3D printing of graphene-based polymeric nanocomposites for biomedical applications

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Abstract

Additive manufacturing techniques established a new paradigm in the manufacture of composite materials providing a simple solution to build complex, custom designed shapes. In the biomedical field, 3D printing enabled the production of scaffolds with patient-specific requirements, controlling product architecture and microstructure, and have been proposed to regenerate a variety of tissues such as bone, cartilage, or the nervous system. Polymers reinforced with graphene or graphene derivatives have demonstrated potential interest for applications that require electrical and mechanical properties as well as enhanced cell response, presenting increasing interest for applications in the biomedical field. The present review focuses on graphene-based polymer nanocomposites developed for additive manufacturing fabrication, provides an overview of the manufacturing techniques available to reach the different biomedical applications, and summarizes relevant results obtained with 3D printed graphene/polymer scaffolds and biosensors.

Keywords: Additive manufacturing, 3D printing, Graphene-based nanocomposites, Biomedical applications

Introduction

Graphene, a one-atom thick two-dimensional carbon material, has remarkable mechanical, thermal, electrical, and physicochemical properties. However, practical applications of graphene are still limited. As a nanofiller for polymer-based composites, graphene can significantly improve the properties at low incorporation levels. These composites are potentially suitable for a wide range of applications in electronics, energy storage, thermal engineering, automotive and healthcare [1].

The biomedical device industry is vast and expected to be valued at 409.5 billion dollars by 2023 [2]. Graphene-based composites are mostly used for the fabrication of 3D structures such as scaffolds for tissue engineering [3] and also for biosensing [4, 5]. Suitable scaffolds have been developed using conventional fabrication methods such as solvent casting, freeze-drying and salt leaching [6–8]. However, control over the scaffold geometry is limited [3] and the production of multifunctional, multi-material scaffold structures is challenging [2]. The recent advancements in additive manufacturing brought about new and exciting possibilities for the fabrication of complex architectures for biomedical applications [3].

Additive manufacturing (AM), also known as 3D printing or rapid prototyping, denotes a group of techniques where a computer-aided design (CAD) model is converted into a 3D object, which is successively built layer-by-layer. This contrasts obviously with the conventional formative (e.g., injection moulding, casting) and subtractive methods (e.g., turning, drilling). AM techniques can fabricate complex structures (eventually not possible to obtain by the other methods) in a relatively short time and with low cost, since they do not require a
shaping features induced revolutionary applications of AM in healthcare, aerospace, automotive, electronics and civil construction [2, 9, 10]. In the medical/biomedical field, AM has been applied in tissue engineering to design personalized scaffolds or artificial tissues and organs for transplants, drug delivery systems, probes and tools to detect specific medical parameters, orthoses, prostheses and implants, anatomical models and medical instruments for diagnostics and surgery [11, 12].

It is estimated that biomedical applications share about 11% of the AM technology market and this percentage is steadily increasing [13]. Indeed, AM is suited to manufacture small quantities of customized prototypes or products, which is particularly useful as patient-customized products are frequently required [14]. The availability of 3D printed prototypes of a patient unique anatomy can improve surgery planning, implant design, and provide specialized medical procedure training. The ability to 3D print not only biomaterials, but also living cells and/or other biological components (e.g. distinct proteins, growth factors), known as bioprinting, enables the creation of biological constructs that can regenerate or augment lost/damaged tissues and organs [2].

The use of polymer nanocomposites combined with the versatility of AM provide great potential to meet the demands of a wide range of clinical needs [2, 3] by creating personalized implants, organ printing, drug delivery devices as well as regenerative scaffolds. In particular, the incorporation of graphene and its derivatives can improve the dimensional accuracy as well as the mechanical, electrical and biological properties of novel biomedical devices [2].

This review discusses the recent advances of using AM techniques and graphene-based polymer nanocomposites to manufacture biomedical products and devices, with emphasis on scaffolds and biosensors. The main 3D printing techniques used with polymers are first introduced. Then, the main preparation methods of graphene-based polymer nanocomposites are reviewed. Finally, representative examples of 3D printed graphene-based scaffolds and biosensors are discussed.

### 3D printing techniques for polymers

AM techniques use computer files describing the 3D geometry of the part or prototype to be produced, to successively build a series of horizontal 2D layers in the vertical direction, usually from the bottom to the top. The techniques vary in terms of the material deposition procedure, the physical form of the material (liquid, solid or powder-based) and the nature of the material (e.g., thermoplastics, thermosets, nanocomposites, fiber composites). The length-scale, dimensional accuracy and surface finish of the printed object will also depend on the AM technique utilized. Moreover, some techniques are better suited to industrial production than others [13, 15].

Generally, the printing process involves the following steps (see Fig. 1):

- a) Definition of the geometry and dimensions of the 3D structure to be manufactured, using computer-aided design (CAD) software;
- b) Conversion of the design into a digital approximation (using triangulations) as a Standard Tessellation Language (STL) file format;
- c) Slicing of the 3D model into layers of specified thickness, using dedicated software;
- d) Use of G-code commands (often via a user-friendly interface) to define how the printer will work during the printing sequence. For example, the in-fill density will determine the porosity of a part;
- e) Transfer of the files to the actual 3D printer;
- f) Layer-by-layer manufacturing of the part, eventually followed by post-processing operations (such as removing support material, sintering, or polishing the surface) [16].

Figure 2 presents three major families of AM techniques used for graphene-based composites. 3D printing of polymer/graphene composite parts has been mainly carried out using extrusion-based and powder-based techniques such as fused deposition modeling (FDM), direct ink writing (DIW) and selective laser sintering (SLS), as well as stereolithography (SLA) [17].

#### Fused deposition modeling

FDM involves the deposition of thin filaments of thermoplastic polymers or composites onto a support, to create a vertical series of horizontal 2D layers of the part under manufacture (see Fig. 3a). The 3D printer is fed with a spool of a previously extruded filament with standard diameter of 1.75 mm or 3 mm, which is pushed into a heated nozzle - generally by means of a pair of counter-rotating gears - where it melts and is extruded as a thinner filament (diameter of approximately 0.1–0.2 mm), which is deposited to fabricate the 3D part. The nozzle moves in the XY direction to build each layer, and vertically (Z-axis) to create other layers [19]. An interesting variant of FDM is Free Form Extrusion (FFE), where a screw extruder replaces the nozzle to produce the thin filament. The extruder can be directly fed by polymers/composites in pellet/powder form, thus widening quite significantly the range of materials that can be printed, considering the relatively limited scope of commercial materials in filament form available for printing. FDM/FFE is by far the most used 3D printing technology [20]. Currently, many 3D printers can handle simultaneously two or more materials. This brings about several advantages:
the production of parts with specific geometrical features; in the case of a geometry consisting of a vertical cylindrical column with a wide disk on top, it is necessary to deposit two materials; one generates the column, while the other will serve as support when printing the disk above; the support material is subsequently removed during post-processing;

- the production of parts with gradient of particular local properties (e.g., soft vs. rigid zones);
- obtaining aesthetical effects, for example by mixing filaments with different colors and in varying proportions in a single nozzle.

Despite of its apparent simplicity, FDM/FFE is governed by a large number of process parameters, which creates requirements in terms of materials properties, particularly rheological and thermal [21]. Operating parameters include extrusion velocity and temperature, printing velocity, build orientation (the rotation of the part in the manufacturing space around the axes of the machine’s coordinate system), deposition sequence (the path taken by the filament during deposition, for example, unidirectional and aligned, unidirectional and skewed, or perpendicular), infill % (0% is hollow while 100% is solid), environment temperature, and support temperature. Flow in the nozzle depends on the melt viscosity of the polymer/composite, which is a function of temperature and shear rate, i.e., extrusion velocity and temperature. Upon exiting the nozzle, the molten filament swells, but is simultaneously stretched axially by the printing head (the filament diameter resulting from these two conflicting effects depends on the viscoelastic response of the material). Deposition involves filament deformation and bonding with contacting filaments previously deposited. Therefore, once deposited, each filament should solidify quickly to minimize its deformation due to its own weight and/or the weight of the material.
that will be deposited above it, thus preserving dimensional accuracy. Conversely, the new filament should remain sufficiently hot during enough time, to ensure adequate bonding (i.e., molecular diffusion) with the neighboring filament(s) [22].

Therefore, process parameters and material properties influence the morphology and properties of printed parts. These are usually significantly anisotropic and may exhibit warping and eventual delamination due to differences in local shrinkage. Adequate bonding between filaments and good porosity control are key factors to obtain quality parts [19, 23]. Fiber and nanofiller matrix reinforcement usually cause improvements in the mechanical properties [23].

Direct ink writing
DIW uses a printing mechanism similar to that of Fused Deposition Modelling but focuses on meso- and micro-scale parts, and generally handles materials with viscosities lower than those of typical polymer melts. The material(s) is (are) supplied under controlled flow rate using a piston (e.g. a syringe), an Archimedes-type screw, or pneumatic force [24] and are heated, so that the nozzle extrudes a liquid “ink” or a “paste” filament that is subsequently printed (Fig. 3b). The pneumatic force system is easy to install and enables pressure adjustment, whereas the screw system is more appropriate for high viscosity materials [18]. If a low heating temperature is used, cells and bacteria can survive during printing, making DIW suitable for biomedical applications. Biopolymers [25–27], thermoplastics [28], hydrogels [29–32], organic/inorganic monomers [33] or graphene-based nanocomposites [34–36] have been printed. Several DIW printers designed for healthcare and food research are currently available in the market, many having a temperature-controlled building platform [17].

As in FDM, the printability of a given material is related to its rheological response. The ink is extruded as liquid, but its viscosity should immediately increase thereafter, so that the printed structure holds without significant dimensional distortion. Hence, rheofluidifying materials, whose viscosity decreases with increasing shear rate, as well as fluids with a yield stress, are recommended. This increase in viscosity can also be induced by cooling, via phase change, or achieved through chemical reactions [37]. Low viscosity (often Newtonian) fluids can also be used in DIW, by extruding into a coagulation liquid where they form a gel. High viscosity inks are thus easier to print, but they tend to clog the nozzle. Proper ink formulation is a key issue in DIW [17].

Stereolithography
SLA consists on the layer-by-layer deposition of a photo-curable liquid resin/monomer, which cures/polymerizes and solidifies by irradiation with a spatially controlled ultraviolet radiation (UV) light or laser beam, forming the desired pattern (see Fig. 4a). Quick solidification is important in order to hold subsequent layers. After printing, the unreacted resin is removed. The thickness of each layer is controlled by the energy of the
light source and exposure time. Post-processing may involve heating or further photo-curing, in order to achieve higher mechanical performance [17].

SLA is one of the earliest additive manufacturing techniques. Although relatively slow, expensive and displaying a limited range of materials for printing, it produces high quality parts, and higher resolution that those produced by extrusion-based techniques, with an accuracy as low as 10 μm [39].

Materials and nanocomposites for SLA must fulfill two important requirements: i) Fast response to the light irradiation (i.e., rapid initiated polymerization) in order to assure swift solidification; ii) sufficient low viscosity, to allow for the dipping of the resin layer. The latter usually limits the level of filler incorporation and demands a homogeneous filler dispersion [17].

Selective laser sintering
SLS employs a laser to progressively sinter thin layers of powdered polymer/composite on top of a platform, to create a 3D part. The manufacturing cycle may start by preheating the powder to a temperature just below the formation of a melt. Then, the laser scans one horizontal slice of the geometry, sintering the particles together to create one solid layer. Next, the platform lowers vertically the equivalent to the thickness of one layer. A roller applies a new layer of powder material on top, and the above steps are repeated until printing is completed. The part cools down inside the printer and then is removed and cleaned [40, 41]. Generally, SLS parts exhibit a slightly rough surface, hence post-processing operations are often carried out [42]. Unlike most other AM techniques, SLS does not require the use of support material to generate parts with overhanging features, since the unfused powder supports the part during printing, thus facilitating the production of particularly complex structures [43]. SLS is widely used for advanced applications, such as scaffolds for tissue engineering. Powder particle size and particle distribution determine the density of the printed part. Normally, powder particles with diameters in the range of 10–100 μm are used to facilitate spreading, as well as play efficiently the role of support material if unfused [44]. Nevertheless, the porosity of the part can also be controlled through a post-treatment by infiltration [45]. A low melting/sintering material temperature also facilitates the use of the laser.

For the 3D printing techniques most used for biomedical applications, Table 1 identifies the most common polymers and applications, together with the main advantages and limitations.

Graphene-based polymer nanocomposites
Graphene and graphene derivative particles
Graphene consists of a single 2D sheet of sp²-hybridized carbon atoms arranged in a crystalline hexagonal lattice. Its properties are estimated to be similar or even higher than those of carbon nanotubes [47]. Graphene is not permeable to gases, has high flexibility and an optical transmittance of 97.7% of the total incident light over a wide range of wavelengths [48]. Yet, graphene has found limited practical applications due to the difficulty and cost to obtain, at a commercial production scale, in the form of a single defect-free carbon sheet. A good method to produce graphene derivatives at large scale has been the exfoliation of graphite, using different procedures, such as depicted in Fig. 5a. The most significant methods have been: i) thermal expansion of graphite intercalation compounds (GICs), ii) chemical oxidation of graphite to obtain graphene oxide (GO) [24], and iii) liquid phase exfoliation (LPE) of graphite using liquids of appropriate surface tension and application of high shear forces [51].
GICs are formed by the insertion of atomic or molecular layers of different chemical species (alkali metal, metal oxide, metal halogenate, etc.) within the graphene layers, which receive or donate electrons to the intercalated species. The graphite obtained with intercalation compounds presents increased interlayer spacing and weakening of the van der Waals interactions between graphene layers. Subjecting GIGs to sufficiently high temperature or microwave radiation, it will further expand due to vaporization of the intercalate, forming expanded graphite (EG) flakes with nano-size thickness [48, 52].

GO is prepared by treatment of graphite flakes with oxidizing agents, bonding oxygen containing functional groups at the graphite surface and edge carbons. In this process the hybridization of part of the graphene carbon changes from sp² to sp³, expanding the spacing between the graphene layers and facilitating the exfoliation of GO single layers in liquid phase, aided by the application of mechanical forces. GO is electrically insulating, the electrical resistivity varying with the degree of oxidation attained [52], and is stable in aqueous solution due to the polar nature of the oxidizing groups. The subsequent reduction of GO produces reduced graphene oxide (rGO) which can partially recover the electrical conductivity, but a high oxygen content and physical damage remains in the rGO structure. GO and rGO are less performing than graphene, but the oxygen containing functional groups at the GO or rGO surface potentiate covalent bonding and strengthen the interface with the polymers, or provide reactivity with biomolecules [24].

LPE of graphite is a cost-effective process that can be up-scaled to mass-production of few layer graphene (FLG), typically based on the application of high shear to graphite suspensions in organic solvents with surface tension near 40 mJ m⁻² [51], ionic liquids, or water-

| AM Technique | Polymer | Resolution | Biomedical Applications | Advantages | Disadvantages |
|--------------|---------|------------|-------------------------|------------|---------------|
| FDM          | Poly(lactic acid (PLA); acrylonitrile butadiene styrene (ABS); polycarbonate (PC); nylon | 100–200 μm | Scaffolds for cell culture, and tissue engineering | Low cost, high strength, composite materials, not using toxic solvents | Nozzle clogging, anisotropy, lower resolution, thermal degradation of polymer |
| DW           | Alginate (ALG); chitosan (CHI); collagen; gelatin; silk | 1–100 μm | Tissue regeneration, wound healing, drug delivery | Structures with different geometries, sizes, and materials | Formulation of inks |
| SLA          | Photocurable resin (epoxy or acrylate-based resin) | 1.2–200 μm | Scaffolds for cell culture, tissue, and organ | High resolution, fast, good cell viability, nozzle free | Cytotoxicity, high cost, material limitation, possible harm to deoxyribonucleic acid (DNA) by UV |
| SLS          | Polycaprolactone (PCL), polylactic acid (PLA) | 100–200 μm | Temporary and degradable rigid implants | Good strength, easy removal of support, no solvents required | High cost, medium resolution, post-processing required |

| a) | Exfoliation of graphite intercalation compounds | Thermal expansion | Oxidation and reduction GO → rGO | Electrochemical activation | Sonication with stabilizer | Sonication with organic solvents |

b) | In situ polymerization | Mixing | Polymerization | Graphene | Monomer | + Curing agent |

| Solvent | Dispersion | Solvent evaporation | Graphene | + Polymer |

| Melt Mixing | Polymer | Melt blending |

Fig. 5 Schematic representation of graphene nanocomposite production: a) Preparing graphene by (1) exfoliation of graphite intercalation materials; (2) solvothermal exfoliation; (3) electrochemical activation uses an applied potential to drive exfoliation; (4) sonication in organic solvents uses direct sonication in various solvents, but without auxiliary stabilizer present; (5) sonication with stabilizer; (6) chemical oxidation to graphene oxide followed by chemical reduction back to reduced single-sheet graphene oxide; b) Dispersion of graphene in the polymer matrix by melt mixing, solution mixing or in situ polymerization. Adapted from [49, 50]
surfactant solutions [53]. The use of organic solvents and ionic liquids is not suited for most biomedical applications, thus LPE of graphite in water solutions of amphiphilic molecules has been favored in this field. LPE produces mainly FLG, with possible formation of a low concentration of monolayer graphene, however requiring the application of high shear rates, larger than $10^4$ s$^{-1}$.

Graphene nanoplatelets (GNPs) can be obtained from graphite through liquid phase exfoliation processes, exposure of acid-intercalated graphite to microwave radiation, ball milling and shear exfoliation. These techniques produce GNPs with varying morphology, within a range of thicknesses, lateral size, aspect ratio and concentration of defects [54, 55]. Commercially available GNPs are typically formed by a combination of single layer, few layer and nano-sized graphite platelets. Thus, its thickness can vary from 0.34 to 100 nm within the same production batch [55].

GNPs exhibit interesting properties such as low density, high aspect ratio, 2D morphology, high electrical and thermal conductivity, mechanical toughness, at low cost. Typical mechanical and physical properties (electrical and thermal conductivity) of graphene and graphene derivatives are presented in Table 2. In this way, GNPs have potential for application in several fields, including thermal interface materials, gas barriers, flame retardants, composites with sensing ability and in the biomedical area [55].

EG and FLG are often used as reinforcement in polymer composites. These nanoparticles are typically produced before composite preparation, and then mixed in the resin or polymer melt using adequate techniques that ensure dispersion of the nanoparticle agglomerates. Only the composites produced in the liquid phase (polymer dilute solutions, monomers for in situ polymerization) may allow the simultaneous LPE of graphite while mixing with the solution.

With the growing interest in graphene-based materials for applications in the medical field, the issue of biocompatibility has gained importance and the studies concerning its cytotoxicity and biocompatibility have multiplied, however there is no scientific consensus on this topic yet [64]. To date, in vitro studies have been performed with bacterial and mammalian cell models, and in vivo studies performed with animals and embryos [65]. The inherent hydrophobic nature of graphene may be responsible for cellular toxicity, since it interferes with the hydrophobic protein-protein interaction in the membrane, interrupting the cell’s metabolism, leading to cell mortality [66]. Functionalized graphene-based nanomaterials have shown reduced cellular toxicity [67] since it can help in pacifying the strong hydrophobic interaction of graphene/GO with cells and tissues [64]. Dextran-functionalized GO was found to accumulate in the reticuloendothelial system of mouse such as the liver and spleen after intravenous injection, and could be cleared from its body within a week without significant toxicity [68]. Besides, rGO has greater cellular toxicity than GO, due to the different surface functional group density and oxidation status [67].

However, it is not possible to make general statements on graphene-based materials safety or toxicity in eukaryotic cells, since it is strongly influenced by several factors such as the production methods, concentration, shape, distribution, surface area, number of layers, lateral dimensions, hydrophobicity, purity, particulate state, surface functionalization [69–71] and even to the type of cells that graphene was exposed to [72]. For instance, Wei et al. reported that pristine GO inhibited the proliferation of bone mesenchymal stem cells (BMSCs) at a high concentration of 10 μg/mL, but enhanced their proliferation at a low concentration of 0.1 μg/mL [73]. Similarly, Zhang et al. investigated the toxic effects of GO nanosheets on BMSCs. A high concentration of GO inhibited cell viability and membrane integrity. It was observed that the toxic effects of GO on BMSCs occur in a dose-dependent manner through the mitochondrial apoptotic pathway and autophagy. Exposure of BMSCs to GO at 0.1, 0.5 and 2.5 μg/mL for 24 h resulted in a slight increase in apoptosis by 5, 7 and 8% [74].

In addition, the in vivo effect of graphene-based materials is also dependent on the dose/time exposure, administration route, and the characteristics of the animals used in the experiment [65, 75]. Thus, conflicting results on the toxicity/biocompatibility of graphene derivatives have been reported. Some studies suggested that graphene-based materials are beneficial [76–81] and

| Properties                  | Graphene-based materials | GO  | rGO   | Graphite | GNPs |
|-----------------------------|--------------------------|-----|-------|----------|------|
| Thermal conductivity [W m$^{-1}$ K$^{-1}$] | ~ 5000 [56] | 3000 [57] | 30–250 [58] | 3000 [52] | 5000 [59] |
| Young’s Modulus [GPa]       | ~ 1000 [56]            | 200 [60]    | 250 [60]    | 20 [61]    | 1000 [62] |
| Electrical conductivity [S m$^{-1}$] | ~ 10$^6$ [63] | Insulator [60] | ~ 667 [58] | Insulator | 10$^6$ [59] |
others mention adverse responses including cytotoxicity [82, 83], tissue fibrosis [84], and inflammatory cell recruitment, usually in lung and liver [83, 85–87].

**Nanocomposites preparation methods**

Most preparation methods of commercial graphene-derivatives are based on the bulk exfoliation of graphite. Given the atomically flat surface of graphene, those materials may be obtained as cohesive agglomerates of individual nanoparticles, stabilized by non-covalent interactions through Van der Waals forces [88]. To fully explore the reinforcing effect and the functional properties resulting from the incorporation of graphene-derivatives into polymer matrices, it is essential to guarantee the uniform dispersion and distribution of the individual nanoparticles, in order to generate a large interfacial area between the composite components [89].

There are three main routes economically viable and scalable to produce well-dispersed polymer composites, as summarized in Fig. 5b. Solution and melt mixing are essentially based on the mechanical action and physical interactions between the polymer and nanoparticles, while in situ polymerization also uses a chemical approach [90].

Solution mixing is especially used for the small-scale manufacture of polymer composites whenever they are soluble in aqueous or organic solvents [91]. The technique entails the dispersion of the nanoparticles in a solvent by mechanical mixing, magnetic stirring, or sonication, followed by the addition of the polymer solution and mixing again by the same methods. The composite is obtained by removing the solvent by precipitation in a non-solvent or by direct solvent evaporation [92]. During mixing, the nanoparticles further separate-expand due to diffusion of the low viscosity polymer solution, and may adsorb polymer molecules at their surface, which helps preventing reagglomeration when the solvent is evaporated.

Melt mixing uses polymer processing equipment, typically internal mixers, or twin-screw extruders, to subject the flowing polymer melt and the graphene-based reinforcement to high hydrodynamic (shear and extensional) forces and complex flow fields. Processing parameters such as temperature, mixing speed, residence time and mixer geometry need to be adequately selected to obtain good dispersion levels [93]. Nevertheless, melt mixing generally attains lower dispersion levels than the remaining methods, although it exhibits strong advantages, such as avoiding the use of solvents, easily scaling to industrial production and high yield. The composite can be readily processed into a final product, or used as a masterbatch, i.e., diluted into the polymer using the same processing method to obtain the desired filler incorporation level.

In situ polymerization involves the preparation of a stable dispersion/suspension of the graphene-derivative particles in the liquid monomer, followed by polymerization. The latter may be initiated by heat, radiation, or by the addition of a catalyst. If the nanoparticles are functionalized with the adequate chemical functions, reaction may occur with the growing polymer chains during polymerization, enhancing interfacial strength and avoiding nanoparticle reagglomeration [88, 91].

For each of the above methods, Table 3 presents a list of polymers used for the manufacture of graphene-based composites, together with their advantages and limitations.

Kim et al. compared the performance polyurethane (PU)/graphene-based nanocomposites produced by different methods using the procedure represented in Fig. 6 [32]. Solvent-based methods yielded the best dispersion, improving the composite properties. Indeed, composites obtained by in situ polymerization exhibited a slightly higher percolation threshold when compared to those prepared by solution mixing. The electrical conductivity of the composites produced by in situ polymerization and solvent mixing was higher relative to that of materials produced by melting mixing, at similar reinforcement concentration [96].

**Properties of 3D printed graphene-based nanocomposites**

3D-printing of graphene-based nanocomposites has been the focus of extensive research, aiming to explore the potential to develop components for advanced applications in electronics, healthcare, energy storage, among others. Table 4 compares results obtained for the properties of 3D printed composites produced with different polymers and using different AM techniques. Despite the advantages of these technologies, some practical issues such as efficiency, cost, feasibility of mass production, still prevents the large-scale adoption of this solution [97].

**3D printing of graphene-based polymer composites for biomedical applications**

**Scaffolds for tissue engineering**

The development of scaffolds with higher mechanical properties suitable for in vivo application requires sophisticated manufacturing methods. Additive manufacturing techniques brought the possibility to process composite scaffolds with fine control of geometry and size, enabling the development of complex and personalized shapes with vast potential to regenerate distinct tissues [98].

With the advances of 3D-printing, the production of 3D biological constructs that mimic the structure and function of native tissues with precise control over the
positioning of both cells or other biological components (proteins, growth factors...) and biomaterials, called biofabrication, is now possible [99, 100]. Bioprinting such constructs comprises several technologies such as SLS, SLA, FDM or inkjet bioprinting, already addressed in section 2. Most of them were originally developed as AM technologies for rapid prototyping, but are included as biofabrication strategies when used for biomedical applications [99, 101].

The bioprinting process can be performed by two different approaches designated as pre-seeding, or direct, and post-seeding, or indirect [102]. The former involves the simultaneous printing of materials and cells combined, while the latter consists of initially printing the material and then co-culturing it with the proper cells. The studies found in literature related with graphene-based 3D-printed scaffolds, which will be overviewed in this section, fall under the category of post-seeding bioprinting.

Recently, Belaid and co-workers [103] developed a 3D-printed scaffold of PLA reinforced by the incorporation of GO at different concentrations (0.1, 0.2 and 0.3 wt%). A composite polymer solution of GO and PLA was cast to form a film that was cut into pieces and introduced into a single-screw extruder, at 200°C. A filament with a diameter of 1.75 mm was obtained and used to print the 3D-scaffolds by FDM. To facilitate bone regeneration, the authors generated scaffolds with a porous interconnected network and a pore size around 300 μm, corresponding to an infill of 70%. With the incorporation of GO, the scaffolds revealed a rougher and hydrophilic surface, when compared to control PLA scaffolds. Tensile tests showed that the mechanical properties were improved with the GO incorporation (30% increase of the Young’s modulus with 0.3% GO) - Fig. 7. Composite scaffolds were also biocompatible as the cell viability, attachment, proliferation, and differentiation assays using MG-63 osteosarcoma cells showed. They promoted cell proliferation and mineralization more efficiently than pure PLA scaffolds which potentially allow bone formation [103].

The need for conductive and mechanically robust, biocompatible, and biodegradable materials in tissue engineering (TE) is constantly increasing. The advantages of electrical conductivity and electrical stimulation (ES) have been recognized and explored in biomedical field [104], since cell adhesion and differentiation are affected by a residual permanent charge on the materials surface. Therefore, it is expected that the use of ES after biomaterial implantation would induce osteointegration. Graphene-based materials have been proposed as additives to produce electrically conducting composites demonstrating that the simple presence of the electrical stimuli of the materials contribute to improve cellular functions [104, 105]. However, as mentioned above, there is no consensus about graphene cytotoxicity and some authors still mention some risks [106]. Wang and colleagues printed PCL-graphene scaffolds loaded with 0.13 wt%, 0.50 wt% and 0.78 wt% of graphene and associated a non-invasive microcurrent therapy to treat rat calvaria critical size bone defects [106]. The blended composite material was cut into small pellets for processing. The melt processing temperature was 90°C, 220 μm of slice thickness, 22 rpm of screw velocity, and 20 mm/s of deposition velocity. The fabrication process was performed at room temperature. A 0°/90° lay-down pattern was used to obtain pores with a regular geometry, a constant filament diameter of 330 μm, and a filament distance of 680 μm. Thereafter, MC3T3 pre-osteoblastic cells were seeded on the scaffolds - Fig. 8. These results showed that cell proliferation was stimulated with the increase of graphene concentration and no significant cytotoxicity was found. Printed scaffolds with 0.78 wt% induced an acceptable level of immune response, revealed by the low levels of tumor necrosis factor-alpha (TNFα) and interleukin-1-beta (IL-1β), suggesting high potential for in vivo applications. The incorporation of graphene and the application of electrical stimulation (10 μA) led to an increase in cell migration,

### Table 3 Preparation methods of graphene-based composites

| Preparation method | Polymer matrix | Advantages | Disadvantages | Ref |
|--------------------|----------------|------------|---------------|-----|
| Solution Mixing    | PC, Polymethylmethacrylate (PMMA); Polyvinyl alcohol (PVA); Polystyrene (PS); Polyurethane (PU) | Inexpensive; Homogeneous dispersion. | Need to remove solvents | [3, 88, 90, 94] |
| Melt Mixing        | PU; Polyethylene terephthalate (PET); PLA; PC | Applicable to polar and non-polar polymers; does not use solvents; adequate for both small and large-scale production. | Less effective in dispersion; possible buckling, rolling, or shortening of graphene sheets. | [90, 91, 95] |
| In Situ Polymerization | PA; Polyethylene (PE); Polypyrrole (PPy); PMMA | Applicable to insoluble or thermally unstable polymers. | Need to remove solvents. | [88, 92, 95] |
Fig. 6 Schematic representation of TPU/Graphene Composite Preparation: (a) After oxidation of graphite functionalized layers of graphene can be obtained by (b) Rapid Thermal Expansion or (c) Organic Modification with Isocyanate in dimethylformamide (DMF) (iGO); (d) Graphite can be mixed with TPU via Melt mixing, or (e) Solvent mixing, followed by solvent removal; (f) In opposite, composites can be produced monomers using In situ polymerization. Black lines represent graphitic reinforcements. TPU hard and soft segments are represented by short blue blocks and thin red curves, respectively [96]
| Technique | Matrix | Filler | Properties | Ref. |
|-----------|--------|--------|------------|------|
| SLA       | Photopolymer (Envision TEC PIC 100 resin) | GO | 0.2wt%GO increased 62.2% and 12.8% the tensile strength and ductility, respectively of the printed composite. | [54] |
| DIW       | Polyaniline (PANI) | | Specific capacitance of the printed supercapacitor: 1329 mF cm$^{-2}$. | [56] |
|           | Polydimethylsiloxane (PDMS) | | The printed device exhibited a resistivity of 1660 $\Omega$ cm at a low GO percolation threshold (0.83 vol.%) | [57] |
|           | Epoxy | Graphene | Printed composites with 10 wt.% of graphene presented flexural modulus of 3.2 GPa and bulk resistivity of $1 \times 10^4$ $\Omega$ m, i.e., five orders of magnitude lower than that of the neat epoxy. | [58] |
| SLS       | PA | | Tensile modulus for PA-rGO compared to PA (221 ± 17 vs 149 ± 26 MPa) and similar tensile strength. | [59] |
| FDM       | ABS | | Electrical conductivity increased to $1.78 \times 10^{-7}$ and $1.05 \times 10^{-3}$ S.m$^{-1}$ with the addition of 0.4 and 5.6 wt% of graphene, respectively | [60] |

Fig. 7 Mechanical properties of PLA and PLA/GO scaffolds produced by FDM: a Young’s modulus, b Tensile stress at break, and c Poisson’s ratio. Reproduced with permission from [103]
leading to new tissue formation, well-organized tissue deposition and bone remodeling, as can be seen in Fig. 9 [106].

In another study, Jakus et al. [107] developed 3D-printable graphene inks and used them to produce electrically conductive, mechanically resilient, and biocompatible scaffolds with high graphene content (20 and 60 vol% of solid). The ink contained graphene flakes and polylactide-co-glycolide (PLGA) and was produced by combining a solution of PLGA and dichloromethane with a graphene dispersion, followed by solvent evaporation. The authors used this ink via extrusion-based 3D printing to create mechanically robust graphene scaffolds. That retained electrical conductivity greater than 800 S/m. These composite scaffolds supported cells adhesion/proliferation, as well as neurogenic differentiation, as illustrated in Fig. 10. On day 14, a distinct difference in cell morphology was observed, where human mesenchymal stem cells (hMSC) on 20 vol% graphene scaffold exhibited a sheet-like morphology, characteristic of adherent cell types such as fibroblasts. In contrast, cells on 60 vol% graphene scaffold presented highly elongated morphologies similar to uni- or multipolar neurons - Fig. 10. In vivo experiments of 30 days in mice showed promising biocompatibility with no evidence for the accumulation of graphene flakes in the kidney, liver, or spleen. Besides, the scaffolds present exceptional handling characteristics being able to apply in fine surgical procedures [107].

Sayyar et al. [108] used different percentages of rGO to reinforce a chitosan -lactic acid matrix and produce conductive hydrogels. These composites were easily processed into 3D scaffolds by extrusion-printing. Scaffolds with 30 layers of each graphene/chitosan dispersion were printed from a 200 μm diameter nozzle, at 0°/90° orientation, a feed rate of 150 mm.min⁻¹ and with a strand spacing of 0.6 mm. The addition of graphene led to significant improvements in terms of mechanical strength. An addition of 3 wt.% of graphene caused an increase over 200% in tensile strength. The resulting structures were seeded with fibroblast (L−293) cells that adhered and proliferated on several layers of the 3D scaffold. These scaffolds revealed to be excellent conducting substrates for the growth of electro-responsive cells [108].

Zhou and co-workers [109] combined GO in different concentrations (0, 0.05, 0.1, 0.25, 0.5 and 1 mg/mL) with a gelatin-methacrylate (GelMA) and poly (ethylene glycol) diacrylate (PEGDA) solution to produce a biocompatible photopolymerizable ink and then 3D-printed scaffolds. GelMA-PEGDA inks with and without GO were placed on the z-control movable platform and printed by SLA, using a table-top stereolithography-based printer that applied UV laser. The printing
parameters were: 200 μm diameter laser beam, 25 μJ intensity output of 20 kHz emitted UV, and 10 mm/s printing speed. Human MSCs spread and extended on these scaffolds, after 5 days of culture. The results showed that the optimal combination was 15% GelMA+PEGDA with GO incorporated at 0.1 mg/mL, as seen in Fig. 11. Moreover, the authors showed that these GelMA-PEGDA-GO scaffolds induced chondrogenic differentiation of Human MSCs and promoted the glycosaminoglycan and collagen levels. Collagen II, SOX 9 and Aggrecan gene expressions associated with chondrogenesis were also greatly promoted on the scaffolds. Such scaffolds are excellent candidates for cartilage regenerative applications.

Composite scaffolds of polyether ether ketone (PEEK)/PVA reinforced with GO was developed by Feng and co-workers using a SLS system envisaging bone regeneration [110]. The resulting scaffolds presented good hydrophilicity and degradability. For a GO loading of 1 wt.%, the strength and modulus of PEEK/PVA scaffolds
increased by 97.16% and 147.06%, respectively. In vitro tests with MG63 cells revealed that the scaffolds promoted cell attachment and proliferation as well as osteogenic differentiation and bone regeneration in vivo, with rabbits [110].

In addition to all outstanding mechanical and electrical properties, graphene can be exploited in the medical field because of its antibacterial activity. Angulo-Pineda et al. explored this property and developed SLA composite scaffolds of PCL filled with conductive thermally reduced graphene oxide nanoparticles at concentration above the percolation threshold. By applying a voltage along of their surface, a bactericidal effect was observed. Moreover, under the same regime of ES, the adhesion and viability of human stem cells were further enhanced when compared with pure PCL scaffolds with and without ES [111]. The antibacterial activity was also observed by Zhang and co-workers in 3D printed Ag-GO nanocomposite scaffolds [112] as well as by Cabral et al., that produced tricalcium phosphate/gelatin/chitosan scaffolds reinforced with rGO displaying antimicrobial activity without compromising osteoblasts’ viability and proliferation [113].

Cryogenic 3D printing is a recently developed biofabrication process that allows the production of scaffolds with a predesigned shape, controllable architecture, and

![Fig. 10 a Images and scanning laser confocal 3D reconstruction projections of live stained (green) and dead stained (red) hMSCs on different scaffolds after 1, 7 and 14 days of seeding. b Number of hMSCs present on scaffolds as a function of composition and days after seeding, according to DNA quantification. Dotted line represents initial cell seeding number. c Neurogenic relevant gene expression of cells on scaffolds with 20 and 60 vol% graphene after 7 and 14 days of seeding, normalized to expression of day 0 (unseeded hMSCs). SEM micrographs of hMSCs on (d) 20 and (e) 60 vol% graphene scaffolds 7 days after seeding. f High-magnification SEM micrograph of cells on day 7, 60 vol% graphene scaffolds, revealing hMSC connecting via a small “intercellular” wire. g Scanning laser confocal 3D reconstruction of live (green) and dead (red) cells on day 14 for 60 vol% graphene scaffolds and (h) detail of cell indicated by yellow arrow in (f). For panels b and c, ** indicates significance of p < 0.05 between compared groups (n = 4); *** indicates significant (p < 0.05) difference over previous time point for the same material group. Reproduced with permission from [107].]
suitable mechanical strength at the relatively low temperature of −32 °C. It allows the incorporation of a large quantity of biomolecules/drugs into scaffolds and the retention of a high level of the biomolecule’s biological activity. It does not require UV light and post-sintering use of a high-power laser, avoiding some typical 3D printing’s disadvantages. Despite being barely explored, the in situ incorporation of GO-loaded drugs/biomolecules into scaffolds by this method shows great potential in TE [114]. A recent and novel bioactive PLGA/β-tricalcium phosphate (β-TCP) composite scaffold, in which GO and a bone morphogenetic protein were loaded in situ, was produced by Zhang et al. using the cryogenic 3D printing method [115]. The composites

![Image](image_url)

**Fig. 11** a Evolution along 5 days of MSCs proliferation on hydrogels with different compositions. b MSCs proliferation on scaffolds with GelMA-PEGDA and different concentrations of GO for 5 days. The photographs represent the corresponding scaffolds. Reproduced with permission from [109].

**Fig. 12** Bone volume and bone surface results obtained by Micro-CT for the in vivo animal tests: a Implantation process; b Micro-CT image of the scaffold; c 3D reconstructed images of the control, PT, PT/P, and PTG/P scaffolds 4 weeks after the implantation (top row) and 12 weeks after the implantation (bottom row); d Bone volume and e bone surface of the newly formed bone in the critical-size defects (p < 0.05). Reproduced with permission from [115].
were produced at various GO concentrations (0, 0.025, 0.05, and 0.1 wt.% with respect to the final mixture) and were designated as ‘PT/P’, ‘0.025PTG/P’, ‘0.05PTG/P’, and ‘0.1PTG/P’, respectively. The PLGA/β-TCP scaffold without peptide was generated as the negative control and denoted as ‘PT’.

Critical-sized calvarial defects were successfully created in rats (Fig. 12). Scaffolds with a diameter of 5 mm were implanted in these animal models to evaluate in vivo bone regeneration. When the implantation time was extended to 12 weeks, more bone formation was observed in the groups with scaffolds than in the control group, particularly in the graphene-based scaffolds. The bone volume (BV) and bone surface (BS) were quantitatively analyzed to confirm the bone regeneration ability - Fig. 12. The scaffolds presented a customized shape with hierarchical porosity and were mechanically comparable to human cancellous bone. GO improved the scaffold’s wettability and mechanical strength as well as tuned the peptide release, promoting bone marrow – derived mesenchymal stem cells ingrowth into the scaffold, enhancing the osteogenic differentiation in a critical bone defect [115].

In summary, several graphene-based composites have been processed by 3D printing to manufacture scaffolds with enhanced biological, mechanical and electrical performance, to regenerate a variety of tissues such as bone, cartilage, or the nervous system. A summary of relevant examples of such scaffolds, as well as their main processing details and properties, is presented in Table 5.

**Biosensors**

The incorporation of conductive materials such as graphene derivatives into polymers to form conductive composites allows the printing of electrodes and circuits, which can be integrated into complex structures, permitting the electrochemical detection of organic compounds and biologically active molecules. Graphene-based composites can be combined with biological receptors such as enzymes, antibodies, and single-stranded DNA. Printed graphene-based biosensors have good reliability and great potential for numerous applications [124].

Palenzuela et al. [125] fabricated ring-shaped electrodes by FDM using graphene/PLA composite filaments, for the electrochemical sensing of picric acid and ascorbic acid. The activation of the graphene-based 3D-printed electrodes consisted of DMF-assisted partial dissolution of the insulating PLA. Different activation times (1, 10, 20, and 60 min) were studied, but immersing the 3D printed electrodes in DMF for 10 min was the optimum time. Picric acid and ascorbic acid could be detected with these electrodes in a wide range of concentrations, from 5 to 350 ppm for the former and 10 to 500 μM for the latter [125].

Cardoso et al. [126] used FDM to produce biosensing platforms from graphene-PLA filaments. The enzymatic biosensor was fabricated on the PLA-graphene surface and used for glucose sensing in blood plasma. The biosensor presented a limit of detection (LOD) of 15 μM L⁻¹, inter-day, and intra-day precision lower than 5%, and adequate recovery values (90–105%) for the analysis of plasma. At the same time, after a surface treatment of the 3D-printed sensor, it also exhibited improved electrochemical properties for the direct detection of nitrite and uric acid [126].

Marzo and co-workers [127] developed 3D-printed enzymatic graphene-PLA electrodes to immobilize horseradish peroxidase (HRP) creating a direct electron transfer biosensor for hydrogen peroxide (H₂O₂) detection. 3D electrodes were first printed by FDM using graphene/PLA filaments and then exposed to a chemical and electrochemical treatment for activation. Electrodes modified with gold nanoparticles enhanced the direct electron transfer between the HRP and the biosensors. Figure 13a depicts the good selectivity to H₂O₂ of DMF-EC/ AuNPs/HRP. The response of these biosensors with and without gold nanoparticles (DMF-EC/HRP and DMF-EC/ AuNPs/HRP, respectively) in human serum was also evaluated (Fig. 13b) as well as their long-term stability up to 7 days, in the presence of 25 (Fig. 13c) and 50 μM of H₂O₂ (Fig. 13d). This work opens possibilities for 3D-printed enzymatic systems to detect other biomarkers without the use of electron mediators and binder polymers [127].

Leigh et al. [128] used a PCL/carbon black (CB) composite filament to manufacture electronic sensors capable of detecting mechanical flexing and capacitance changes, using a low-cost 3D printer. The filler ratio was defined considering the percolation threshold and the melt viscosity of the composite, the final chosen loading of CB was 15 wt%. Through resistivity tests, the authors verified the existence of a piezoresistive behaviour, enabling its use in the field of biomechanics. The sensors also have a capacitive behaviour when printed as part of an interface device or embedded inside a “smart” vessel, making it possible to detect the presence and quantity of a liquid inside [128].

Lee and co-workers [129] produced PCL/graphene (10 wt%) electrodes using a KIMM SPS1000 bioplotter extrusion printing system. These electrodes presented robustness, flexibility, biodegradability, and conductivity required for the electrochemical studies. Before the electrochemical experiment, the electrode was immersed in DMF solvent for 10 min to remove the PCL that blocked the graphene layer, allowing an ideal contact between the graphene and the electrolytes. Biocompatibility was demonstrated by the electrochemical response derived from the diatom
| AM technique | Scaffold | Printing parameters | Cellular behavior | Mechanical properties | Ref |
|--------------|----------|---------------------|------------------|-----------------------|-----|
| Extrusion-based 3D-printing | PCL + Graphene (0.13 and 0.78 wt.%); Coating with P1-latex protein | Pattern: 0/90 lay down; 90°C; slice thickness: 220 μm; 22 rpm; speed: 20 mm/s | Human adipose-derived stem cells (ADSCs) adhered/spread and presented a spindle-like morphology; osteogenic differentiation; Scaffolds (0.78 wt.%): higher viability/spread | Compressive strength (C_s): 80 MPa; C_s (0.78 wt.%): 130 MPa | [116] |
| | PCL + Graphene (0.5 and 0.78 wt.%); Coating with P1-latex protein | Pattern: 0/90 lay down; 90°C; slice thickness: 220 μm; 22 rpm; speed: 20 mm/s | ADSCs adhered/spread and presented a spindle-like morphology; osteogenic differentiation; Scaffolds (0.78 wt.%): higher viability/spread | No mechanical results for scaffolds | [117] |
| | PCL + Modified graphene nanoplatelets (0.5 wt.%); Coating with P1-latex protein | Pattern: 3 layers arranged in at 90°; Layers height: 0.15 mm; 190°C; Speed: 15 mm/s | Adhesion and proliferation of human chondrocytes; Higher cell proliferation in 3D scaffold comparing to polystyrene positive control; Graphene did not increase the cellular toxicity | | [118] |
| | PCL + GO (0.1 and 0.5 wt.%); Coating with P1-latex protein | Cuboidal shape 22 layers; Layer thickness: 2.2 mm; 100°C; Speed: 1 mm/s | Murine preosteoblast cells with higher proliferation and osteogenic differentiation in scaffolds with 0.5 wt.% GO | | [119] |
| | Poly (trimethylene carbonate) (PTMC) + Graphene sheets from rGO (3 wt.%); Coating with P1-latex protein | 7 layers; 1 x 1 cm 60–150°C; Nitrogen pressure: 100–200 kPa; cross-linking by UV irradiation for 10–15 min | MSCs with good attachment and viability; Addition of graphene did not alter cell number; Electrical stimulation did not compromise MSCs and the osteogenic markers were upregulated | | [104] |
| | Chondroitin sulfate/ALG/gelatin+GO (1 mg ml⁻¹); Coating with P1-latex protein | 30 x 30 x 1 mm³ mesh-like inner pattern: 1.5 mm of thread spacing; Speed: 50 mm s⁻¹; Pressure: 1 bar; extrusion needle tip: 25 G; Petri dish at 2°C; UV: 9 mW cm⁻² for 5 min | hMSCs cells adhered and spread; Composites presented cells with great proliferation, alignment and distribution; Chondrogenic differentiation | | [121] |
| SLS | PVA+ GO (2.5 wt.%); Coating with P1-latex protein | Laser power: 5 W scan speed: 400 mm.min⁻¹ spot diameter: 1.6 mm scan spacing/layer thickness: 2.7/0.1–0.2 mm, respectively | Human osteoblast-like MG-63 adhered and spread; The addition of GO to PVA led to higher cell growth and proliferation comparing to pure PVA scaffold. | | [123] |

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microalgae grown on the PCL/graphene substrate. The produced electrode offers a great potential in electrical stimulation to promote tissue formation, as well as in bioelectronic applications [129].

Conclusions

The manufacture of 3D printed devices is an emerging multidisciplinary field, namely in the biomedical area. In this review, the whole 3D printing process was presented, from the preparation of composites to the most used techniques for printing graphene-based polymer composites for biomedical applications. The properties of the reported graphene-based composite structures have shown significant improvement over polymer matrices without reinforcement, and enhancement of cell response.

3D printing demonstrated the versatility to build complex shapes and custom designed parts, revolutionizing the development of complex structures with target properties, as compared to conventional manufacturing methods. The processing of graphene-based polymer composites with the recent AM technologies has enormous potential for several biomedical applications, namely to engineer distinct tissues such as bone, cartilage, or the nervous system, as demonstrated by the examples discussed in section 3, and also other tissues such as tendon, ligament, spinal cord or cardiac tissue, which could potentially benefit from the electrical properties conferred by graphene. The development and integration of biosensors is also a growing application of such 3D printed graphene-based nanocomposites, with promising results, as shown in this review.

Abbreviations

AM: Additive manufacturing; CAD: Computer-aided design; STL: Standard Tessellation Language; FDM: Fused deposition modeling; DW: Direct ink writing; SLA: Stereolithography; SLS: Selective laser sintering; FFE: Free form extrusion; UV: Ultraviolet radiation; PLA: Polylactic acid; ABS: Acrylonitrile butadiene styrene; PC: Polycarbonate; ALG: Alginate; CHI: Chitosan; DNA: Deoxyribonucleic acid; PCL: Polycaprolactone; PA: Polylamide; GICs: Graphite intercalation compounds; GO: Graphene oxide; rGO: Reduced graphene oxide; EG: Expanded graphite; GNP: Graphene nanoplatelets; BMSCs: Bone mesenchymal stem cells; PMMA: Polymethylmethacrylate; PVA: Polyvinyl alcohol; PS: Polystyrene; PU: Polyurethane; PET: Polyethylene terephthalate; PE: Polyethylene; PPy: Polypyrrole; D; Dimethylformamide; PANI: Polyaniline; PDMs: Polydimethylsiloxane; TE: Tissue engineering; ES: Electrical stimulation; TNF-α: Tumor necrosis factor alpha; IL-1β: Interleukin-1beta; NBR: Natural bone regeneration; PLGA: Polylactide-co-glycolide; hMSCs: Human mesenchymal stem cells; GelMA: Gelatin-methacrylate; PEGDA: Poly (ethylene glycol) diacrylate; PEEK: Polyetherether ketone; B-TCP: β-tricalcium phosphate; BV: Bone volume; BS: Bone surface; ADSCs: Adipose-derived stem cells; Cs: Compressive strength; PTMC: Poly (trimethylene carbonate); TEGDA: Triethylene glycol dimethacrylate; LOD: Limit of detection; HRP: Horseradish peroxidase; DMF-EC/AuNPs/HRP: 3D electrode with chemical/electrochemical treatment and gold nanoparticles; DMF-EC/HRP: 3D electrode with chemical/electrochemical treatment without gold nanoparticles; UA: Uric acid; AA: Ascorbic acid; D: Dopamine; CB: Carbon black

Fig. 13 a Selectivity assay of DMF-EC/AuNPs/HRP for H$_2$O$_2$ analysis in the presence of 50 μM of uric acid (UA), ascorbic acid (AA), and dopamine (D). b Accuracy assay of DMF-EC/AuNPs/HRP and DMF-EC/HRP for H$_2$O$_2$ detection. Long-term stability response of 3D DMF-EC/HRP and 3D DMF-EC/AuNPs/HRP biosensors for 25 (c) and 50 (d) μM H$_2$O$_2$ detection. Reproduced with permission from [127].
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Authors’ contributions

MS and ISP – literature search, manuscript drafting, figure preparation; JAC, MCP and NMA – conceptualization and manuscript revision. All authors read and approved the final manuscript.

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Availability of data and materials

Data sharing is not applicable to this article as no datasets were generated or analysed during the current study.

Declarations

Competing interests

The authors declare that they have no competing interests.

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