 14.7% (195 MPa) | 11.8% (85 HV) | 9.53% (13.4%) | –                      | [17] |
| Al     | GNS 0.30 wt%  | BM of Al1Al dispersion in PVA aqueous solution | MM of PVA / Al and GNSs in H2O | HP hot extrusion | 62% (269 MPa)          | –             | –        | 13%    | –                      | [73] |
| Al     | GO 0.20 wt%   | GO sonication in H2O Magnetic stirring (MS) of Al in IPA | Magnetic stirring CP sintering | – | – | 29% | – | – | – | [59] |
| Al-1.5 wt% Sn | GO 1.00 wt% | MS of Al in ethanol GO sonication in ethanol / H2O | MS Sonication CP sintering | 26% (120 MPa) | – | 43% (+58 HV) | – | – | [129] |
| Al     | GNS 0.15 wt%  | Dispersion of GNSs in EC / IPA Dispersion of Al in EC / IPA | MS | Filtration / CP sintering | – | – | 43% (377 HV) | – | – | [58] |
| Al     | GNP 1.00 wt%  | GNP sonication in acetone | Sonication | SPS | 54.8% (105 MPa) | 84.5% (65 MPa) | 21.4% | – | – | [24] |
| Al6061 | Gr 0.50 wt%   | GNP sonication in acetone | Sonication | CP sintering | – | – | 26% (+84 HV) | – | – | [44] |
| AA5052 | GNP 3.00 vol% | – | – | FSP | 35.7% (224 MPa) | 112.4% (148.7 MPa) | 53% (84 HV) | – | 20% (28.5%) | [98] |
| Al     | MLG 40.0 at%  | – | – | FSP (8 passes) | – | 75% (147 MPa) | 88% (94 MPa) | – | 26% | [139] |
Table 4: ...continued

| Matrix | Reinforcement | Pretreatment | Mixing | Compacting | % Increase | Strength | Hardness | Others | % Elongation (original) | Ref. |
|--------|---------------|--------------|--------|------------|------------|----------|----------|--------|------------------------|------|
|        |               |              |        |            | TS        | YS       |          |        |                        |      |
|        |               |              |        |            |            |          |          |        |                        |      |
| Al2009 | GNP 1.00 wt%  | –            | FSP    | –          | 23.3%      | 30.5%    | –        | –      | –                      | [120]|
|        |               |              |        |            | (514 MPa)  | (398 MPa)|          |        |                        |      |
|        |               |              |        |            |            |          |          |        |                        |      |
| AA5052 | GNP 3.00 vol% | FSP          | –      | –          | 18.6%      | 112%     | 40%      | –      | 20%                    | [154]|
|        |               |              |        |            | (244 MPa)  | (148.7 MPa)|          |        |                        |      |
|        |               |              |        |            |            |          |          |        |                        |      |
| Al     | GNS 2.50 vol% | Magnetic stirring of Al and Cu in ethanol | In situ synthesis of Gr by CVD with methane as C source | VHP / sintering | 200%      | –        | –        | –      | –                      | [143]|
|        |               |              |        |            | (318 MPa)  |          |          |        |                        |      |
|        |               |              |        |            |            |          |          |        |                        |      |
| Al     | GNS 2.00 wt%  | –            | One step in-situ synthesis in molten Al under alkali halides melt | –          | 16%       | 51.6%    | 262%     | 45.1%  | 26.2%                  | [140]|
|        |               |              |        |            | (48.1 MPa)| (93.78 MPa)| (57.19 HV)| (87.9 GPa)| (13.04)|                        |      |
Table 5: Summary of Reported Gr-Reinforced Cu Matrix Composites Studies.

| Matrix | Reinforcement | Pretreatment | Mixing | Compacting | % Increase | Strength | Hardness | Others | % Elongation (original) | Ref. |
|--------|---------------|--------------|--------|------------|------------|----------|----------|--------|------------------------|------|
| Cu     | rGO 2.5 vol%  | Sonication of GO collide in DI water | MLM    | SPS        | 32.4%      | (748 MPa) | –        | –      | –                      | [135]|
| Cu     | FLG 0.5 vol%  | Dip FLGs in Cu(OAc)2 Dry and calcination | PM     | HP         | –          | 15.2%    | (612 MPa) | –      | –                      | [101]|
| Cu     | GNS 0.50wt%   | GO sonication in ethanol | BM     | Hot press sintering (HPS) | ≈24% (≈230 MPa) | –        | –        | –      | ≈24% (≈20%)            | [49] |
| Cu     | rGO & GNP 1.00 wt% | – | BM | HP | – | – | HV(rGO/Cu) > HV(GNP/Cu) > HV (Cu) | – | [63] |
| Cu     | MLG 1.00 vol% | – | Sonication ethanol BM in HRDSR | – | 16.4% (425.5 MPa) | 14% | (360.5 MPa) | – | 16.4% (24.1%) | [100] |
| Cu     | Graphite 0.5 vol% | – | Attrition milling | HP | – | – | – | 14.4% (841 MPa) | CYS | – | [101] |
| Cu     | GO 0.20 wt%   | Treating Cu with PVA | MM     | Heating/CP/sintering | – | – | – | 10% (280 MPa) Compressive strength (CS) | – | [66] |
| Cu     | rGO 0.30 wt%  | Treating Cu with PVA | MM     | Heating/SPS | 22% drop (163 MPa) | 64.2% | (156 MPa) | – | 73% (208 MPa) CS | – | [71] |
| Cu     | GNP 0.30 wt%  | Treating GNP with PVP Treating Cu with PVA | MM | Heating/SPS | 10.5% drop (187 MPa) | 81.3% | (172 MPa) | – | 90% (228 MPa) CS | – | [71] |
| Cu     | rGO 5.00 wt%  | MM of GO in H2O with N2H4·H2O addition | MM | CP/SPS | – | 35.3% | (121.2 MPa) | – | 21.5% | [125] |
| Cu     | GNP 0.50 wt%  | GNP activation in SnCl2 and PtCl2 GNP electroless plating with Ni | MM | SPS | – | 64.5% | (181 MPa) | – | – | – | [127] |
| Matrix | Reinforcement | Pretreatment | Mixing | Compacting | % Increase in Strength | Hardness | Others | % Elongation (original) | Ref. |
|--------|--------------|--------------|--------|------------|-----------------------|----------|--------|------------------------|------|
| Cu     | GNS 1.00 vol%| In-situ chemical reduction Ni-decorated GNSs | Sonication | SPS        | 96% increase in TYS (268 MPa) | –        | 61% (132 GPa) YM | –          | [113] |
| Cu     | GNP 0.13 wt% | GNP sensitization in PVP, activation in SnCl2 and PdCl2, electroless plating with Ni | Sonication | SPS        | 12% increase in UTS (232 MPa) | –        | –      | 60%        | [2]   |
| Cu     | GNP 0.60 vol%| –            | MLM (sonication of GNPs in Cu(NO3)2·3H2O) | SPS        | –                     | 118.3% (310 MPa) | 75% (1.75 GPa) | 65% (147 GPa) YM | [97] |
| Cu     | GNP 0.10 wt% | GNP activation in SnCl2 and PdCl2 | MLM (sonication of Cu acetate and GNPs in H2O) | SPS        | –                     | 52.9% (208 MPa) | –      | –          | [134] |
| Cu     | rGO 0.10 wt% | GO sonication in H2O | MLM (sonication of Cu acetate and GO in H2O) | SPS        | –                     | 25% (170 MPa) | –      | –          | [134] |
| Cu     | rGO 2.50 vol%| Sonication of GO in H2O | MLM (sonication of Cu acetate and GO in H2O) | SPS        | 30% (335 MPa) | 80% (284 MPa) | –      | 30% (131 GPa) YM | [72] |
| Cu     | GNR 1.00 vol%| GNR sonication in ethanol MS of Cu in ethanol | Stirring | SPS/hot rolling | – | 55% (216 MPa) | – | 6.4% | – | [52] |
| Cu     | GO 0.3 wt%   | Sonication of GO in H2O and stirring of Cu in CTAB and H2O | Stirring of GO and Cu | CP/sintering | – | – | 18.5% (51 HV) | – | – | [64] |
| Cu     | GO 0.30 wt%  | GO sonication in H2O MS of Cu in CTAB/H2O | MS | CP/sintering | 13.5% (210 MPa) | ≈18.6% (≈51 Hv) | – | – | [51] |
| Cu     | GNP 0.5 wt%  | GNP sonication in ethyl alcohol | MS | CP/sintering/HP | – | – | ≈44.7% (≈97 HV) | – | – | [126] |
| Cu     | GNS 7.50 vol%| GNS sonication in acetone | MS | HP          | – | – | 44% (97 HV) | – | – | [126] |
Table 5: ...continued

| Matrix | Reinforcement | Pretreatment | Mixing | Compacting | % Increase Strength | Hardness | Others | % Elongation (original) | Ref. |
|--------|---------------|--------------|--------|------------|---------------------|----------|--------|------------------------|------|
| Cu     | Gr            | –            | CVD    | –          | –                   | –        | –      | –                      | [122]|
| Cu     | GNP 10 vol%   | –            | VM     | SPS        | 26% (265 MPa)       | –        | –      | –                      | [93] |
| Cu-0.2 at.%Cr & rGO 2.50 vol% | –            | VM          | Annealing/HP | 82% (352 MPa) | –       | –      | –                      | [128]|
| Cu     | GNF 3.6vol%   | –            | Mixing by pestle and mortar | CP/microwave sintering | –       | –      | 93.5% (89 HV) | – | [137]|
| Cu     | GNP           | –            | Coating GNP with TiC and VC via a controlled reaction in molten salts | MLM | Compaction via SPS | 40% (470MPa) for TiC-GNP/Cu | – | – | 133% increase for VC-GNP/Cu | [96]|
| Cu     | Gr (0.50 wt%) | Ball milling of Cu and PMMA | In-situ growth of Gr in Cu by CVD with PMMA | HPS | – | 35.7% (308 MPa) | 233.3% (290 MPa) | – | – | – | [88]|
| Cu     | Gr 0.95 wt%   | BM Cu and PMMA | In-situ growth of Gr in Cu by CVD with PMMA | HP | – | 274% (274 MPa) | 177% (244 MPa) | 16.2% (143 HV) | – | – | [142]|
| Cu     | GNS           | GNS sonication in acetone | – | ARB (6 cycles) | 71.3% (496 MPa) | – | – | – | 5% (43%) | [123]|

Legend:
- TS: Tensile Strength
- YS: Yield Strength
- Others: Additional properties or notes
### Table 6: Summary of Reported Gr-Reinforced Mg Matrix Composites Studies.

| Matrix   | Reinforcement | Pretreatment | Mixing                  | Compacting                  | % Increase Strength | Hardness | Others | % Elongation (original) | Ref. |
|----------|---------------|--------------|-------------------------|-----------------------------|---------------------|----------|--------|------------------------|------|
| AZ31 Mg  | GNP 0.30 wt%  | –            | BM                     | Sintering / hot extrusion   | 3.5% drop (275 MPa) | 11.3% drop (173 MPa) | 22.4%   | 21.7% (14.5%)          | [48] |
| Mg       | GNP 2.00 vol% | –            | BM                     | SPS                        | –                   | –        | 36.9% (63 HV)          | –    | –                      | [109]|
| AZ31Mg   | rGO 0.40 wt%  | Sonication of GNPs and alloy in hexane | MM BM | CP                  | –                   | –        | 8.6% drop CS           | –    | –                      | [65] |
| Mg-10 wt% Ti | GNP 0.18 wt% | MM of Mg and Ti in ethanol GNP sonication in ethanol | MM | CP / sintering / hot extrusion | 41.1% (230 MPa) | 22.13% (160 MPa) | –      | 14% (3.2%)             | [104]|
| Mg-1 wt% Al | GNP 0.30 wt% | –            | MM                     | CP / sintering / hot extrusion | 139.8% (446 MPa) | 49.6% (178 MPa) | 34.15% (55 HV) | 16.9% (9.7%) | [105]|
| Mg-1 wt% Al | GNP 0.18 wt% | MS of Al Sonication of GNPs MM of Mg (All in ethanol) | MM | CP / sintering / hot extrusion | 30.3% (254 MPa) | 17.3% (190 MPa) | –      | 319% (15.5%) Failure strain | [130]|
| Mg       | GNP 0.30 wt%  | MM of Mg in acetone GNP sonication in acetone | MM | CP / sintering / hot extrusion | 8% (238 MPa)  | 5% (197 MPa)  | 19.3% (68.5 HV) | 10.6% (14.6 GPa) | 3.11% | [106]|
| Mg-1 wt% Al-1 wt% Sn | GNP 0.18 wt% | Sonication of GNPs in ethanol Mixing of alloy in ethanol | MM | CP / sintering / hot extrusion | 14% (269 MPa) | 29.2% (208 MPa) | –      | 10.9% (16.7%) | [107]|
| Mg       | GNP 0.50 wt%  | Sonication of ethanol GNP sonication in ethanol | Sonication MM | HP / sintering | –                   | –        | 20.6% (48.5 HV) | –    | –                      | [108]|
| Mg       | GNP 1.20 vol% | –            | –                      | Sonication / solid state stirring | –                   | –        | 78% (66 kg mm-2) | –    | –                      | [103]|
| ZK60 Mg  | GNP 0.05 wt%  | Sonication of ethanol GNP sonication in ethanol | Sonication Hot extrusion to rod facile melt stirring | Solidification of hot melt Hot extrusion | –                   | 62% (256 MPa) | –      | –                      | [138]|
| Mg-62Zn  | GNP 1.50 wt%  | –            | –                      | DMD                        | Casting of molten metal Hot extrusion | 13.4% (313 MPa) | 34.6% (214 MPa) | 43.3% | 21% (17%)              | [110]|
| AZ61 Mg alloy | GNP 3.00 wt% | –            | –                      | DMD                        | Casting of molten metal Hot extrusion | 11.7% (335 MPa) | 26.1% (232 MPa) | 15.9% | 10.7% (11.5%) | [111]|
Table 7: Summary of Reported Gr-Reinforced Other Metals Matrix Composites Studies.

| Matrix | Reinforcement | Pretreatment | Mixing | Compacting | % Increase Strength | TS | YS | Hardness | Others | % Elongation (original) | Ref. |
|--------|---------------|--------------|--------|------------|---------------------|----|----|----------|--------|------------------------|------|
| Ni     | GNP 1.00 vol% | –            | BM     | SPS        | –                   | 131.2% (370 MPa) | –  | 112.85% (165 HV) | –      | 43% (50%)               | [46] |
| Ni     | MLG (sucrose as C source) | –            | MM     | CP / sintering | 311% (370 MPa) | –  | 69.8% (107 HV) | –      | 25.5% (36.7%)          | [89] |
| Ni     | GNS 1.00 vol% | GNS synthesized via CVD using NaCl template | Impregnation and reduction | SPS | 26% (546 MPa) | 188.4% (474 MPa) | –  | –  | –      | 25.5% (36.7%)          | [90] |
| Ni     | rGO 1.50 wt%  | –            | MLM    | SPS        | 95.4% (948 MPa) | 328% (826 MPa) | –  | –  | –      | 12.1% (34.2%)          | [67] |
| Ni     | rGO           | –            | Electrodeposition | –  | 25% (864 MPa) | –  | –  | –      | –      | 20.6% (15.1%)          | [61] |
| Ni     | GNP 5.00 wt%  | MS of GNP   | Coating AISI410 with GNPs, PVA, and Ni in water | LS | –  | –  | 175.1% (605.2 HV) | –      | –      | [112] |
| Ti     | GO 2.50 wt%   | Sonication of GO | Coating AISI410 with GO and Ti in water | LS | –  | –  | 312.2% (742 HV) | –      | –      | [60] |
| Ti     | GNS 2.50 wt%  | Sonication of GNS | Coating AISI410 with GNSs and Ti in ethanol | LS | –  | –  | 161.1% (470 HV) | –      | –      | [153] |
| Ti     | MLG 0.20 wt%  | –            | BM     | SPS / hot rolling | 91.7% (1050 MPa) | 134.9% (1010 MPa) | –  | –  | 5% (27%) | [114] |
| Ti     | MLG 0.50 wt%  | Sonication of MLG in water | Sonication BM | SPS | –  | 94.1% (918 MPa) | 71.4% (15.39 HV) | 6.4% (264.3 GPa) YM | 10% (30%) | [131] |
| Ti     | GNP 0.10 wt%  | GNP sonication | Sonication BM | SPS / hot rolling | 54.2% (887 MPa) | 57.1% (817 MPa) | –  | –  | 10% (30%) | [115] |
| Ti     | MLG 0.80 wt%  | BM of MLG MM of Ti | BM | SPS / hot rolling | –  | 280% (2 GPa) | 96% | 16% YM | –      | [116] |
| Ti6Al4V| GNF 0.50 wt%  | –            | MM     | HIP Forging | 12.3% (1058 MPa) | 20.1% (1021 MPa) | 14.7% (125 GPa) YM | 9.3% (9.4%) | [117] |
On the high concentration side of Gr reinforcement, several studies have reported its effect on the mechanical properties. One study reported the use of 10 wt%, 15 wt%, and 20 wt% GNPs as reinforcement in the AA2900 alloy [132]. In this study, the Gr-Al composite failed and broke during a standard Brinell hardness test because of the large amounts of Gr particles used. This led to a switchover in the roles of Gr and the metal alloy, where the reinforcement behaved as the matrix and the matrix became the reinforcement. Chu et al. [93] reported interesting results for the GNP-Cu composite. Adding 10 vol% GNPs led to a 26% increase in strength, which significantly dropped to −7.1% when the GNP addition was increased to 20 vol%. The TEM results suggested that GNP debonding occurred during tensile testing for both 10 vol% and 20 vol% GNP additions. At lower volume fractions, the enhancement in strength was attributed to grain refinement and dislocation strengthening. However, the enhancement was hindered by the poor Cu-GNP bonding, which weakened the load transfer ability and resulted in the unsatisfactory strength enhancement of the composites. At 20 vol% GNP content, the GNP delamination played a dominant role over grain refinement and dislocation strengthening, which led to low load transfer efficiency and caused weakening in the composite strength.

Rashad et al. [110] studied the effect of GNP concentration on the mechanical behavior of Mg–6Zn prepared via DMD and hot extrusion. Vickers hardness was measured in the directions longitudinal and perpendicular to the extrusion direction. The hardness values measured in the longitudinal direction were higher than those in the perpendicular direction at any GNP content [110]. However, the effect of changing the GNP content on the hardness measurements was related to the direction in which the hardness was measured. In the longitudinal direction, the hardness increased only until the GNP content was 0.5 wt%, after which the hardness started to decrease. In the perpendicular direction, however, the hardness increased continuously with increasing GNP content up to 1.5 wt%.

**Effect of Graphene Type**

The reinforcement materials (type of Gr) play an important role in the synthesis of GRMMCs. Various reports have shown that the thickness, number of layers, lateral size, and chemical structure of the Gr reinforcement directly affect the mechanical behavior of GRMMCs [51, 58, 63, 71, 87, 124, 126, 129, 134, 140].

The effect of the chemical nature of the Gr derivative used as a reinforcement has been reported by several studies [58, 71, 87, 129, 134] in which authors have compared the use of unfunctionalized Gr, such as GNPs, with functionalized Gr, such as GO. These studies reported contradictory results as to which Gr derivative exhibited better performance as a reinforcement in GRMMCs. Jiang et al. [71] compared the use of GNPs and rGO in a Cu matrix. The results demonstrated that GNP outperformed rGO and offered a better performance as a reinforcement. The GNP-Cu and rGO-Cu composites achieved a 90% and 72% increase in yield strength, respectively, compared with pure Cu. According to the author’s remarks, the absence of oxygen-containing functional groups in GNPs caused fewer structural defects than that in the rGO. Thus, GNP performed better as a reinforcement and enhanced the strength of the metal composite by blocking the dislocation motion. Similarly, Shao et al. [87] and Liu et al. [58] reported that unfunctionalized Gr performed better than rGO as a reinforcement; it achieved higher increases in hardness and strength of the composite for the same content of reinforcement added. In contrast, high mechanical properties of the MMCs reinforced with functionalized Gr were reported in several studies. Zhang et al. [134] produced GNP-Cu and rGO-Cu composites via MLM synthesis. The tensile behavior of both composites at different reinforcement contents is shown in Figure 19. The GNP-Cu composite exhibited higher strength at a lower GNP content, while the rGO-Cu composite exhibited a continuous increase in strength and a higher mechanical performance with increasing rGO content. This is because GNP has fewer structural defects than rGO, which enabled the GNPs to disperse uniformly and achieve better stress transfer and higher strength at a low reinforcement content. Increasing the reinforcement content led to the agglomeration of GNPs and

![Figure 19: Tensile stress-strain behavior of GNP-Cu and rGO-Cu composites at different reinforcement content. Reproduced from Reference [134].](image-url)
Figure 20: (a) Hardness vs. temperature curves of copper matrix composites. (b) Schematic diagrams of the hindrance of atom diffusion generated by graphene and graphite. Reproduced from Reference [124].

hence increased porosity, which restricted stress transfer and decreased mechanical strength. However, no aggregates were found in the rGO-Cu composite at a higher reinforcement content because of its hydrophilic functional groups, which aided in forming stronger interfacial bonding between GO and the Cu matrix. Similar observations and reasoning were given by Asgharzadeh et al. [129].

The thickness and lateral size of Gr have been shown to affect the mechanical behavior of GRMMCs [63, 124, 126]. Dutkiewicz et al. [63] used two grades of GNP s varying in thickness and in lateral dimensions in the Cu matrix. The composite contained GNP s with 2- to 4-nm thickness and size less than 100 nm showed a 50% improvement in hardness compared with that with GNP s of 10- to 20-nm thickness and lateral size less than 14 µm. In addition, SEM images revealed a highly uniform microstructure of the composite with finer GNP s than those containing coarse GNP s. Wang et al. [124] studied the effect of temperature on the hardness of GNP-Cu and graphite-Cu composite (referred to as Cu-GN and Cu-GP, respectively); see Figure 20a. The authors concluded that the hardness of both composites with the same reinforcement content is the same in the range between room temperature and 450°C. At 600°C, however, the hardness of GNP-Cu was found to be twice that of graphite-Cu. They attributed that to the higher efficiency of 2D GNP s to hinder atom diffusion and grain boundary motion at higher temperatures than when using the discontinuous interphase formed by the isolated graphite particles. The mechanism of hindering the grain boundary motion is schematically shown in Figure 20b.

Layered GRMMCs

In addition to bulk GRMMCs with homogenously and randomly distributed Gr reinforcements, layered structured GRMMCs have been investigated and have shown promising results [135, 157–159]. Kim et al. [122] reported the synthesis of a Gr-Cu nanolayered composite that consisted of alternating layers of Cu and monolayers of Gr. They used CVD to grow the monolayers of Gr before transferring it onto the deposited Cu layer to fabricate the Cu-Gr–layered structure. Nanopillar compression tests revealed that the nanolayered composite exhibited an ultra-high compression strength of 1.5 GPa when the repeat layer spacing was 70 nm. TEM investigations indicated a higher density of dislocations was present in the upper Cu layer, while the lower Cu layer showed dislocation starvation. This result was attributed to the strong Cu–Gr interface, which acted as a barrier to dislocation propagation across the interface. The authors explained that Gr acted as an impermeable interface to dislocations, preventing its propagation to the lower layers. This was a result of Gr’s high intrinsic mechanical strength, which prevented its rupture and shearing. The compression tests revealed that Gr was not sheared during deformation up to a total compressive strain of 23%.

Yang et al. [135] synthesized another high-strength layered composite using MLM and SPS techniques. The authors investigated the effects of certain synthesis parameters, such as pH and temperature, on the structure and tensile properties of a rGO-Cu composite. Their analysis suggested that changing the pH value during the MLM process
and the reaction temperature highly influenced the structure of the composite and its tensile properties. The tensile strength of the 2.5 vol% rGO-Cu composite was reported to be as high as 748 MPa for the composite fabricated at a pH of 13.6 and a reaction temperature of 20°C. The authors concluded that at high pH values and low reaction temperatures, the composite attained a hierarchical layered structure that was damaged when the synthesis was conducted at a lower pH and higher temperatures. These results highlight the importance of structural design and suggest that the formation of a hierarchical layered structure can play a crucial role in enhancing the mechanical properties of GRMMCs.

3.3.1.2 The Ductility and Deformation Mechanisms

Attaining high strength in GRMMCs without compromising good ductility is a challenge that has yet to be overcome. Mechanical deformation behavior in GRMMCs was reported to vary based on the metal matrix. The majority of the Gr-Al studies have reported a decrease in the ductility of the composites compared with the initial ductility of the metal [11, 18, 20, 24, 25, 27, 53–57, 70, 94]. The deterioration in ductility has been attributed to the pinning of dislocations by the reinforcement [24] and to the agglomeration of the reinforcement along the grain boundaries with increasing Gr content, resulting in weak interface bonding [11, 57]. Nonetheless, some studies have reported promising results in ductility at certain Gr concentrations along with an enhanced strength of the composite [11, 20, 27, 57, 139, 140].

Recently, Dixit et al. [139] reported a high ductility of 26% tensile elongation for their MLG-Al composite fabricated via FSP. The high ductility was attributed to the good bonding between the Al matrix and Gr reinforcement. In addition, the TEM analysis of a quasi-static crack growth proved that Gr was acting against propagation during the ductile fracture test. Figure 21a shows a Gr sheet arresting a microcrack that was developed upon loading. Yan et al. [118] synthesized 0.5 wt% GNF-reinforced aluminum matrix composite by ball milling followed by HIP and extrusion. An increase in the tensile strength from 373 MPa to 467 MPa was reported for pure Al and Gr-Al composite, respectively, without affecting the uniform elongation and ductility of the metal. This finding was attributed to the wrinkled structure of Gr that was preserved even after milling. Upon plastic deformation, the wrinkled structure of Gr straightens and flattens, thereby preserving the good ductility of the metal. Li et al. [53] observed wrinkling of GNP sheets after fabrication, which included ball milling. However, the 0.2 wt% GNP-Al witnessed a reduction in tensile elongation from 11% to 4% due to the formation of microcracks; see Figure 21b. The authors explained that the wrinkles and folds in the GNPs would partially unfold with initial stretching. However, the GNPs couldn’t keep up with the ductile Al, which further deforms with increasing the load, leading to the rapid expansion of the microcracks at the interface and deterioration of the composite’s elongation. In contrast, promising results have been reported by Yolshina et al. [140], who synthesized Gr-Al composites through a one-step process that took place directly in molten aluminum under alkali halides melt. The compos-
ites exhibited simultaneous increase in hardness, elastic modulus, tensile and yield strength with increasing the Gr content up to 2 wt%. Additionally, the authors reported an increase in tensile elongation to reach 26.2% for their 2 wt% Gr-Al composite compared with 13% elongation for pure Al.

Similar to Al, the ductility of pure, ductile, malleable, and easily processed Cu has been reported to drop significantly when adding Gr [49, 51, 71, 93, 97, 123, 125, 127, 134]. The deterioration in ductility has been attributed mainly to the pinning of dislocations by the reinforcement [71] and the agglomeration of the reinforcement along the grain boundaries with increasing Gr content, resulting in weak interface bonding [97]. In addition, several studies have attributed the deterioration in ductility to cracking, which tends to initiate in the grain boundaries where Gr is mostly located and/or at the interface between Gr and Cu, before cracks extend and propagate to the matrix [51, 71, 123]. Nevertheless, unprecedented enhancement in ductility of Gr-Cu compared with pure Cu was reported by Yang et al. [52]. They achieved a 39% increase in ductility over pure Cu for the 3 vol% Gr-reinforced Cu matrix composites prepared via the facile solution-based method followed by SPS and hot rolling. It is crucial to note that the study utilized GNR rather than rGO or GNSs; hence, the promising enhancement in ductility had been associated with the type of Gr derivative used and its distribution. The authors attributed the ductility to the uniform distribution of the GNRs inside the grains, which provided rigid interfacial areas that acted both as dislocation initiation sites and dislocation sinks pinning dislocation motion at the same time. This allows for achieving high strength by dislocation pinning without sacrificing the good ductility of the matrix by generating new dislocations. In addition, the intergran uniform distribution of the GNRs avoided aggregation at the grain boundaries and alleviated the tendency to induce large stress concentrations and microcrack generations at grain boundaries. This behavior has been reported several times with the use of GNSs [51, 71, 123]. Other studies maintained decent ductility or tensile elongation percentages along with the associated strength enhancement of the composites. Jiang et al. [2] produced Ni-plated GNP-Cu composite and reported a 60% elongation, which is better than that of unreinforced Cu. Yue et al. [49] produced rGO-Cu by ball milling and HP using varying GO content. The optimized GO concentration was reported to be 0.5 wt%, which resulted in a 28% increase in UTS over pure Cu and a 24% tensile elongation. This had been attributed to the good interface bonding as a result of the successful distribution of rGO nanosheets in copper. The authors explained that with low GO content, GO sheets can hinder crack propagation by shielding the cracks. This is facilitated by their high aspect ratio and large contact area with the matrix. However, increasing the GO content more than 0.5 wt% resulted in rGO aggregation, providing sites for composite cracking. The same behavior and justification had been reported by Zhang et al. [134] for a 0.1 vol% rGO-Cu composite. SEM images showed the ductile behavior of the composite prepared by molecular-level mixing and SPS. Dimples were clearly distributed on the fracture surface, demonstrating a typical plastic deformation fracture behavior. The ductility behavior of the samples in References [2, 49, 52], and [134] are shown in Figures 22a, 22b, 22c, and 22d, respectively.

Unlike the face-centered cubic metals (Al and Cu) discussed previously, in which the increase of reinforced mechanical strength and hardness is mostly accompanied by a deterioration in ductility, the Gr-reinforced Mg matrix composites showed a significant improvement of ductility and strength compared with pure Mg or its alloys processed at the same conditions [48, 104, 105, 110, 111]. Rashad et al. [104, 105, 110] reported a significant increase in the ductility of the Mg matrix composites by adding GNPs. They reported an increase in tensile fracture strain from 17% for Mg-6Zn to 21% for Mg-6Zn-1.5 wt% GNPs [110], from 9.7% for pure Mg to 16.9% for Mg-1Al-3 wt% GNPs [105], and from 3.2% for pure Mg to 14% for Mg-10Ti-0.3 wt% GNPs [104]. SEM images of the tensile fracture surfaces of Mg-6Zn-1.5 wt% GNP are shown in Figure 23, where ductile fracture features are represented by dimples and tear ridges. An improvement in ductility has been attributed to the high surface area of GNPs and good adhesion to metal matrix [104] as well as the uniform dispersion of GNPs throughout the matrix [110].

### 3.3.2 Thermal and Electrical Properties

#### 3.3.2.1 Thermal Behavior

Different parameters can affect the thermal properties of Gr-reinforced metals, such as the composite’s thermal conductivity, thermal diffusivity, and CTE. Yolshina et al. [140] studied the effect of Gr content on the melting point of the Gr-Al composites. They reported a decrease in the melting point of the Gr-Al composite with an increase in the Gr content. Enhancements in the thermal conductivity of metal matrices after adding Gr have been reported for Cu [51, 160, 161] and Al [50]. Reddy et al. [50] investigated the effect of rGO content on the thermal conductivity (TC) of a rGO-Al composite. The TC is reported to increase from 48.5 W/mk to 58.5 W/mk for Al and 0.5 wt% rGO-Al composite, respectively. Gao et al. [51] reported an increase of the TC
Figure 22: Tensile engineering stress-strain curves for (a) GNR-, (b) GNP-, and (c) GNS-reinforced Cu matrix composites. (d) SEM of fracture surfaces of 0.1 vol% RGO/Cu composites. Reproduced from Reference [52].

Figure 23: Tensile fracture images of as extruded (a) Mg–6Zn alloy, (b) Mg–6Zn–0.5GNP, and (c) Mg–6Zn–1.5GNP composites. Reproduced from Reference [110].
of their Gr-Cu composite with an increase in Gr content to reach a maximum value of 396 Wm$^{-1}$K$^{-1}$ at 0.3 wt%, after which a decrease in the TC was measured. The enhancement of the TC was ascribed to various reasons, such as the higher TC of Gr over pure Cu as well as the electrostatic self-assembly synthesis technique, which allowed for a uniform distribution of Gr in Cu and resulted in a lower interface thermal resistance. However, a higher Gr content resulted in Gr agglomeration in the composite and, hence, formed structural defects such as porosities and voids. These structural defects interfered with the phonon transfer between the Cu matrix and the Gr reinforcement, leading to a decrease in the TC of the composite [51, 97]. The effects of Gr alignment in the Gr-Cu composite on the TC of the composite are reported in References [160–162]. In these studies, the experimental and modeling results proved that achieving the highest possible TC requires the GNP alignment to be highly aligned in the metal composite. The studies concluded that the TC of GRMMCs is highly anisotropic, such that the enhancement achieved is different based on the conductivity's direction—whether perpendicular or parallel to the direction of alignment. In addition, the enhancement of the TC in one direction of the alignment was attained at the expense of the conductivity in the other direction [160, 161]. Chu et al. [161] controlled the alignment of GNS in the Cu-GNS composites via vacuum filtration and SPS. They reported a substantially higher in-plane (perpendicular) TC than through-plane (parallel) TC with a 35% increase in the in-plane TC than that of pure unreinforced Cu.

The thermal diffusivity of Gr-Cu composites has been studied. Chen et al. [97] and Si et al. [96] reported a decrease in the thermal diffusivity after adding GNP to the Cu matrices. Chen et al. [97] reported a constant deterioration of thermal diffusivity measured in both the horizontal and parallel consolidation directions with increasing the GNP content. This behavior was attributed to the poor bonding and large mismatch in the CTE between Gr and Cu, which formed voids and increased the interfacial thermal resistance. Si et al. [96] explained that the increase in thermal resistance is due to the poor interfacial structure of Gr-Cu composites. This led to poor electron-phonon coupling, which is considered to be the main thermal transport mode in Gr-Cu composites. The CTE for Gr-Cu composites has also been studied [101, 124, 161]. It was reported to decrease after adding Gr. Wang et al. [124] attributed this decrease in CTE to the distribution of Gr in the composite and the effective drag force it applied on the grain boundary motion, which hindered grain growth at elevated temperatures.

### 3.3.2.2 Electrical Behavior

The effects of varying Gr content on the electrical conductivity of GRMMCs have been reported. The electrical conductivity was reported to decrease with increasing the Gr content [2, 52, 63, 96, 97, 137]. However, several studies have suggested that the excellent electrical conductivity of metals can be maintained after adding Gr. Li et al. [53] studied the effects of adding GNP on the electrical conductivity of Al and reported a trivial 0.7% reduction in electrical conductivity for their 0.2 wt% GNP-Al composite. This finding was attributed to the low GNP content, suggesting that the Al matrix dominates the conductivity behavior and that phase interface scattering between Al and GNP is very limited. A similar behavior was reported by other studies, where the electrical conductivity of Gr-Cu composite is trivially reduced and high conductivity is still achieved [63, 71, 72, 96, 97, 124, 135, 137]. The decrease in the electrical conductivity of Cu was justified by the presence of the Gr-Cu interface, resulting in a reduction of the electrons’ mean free path as well as the scattering of electrons at the interface [71, 97].

Recent attempts have successfully reported enhanced electrical conductivity of Gr-Cu composites over pure Cu [52, 137, 163]. Yang et al.’s [52] results proved that enhancing the strength of metal composites by adding Gr does not have to be at the expense of electrical conductivity. Their composite achieved a 5% enhancement in the electrical conductivity of GNRs over pure Cu for 1 vol% addition of GNRs. When the GNR content was increased to 3 vol%, the electrical conductivity dropped below that of 1 vol%. However, this value is still 2.7% higher than that of pure Cu. The enhanced electrical conductivity was attributed to the unique geometry of the GNRs as well as its direction in the metal matrix, which was controlled via the hot-rolling process. In addition, the authors suggested that the clean and closely bonded interface reduced the electron scattering and, hence, minimized interfacial impedance. Last, the addition of GNRs in this study resulted in extra paths for electron transport. The electrical conductivity of Ayyappadas et al.’s [137] GNF-Cu microwave-sintered composite was reported to increase from 92% to 94% compared with pure Cu when the addition of GNFs was 0.9 vol%. This result was attributed to the homogenous distribution of the highly conductive GNFs in the matrix. Increasing the Gr content to 1.8 vol% led to maintaining a 92% electrical conductivity, which is similar to that of pure Cu. However, further increasing the GNFs content to 3.6 vol% achieved an 86% electrical conductivity, which is 6% less than that of pure Cu. The decrease in electrical conductivity was correlated to the introduction of porosities in the compos-
ite with increasing the GNF content, which interrupted the continuity of the conductive path necessary to achieve high electrical conductivity values. Recently, more promising results were reported by Cao et al. [163] for a layered Gr-Cu composite that was synthesized using CVD and hot pressing. The layered composite achieved the highest electrical conductivity for a Gr-Cu composite of a 117% International Annealed Copper Standard (IACS) with the addition of only 0.008 vol% Gr. The authors used first-principles calculation results based on the density functional theory to understand the resultant significant electrical conductivity. They attributed the high electrical conductivity of the embedded Gr to a doping effect that resulted in an increased carrier density and an unaffected, very high carrier mobility in Gr. Furthermore, the authors investigated the effect of changing the number of Gr layers on the electrical behavior of the composite through a careful control of the carbon source concentration and the growth time during the CVD process. The results suggested that the electrical conductivity in the Gr layers decreases with increasing the number of layers.

The effects of varying the Gr type in the composite on the electrical behavior of Gr-Cu composites have also been noted [63, 71, 124]. Wang et al. [124] compared the electrical conductivity when using GNPs and graphite in Cu composites. Compared with the 74.1 IACS% reported of graphite-Cu, the GNP-Cu composite was able to maintain 95.9% IACS. Jiang et al. and [71] and Dutkiewicz et al. [63] investigated the effect of reinforcing Cu with rGO and GNPs on the electrical conductivity and resistivity, respectively. Jiang et al. [71] reported a better electrical performance for the GNP-Cu with 84.2% IACS compared with 73.4% IACS for the rGO-Cu composite. Dutkiewicz et al. [63] reported a better electrical performance for rGO-Cu with only a 28% increase in electrical resistivity, compared with 57% for the GNP-Cu composite. While Dutkiewicz et al. [63] gave no justification, Jiang et al. [71] ascribed their results to the already better inherent electrical performance of GNPs compared with GO due to the inevitable defects in GO sheets after oxidation.

### 3.3.3 Surface Properties

#### 3.3.3.1 Corrosion Behavior

The effects of the added reinforcement on the corrosion-resistant behavior of Gr-metal composites have been reported for Al [11, 140], Mg [108], and Cu [164–166]. The influence of the added reinforcement on the corrosion resistance is a main challenge in the fabrication of metal composites, especially in Al composites. Al is known for its corrosion-resistant properties due to its protective oxide film formed on the surface. The addition of another phase to Al, such as Gr, can cause discontinuities in the protective film, creating corrosion-initiating sites. Rashad et al. [11] and Yolshina et al. [140] studied the corrosion behavior of the Gr-Al composite after immersing the composite in 3.5 wt% NaCl solution. In this study [11], the potentiodynamic polarization measurements for the composite synthesized via powder metallurgy and hot extrusion showed an increase in the corrosion rate after adding GNPs. The authors suggested that the cathodic nature of Gr compared with the Al matrix leads to galvanic corrosion in the presence of the electrolyte. Yolshina et al. [140] reported similar observations for the Gr-Al synthesized by a one-step in-situ synthesis in molten Al under an alkali halides melt. The corrosion rate increased with increasing GNP concentration compared with the corrosion rate of pure Al. However, the authors suggested that the increase in corrosion rate from 0.0016 g/m²·h to 0.0048 g/m²·h for pure Al and 2 wt% GNP-Al composite, respectively, is minimal and that the composite is still classified as a class 4 corrosion-resistant material. SEM images of clean and corroded pure Al and 0.5 wt% Gr-Al composites are shown in Figure 24.

More recently, Jin et al. [164] studied the corrosion behavior of a Gr-Cu composite prepared by growing Gr on Cu by heating PMMA-coated Cu at 900°C followed by HP of the powder flakes to form a bulk composite. The corrosion rate of the composite was reported to decrease by 50% in NaCl. Several tests were conducted to evaluate the enhanced corrosion behavior of the synthesized Gr-Cu composite including etching behavior test in FeCl₃ solution, salt spray corrosion test, and electrochemical corrosion test. According to the results, Gr clearly showed anticorrosive advantages, but its performance was anisotropic; its orientation in the Cu matrix influenced the corrosion pathway. In addition, it was concluded that the defects present in the Gr structure are detrimental to its corrosion protection behavior and act as sources that promote corrosion. Finally, the tests confirmed that the corrosive media had to penetrate more Gr layers if the corrosion proceeded in the in-plane direction, offering higher corrosion resistance than when the corrosion proceeded in the cross-plane direction where more Cu channels were present. These results demonstrate the potential applications of such composites, especially when the enhancement in corrosion behavior is associated with substantial mechanical hardening and good electrical conductivities.
3.3.3.2 Tribological Behavior

Graphene and its derivatives are expected to have a positive influence on the tribological properties of MMCs, such as wear and friction behavior. This is attributed to its self-lubricating nature that stems from the squeezing of the Gr sheets in between sliding surfaces and its loadbearing capability, which reduces the wear rates of metal matrices [47].

The effects of changing the Gr content on wear rates and coefficient of friction (COF) have been investigated [47, 99]. It has been reported that the wear rates and COF decrease for Gr-Al composites compared with plain Al-based matrices [47, 99] and for Cu matrices [96, 97, 137] up to certain Gr content, after which an increase in wear rates and COF is reported [64, 126]. El-Ghazaly et al. [47] studied the tribological properties of hot-extruded GNP-reinforced AA2124. They reported superior wear behavior of the composite compared with the alloy. The authors reported a 34% reduction in the wear rate and a 22.5% reduction in the COF for the 0.3 wt% GNP-AA2124 composite.

Figure 24: Before and after SEM images of (a, b) pure Al and (c, d) 0.5 wt% GNP-Al composite. Reproduced from Reference [140].
which were delaminated between the rubbing surfaces. When small amounts of GNPs, below 0.8 vol%, were added, the wear rates meant exposing more GNPs to the surfaces and hence were smeared out uniformly on the sliding surfaces, providing a continuous graphene layer between the rubbing surfaces and hence increasing wear resistance.

Tabandeh-Khorshid et al. [99] reported a decrease in COF and an increase in wear rate with increasing the GNP content from 0.1 wt% to 1 wt%. They suggested that higher wear rates mean exposing more GNPs to the surfaces and thus higher lubrication during sliding, which led to a lower COF. Chen et al. [97] reported that the COF did not change when small amounts of GNPs, below 0.8 vol%, were added to Cu. However, increasing GNPs to 4.0 vol% led to a 60% reduction in COF over pure Cu. The trivial decrease in COF at a lower Gr content was justified by the quick wearing of the Gr thin film that forms at the worn surfaces. In contrast, increasing the Gr content formed a thick continuous film that prohibited oxidation and reduced friction. Si et al. [96] reported a drastic reduction in wear rates of the Gr-Cu composite with GNP concentrations starting from 0.9 vol%, with a minimum wear rate of 0.28×10 ^{-5} g/m for their 3.6 vol% GNP-Cu composite at a load of 30 N. This was attributed to the interlayer shearing mechanism of Gr that prevented metal-to-metal contact and led to lower wear rates and COFs.

Gao et al. [64] and Li et al. [126] reported that there is a critical Gr content above which the tribological behavior of the Gr-Cu composite deteriorates. In their work [64], the COF is first decreased to 65% lower than that of pure Cu by adding 0.3 wt% GNP, and then it increases when the GNP is increased to 0.5 wt%. The authors explained that at a high GNP content, GNPs were located at the grain boundaries, which led to the poor sintering ability of the 0.5 wt% GNP-Cu composite and resulted in Cu grains separating during the friction test. Li et al. [126] reported a similar behavior for the wear rate of the GNS-Cu composite, which decreased from 13.6×10^{-4} to 2.3×10^{-4} mm^3/ma as the volume fraction increased from 2.5 vol% to 7.5 vol%, but later it increased as the GNS content increased to 10 vol%. This was justified by the poor mechanical behavior and the load-bearing ability of the 10 vol% GNS-Cu composite, where the hardness dropped from 97.4 Hv to 56.8 Hv when the GNS content increased from 7.5 vol% to 10 vol%, respectively. Similarly, the relationship between the wear behavior and the mechanical properties was referred to and addressed by El-Ghazaly et al. and Si et al. [47, 96].

### 4 Summary

This review focused on the three main pillars in bulk GR-MMC studies: synthesis, structure, and properties. First, the variety of techniques used to produce GRMMC have been summarized and categorized based on the steps involved in the synthesis: pretreatment of metal and Gr, mixing, and postsynthesis/consolidation. Regardless of the synthesis route chosen, a synthesis technique was assumed to be successful when the composite attained two main structural features: a homogenous distribution of Gr within the metal matrix and a defect-free clean matrix-reinforcement interface. These two features facilitate mechanical load, electron, or phonon transfer between the metal matrix and the Gr reinforcement, all of which directly affect the enhanced response and behavior of the GR-MMCs. In addition, studies showed that different metals have different interactions with Gr, and many challenges persist regarding the interfacial interaction formation. For example, the lack of second-phase formation had been attributed to the poor solid solubility between Cu and C in Cu-Gr systems, and to the poor wettability between Mg and Gr in Mg-Gr systems. Nonetheless, the formation of a second phase between Al and C in Al-Gr systems has been highly investigated and is considered a point of debate among researchers (i.e., whether the formation of the AlC phase is caused by the exposure of the composite to high temperatures during synthesis or is initiated by the disorder and defect points in the graphitic structure regardless of temperature). Furthermore, the role such phase played in the behavior of the composites is also a point of debate. Some studies suggested that it caused detrimental effects on the mechanical behavior of Al-Gr composites due to the structural damage involved with the formation of AlC, while other studies suggested that it enhanced the mechanical strength and hardness by pinning dislocations.

Impressive enhancements in the tensile and compressive strengths and hardness have been reported in studies utilizing relatively small concentrations of Gr, regardless of the metal type. In these studies, there were optimized concentrations of Gr, above which an observed deterioration in the behavior was reported. Such behavior had been attributed to the difficulty in homogeneously distributing larger reinforcement content in the matrix, lead-
ing to agglomeration and poor contact between the reinforcement and the matrix. The ductility or mechanical deformation behavior in GRMMCs was reported to vary based on the metal type. In Al-Gr and Cu-Gr systems, most studies reported a decrease in the ductility of the composites compared with the initial ductility of the metal. In Mg-Gr systems, a simultaneous improvement in ductility and strength compared with pure Mg or its alloys processed at the same conditions has been reported. Another point of investigation highlighted in this review is the effect of the Gr derivative type and morphology on the overall behavior of the GRMMCs. Various reports have shown that the thickness, number of layers, lateral size, and chemical structure of the Gr reinforcement has directly affected the mechanical behavior of GRMMCs. So far, the main interest has been comparing the use of unfunctionalized Gr, such as GNPs, with functionalized Gr, such as GO. Some researchers have suggested that GNPs performed better as reinforcement due to the absence of oxygen-containing functional groups in GNPs, which would result in lower structural defects. Other researchers concluded that the hydrophilic functional groups in GO aided in stronger interfacial bonding between GO and the metal matrix, allowing the composite to withstand higher concentrations of the reinforcement without aggregation.

With the technological advancements in every known area, the demand for high-performance materials is more pressing than ever. Combining Gr into bulk metals promises enhanced mechanical performances and superior strength-to-weight ratios, as well as the expected enhancements in thermal and electrical performances. The field of GRMMCs is still in its early stages, but the increasing number of publications on this topic annually and the reported results prove both its merit and future. At the end of this review, it is important to highlight that inconsistent experimental findings with regard to the behavior and response of GRMMCs are reported and explained throughout the literature. Discrepancies arise largely from the large number of variables involved in different studies, such as the different Gr derivatives used, the metal type investigated, the handling of the materials, the synthesis technique of choice, and the amount of Gr reinforcement used. Thus, a more systematic approach toward producing and characterizing these composites is required to achieve a more comprehensive understanding of the behavior and response of these composites.

Acknowledgement: This work was made possible by NPRP Grant no. NPRP9-180-2-094 from the Qatar National Research Fund (a member of the Qatar Foundation). The statements made herein are solely the responsibility of the authors.

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