Applications of Graphene and Its Derivatives in Bone Repair: Advantages for Promoting Bone Formation and Providing Real-Time Detection, Challenges and Future Prospects

Abstract: During continuous innovation in the preparation, characterization and application of various bone repair materials for several decades, nanomaterials have exhibited many unique advantages. As a kind of representative two-dimensional nanomaterials, graphene and its derivatives (GDs) such as graphene oxide and reduced graphene oxide have shown promising potential for the application in bone repair based on their excellent mechanical properties, electrical conductivity, large specific surface area (SSA) and atomic structure stability. Herein, we reviewed the updated application of them in bone repair in order to present, as comprehensively, as possible, their specific advantages, challenges and current solutions. Firstly, how their advantages have been utilized in bone repair materials with improved bone formation ability was discussed. Especially, the effects of further functionalization or modification were emphasized. Then, the signaling pathways involved in GDs-induced osteogenic differentiation of stem cells and immunomodulatory mechanism of GDs-induced bone regeneration were discussed. On the other hand, their applications as contrast agents in the field of bone repair were summarized. In addition, we also reviewed the progress and related principles of the effects of GDs parameters on cytotoxicity and residues. At last, the future research was prospected.

Keywords: graphene, graphene oxide, bone repair, real-time detection, signaling pathways

Introduction
In recent years, due to the increasing incidence of bone defects caused by different reasons, such as post-traumatic, degenerative, neoplastic or congenital, bone transplantation has become the second largest tissue transplantation after blood transfusion.\(^1\)\(^-\)\(^3\) Although autogenous bone transplantation is still considered as the gold standard for bone repair, the application of this method is limited owing to the lack of autogenous materials and the occurrence of secondary defects at the bone donor site.\(^4\)\(^-\)\(^6\) Consequently, there has been an urgent need for the development and application of bone repair materials. During continuous innovation in the preparation, characterization and application of various bone repair materials for several decades, nanomaterials have exhibited some unique advantages. For example, the structure of some nanomaterials is similar to that of the natural bone, and they have special characteristics such as outstanding mechanical properties, desired electrochemical capacities, increased surface area and satisfactory wettability,\(^7\)\(^-\)\(^8\) which
could not only provide structural support for cells, but also regulate their proliferation, differentiation and migration, and ultimately improve the effect of bone repair.9,10

Currently, as a representative two-dimensional nanomaterial, graphene and its derivatives (GDs) have been shown promising potential for the application in bone repair based on their excellent mechanical properties, electrical conductivity, large specific surface area (SSA) and atomic structure stability.11,12 It has been demonstrated that GDs can provide sufficient mechanical reinforcement to bone repair scaffolds,13 bring desired electrical stimulation to cell osteogenic activities and bone formation,14 and be conducive to the adsorption of active substances.15 Moreover, the hexagonal single-layer carbon atomic structure of graphene makes strong structural stability, which is not easy to be destroyed during complex scaffold preparation process with organic solvents and at the implantation sites under physiological environments.16 But, pure graphene particles are difficult to form a 3D scaffold themselves and have poor fluidity to inject into the body, so they have been normally compounded with other substrates to use in bone repair.17 However, because pure graphene particles are easy to agglomerate due to strong van der Waals force, which makes it difficult for GDs to uniformly disperse into composites,18 they usually need to undergo functionalization before use.19,20 In most cases, the preparation of graphene oxide (GO) is the first step for their functionalization.21,22 On the basis of GO, it is possible for carboxyl,23 amino,24 hydroxy25 and other functional groups to be introduced not only to improve the fluidity and dispersion of graphene, but also to further endow it with new functions, among which is that various active substances can be loaded to improve bioactivities.26,27 At present, the functionalized GDs have been compounded with various substrates such as metals,28 inorganic,29 natural polymer30 and synthetic polymer31 to prepare scaffolds,32,33 coatings,34 membranes35 and injectable hydrogels.36 Lots of studies in vitro and in vivo have demonstrated that the GDs-containing composites could regulate the extracellular micro-environment to effectively promote bone regeneration. It has been shown that GDs could not only promote the adhesion, proliferation and mineralization of osteoblasts,37,38 but also maintain the activity of bone marrow mesenchymal stem cells and those derived from soft tissues and induce them to differentiate into osteogenic cells.39,40 Further studies into related signaling pathways demonstrated that the GDs achieved the osteogenic performances by activating some specific signaling pathways, such as Wingless/Integrated (Wnt)/β-catenin,41 Phosphatidylinositol kinase (PI3K)/

Protein kinase B (Akt)/Glycogen synthase kinase-3 (GSK-3β)/β-catenin,42 Mitogen-activated protein kinases (MAPK)43 of mesenchymal stem cells, and Wnt and bone morphogenetic protein (BMP)44 of monocytes and macrophages. Moreover, in vivo experiments of the skull and radius defect models of various animals such as murines,45 rabbits46 and canines47 have demonstrated that the GDs could significantly promote bone repair.

In addition, it has been indicated that GDs could be applied to real-time detection of bone repair process. After GDs were modified with functional groups or combined with fluorescent/magnetic molecules, the imaging efficiency of the GDs-containing composites with various detection equipment such as computed tomography (CT),48 ultrasound49 and nuclear magnetic resonance (NMR)50 could be significantly improved, by which the structural integrity,51 tumorigenicity,52 degradability,53 and mineralization54 of the composites could be real-timely evaluated, which was definitely helpful for further and deeper study of the bone repair process. Compared with most of other substrate contrast agents, GDs could be more conducive to the quantitative control of loaded fluorescent markers or paramagnetic substances55 based on their excellent conductivity and large SSA. Moreover, the imaging performances of GDs could be further changed and optimized by functionalization.56

Although GDs have exhibited attractive advantages in the bone repair application, as shown above, there are still some concerns, the main two aspects of which are their potential toxicity and possible residue in vivo. The current related studies demonstrated that the potential toxicity of GDs was dependent on their size, concentration, functionalization and formed structure, etc. For example, the graphene nanosheets with smaller size (<100 nm) had bigger potential toxicity than those with larger size (>400 nm).57 The concentration of GDs greater than 10 μg/mL might inhibit the proliferation of bone marrow mesenchymal stem cells.58 Functionalization could decrease toxicity potential of GDs to some extent.59 In addition, the GDs in sheet structures60 or compounded with other materials61 could bring lower toxicity risk because of decreased possibility in penetration into the cell membrane. On the other hand, the degradation of GDs has been shown to be closely related to their layer number, introduction of other atoms, binding of functional groups, and compounding with degradable polymers. For example, improving the dispersion of GDs was beneficial to increase the contact between GDs and enzyme, thus promoting the degradation
of GDs. Single-layer or low-layer GDs could be relatively more easily degraded by specific enzymes around the implantation site, such as peroxidase. Covalent bonds of some specific functional groups such as phosphate groups could regulate the biodegradation of GDs by improving the interactions between the GDs and enzymes. Moreover, compounding with some degradable polymers such as polycaprolactone (PCL) was demonstrated beneficial to the excretion of GDs with the degradation products.

Herein, we reviewed the updated application of GDs in bone repair in order to present as comprehensively as possible their specific advantages (Figure 1), challenges and possible solutions. Finally, future prospects were proposed.

**Utilization of Advantages of GDs in Bone Repair Materials**

As shown above, GDs have exhibited many advantages that present great potential to directly or indirectly promote bone repair (Figure 2). In this section, how the various advantages of GDs have been utilized in bone repair materials was discussed. Especially, the effects of further functionalization or modification of GDs were emphasized.

**Outstanding Mechanical Properties**

It has been well recognized that the mechanical properties of bone repair materials have important effects on their performances. In this subsection, the main factors that have been shown to significantly influence the reinforcing effect of GDs were focused.

Graphene is a monolayer carbon atom composed of sp² hybrid C-C bond, which has excellent mechanical properties because of its huge bond energy. At present, graphene is considered to be the strongest substance and harder even than diamond, Young’s modulus, stiffness and fracture strength of which could be up to 1.0 TPa, 2.0 TPa and 130 GPa, respectively. Current studies have...
demonstrated GDs possess an outstanding potential to significantly reinforce bone repair materials, the effect of which is mainly dependent on the layer number, content and uniformity of the GDs in the matrix, and the binding mode between GDs and the matrix.

Firstly, it should be noticed that the increase of graphene layers could normally lead to the decrease of their mechanical properties, which was mainly due to the fact that there was no covalent bond connection between the layers, but only a relatively weak van der Waals force.\textsuperscript{75–77} For example, Eqra et al\textsuperscript{75} reinforced epoxy resin with <10 layers of graphene (GEC10) and <30 layers of graphene (GEC30), the results of which showed that when the content of graphene was 0.1 wt.%, the tensile strength of GEC10 and GEC30 samples both reached the maximum, and the peak tensile strength of GEC10 samples was 4.8% higher than that of GEC30 samples.

Another aspect that needed to be highlighted was that with the increase of GDs content in the matrix, the mechanical properties of the composites would normally firstly increase and then decrease, which was mainly caused by agglomeration of GDs and the change of reinforcing mechanism.\textsuperscript{78–80} For example, Tang et al\textsuperscript{78} introduced graphene into barium titanate/poly(methyl methacrylate) composites in order to obtain bone repair materials with both biological activity and satisfactory mechanical properties. As shown in Figure 3A–F, the results showed that the introduction and dispersion of graphene had a significant influence on the reinforcing effect. When the graphene content was 0.5 wt.%, the compressive strength of the composite increased from 83.5 MPa to 89.5 MPa. However, with the further increase of graphene content, the compressive strength of the composites decreased gradually. Due to the accumulation of graphene, there was a phenomenon of stress concentration in the composites, which increased the possibility of their failure. When the content of graphene increased to 2.5 wt.%, the agglomeration of graphene accelerated the decrease of compressive strength of the composites. The results demonstrated that GDs could prevent the propagation of cracks, thus significantly improving the mechanical properties of the matrix. However, when the graphene content increased to some extent, the dispersion of graphene began to decrease obviously observed in the corresponding cross-section micro-morphology (Figure 3B–F), which led to the formation of cohesion and stress concentration, and increased the possibility of failure of the composite. On the other hand, Gao et al\textsuperscript{79} synthesized B\textsubscript{4}C composite ceramics with different graphene content from 0.4 wt.% to 1.2 wt.% by pressureless sintering, and they found that with the increase of graphene content in the matrix, the mechanical properties of the composites increased at first and then decreased, and got to the biggest value when the graphene content was 0.8 wt.% (Figure 3G). Most importantly, they found that different content of graphene led to two different reinforcing mechanisms. For example, the whole graphene layer would be pulled out when the composite with the graphene content of 0.4 wt.% was broken (Figure 3H). The fracture only needed to overcome the adhesion between graphene and B\textsubscript{4}C matrix. However, when the graphene content is 0.8 wt.%, the fracture first occurred in the outermost layer of graphene, and then the inner layer of graphene was pulled out (Figure 3I). The fracture needed to overcome both the resistance of graphene and the bonding force between graphene and B\textsubscript{4}C matrix, thus absorbing more fracture energy and preventing crack propagation. As shown in Figure 3G, with the increasing of graphene content, the porosity of the composites decreased at first and then increased. B\textsubscript{4}C substrate and graphene combined most closely when the graphene content was 0.8 wt.%. When the graphene content was 1.0 wt.%, isolated graphene sheets appeared in the interstice of grains, and this phenomenon was more serious when the graphene content was 1.2 wt.%. So, it was indicated that the different addition content of graphene led to the different changes of the microstructure of the composites, which further led to different reinforcing mechanisms of the graphene, and thus affecting the final reinforcing effect.

Moreover, the functionalization of GDs could also significantly influence the reinforcing effect mainly from two aspects, one of which was that enhanced binding between GDs and matrix might be achieved after the functionalizations,\textsuperscript{81,82} and the other of which was that appropriate functionalizations could improve dispersion of GDs in the matrix, thereby getting better structural uniformity.\textsuperscript{83,84} For example, Ionita et al\textsuperscript{81} prepared porous polyvinyl alcohol Gel-polyvinyl alcohol (Gel-PVA) /GO materials for bone repair by freeze-drying method. Their subsequent results showed that addition of the GO could enhance the compressive strength of the material by 100%. One of the main reasons to achieve such a satisfactory reinforcing effect might be that the covalent binding between -OH on the end group of GO and -COOH on the PVA chain led to the helical recombination.
of the gel chain and improved the crystallinity of the PVA domain, which further led to more uniform structure in the composites. On the other hand, Zhang et al. found that polydopamine functional reduced graphene oxide (PDA-rGO) had a significantly better reinforcing effect than GO in waterborne polyurethane (WPU) matrix mainly because polydopamine (PDA) on the surface of GO prevented the accumulation and aggregation of GO layers, resulting in the uniform dispersion of PDA-rGO in the WPU matrix, and the functional groups such as hydrophilic amine, imine and catechol on the surface of PDA-rGO could react with the -NCO group on the WPU molecular chain to form relatively strong covalent and hydrogen bonds, thus improving the load transfer efficiency from matrix to graphene sheet.

Excellent Electrical Properties
Current studies have shown that the improvement of electrical conductivity of bone repair materials is beneficial to enhance their osteogenic activities. Graphene possesses excellent electrical conductivity because its each carbon atom can form a π-bond structure with its neighboring carbon atoms on the pz orbital perpendicular to the monolayer plane and it has large SSA. The conductivity of graphene monolayer is up to $10^4$ S/cm, and the electron mobility is up to 15,000 cm$^2$/V·s at room temperature.

Figure 3 (A–F) Mechanical properties of graphene/barium titanate/polymethyl methacrylate composites with different graphene content: (A) compressive strength, in which the inset image showed the compressive strength curve of the composites, (B) crack propagation (0.5 wt.% graphene), and (C) scanning electron microscope (SEM) images of fracture morphology (0.5 wt.% graphene), in which red arrow indicated graphene, (D) SEM images of crack morphology (0.5 wt.% graphene), in which the red arrow indicated graphene extracted during crack propagation, (E) SEM images of crack propagation (1.5 wt.% graphene), and (F) SEM images of fracture morphology, in which red arrow showed graphene (1.5 wt.% graphene). (G–I) Mechanical properties of B$_4$C composite ceramics with different graphene content: (G) effect of graphene addition on fracture toughness and bending strength of composites, (H) SEM images of the fracture morphology of the composite with 0.4 wt.% graphene, which showed that the whole graphene layer was pulled out, and (I) SEM image of the fracture morphology of the composite with 0.8 wt.% graphene, which showed that the fracture of graphene first occurred in the outermost layer of graphene, and then the inner layer of graphene was pulled out. Reprinted from Ceramics International, Vol 45(5), Tang YF, Chen L, Duan ZH, Zhao K, Wu ZX, Graphene/barium titanate/polymethyl methacrylate bio-piezoelectric composites for biomedical application, Pages No.6567–6576, Copyright (2020), with permission from Elsevier (A–F). Reproduced from Gao DZ, Jing J, Yu JC, et al. Graphene platelets enhanced pressureless-sintered B4C ceramics. R Soc Open Sci. 2018;5 (4):171837. (G–I).
In this subsection, the main utilization ways of excellent electrical conductivity of GDs in bone repair materials were summarized. One was to compound GDs with other materials to prepare 2D/3D composites with satisfactory electrical conductivity, which could promote osteogenic activities of cells in vitro and bone formation in vivo under external electrical stimulation. GDs could be introduced into the matrix as an electrode to promote bone tissue regeneration under external electrical stimulation. Compared with traditional metal electrodes, graphene-containing composites had lower impedance and higher charge injection capacity. For example, Li et al \(^92\) prepared GDs-cellulose (G-C) scaffold and assembled an electrode device on this basis, which provided a platform for support and osteogenic differentiation of adipose-derived stem cell (ADSC). The results showed that the charge storage capacity of G-C electrode (0.966 mC/cm\(^2\)) was higher than that of gold electrode (0.462 mC/cm\(^2\)). Compared with the gold electrode, the G-C electrode showed lower impedance at low frequency (0.1–150 Hz), which indicated that it had higher energy efficiency and higher charge injection ability (Figure 4A and B). Furthermore, the results of in vitro experiments showed that the proliferation, mineral deposition and alkaline phosphatase (ALP) expression of adipose-derived stem cells (ADSCs) cultured with G-C electrode increased after electrical stimulation. Moreover, GO could be functionalized to further improve its electrical conductivity. For example, Jin et al \(^94\) introduced GO into the polyvinyl chloride/poly(lactic-co-glycolic acid) (PVC/PLGA) scaffold by electrospinning, and then GO-PVC/PLGA

**Figure 4 (A and B)** Electrochemical characteristics of Au and graphene-cellulose scaffold electrodes: (A) cyclic voltammogram, and (B) electrochemical impedance spectroscopy (EIS); (C and D) Conductivity of polyvinylidene fluoride (PVDF)/GO composite scaffolds: (C) open-circuit voltage, and (D) short-circuit current. Reprinted from Materials Science and Engineering: C, Vol 107, Li JF, Liu X, Crook JM, Wallace GG, Electrical stimulation-induced osteogenesis of human adipose derived stem cells using a conductive graphene-cellulose scaffold, Pages No.110312, Copyright (2020), with permission from Elsevier (A and B). \(^92\)Reprinted from Materials & Design, Vol 190, Shuai CJ, Zeng ZC, Yang YW, et al, Graphene oxide assists polyvinylidene fluoride scaffold to reconstruct electrical micro-environment of bone tissue, Pages No.108564, Copyright (2020), with permission from Elsevier (C–D). \(^96\)
composite scaffolds were immersed in hydrogen iodide solution to obtain rGO-PVC/PLGA scaffolds. The results showed that compared with GO-PVC/PLGA scaffolds, rGO-PVC/PLGA scaffolds had higher capacitance and better conductivity at a suitable voltage, which was due to the larger SSA of rGO-PVC/PLGA and the formation of more conjugated π-bonds on the surface of rGO. In another case, Silva et al\textsuperscript{95} introduced cyclic amine into graphene by 1,3-dipolar cycloaddition reaction, and then functionalized graphene was compounded with chitosan (CHI) and sodium alginate (ALG) to fabricate biofilms. The results showed that the introduction of functionalized graphene significantly reduced the resistivity of the thin films from 3.00 $\times$ 10\textsuperscript{12} $\Omega$ m to 2.25 $\times$ 10\textsuperscript{9} $\Omega$ m.

The other method was to use the excellent electrical conductivity of GDs to improve the performances of piezoelectric biomaterials that could bring desired electrical stimulation for new bone formation under biomechanical environment in vivo.\textsuperscript{96,97} In addition to the excellent electrical conductivity of GDs themselves, GDs could also enhance the electrical output performance of the piezoelectric biomaterials by changing the phase structure of the matrix. For example, Shuai et al\textsuperscript{96} introduced GO into polyvinylidene fluoride (PVDF) to fabricate composite scaffolds for bone repair. The results showed that the open-circuit voltage and short-circuit current of the composite increased at first and then decreased with increasing of GO content. Compared with pure PVDF, the scaffold with 0.3 wt.% GO showed the best electrical output performance, and its maximum output voltage (~8.2 V) and maximum current (~101.6 nA) were increased by 82.2\% and 68.2\%, respectively (Figure 4C and D). The main reason of which might be that there was a strong hydrogen bond between the oxygen-containing functional groups of GO and the fluorine groups of PVDF, which improved the interfacial bonding, and GO could induce the phase transition with higher energy conversion efficiency. However, when the content of GO was exorbitant, the electrical output capacity of the composites decreased, which might be due to the agglomeration of GO in the composites. Therefore, the content and dispersion of GDs could have a significant impact on the electrical output properties of the composites. Subsequently, in vitro experiments demonstrated that the introduction of GO could significantly promote the proliferation and ALP expression of MG-63 cells. The reason might be that GO had stronger electromechanical conversion ability, resulting in stronger electrical stimulation under the same external stress, and finally effectively promoting the behavior of cells.

Large Specific Surface Area (SSA) and Easy Modification

The large SSA of GDs could not only greatly improve cell adhesion, but also make GDs to be easily further functionalized and incorporated with other active materials to obtain bone repair materials with better performance. The functionalization of GDs could not only improve different properties in the matrix, but also used as a “bridge” for binding with active substances.\textsuperscript{98,99} On the other hand, GDs could be further endowed with new functions by binding with active substances.\textsuperscript{100–102}

In this subsection, we reviewed the different functional groups of GDs modification in bone repair materials and the progress had been made in the properties and functions of the composites. Then, GDs were bound with different active substances such as peptides and drugs, and which functions of bone repair materials were improved by in vitro and in vivo experiments.

The common functional methods of GDs used in bone repair were carboxylation,\textsuperscript{103–105} silanization,\textsuperscript{106,107} hydroxylation,\textsuperscript{25} dopamine modification\textsuperscript{108,109} and amination,\textsuperscript{24,110} as shown in Table 1. These functional modifications improved the different properties of the composites, such as binding other ions, dispersibility, hydrophilicity, adsorption of serum proteins and attraction to osteoblasts.

Firstly, the surface of GDs was negatively charged by carboxyl modification, which was more beneficial to improve the properties of the composites by combining cations. For example, Chen et al\textsuperscript{103} prepared carboxylated graphene oxide (GO-COOH) by NaOH/CH\textsubscript{2}COOH and further covalently bound with ZnO (Figure 5A). The results showed that the surface of GO-COOH was negatively charged, which contributed to the stable bonding with Zn\textsuperscript{2+}. Subsequently, the results of in vitro experiments showed that ZnO/GO-COOH composite significantly increased ALP activity, osteocalcin production and extracellular matrix mineralization in MG-63 cells. In addition, ZnO/GO-COOH nanocomposites had a bacteriostatic effect on Streptococcus mutans.

Secondly, silanization improved the dispersion of GDs, resulting in stronger adhesion between GDs and the matrix after polymerization. Paz et al\textsuperscript{106} prepared the silylated graphene by (3-methacryloxypropyl)
trimethoxysilane (MPS) and then introduced it into polymethyl methacrylate (PMMA) bone cement to prepare composite materials (Figure 5B). The results showed that the compressive strength, bending strength and fracture toughness of the composites containing silylated graphene were increased by 12%, 13.7% and 28%, respectively, compared with those containing original graphene.

### Table 1 Functionalization of GDs for Bone Repair

| Functionalization | Preparation Method | Properties | Functions | References |
|-------------------|--------------------|------------|-----------|------------|
| Carboxylation     | NaOH/ClCH<sub>2</sub>COOH or KMnO<sub>4</sub>/H<sub>2</sub>SO<sub>4</sub> | Binding more positive amino acids | Improving mineralization of MG63 cells | [103–105] |
| Silanization      | Silane coupling method (hydrolysis, blending, washing, freeze-drying) by (3-methacryloxypropyl) trimethoxysilane (MPS) | Improving the dispersion of graphene; enhancing the compressive strength and bending strength of the materials | Lower cytotoxicity | [106,107] |
| Hydroxylation     | 1. KMnO<sub>4</sub>/H<sub>2</sub>SO<sub>4</sub> 2.Sheared and emulsified in NaOH | Better mechanical property and electrical conductivity | Improving the proliferation of rat adipose stromal cells | [25] |
| Dopamine          | Chemical vapour deposition method | Significantly improving the binding ability and release ability with BMP-2 | Improving the exogenous BMP2-induced osteogenic differentiation | [108,109] |
| Amination         | Sulfoxide chloride (SOCl<sub>2</sub>)/ethylene diamine/dimethylformamide (DMF) | Better thermal stability and higher mechanical properties | Stronger osseointegration and lower cytotoxicity | [24,110] |

**Figure 5** Schematic diagram of GDs modified by various functional groups: (A) carboxylation, (B) silylation, (C) dopaminetization, and (D) amination. Reproduced with permission from Colloids and Surfaces B: Biointerfaces, Vol 147, Chen JY, Zhang X, Cai H, et al, Osteogenic activity and antibacterial effect of zinc oxide/carboxylated graphene oxide nano-composites: preparation and in vitro evaluation, Pages No.397–407, Copyright (2016), with permission from Elsevier (A); Reprinted from Materials Science and Engineering: C, Vol 104, Paz E, Ballesteros Y, Forró I, Dunne NJ, Del Real JC, Graphene and graphene oxide functionalisation with silanes for advanced dispersion and reinforcement of PMMA-based bone cements, Pages No.109946, Copyright (2019), with permission from Elsevier (B), Reprinted (adapted) with permission from Jia ZJ, Shi YY, Xiong P, et al. From solution to biointerface: graphene self-assemblies of varying lateral sizes and surface properties for biofilm control and osteodifferentiation. ACS Appl Mater Interfaces. 2016;8(27):17151–17165. Copyright (2016) American Chemical Society (C); Reprinted (adapted) with permission from Sharma R, Kapusetti G, Bhong SY, et al. Osteoconductive amine functionalized graphene-poly(methylmethacrylate) bone cement composite with controlled exothermic polymerization. Bioconjugate Chem. 2017;28(9):2254–2265. Copyright (2017) American Chemical Society (D).
Thirdly, through hydroxylation modification, the surface of GDs was negatively charged, thus improving their hydrophilicity. Sun et al\textsuperscript{25} prepared hydroxylated graphene by two-step methods of K\textsubscript{2}MnO\textsubscript{4}/H\textsubscript{2}SO\textsubscript{4} and NaOH shear emulsification. In vitro experiments showed that the hydroxylation of graphene significantly promoted the proliferation of ADSCs. The hydrophilicity of hydroxylated graphene provided a better microenvironment for cell adhesion and proliferation.

Fourthly, dopamine modification could improve the adsorption of GDs to serum protein. The catechol groups on polydopamine (PDA) could covalently bind to the amino group on the protein through o-benzoquinone-amine coupling, promoting more serum proteins to be anchored to the surface of the composite, thus establishing a favorable extracellular matrix for rapid cell recruitment, stable adhesion and effective cytoskeleton construction. Jia et al\textsuperscript{108} covalently bound dopamine (DA) and GO by the method of evaporation-assisted electrostatic assembly to prepare rGO-PDA, and then rGO-PDA was introduced into titanium substrate to prepare composites (Figure 5C). In vitro experiments showed that rGO-PDA could significantly increase the adhesion, growth and ALP expression of MC3T3-E1 cells compared with GO.

Moreover, through amination, the positive charge was introduced on the surface of GDs, which increased the attraction of GDs to the negative charge of carboxylate and phosphate ions on the surface of the osteoblast membrane. Sharma et al\textsuperscript{24} synthesized amine-functionalized graphene by Sulfoxide chloride (SOCl\textsubscript{2})/Ethylene diamine/Dimethylformamide (DMF) and blended it with PMMA bone cement matrix to prepare bone cement composites (aminated graphene/PMMA) (Figure 5D). In vitro and in vivo experiments showed that compared with pure graphene and GO, the aminated graphene promoted the osseointegration of the composites and reduced its cytotoxicity, and the aminated graphene/PMMA composites were calcified earlier after implantation in rabbits.

On the basis of GDs functionalization, the binding with active molecules such as genes,\textsuperscript{111,112} peptides,\textsuperscript{26,113,114} activity factor\textsuperscript{115,116} and drugs\textsuperscript{117–119} contributed to the continuous release of active molecules, as shown in Table 2. The active molecules could enhance a variety of functions of bone repair materials for a long period, such as proliferation, growth, osteogenic differentiation and antibacterial.

However, the mechanism and effect of functionalized GDs on different active molecules were different. The functionalized GDs were used as a carrier for gene transfection into cells, the charge on the surface of which was one of the most important factors affecting the transfection efficiency. For example, Liu et al\textsuperscript{111} blended graphene with oleic acid at first and then covalently bound them with polyamidoamine (PAMAM) to prepare graphene-oleate-PAMAM composites. The results showed that the covalent modification of graphene by PAMAM greatly improved the transfection efficiency of green fluorescent protein gene (GFP). The transfection efficiency of graphene-oleate-PAMAM on HeLa cells was up to 18.3\%, which was 13 times that of GO. Because the primary amine group on the surface of PAMAM had a large amount of positive charge, graphene-oleate-PAMAM could effectively immobilize gene molecules and obtaining high transfection efficiency.

In another case, the improvement of the bioactivity of GDs-containing composites was not only due to the interaction between GDs and active molecules, but also related to the charge on the surface of the carrier. For example, Eckhart et al\textsuperscript{26} covalently bound polylsine with GO-COOH to prepare composites by acylation. The results showed that the covalent binding improved the electrical conductivity of the composites, which might be due to the strong interaction with the covalently bound peptides between the adjacent graphene layers, such as hydrogen bonding and chain entanglement, which led to the closeness between the graphene layers. On the contrary, the electrical conductivity of the composite prepared by non-covalent binding of peptide and GDs was lower than that of pure graphene, which might be due to the phase separation between peptide and GDs. The phase separation blocked the electron migration, so that it reduced the electrical conductivity of the composites. On the other hand, the terminal group of polylsine had a positive charge under the condition of physiological pH, which was consistent with the principle that charge affects cell function. Therefore, these factors led to the enhancement of the adhesion of PC12 cells on the composites prepared by covalent binding.

Based on the above principle, including the interaction between GDs and active molecules, as well as the charge on the surface of the carrier, GDs binding with active molecules could promote stem cell proliferation, migration and differentiation. For example, Zhang et al\textsuperscript{115} introduced Fe\textsuperscript{3+} into GO through -COOH to prepare GO-Fe\textsubscript{3}O\textsubscript{4} magnetic nanoparticles through FeCl\textsubscript{3}/NaOH reaction. Then, BMP-2 was fixed to the nanoparticles, which enter the stem cells through endocytosis. The results showed that BMP-2 enhanced the proliferation, migration and
differentiation of dental pulp stem cell (DPSCs) into osteoblasts around the cell membrane to form the new bone. Compared with traditional Fe$_3$O$_4$ nanoparticles, GO provided many carboxyl groups that could bind to growth factors. With the help of the magnetic nanoparticles, BMP-2 was integrated into cells and induced into the bone.

In addition, covalent binding was conducive to the continuous release of active molecules, so it could maximize the function of active molecules. For example, Unnithan et al$^{117}$ crosslinked simvastatin (SV) with GO by EDC, and then introduced them into chitosan (CS) and hydroxyapatite (HA) to prepare SV-GO-CS-HA composite scaffolds. The release curve of SV showed that drugs could be released from CS-HA-GO scaffolds in a controlled manner, and the drug loading rates of 0.5% and 1% SV-loaded on GO-CS-HA scaffolds were 68% and 82%, respectively. In vitro experiments showed that SV significantly enhanced the osteogenic and biomineralization ability of MC3T3 cells. Moreover, GO initiated the

| Types of Active Substance | Examples of the Active Substance | Preparation Method | Properties | Functions | References |
|--------------------------|---------------------------------|-------------------|-----------|-----------|------------|
| Genes                    | Green fluorescent protein gene (GFP) | The PEGFP-N1 plasmid containing GFP gene was adsorbed on the surface of graphene-oleic acid-polyamidoamine (PAMAM) composites | Good dispersion and stability in aqueous solutions | Improving the efficiency of gene transfection | $^{111}$ |
| MiR-7b                   | Blending of MiR-7b plasmid and GO-PEI complex in aqueous solution to prepare GO-PEI-miR-7b composites | / | Excellent transfection efficiency in bone marrow macrophages. Blocking the cell-cell fusion of osteoclasts and promoting bone regeneration. | $^{112}$ |
| Peptides                 | Polysine                         | Self-assembled    | Mechanically robust | Enhancing adhesion and neurite outgrowth of PC12 cells | $^{26}$ |
| Motif-specific peptide (LLVFGAKMLPHHGA) | The peptides were adsorbed on the surface of the scaffolds by impregnation | Adjustable shape, superlow weight and high porosity | Easy to mineralize | $^{113}$ |
| Arginine-glycine-aspartic acid (RGD) peptide | RGD peptides were self-assembled onto the surface of GO/silica nanoparticles composites on indium tin oxide (ITO) substrates. | Excellent electrical conductivity | Promoting osteogenic differentiation of adipose-derived mesenchymal stem cells (ADSCs) | $^{114}$ |
| Activity factors         | Bone-morphogenetic-protein-2 (BMP-2) | The BMP-2 factor was adsorbed on the surface of GO-Fe$_3$O$_4$ nanoparticles by impregnation. | / | Promoting osteogenic differentiation of dental pulp stem cells (DPSCs) | $^{115}$ |
| Human recombinant bone morphogenetic protein-2 (rBMP-2) | The rBMP-2 factor was adsorbed on the surface of heparin modified graphene scaffold by impregnation. | Significantly prolonged the release of factors. | Promoting osteoblast differentiation of C2C12 cells | $^{116}$ |
| Drugs                    | Simvastatin (SV)                 | SV was crosslinked with graphene oxide-chitosan-hyaluronic acid composite scaffold by EDC/NHS method. | Higher mechanical support for scaffolds | Higher mineralization activity of MC3T3 cells | $^{117}$ |
| Dexamethasone (DEX)      | DEX was used to prepare DEX-phosphate monoester-GO-chitosan composite scaffold by covalent bonding of phosphate ester bond. | A sustaining release for longtime in scaffolds in different pH conditions. | Excellent biocompatibility | $^{118}$ |
| Vancomycin (VA)          | VA was adsorbed on the surface of rGO-nano hydroxyapatite composite scaffold by impregnation. | Persistent pathogen resuscitation inhibition | Improving cell adhesion and osteogenic differentiation of Bone marrow mesenchymal stem cells (BMSCs) | $^{119}$ |
effective interaction between the scaffolds and mineral ions through the existence of functional groups such as -COOH, -OH and epoxy groups on the scaffold surface, thus promoting the formation of HA particles.

Improving Functions of Bone-Related Cells in vitro and Bone Regeneration in vivo

GDs had been considered as the next generation of bioactive nanomaterials, which could not only significantly improve the functions of bone-related cells such as adhesion, proliferation, osteogenic differentiation and mineralization in vitro, but also promote bone regeneration in vivo.

Many in vitro experiments showed that introducing a small amount of GDs into substrates could significantly improve the related functions of osteoblasts such as adhesion and proliferation. For example, Liang et al prepared nano-hydroxyapatite/poly(lactic-co-glycolic acid)/graphene oxide (nHAC/PLGA/GO) three-dimensional porous scaffolds by freeze-drying, then co-cultured them with MC3T3-E1 cells. The results showed that nHAC/PLGA/GO (1.5 wt.%) scaffold could promote the adhesion and proliferation of osteoblast (MC3T3-E1) more significantly than nHAC/PLGA scaffold at days 3, 5, and 7, and that the cells were larger and more stretched. In another case, Li et al prepared rGO coating on the surface of Ti6Al4V alloy and studied the effect of rGO coating on the growth and osteogenic differentiation of MC3T3-E1 cells, the results of which showed that the rGO coating significantly promoted the growth and osteogenic differentiation of MC3T3-E1 cells on Ti6Al4V implants at days 3 and 7. In addition, the quantitative reverse transcription-polymerase chain reaction (RT-qPCR) results demonstrated that bone sialoprotein (BSP), osteocalcin (OCN) and runt-related transcription factor 2 (Runx2) expressions of the cells cultured on the rGO-coated substrate at day 7 were significantly higher than those of the cells cultured on the original Ti6Al4V substrate (p<0.05).

Moreover, GDs could regulate the osteogenic differentiation of stem cells from different sources because they could provide stem cells with a unique physical framework comparable to natural extracellular matrix (ECM). For example, Newby et al prepared GO nanoparticles with oxygen content of 6–10%, then co-cultured them with human mesenchymal stem cells (MSCs), and evaluated the effect of graphene nanoparticles on osteogenic differentiation of these stem cells by alizarin red staining and quantitative analysis. The results showed that the calcium content of MSCs co-cultured with GO nanoparticles was significantly higher than that of cells on the control surface. Interestingly, the up-regulation was observed in the absence of any osteoinductive agents (dexamethasone, β-glycerophosphate or ascorbic acid), which indicated that GO nanoparticles could spontaneously induce calcium accumulation in MSCs. In another case, Krukiewicz et al prepared GO/PMMA composites by pressing machine and artificial smearing, respectively. Then, the osteogenic differentiation efficiency of human primary mesenchymal stem and progenitor cells (hMSPCs) cultured on GO/PMMA composite was studied by three differentiation markers, namely ALP, secreted protein acidic and rich in cysteine (SPARC), and bone morphogenetic protein-2 (BMP-2), the results of which showed that both GO/PMMA composites were effective in inducing osteogenic differentiation and were superior to unmodified PMMA. Moreover, the GO/PMMA composite prepared by artificial smear had the highest efficiency of inducing osteogenic differentiation of the cells, which might be due to a large number of voids and pores on its surface.

On the other hand, in vivo experiments on various animal models showed that GDs could not only promote bone repair but also significantly enhance ectopic bone formation, indicating the great potential of GDs as bone repair material. For example, Zou et al prepared graphene and Laponite (GL) composite nanoparticles, then mixed the nanoparticles with MSCs infected with recombinant adenovirus-bone morphogenetic protein-9 (Ad-BMP9), and implanted the mixture under the skin of athymic nude mice. The results showed that graphene-containing composite nanoparticles could significantly enhance BMP9-induced ectopic bone formation from MSCs than pure GL nanoparticles. In another case, Nie et al prepared 3D porous rGO composites by self-assembly of GO and nHA and then constructed biomimetic scaffolds by freeze-drying to repair bone defects. In vivo experiments showed that after 6 weeks of implantation, the porous scaffold containing rGO successfully cured rabbits round skull defect with a diameter of 4 mm, which was faster than that of pure nHA scaffolds. Computed tomography (CT) and histological analysis showed that the introduction of rGO could significantly
improve the collagen deposition, cell proliferation and new bone formation in the defect model than nHA scaffolds.

**Composites with Different Substrates for Bone Repair Materials**

GDs could be introduced into different kinds of substrates, such as metals, inorganic nonmetals, natural polymers and synthetic polymers to prepare bone repair materials with improved performance, as shown in Table 3. In this subsection, the updated progress on research and development of those materials was reviewed. Especially, the functions of GDs themselves and the synergistic effects with other materials in vitro and in vivo were emphasized.

GDs played a very important role in the preparation and development of load-bearing bone materials with excellent functions. Metal substrate provided hard mechanical support for load-bearing bone materials, but it also had the problems of fast corrosion rate, stress shielding and low bioactivity. Therefore, GDs with the excellent properties were used to improve the in vitro/in vivo function of the composites, and the synergistic effect of GDs and substrate provided a good microenvironment for further improving the performances of the composites.

In the past, GDs were mostly introduced into the metal substrates in the form of coatings. However, in order to obtain the microstructure closer to the natural bone, GDs were introduced into the metal substrate to prepare a three-dimensional structure. For example, Qiu et al introduced GO into titanium substrates treated by NaOH/3-aminopropyltrimethoxysilane (APS) method. Subsequently, GO was reduced to rGO by alkaline thermal method to prepare three-dimensional porous nanocomposites. The results showed that nanocomposites could promote ALP activity, extracellular matrix (ECM) mineralization and collagen secretion of rat bone marrow mesenchymal stem cells (rBMSCs). Moreover, the expression of osteogenesis-related genes such as ALP, bone morphogenetic protein-2 (BMP-2), osteocalcin (OCN) and osteopontin (OPN) increased on the surface of nanocomposites. The improvement of function might be attributed to the unique three-dimensional structure formed through the cross-linking of silanol-based (SiOH) introduced on the surface of titanium substrate, which affected the interactions of cell-cell and cell-material. Furthermore, the reduction of GO to rGO resulted in more aromatic structures on the surface of the composites, which have a strong non-covalent interaction with biomolecules.

In another case, Lyu et al constructed layered ZnO nanotubes/silk fibroin/graphene oxide nanostructures (SF/GO-ZnO) on pure zinc substrates by anodizing and SF/GO self-assembly. Subsequently, SF/GO-Dex-ZnO was prepared by coating the osteogenic drug dexamethasone (Dex) to further improve the function of the composites. The results showed that SF/GO-Dex-ZnO had excellent bactericidal and osteogenic activity, which might be due to the unique structure of GO and the active oxygen produced during the degradation of ZnO. Among them, the “sharp” of GO nanosheet structure destroyed the cell membrane of bacteria through direct mechanical contact with bacteria. In addition, on the composite with SF/GO coating, the cells stretched in different directions, while the cells on the pure zinc and the cells on anodized zinc were elongated, which showed that SF/GO coating had a positive effect on promoting cell proliferation. The reason for the above results might be that the composition of SF was similar to that of collagen, the main component of extracellular matrix, and GO had a higher adsorption capacity for serum protein, resulting in a higher density of adhesion molecules could be used for cell adhesion and growth. Therefore, SF and GO jointly promoted the early diffusion and proliferation of cells.

In the composition of bone, inorganic substances such as hydroxyapatite accounted for a large proportion, so whether it could stimulate the biomineralization of inorganic substances in natural bone was regarded as an important index in the process of bone repair. In the past, inorganic/organic composite scaffolds based on polymers were a common choice for bone repair, but the agglomeration of inorganic particles affected their overall stability and bioactivity. Therefore, the introduction of GDs could solve the problems. For example, Zhao et al prepared GO/chitosan/nano-hydroxyapatite particles (GO/CS/nHAP) scaffolds by introducing nHAP into CS and GO covalent binding network matrix by in situ one-step bionics technology. The scaffolds had excellent physical and chemical properties, such as three-dimensional porous bone structure, mechanical properties, biodegradability and water absorption. Furthermore, in vitro experiments demonstrated that compared with the defect in CS/nHAP group, the defect in GO/CS/nHAP group was almost filled with new bone tissue and formed a complete structure with the natural bone, the reasons for which included the following aspects.

Firstly, the network substrate of GO/CS effectively regulated the uniform dispersion of nHAP on the nanometer scale and promoted the formation of bone-like organic-inorganic hybrid structure. Secondly, the covalent binding of GO/CS strengthened

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**Table 3**

| Material | Properties | Applications |
|----------|------------|--------------|
| GO       | Excellent mechanical strength and bioactivity | Bone repair materials |
| CS       | Good stability and bioactivity | Bone repair materials |
| nHAP     | Stable mechanical properties | Bone repair materials |

**Figure 3**

3-aminopropyltrimethoxysilane (APS) method.
| Types of Substrates      | Examples             | Preparation Method                                                                 | Structure                  | Properties                                | Functions                                                                                       | References |
|-------------------------|----------------------|-------------------------------------------------------------------------------------|----------------------------|-------------------------------------------|------------------------------------------------------------------------------------------------|-----------|
| Metal                   | Titanium             | Silane coupling using NaOH/APS                                                      | 3D porous nanosheets       | Super hydrophilic, rough surface         | Promoting ALP activity, ECM mineralization and collagen secretion of BMSCs                      | [130]     |
|                         |                      | Zinc/GO was self-assembled and coated on anodized zinc substrate                     | The GO coating was attached to the Zn surface where many ZnO nanotubes were formed. | /                                         | Excellent bactericidal activity and osteogenic activity                                        | [131]     |
|                         |                      | Magnesium                                                                          | Micron particles           | Improving hardness and corrosion resistance | /                                                                                              | [132]     |
| Inorganic substance     | Hydroxyapatite       | Preparation of GO/chitosan/nHAP by in-situ synthesis and microwave-assisted method | 3D porous scaffold         | Better dispersion and remarkable strength | Significantly improving MC3T3-E1 cells viability                                              | [133]     |
|                         | Calcium phosphate    | Liquid-phase and powder-phase mixing                                                | Rough microporous          | Higher stability, injectability and rheology | Increasing the survival rate of BMSCs slightly                                                | [134]     |
|                         | Calcium carbonate    | CaCl\(_2\)/rGO and Na\(_2\)CO\(_3\) aqueous solution reaction                   | Hollow microsphere          | Sensitive release characteristics of pH    | No cytotoxicity                                                                              | [135]     |
|                         | Tricalcium silicate  | Liquid-phase and powder-phase mixing                                                | Spherical particles with rough and porous surface | Higher compressive strength and lower pH value | Excellent ability of MC3T3-E1 cells proliferation and antibacterial                          | [136]     |
| Natural polymer         | Chitosan             | In situ crystallization process (acetic acid, Ca\text{(NO}_3\text{)}_2\text{, and K}_2\text{HPO}_4 aqueous solution) | 3D porous bone-like hierarchical structure | Biodegradability and appropriate water absorption and water retention | Excellent biological activity and ability to induce bone regeneration in situ | [137]     |
|                         | Cellulose            | Dispersion deposition-drying                                                        | GO-coated cellulose tissue paper | Lower impedance and higher charge injection capacity | Promoting the proliferation and osteogenic differentiation of ADSCs | [138]     |
|                         | Collagen             | Sol-gel method                                                                      | Highly porous and interconnected aerogels | Excellent hydrophilicity and improving compression modulus | Better biomineralization and cytocompatibility                                                | [139]     |
| Synthetic polymer       | Polycaprolactone     | 3D printing                                                                         | 3D porous structures       | Significantly higher electrical conductivity | Higher mineralization nucleation rate in vitro                                               | [140]     |
|                         | (PCL)                | Polycaprolactone (PCL)                                                               | 3D porous structures       | Significantly higher electrical conductivity | Higher mineralization nucleation rate in vitro                                               | [140]     |
|                         | Polyethyleneimine    | PEI were crosslinked with GO by EDC/NHS                                             | Nanofillers                | Improving the ultimate tensile strength and elastic modulus | Promoting the effective interfacial adhesion and compatibility with the matrix | [141]     |
|                         | (PEI)                | UV crosslinking using 3D printing technology                                         | 3D porous structures       | Better tensile strength and bending strength | /                                                                                             | [142]     |
|                         | Polyurethane (PU)    | Electrospinning                                                                      | Nanocomposite fiber        | Increasing tensile strength and modulus of elasticity | Promoting Saos-2 cells proliferation                                                         | [143]     |
the interfacial interaction between organic and inorganic materials, and it enhanced the stability of GO/CS/nHAP scaffolds in the degradation process. Thirdly, the sheet structure of GO maintained a certain space between the composite fiber units and improved the hydrophilicity of GO/CS/nHAP scaffolds. Finally, the abundant functional groups on GO (such as -COOH, -OH and epoxy groups) and CS (such as -NH$_2$) promoted the in situ recruitment, proliferation and differentiation of endogenous stem cells at the injured site, thus realizing endogenous bone tissue regeneration.

Polymers had been widely used as bone repair materials because of its many advantages, such as easy to prepare into three-dimensional porous structure, degradability and similar mechanical properties to the implanted tissue. However, many pure polymers such as poly (caprolactone) (PCL), polylactic acid (PLA) were hydrophobic solids and lacked functional groups, resulting in low cellular response. GDs could be uniformly dispersed into the polymer matrix by membrane lamination, fiber bonding, 3D printing and other methods to make up for these shortcomings. For example, Unagolla et al$^{140}$ prepared PCL-GO composite scaffolds by 3D printing, which could independently control the size and pore diameter of the implants. The morphology observation showed that the surface of the pure PCL scaffold was rough and irregular, while the surface of the scaffold containing GO was smooth and there were GO particles on the surface, which indicated that the introduction of GO improved the fluidity of the composites. The ALP activity assay confirmed that GO increased ALP activity of composite scaffolds (Figure 6A). Subsequently, as shown in Figure 6B and C, the calcium and phosphorus analysis showed that compared with simple PCL scaffolds, GO promoted the mineralization of scaffolds. The reason of which might be that the functional groups and electrical conductivity of GO improved the function of the composites. Furthermore, Western blot analysis demonstrated that the porous structure of GO and PCL composite scaffolds promoted the secretion of bone morphogenetic protein-2 and osteopontin, thus stimulating its osteogenesis.

**Mechanism of GDs-Induced Bone Regeneration**

**Possible Signaling Pathways Involved in GDs-Induced Osteogenic Differentiation of Stem Cells**

The important index to evaluate the performance of bone repair material was whether it could recruit reactive osteoblast progenitor cells and eventually differentiate into osteocytes in the process of bone regeneration. Stem cells from different sources could be used as progenitor cells in the process of bone formation, so their recruitment and osteogenic differentiation were key factors for effective bone regeneration. GDs were proved to have the ability of osteogenic induction,$^{144–152}$ as shown in Table 4. In this subsection, the signaling pathways involved in GDs-induced bone regeneration were discussed, including which signaling pathways of osteogenic differentiation of stem cells from different sources were affected by GDs and the effect of GDs-related parameters on the signaling pathway.

Currently, different types of stem cells were studied, including bone marrow mesenchymal stem cells (BMSCs), adipose-derived stem cells (ADSCs), stem cells from human exfoliated deciduous teeth (SHEDs) and tonsil-derived mesenchymal stem cells (TMSCs). Cytobiological assay and Ribonucleic Acid (RNA) sequencing were used to evaluate the signaling pathway of GDs-induced stem cells to differentiate into osteoblasts. These studies were systematically summarized in Table 4, and some possible rules were found.

Firstly, when GDs were used alone and GDs were combined with other materials, the signaling pathways involved in GDs-induced bone regeneration were different, which might be due to the change of the surface charge and groups after GDs were introduced into the matrix. For example, Qi et al$^{144}$ prepared sericin methacryloyl/GO (SMH/GO) composite hydrogel as bone repair material by photo-crosslinking. The results of in vivo experiments showed that the hydrogel containing GO had a better bone regeneration effect, and the structure and function of rat skull defect model were repaired within 12 weeks. Subsequently, the results of cytotiological assay and RNA sequencing showed that the expressions of key genes related to bone marrow mesenchymal stem cell migration (such as Bmp6, Tgf-β1, Mmp2, Mmp3, Mmp8, Mmp10 and Mmp12) and osteogenic differentiation-related genes (such as Tgf-β1, Smad1, Bmp6, Fgf7, Mapk3, Mapk6 and Gs3k) were the highest in SMH/GO hydrogel group. The up-regulated expression of these genes indicated that SMH/GO hydrogel might promote the migration and osteogenic differentiation of BMSCs by regulating biological processes such as “positive regulation of cell migration”, “positive regulation of osteoblast differentiation” and “positive regulation of myeloid cell differentiation”. Moreover, through signaling pathway analysis, they revealed the possible mechanism of bone regeneration, SMH/GO hydrogel regulated the migration and osteogenic
differentiation of BMSCs by activating MAPK, tumor necrosis factor (TNF) and chemokine signals, thus promoting bone regeneration. In another case, Halim et al. studied the signaling pathways of GO controlling the differentiation of stem cells into osteoblasts. GO was made into suspension with a concentration of 0.1 μg/mL and...
then co-cultured with BMSCs. The results showed that GO had no effect on the bioactivity of BMSCs, and it slightly promoted the proliferation of BMSCs. Moreover, the osteoinductive effect of GO was realized by activating c-Jun N-terminal kinase (JNK) and Forkhead transcription factor 1 (FoxO1) signaling pathways.

Secondly, the content of GDs in the composites might affect the signaling pathway of stem cell osteogenic differentiation. For example, Wu et al. incorporated an appropriate amount of graphene into polylactic-glycolic acid copolymer films, and then they discussed the possible effect of PI3K/AKT/GSK-3β/β-catenin signal pathway on

**Table 4 Study on the Signaling Pathways Involved in GDs-Induced Osteogenic Differentiation of Stem Cells**

| Signaling Pathways | Materials Structure | Stem Cell Species | Method | References |
|--------------------|---------------------|-------------------|--------|------------|
| Wnt/β-catenin      | GO: coating β-TCP 3D scaffolds | BMSCs | The cells were cultured on GO coating, which was prepared by immersion in aqueous suspension and heat treatment for 1, 3, 7 and 14 days. | [41] |
|                   | GO/Fe$_3$O$_4$ Nanoparticles | BMSCs | The GO/Fe$_3$O$_4$ nanoparticles prepared by chemical coprecipitation of ferric trichloride and ammonia were added to the cell suspension, and cultured for 1, 3, 5, 7, 14 and 21 days. | [58] |
|                   | Graphene oxide quantum dots (GOQDs) Suspension | Stem cells from human exfoliated deciduous teeth (SHEDs) | GOQDs was dispersed by ultrasound and cultured in cell suspension for 1, 3, 5, 7 and 14 days. | [149] |
| PI3K/Akt/GSK-3β/β-catenin | GO/poly(lactic-co-glycolic acid) (PLGA) Films | BMSCs | The cells were cultured on GO/PLGA films, which were prepared by acetone solution and mold casting for 1, 3, 7 and 14 days. | [42] |
| MAPK and its subspecies (ERK1/2, p38) | GO/Srtrontium (Sr)/collagen Porous scaffolds | Adipose-derived stem cells (ADSCs) | The cells were cultured on porous scaffolds, which were crosslinked by EDC/NHS and freeze-drying for 1, 3 and 14 days. | [43] |
|                   | GO/sericin methacryloyl (SerMA) Porous hydrogels | BMSCs | The cells were cultured on GO/SERMA porous hydrogels, which were prepared by UV cross-linking method for 1 day. | [144] |
|                   | GO/copper (Cu)/calcium phosphate(CaP) GO/CaP: coating CaP: porous scaffolds | BMSCs | The cells were cultured on GO/CaP coating, which was synthesized in situ by CuSO$_4$ and ascorbic acid for 1, 3, 7 and 14 days. | [145] |
|                   | Graphene/CaP/chitosan/magnesium (Mg) alloy Graphene nanosheets/ CaP/CS: porous coating | BMSCs | The cells were cultured on graphene/CaP/CS porous coating, which was prepared by electrophoresis deposition (EPD) for 1, 2, 8, 16 and 24 days. | [147] |
|                   | Graphene/single-walled carbon nanotube (G/SWCNT) hybrids Nanoparticles | BMSCs | G/SWCNT hybrids nanoparticles prepared by chemical vapor deposition processing were added to the cell suspension, and cultured for 1, 3, 7 and 14 days. | [150] |
| JNK and FoxO1     | GO Suspension | BMSCs | GO was dispersed by ultrasound and cultured in cell suspension for 1, 2, 3, 7 and 14 days. | [146] |
| Focal adhesion kinase (FAK) | GO/chitosan/ nanohydroxyapatite (nHA) Porous scaffolds | ADSCs | The cells were cultured on GO/chitosan/nHA porous scaffolds, which were crosslinked by EDC/NHS and freeze-drying for 1, 3, 5, 7 and 14 days. | [148] |
|                   | GO/polyethylene glycol (PEG) Porous hydrogel scaffolding | Tonsil-derived mesenchymal stem cells (TMSCs) | The cells were cultured on GO/PEG porous hydrogel scaffolds, which were coupled by APS and freeze drying for 1, 4, 7 and 14 days. | [151] |
|                   | GO/polyethylene glycol-acrylate (PEGA) Hydrogels | ADSCs | The cells were cultured on GO/PEG hydrogels, which were prepared by UV cross-linking method for 1, 7, 14 and 21 days. | [152] |
osteogenic differentiation of BMSCs induced by graphene. The results showed that the expression of total Akt and phosphorylated-Akt (p-Akt) was up-regulated with the increase of graphene nano-tablet concentration to 0.5 wt.%. However, when the concentration of graphene nanoplates further increased to 1.0 wt.%, the expression of Akt and p-Akt decreased. The above results suggested that 0.5 wt.% GO-PLGA showed better osteogenic induction than other graphene groups. Therefore, the content of GDs could affect the signaling pathway of stem cell osteogenic differentiation by regulating the expression of related proteins.

Finally, the composites containing GO were prepared into porous structures, which tended to activate the signal pathways of MAPK and its subspecies (extracellular regulated protein kinases 1/2 and p38). For example, Zhang et al. prepared GO/Cu solution by in-situ synthesis of CuSO₄ and ascorbic acid, then coated them on the surface of calcium phosphate (CaP) substrate to fabricate GO/Cu/CaP composite scaffolds. Western blot analysis demonstrated that GO up-regulated the expression of Hif-1α in BMSCs and secreted bone morphogenetic protein-2 by activating 1/2 (ERK1/2) signal pathway (a subclass of MAPK).

**Immunomodulatory Mechanism of GDs-Induced Bone Regeneration**

In the process of bone repair, immune cells were the first cells to contact and react with the implant. These cells produced growth factors by regulating acute inflammation, which promoted the differentiation of mesenchymal stem cells and played an important role in maintaining bone homeostasis and promoting bone repair. The studies had confirmed that inflammatory cytokines such as Interleukin-4 (IL-4), IL-10 and IL-13 could promote osteogenesis, while tumor necrosis factor α (TNF-α), IL-6 and IL-1 had the opposite effect. GDs had been shown to stimulate immune cells to produce different responses. In this subsection, we reviewed the immunomodulatory mechanism of GDs-induced bone regeneration, which was normally in two ways.

One was that GDs stimulated macrophages to transform into M1 phenotype and up-regulated the expression of a variety of pro-inflammatory cytokines, thus creating a good immune microenvironment for osteogenesis. Subsequently, the immune microenvironment could stimulate stem cells to differentiate into osteoblasts. For example, Xue et al. studied the effect of GO on the inflammatory response of mouse macrophage line RAW-264.7 cells and found that the expression of proinflammatory cytokines IL-6, IL-1 β, Interferon γ (IFN-γ), Oncostatin M (OSM) and TNF-α was up-regulated in RAW 264.7 cells treated with GO. Subsequently, they used the supernatant of GO/RAW 264.7 as the conditioned medium to culture BMSCs, the results of which showed that the expressions of myeloiddifferentiationfactor88 (MYD88), BMP2, vascular endothelial growth factor (VEGF) and OSM of BMSCs were up-regulated in this conditioned medium, indicating that GO stimulation was beneficial to the formation of immune microenvironment of BMSCs osteogenesis. Finally, they researched the immunomodulatory mechanism of GO in the osteogenesis of BMSCs. The results demonstrated that the immune microenvironment induced by GO stimulated the osteogenic differentiation of BMSCs through OSM and nuclear factor kappa-B (NFκB)-VEGF signaling pathways.

The other was that GDs activated monocytes, which could increase the release of osteogenic factors and induce stem cells to differentiate into osteoblasts. For example, Bordoni et al. prepared calcium phosphate (CaP)-GO nanoparticles in phosphate solution/GO/cyclohexane/ethanol system by emulsion synthesis, and verified their activating effect on monocytes. They found that the expression of proinflammatory cytokines such as IL-6, TNF-α was up-regulated in the cell culture medium, which confirmed that monocytes were activated. Then, the nanoparticles were co-cultured with monocytes and BMSCs to compare the effects of the introduction of CaP and CaP-GO on the osteogenic differentiation of BMSCs. The results showed that GO stimulated monocytes to produce higher osteogenic factors (OSM), which further up-regulated the Wnt and BMP signaling pathways of BMSCs, and finally promoted the osteogenic differentiation of the BMSCs.

**Application of GDs in Real-Time Detection of Bone Repair Process**

In the field of bone repair, GDs were used not only as the bone repair material, but also as the contrast agent for in vivo imaging. By being functionalized or compounded with other materials, GDs could be applied to real-time detection of bone repair processes to detect the changes in the structure, tumorigenicity, degradation and mineralization of bone repair materials in complex environments in vivo (Figure 7 and Table 5). In this section, we summarized the related principles and advantages of GDs as a contrast agent.

GDs were used as the contrast agent from these three aspects. Firstly, it had a photoelectric effect to convert the
absorbed electromagnetic energy into sound waves, thus being used in CT and ultrasound. In most cases, GDs were usually prepared in composites for these applications. For example, Kundu et al.\textsuperscript{48} blended GO with AgNO\textsubscript{3}/NaBH\textsubscript{4}/human serum albumin (HSA) and AgNO\textsubscript{3}/NaBH\textsubscript{4}/bovine serum albumin (BSA) system to prepare silver nanocluster (AgNCs)/GO composites as contrast agents and fluorescent markers. The results of CT scanning showed that the introduction of GO enhanced the imaging effect of the composites in Swiss albino mice (Figure 8A and B). Moreover, GO ensured the sustained release of AgNCs in vivo as a stable fluorescent marker and reduced the cytotoxicity of the composite. In another case, Toumia et al.\textsuperscript{49} coated graphene on the surface of polyvinyl alcohol (PVA) microspheres to prepare graphene/PVA (G/PVA) composites, which were endowed with the function of graphene to convert absorbed radiation into ultrasound waves for contrast agents and based on the nano-microsphere structure, drugs could be loaded for bone repair. The results of in vivo experiments showed that G/PVA as a contrast agent had excellent photoelectric sensitivity and ultrasonic echo ability. In addition, G/PVA could be degraded in vivo and had no cytotoxicity.

Secondly, GDs could obtain magnetism through functionalization to meet the electrophysiological requirements and be used in nuclear magnetic resonance imaging.\textsuperscript{56,158,159} For example, Enayati et al.\textsuperscript{159} prepared rGO by γ-ray irradiation to study the effects of oxygen functional group content and structural defects on the magnetic properties of rGO. The results showed that under the action of γ-ray irradiation, the epoxy groups on the surface of GO were dislodged, resulting in deoxidation defects, so the magnetic properties of rGO were enhanced. Moreover, the degree of oxidation on the surface of GO would also affect its magnetic properties after reduction. The stronger the degree of oxidation, the higher the content of epoxy groups in GO, correspondingly, the higher the defect density of rGO, the stronger the magnetic properties. In another case, Hu’s study confirmed that fluorinated graphene oxide (FGO) had magnetic properties due to the existence of C-F bond, so the FGO could be used as a carbon-based contrast agent for magnetic resonance imaging (MRI) without other magnetic nanoparticles.\textsuperscript{56} Moreover, compared with the distribution of proton (\textsuperscript{1}H) in the human body, the distribution of \textsuperscript{19}F was rarer. Therefore, FGO had a higher signal-to-noise ratio compared with the traditional MRI contrast agents.

Thirdly, GDs could be combined with magnetic particles and then introduced into bone repair materials, which could not only make use of the excellent properties of GDs to accelerate bone regeneration, but also enabled that magnetic particles were evenly dispersed and tracked by nuclear magnetic equipment.\textsuperscript{53,160–162} For example, Chen et al.\textsuperscript{53} prepared a composite scaffold containing gelatin/PCL/ultra-small paramagnetic iron oxide/GO (Gel/PCL/USPIO/GO) by freeze-drying. The negatively charged GO could effectively combine with the positively charged USPIO, to ensure that the USPIO provided a stable imaging effect for the scaffolds. The results of MRI detection showed that the scaffolds still had clear visual imaging after 7 days of degradation in vitro (Figure 8C). Furthermore, GO could stimulate the proliferation and osteogenic differentiation of BMSCs and provided mechanical support for composite scaffolds. In another case, Pramanik et al.\textsuperscript{162} prepared Fe\textsubscript{3}O\textsubscript{4}/rGO/poly(hydroxybutyrate-co-valerate) (PHBV) nanocomposite scaffolds. The results showed that the large SSA and lots of surface charges of the rGO enhanced the capture of the graphite lattice by the magnetic particles. Moreover, the rGO could make Fe\textsubscript{3}O\textsubscript{4} nanoparticles uniformly distributed on its surface, thus realizing the imaging of Fe\textsubscript{3}O\textsubscript{4} nanoparticles in MRI. Therefore, the degradation process of PHBV-based composites could be...
detected in real time by MRI. In addition, the introduction of rGO enhanced the mechanical strength and electrical conductivity of the composite scaffolds.

**Some Concerns About the Application of GDs in Bone Repair**

There were still some concerns about the application of GDs in bone repair materials, including their potential cytotoxicity and possible residues in vivo. In this section, on the one hand, we summarized the mechanism and related factors of GDs-induced cytotoxicity. On the other hand, we reviewed the possible strategies to avoid or decrease GDs-induced cytotoxicity, mainly including biological wrapping and biodegradation.

### Main Factors of GDs-Induced Cytotoxicity

The relative studies had shown that GDs might induce cytotoxicity by depleting mitochondrial membrane potential (MMP) and increasing intracellular reactive oxygen species (ROS), thus triggering apoptosis. Moreover, the potential cytotoxicity of GDs was related to their concentration, size, shape, and degree of functionalization.

Firstly, some studies had confirmed that GDs had concentration-dependent cytotoxicity. When the concentration of graphene in the solution was too high, it could cause a high level of oxidative stress by accumulating on the membrane, resulting in physical destruction of cells in the way of membrane penetration, or cell vitality was endangered through the interaction between the

### Table 5 Application of GDs Composites in Real-Time Bone Detection

| Testing Equipment | Composites | Combination Mode/Modification Method | The Used Properties of Graphene | Functions | References |
|-------------------|------------|--------------------------------------|---------------------------------|-----------|------------|
| Computed tomography | AgNCs/GO  | AgNCs adsorbed on graphene | Photoelectric effect | Significant contrast enhancement of bone tissues in mice models | [48] |
|                   | BaGdF<sub>5</sub>/GO | BaGdF<sub>5</sub> NPs were directly grown on the surface of GO nanosheets by a solvothermal method | Photoelectric effect | Effective imaging of tumor model in vivo | [52] |
| Ultrasonic        | Graphene/PVA | Graphene was bound to the surface of PVA microspheres by EDC/NHS. | Photoelectric effect | Performing real-time multiplex photoacoustic imaging in vivo | [49] |
| Magnetic resonance imaging | Ultra-small paramagnetic iron oxide (USPIO)/GO | Covalent binding between amino groups on USPIO and carboxyl groups on GO | Excellent electrical conductivity | Real-time detection of implant degradation | [53] |
|                   | Fluorinated graphene oxide (FGO) | Fluorinated graphite polymer (FG) is dispersed in a 9:1 mixture of concentrated sulfuric acid and concentrated phosphoric acid. | Easy to paramagnetic modification | It has higher retention time and imaging efficiency than the traditional metal contrast medium. | [56,158] |
|                   | Reduced graphene oxide (rGO) | Reduction of GO by γ-ray irradiation | Easy to paramagnetic modification | Higher imaging efficiency | [159] |
|                   | Graphene quantum dots/Gd-DOTA | Graphene quantum dots were crosslinked with Gd-DOTA by EDC/NHS. | Excellent electrical conductivity, low toxicity and high water solubility | Improving the contrast enhancement of magnetic resonance imaging | [160] |
|                   | Gd<sup>3+</sup>/Graphene nanoribbons | The reagent containing Gd<sup>3+</sup> was blended with graphene, and the mixture was covalently bound with PCP. | Large specific surface area | Labeling porcine bone marrow mesenchymal stem cells for magnetic resonance imaging | [161] |
hydrophobic surface of graphene and the cytoplasmic matrix. For example, Pang et al.\textsuperscript{163} found that graphene had little cytotoxicity when the concentration of graphene was in the range of 5–10 μg/mL. In vitro studies had demonstrated that when the concentration of graphene or GO was 10 μg/mL, the adhesion and morphology of BMSCs were good. When the concentration of graphene was more than 50 μg/mL, the cell size became smaller, its surface secretion increased and the microvilli extended. When the concentration of GO was more than 50 μg/mL, BMSCs shrunk and deformed, and most of the cells died. These results also suggested that at the same concentration, the cytotoxic effect of GO on BMSCs was higher than that of graphene. This might be due to the stronger adsorption capacity of GO, which could adsorb serum proteins to form protein inclusion bodies.

Secondly, size had an important effect on the cytotoxicity of GDs.\textsuperscript{165–167} For example, Wychowaniec et al.\textsuperscript{165} prepared GO paper using two different sizes of GO sheets (4.9 μm and 151.6 nm). The results of in vitro experiments showed that GO paper prepared with smaller size GO sheets had higher cytotoxicity, which might be due to the different mechanisms of interaction between GO sheets of different sizes and cells. The larger GO sheets first attached to the cell surface, then the membrane invagination and pseudopodia extension, and finally entered the

Figure 8 (A) Preprocessing CT images of Swiss albino mice were treated with AgNCs/GO composites. (B) Postprocessing CT images of Swiss albino mice were treated with AgNCs/GO composites, including top view, right view, front view and 3D view. (C) magnetic resonance imaging (MRI) contrast of Gel/PCL/USPIO/GO scaffolds with and without BMSCs in vitro. Reproduced with permission of American Chemical Society from Kundu N, Mukherjee D, Maiti TK, Sarkar N. Protein-guided formation of silver nanoclusters and their assembly with graphene oxide as an improved bioimaging agent with reduced toxicity / Phys Chem Lett. 2017;8(10):2291–2297. (A and B). Further permissions related to the material should be directed to the ACS.\textsuperscript{48} Reproduced with permission of IOP Publishing, Ltd from Chen J, Hu HL, Feng LB, et al. Preparation and characterization of 3D porous conductive scaffolds with magnetic resonance enhancement in tissue engineering. Biomed Mater. 2019;14(4):045013; permission conveyed through Copyright Clearance Center, Inc (C).\textsuperscript{53}
cells mainly through phagocytosis, while the smaller GO sheets mainly entered the cells through molecular sieve-mediated endocytosis. In another case, Gurunathan et al. studied the cytotoxic effects of GO with sizes of 100 nm and 20 nm (GO-100 and GO-20) on Leydig (TM3) and Sertoli (TM4) cells. The results showed that the cell viability and cell proliferation induced by GO-20 were significantly lower than those induced by GO-100, the reason of which might be that GO-20 could significantly increase intracellular ROS, induce the significant decreasing of MMP, up-regulate the expression of pro-apoptotic genes, and down-regulate the expression of anti-apoptotic genes. Moreover, they also found that the cytotoxicity of GO-100 and GO-20 might be due to the regulation of EGFR/AKT signal transduction pathway of the TM3 and TM4 cells.

Thirdly, the various shapes of GDs also led to the complexity of their cytotoxicity. For example, Wu et al. prepared rGO using Na2S as a reductant and compared the cytotoxicity of GO and rGO with different reduction degrees on primary bone marrow-derived macrophages (BMDMs). The results of in vitro evaluation showed that the cytotoxicity of rGO was higher than that of GO, the main reason of which was found not the change of composition caused by reduction, but the morphology change from the flake of GO to the polygonal crimped shape of rGO. Therefore, rGO with the structure stimulated more ROS production than GO in BMDMs. Moreover, the observation of cell morphology showed that GO enlarged the cell body and prolonged the processes of BMDMs, while rGO caused the size of cells to become smaller and the length of processes shortened, which might be due to the fact that rGO was more likely to condense in the cytoplasm and cause higher cytotoxicity to macrophages. In another case, Dervin et al. prepared two kinds of rGO (H-rGO and AA-rGO) using hydrazine and ascorbic acid as reductants, respectively and compared the cytotoxicity of GO, H-rGO and AA-rGO to A549 human lung epithelial cancer cells. Transmission electron microscope (TEM) observation demonstrated that the shape of H-rGO remained almost unchanged after hydrazine treatment, while the plane structure of AA-rGO was distorted and sharp protuberances appeared on the edge after ascorbic acid treatment. The results of in vitro experiments showed that the cytotoxicity of AA-rGO was higher than that of GO and H-rGO. The reason for these results might be that the irregular shape of AA-rGO reduced the integrity of cell membrane, resulting in the loss of cell structure and stability, and finally led to cell death. The above results indicated that the shape of GDs had an effect on cytotoxicity.

Main Means of Avoiding or Deceasing GDs-Induced Cytotoxicity

GDs might cause potential harm if they stayed in the human body for a long time. Until now biological wrapping and biodegradation could be used as the main feasible strategies to avoid or decease GDs-induced cytotoxicity. GDs could be probably wrapped by regenerated bone tissues with the time going on after implantation, resulting in the avoiding or decreasing diffusion of the residues into other tissues or organs. For example, Zhou et al. prepared three-dimensional porous HA/rGO scaffolds and implanted them into the 6 mm critical-size segmental diaphyseal bone defect of rats to repair the defect. The results showed that rGO was significantly wrapped in regenerated bone tissue at month 6 after the implantation, which was accomplished mainly by the following steps: growing of the new bone into the scaffold, cracking of the substrate under compression, being exposed of GDs that further accelerated bone regeneration, and eventually being wrapped of GDs by the newly regenerated bone. Histological results demonstrated that there was no obvious deposition of rGO and abnormality in the heart, liver, spleen, kidney, lung and pancreas of the animals. The above results might be related to factors such as the pore size, porosity, GDs content, substrate degradability of the scaffolds.

On the other hand, the biodegradation of GDs was mainly through the action of some specific enzymes, such as myeloperoxidase (MPO) and horseradish peroxidase (HRP). These enzymes mainly interacted with hydrogen peroxide (H2O2) produced during metabolism in the body. The H2O2 was catalyzed by the enzyme to produce strong oxidant hypochloric acid and reaction free radical intermediate, which cleaved the hydroxyl and epoxy groups connected to C atoms on GDs to form porous amorphous graphene. On the basis of the above principles, the effects of dispersion, layer number, oxygen-containing functional group content and functionalization of GDs on their biodegradation rate were studied. For example, Kurapati et al. found that the better the dispersion of GO, the easier it was to be degraded by MPO. These effects included electrostatic repulsion between GO layers leading to better dispersion of GO to form stable colloids, and MPO tended to interact with the negative charge on...
the surface of GO, thus inducing GO degradation at the position of carbon atoms connected to oxygen-containing groups. Subsequently, they also found that monolayer graphene (MLG) and few layer graphene (FLG) could be degraded by MPO in the presence of H₂O₂ and NaCl without chemically functionalized and stable surfactants.⁶³ The degradation rate of MLG by MPO was slower compared with that of FLG, which was probably because there were fewer defects and low content of oxygen-containing functional groups on the surface of MLG, so that there were relatively few contact sites between MLG and MPO.

Because the functional groups on the surface of GDs could affect their degradation, the regulation effect of functionalization on the degradation rate of those had been studied. For example, Li et al.¹⁷² prepared functionalized GO by polyethylene glycol (PEG) and bovine serum albumin (BSA). The results of in vitro experiments showed that both peg-modified and BSA-functionalized GO could not be degraded by HRP, which might be due to the steric hindrance provided by PEG and BSA, thus preventing the interaction between HRP and GO surface. Subsequently, they functionalized GO with disulfide bonds to form GO-SS-PEG and found that GO-SS-PEG could be degraded in mice, which possibly because GO-SS-PEG interacted with macrophages in the body destroyed the disulfide bonds in GO-SS-PEG, thus promoting the contact of the enzyme with GO.

Above all, it is possible for GDs to be biologically wrapped in vivo to decrease their negative effects on other tissues or organs. Moreover, another one of the effective methods to further reduce and avoid the cytotoxicity of GDs could be to introduce GDs evenly into the substrate, which not only avoided the direct contact between GDs and cells to reduce the probability of cell destruction, but also attracted the biological enzymes through the charge on the surface of GDs and substrate, thus accelerating the degradation of GDs.

**Summary and Prospect**

It has been proved that GDs, as a bone repair material, have the advantages of excellent mechanical properties, good electrical conductivity, large SSA, easy modification and so on. Among them, the mechanical reinforcement of GDs on bone repair composites is mainly affected by their content. With the increase of GDs content, the mechanical properties of composites have a trend of increasing at first and then decreasing, mainly through agglomeration and changing the mechanical mechanism of pullout effect. Then, the good electrical conductivity of GDs can not only improve the electrical conductivity of bone repair materials directly through the characteristics of low impedance and high charge, but also indirectly enhance those by changing the phase structure of the matrix, thus transmitting electrical signals to improve the bone regeneration behavior of cells and tissues. Subsequently, through the modification of different functional groups, the properties of GDs such as dispersibility and hydrophilicity are significantly improved in the composites, thus enhancing the functions related to bone repair and the binding ability of GDs to active substances.

Moreover, GDs can combine with various matrices in different ways, endowing the matrix with better properties, resulting in further improving the functions of cells such as adhesion, proliferation, osteogenic differentiation and osteoblast-related mineralization. In addition, some studies have reported that the signaling pathways involved in GDs-induced bone regeneration might be affected by the content of GDs, functional groups and the structure of composites. In particular, some possible rules have been found. For example, the porous structure of composites containing GDs is tending to activate the MAPK signal pathway. Furthermore, it has been confirmed that GDs can regulate immunity to induce bone regeneration. After implantation, GDs first interact with immune cells to regulate the expression of a variety of pro-inflammatory cytokines, thus stimulating stem cells to differentiate into osteoblasts.

On the other hand, GDs can also be used as a contrast agent in the field of bone repair. Using photoelectric effect, composites containing GDs can be used for CT and ultrasonic testing. In another case, the magnetic properties of GDs-containing composites can be changed by functional modification of GDs or compounding with GDs and magnetic particles, which can be used in nuclear magnetic resonance. Finally, we also summarized the factors related to the mechanism of cytotoxicity induced by GDs. It has been confirmed that the cytotoxicity of GDs can be reduced or even avoided by controlling some parameters of GDs, such as the concentration, size and shape of nanosheets. Additionally, most studies have found that biodegradation or biological wrapping could be used as the feasible strategy to solve the residues of GDs.
Many studies have confirmed that GDs have a broad application prospect in the field of bone repair, but there are still some challenges to be solved before they are used in clinics, including the following aspects.

Firstly, the uniform dispersion of GDs in the matrix has been found to significantly affect the performance of bone repair materials, but the evaluation is only qualitative, lacking common evaluation criteria.

Secondly, for the use of GDs to reinforce bone repair materials, although effects of many parameters such as the number of layers, content and functional modification have been investigated, further studies on the influence of the matrix porosity together with those parameters are necessary because most of the current bone repair materials are porous. On the other hand, a lot of work has been done to improve the combination of the GDs and matrices and enhance the reinforcing effectiveness, including the functionalization of GDs. However, the corresponding systematic research is still lacking, such as that on the effects of different composition means of the GDs and matrices including various kinds of covalent bindings and non-covalent combinations. Moreover, how the treatments affect the mechanical properties of GDs themselves has not been revealed.

Thirdly, it has been proved that the excellent conductivity of GDs can be used to conduct electrical signals, thus promoting the bone regeneration behavior of cells and tissues. At present, the effects of GDs parameters such as content, defect density and modification of the specific functional group on the electrical conductivity of the composites have been investigated. In future, the effect of interface interaction between GDs and matrices on the electrical conductivity of composites should be systematically studied. For example, the effect of GDs binding with different matrices on the electrical conductivity of composites, or the effect of GDs binding and the same matrix on the electrical conductivity of composites. Moreover, the effects of electrical conductivity of GDs composites on osteogenic function and bone tissue regeneration behavior should be studied in vitro and in vivo without external electric field. In addition, the addition of conditions to promote bone repair has become a more recognized method. For example, piezoelectric materials containing GDs have been prepared, which can convert external stimuli into electrical signals, thus promoting bone regeneration through the conductivity of GDs. In future research, it is necessary to develop more functional composites based on GDs, which can convert various external stimuli such as magnetic stimulation and light stimulation into electrical signals, so as to further stimulate the osteogenesis-related functions of cells.

Fourthly, some progress has been made in the mechanism of GDs-induced osteogenesis and immune regulation. In order to achieve controllable regulation of specific signal pathways, it is necessary to further study how the parameters such as the size, shape, microstructure of GDs and the preparation technology of composites with different matrices affect cell-induced osteogenesis and immunomodulatory mechanism.

Fifthly, GDs has been used as a contrast agent in the field of bone repair to achieve real-time detection of the bone repair process, but the effects of GDs parameters such as size, defect density and the method of assembly with the substrate on the imaging effect need to be studied systematically.

Sixthly, in terms of the potential cytotoxicity of GDs, the effects of some parameters such as concentration, size and shape of GDs on cytotoxicity have been investigated to some extent. In future, it is necessary to deeply study the influence of those parameters on the degree and detailed process of damage to cells and related signaling pathways by utilizing some special techniques or methods, such as high-content cell imaging analysis system and freezing electron microscope. On the other hand, the used animal models are mainly rats, lacking large animal models such as goats, canines and monkeys. Moreover, due to the limitation of experimental time, there is a lack of chronic toxicity studies for more than 6 months, such as those on the possible effects of GDs on animal reproduction and carcinogenesis.

In addition, the biodegradation of GDs should be deeply explored, such as how the changes in the size, crimp and folding of GDs affect their cytotoxicity or inflammation in the process of enzymatic degradation, as well as the study of different metabolic pathways of GDs, such as urine, sweat and feces. These technical breakthroughs will clear the way for GDs to be used in clinics earlier.

Acknowledgments
The authors acknowledge financial support from the National Natural Science Foundation of China (No. 31771042), Fok Ying Tung Education Foundation (No. 141039), the Fund of Key Laboratory of Advanced Materials of Ministry of Education (No. 2020AML10), International Joint Research Center of Aerospace Biotechnology and Medical Engineering, Ministry of Science and Technology of China, the 111 Project (No. 2020A07010101).
B13003), and the Academic Excellence Foundation of BUAA for PhD Students.

**Disclosure**
The authors declare no conflicts of interest for this work.

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