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# REVIEW

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### 1. Introduction

The nanotechnology field has been expanding with nanomaterials' development regarding their different structural, morphological, and chemical characteristics, guaranteeing enormous versatile applications. One of these features includes the ability to link molecules of different natures and functionalities that can be used for diverse applications for various products: pharmaceutical nanoformulations, ointments/creams for therapeutic or aesthetic purposes, curative films, active packaging, food additives, food packaging, and others.<sup>1–3</sup> Considering these cutting-edge nanotechnology applications, food packaging is of great interest since several physical and chemical properties can be tuned through the formulation of nanocomposites.<sup>1–3</sup> The application of nanomaterials in the food industry – mainly in the packaging market – increased its market value from \$20.4 billion in 2006 to more than

# Nanocomposites based on the graphene family for food packaging: historical perspective, preparation methods, and properties

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Nanotechnology experienced a great technological advance after the discovery of the graphene family (graphene – Gr, graphene oxide – GO, and reduced graphene oxide-rGO). Based on the excellent properties of these materials, it is possible to develop novel polymeric nanocomposites for several applications in our daily routine. One of the most prominent applications is for food packaging, offering nanocomposites with improved thermal, mechanical, anti-microbial, and barrier properties against gas and water vapor. This paper reviewed food packaging from its inception to the present day, with the development of more resistant and intelligent packaging. Herein, the most common combinations of polymeric matrices (derived from non-renewable and renewable sources) with Gr, GO, and rGO and their typical preparation methods are presented. Besides, the interactions present in these nanocomposites will be discussed in detail, and their final properties will be thoroughly analyzed as a function of the preparation technique and graphene family-matrix combinations.

\$3 trillion in 2020.<sup>4</sup> Moreover, nanotechnology contributes significantly to reducing food lost amount and thus offers an alternative for the increasing food demand as the worldwide population increases. Food packaging acts as a physical barrier against the interaction of food and the environment and thus preventing its contamination with microorganisms, dust, or even spoiling as a consequence of moisture, light, and shock. Considering that, several strategies for food packaging have been developed over the centuries, most notably in the last century, in which materials chemistry exerted a pivotal role in the search for smart packaging. The combination with conventional polymers used for packaging with other materials offered a wide range of new materials (composites) with improved properties.<sup>4</sup>

In this way, graphene (Gr), graphene oxide (GO), and reduced graphene oxide (rGO)5-23 are examples of these nanomaterials which are used in composites formulations with polymers, such as polypropylene (PP),<sup>24</sup> polystyrene (PS),<sup>25</sup> polyethylene terephthalate (PET),26 polyvinyl alcohol (PVA),27 poly lactic acid (PLA),<sup>28</sup> polyhydroxybutanoate (PHB),<sup>29</sup> chitosan (CS),<sup>30</sup> starch (ST),<sup>31</sup> cellulose<sup>32</sup> and others polymeric matrix.<sup>33–36</sup> The interaction between Gr or GO with these polymers occurs through chemical bonds or molecular interactions and thus improves chemical,37 mechanical, their thermal, optical, antimicrobiological, and barrier properties.16,19,38-45

In light of the considerations mentioned above, this review will focus on the recent developments of the graphene family (graphene, graphene oxide, and reduced graphene oxide)-based

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#### Review

nanocomposites used as food packaging systems. We will provide an overview of the historical use of food packaging materials and present the most common preparation methods of graphene-based nanocomposites – with several fossils and renewable-derived polymeric matrices – their physicochemical interactions, final properties as a food packaging material, and most common applications found in the market.

## 2. Food packaging

#### 2.1. Historical perspective

The first types of food packaging arose between prehistory and ancient history. People stored and protected their food in plant leaves, clay artifacts, seashells, baskets, gourds, skin bags, and animal organs (stomach and bladder), coats, hooves, and horns.<sup>46</sup> Years later, in A.D. 751, paper became known and made throughout Europe, but its use for food packaging became famous many years later. The development of food and beverage storage techniques was essential to transition from nomadic activities to more complex societies.<sup>46</sup>

The discovery of novel technologies applied to food packaging is of great importance for preserving and maintaining the shelf life of foods in general. Many techniques have been employed to manufacture packaging and have continued to change over the centuries. The correct selection of the type of packaging for each food is essential for maintaining the integrity and quality of the food, as they will keep the food protected, fresh and healthy since they leave the industries until they reach the consumer's table. Several types of material have been used

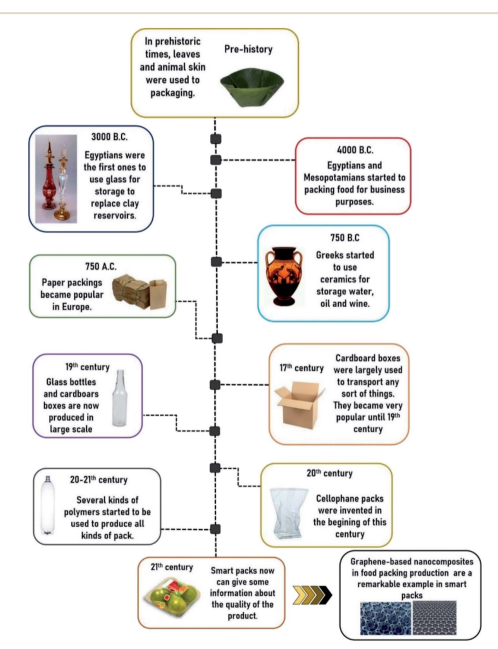


Fig. 1 Historical perspective of food packaging

to pack food, such as glass, aluminum, tin-free steel, paper, cardboard, and rigid and flexible plastics.<sup>47</sup>

The use of glass for food storage dates back to 3000 B.C. The glass containers have characteristics to be impermeable to gases and vapors, keeping food and drink fresh, and offer the possibility of storage for long periods without interfering with the food taste. The main disadvantage of using glass containers is their difficulty transporting because they are fragile and bulky. The glass package may break if there are impacts, thermal shocks or if the internal pressure increases.<sup>47,48</sup>

Metal packaging is generally produced from aluminum, tinfree steel, and tinplate. These materials are versatile and have good protection and barrier properties, and also possess a high decorative potential and more resistance, being lighter and more malleable than glass packaging.<sup>47–49</sup>

In that sense, aluminum packaging is resistant to corrosion because it is naturally coated with aluminum oxide, creating an effective barrier to gases and vapors, chemical attacks, variations in humidity and temperature, odors, light, and microorganisms.<sup>47–49</sup> Steel packages without tin or chrome steel coated with oxide or electrolytic chromium require the coating with organic material to be completely resistant to corrosion.<sup>47–49</sup>

Paper-based packaging is also important for the food industry and is widely applied. Paper and cardboard use as food packaging began in the 17th century and expanded in the late 19th century.<sup>47,50</sup> Kraft paper is the most resistant paper packaging used to pack vegetables, dried fruits, sugar, and flour. Also, glassine is a paper that undergoes a more extreme hydration process than the previously mentioned and produces a denser, smoother, and shiny paper used to pack fried foods, baked goods, and fast foods. Currently, laminated papers are coated or not with cellulose and kraft sulfite. They can also be laminated with plastic or aluminum to improve their gas and moisture barrier properties, substantially increasing the cost of paper.<sup>47,48,50</sup>

The rise of modern food packaging began in the 19th century after the industrial revolution, with the studies of can-food by

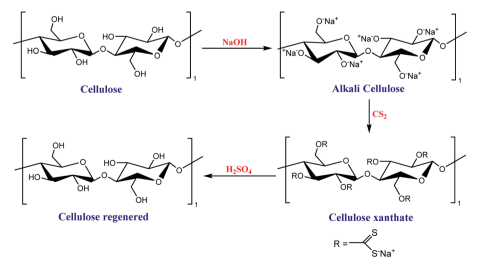
Appert, based on the food microbiology of Pasteur, Prescott, and Underwood.<sup>51</sup> In this way, Fig. 1 shows the types of food packaging used by humanity over the centuries.

In the early 20th century, some food packaging appeared from unexpected sources due to unsuccessful attempts, such as the transparent tablecloths by Jacques Brandenberger, a Swiss textile engineer. However, cellophane on regenerated cellulose kind was created through this unsuccessful attempt.<sup>51</sup>

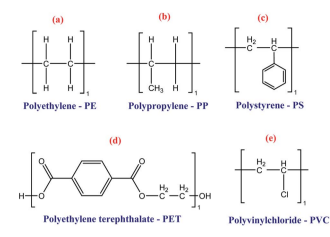
For cellophane production, cellulose is first dissolved in sodium hydroxide solution. It is treated with carbon sulfide producing viscose (cellulose xanthate), which is extruded in a sulfuric acid and sodium sulfate solution to convert the viscose into regenerate cellulose. The regenerated cellulose (cellophane) goes through several baths: sulfur removal, bleaching, and glycerin to avoid fragility,<sup>52,53</sup> as shown in Scheme 1. Cellophane is transparent, with a high permeability barrier to oils, greases, and air; it is used to pack sweets, generally.<sup>54</sup>

During the first (1914–1918) and the second (1939–1945) World wars, there was a lot of innovation in food packagings, such as oil products and wax for dry cereals and cookies. In this way, plastics like polyethylene (PE) and polyvinylidene chloride (PVDC) emerged as aseptic packaging and flexible metal cans for bottling beer. The development of polymers such as polyethylene (PE), polyethylene terephthalate (PET), polypropylene (PP), and polyvinyl alcohol allowed a partial replacement of glass, metal, and rigid plastic packaging by these new flexible plastic packaging.<sup>48,50,51,55</sup> Since the end of the 20th century, many polymers produced from microorganisms, plants, and animals have been studied. Furthermore, in the 20th century, intelligent packaging capable of controlling oxygen diffusion, being a breath mediator, scent controller, and antimicrobial was developed.<sup>48,50,51,54–56</sup>

In 2007 Kalaitizidou and co-workers<sup>24</sup> were the first group of researchers to create a polymer of polypropylene and graphene. They observed that the addition of graphene to the polymer matrix improved the physical, thermal, and mechanical



Scheme 1 Cellophane production route.



**Fig. 2** Structures of: (a) polyethylene – PE, (b) polypropylene – PP, (c) polystyrene – PS, (d) polyethylene terephthalate – PET and (e) poly-vinyl chloride – PVC.

properties and that the polymeric nanocomposite could be used to produce food packaging. Item 3.4 presents more details about graphene-based nanocomposites applied to food packaging.

Considering the pivotal role that polymers-based packaging possesses in our modern society, the following sections will cover the non-renewable and renewable polymers most used in food packaging.

# 2.2. Non-renewable-based polymers applied to food packaging

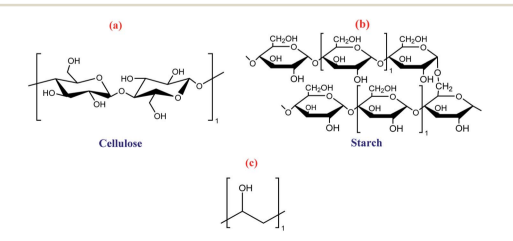
Since the 20th century, the main polymers used in food packaging have been polyethylene, polyethylene terephthalate, polystyrene, polypropylene, and polyvinyl because of their high applicability and availability low molecular weight, and low cost.<sup>51</sup> Most of these materials have some drawbacks to be used as food packaging. In most cases, they are not biodegradable and take several years to completely degrade by nature.<sup>33,48,50</sup> **2.2.1. Polyethylene.** Polyethylene (PE) is one of the most used polyolefins-based polymers in the manufacture of food packaging.<sup>47,56</sup> PE, Fig. 2(a), is produced from the polymerization reaction of ethylene monomers. Two polyethylene types called low and high density can be made, LDPE and HDPE, respectively. LDPE is widely used in food packaging due to its easy processing and extensive ability to coat various substrates; it has light color, lightness, flexibility, and low cost.<sup>57</sup> Moreover, LDPE has low permeability to water vapor, although it does not present barriers to protect gases, volatile compounds, and other gases. It is used to produce grocery bags, food packaging for frozen food, papers and laminates coating, bread bags, *etc.*<sup>47,50,57</sup> On the other hand, HDPE has high resistance, a higher gas permeability barrier, and thermal resistance.<sup>47,50</sup>

**2.2.2. Polypropylene.** Polypropylene (PP) is also one of the most used olefin-based in food packaging manufactured through the polymerization reaction of propylene monomers Fig. 2(b). PP has a strong barrier to protecting against humidity, water vapor, and gases and is resistant to chemicals. In this sense, it is resistant to impacts and ruptures, making it suitable food packaging to be used even in a microwave oven.<sup>47,50,57</sup>

**2.2.3. Polystyrene.** Polystyrene (PS) is formed by the polymerization reaction between the styrene monomers, Fig. 2(c).<sup>47,58</sup> PS is used to manufacture cups, cutlery, disposable plates, food trays, and bottles. The expanded PS is very light, rigid, and opaque, thus acquiring excellent barrier and impact properties, making it suitable to protect fragile products.<sup>50,53,60</sup>

**2.2.4. Polyethylene terephthalate.** Polyethylene terephthalate (PET) is a polyester formed from the polymerization of terephthalic acid (TA) and ethylene glycol (EG), Fig. 2(d). PET is thermally resistant to temperature and impact, also owing good barrier of protection against  $O_2$  and  $CO_2$ .<sup>58</sup> PET is a light, transparent material with an amorphous structure and, therefore, widely used in food packaging and soft drink bottles since it retains  $CO_2$ .<sup>47,50,57,59</sup>

2.2.5. Polyvinyl chloride. The polymerization among vinyl chloride monomers produces polyvinyl chloride (PVC),



Poly(Vinyl alcohol) - PVA

Fig. 3 Structures of: (a) cellulose, (b) starch and (c) poly(vinyl alcohol) - PVA.

Fig. 2(e),  $^{60}$  generating an amorphous, transparent, ductile, striated, dense, and brittle material that can be heat sealed.  $^{47,50,57}$ 

#### 2.3. Renewable-based polymers applied to food packages

Several polymeric materials from renewable feedstocks are already accessible, representing an evolution towards a more sustainable society, allowing at least a partial replacement of fossil-based polymers. These polymers can be classified into four groups: polymers from vegetal sources (cellulose and starch); polymers from animal sources (chitin and chitosan); polymers from microbiological sources (polyhydroxyalkanoates – PHAs); and polymers chemically synthesized from agricultural raw materials (poly(vinyl alcohol – PVA; polylactic acid – PLA; biopolyethylene – bioPE; biopolyethylene terephthalate – bioPET).<sup>54,56</sup>

**2.3.1.** Cellulose. Cellulose is one of the most abundant fractions present in lignocellulosic biomass and can be used as starting material for several food packaging purposes. The chemical structure of cellulose, Fig. 3(a), is responsible for its remarkable properties. An equatorial position of the hydroxyls of its structure allows H bonds to occur – allowing a high degree of organization and crystallinity.<sup>34,35,54,56,61</sup>

Cellulose is highly hydrophilic, insoluble in most solvents, and non-thermoplastic, *i.e.*, it does not melt and cannot be moldable at a specific temperature. Besides, several functional groups, such as carbonyl and carboxyl, can be intercalated in its structure.<sup>34,35,54,56,61</sup> The changes made in the cellulose structure usually involve the intrinsic hydroxyl groups through esterification and etherification reactions, thus changing their physicochemical properties.

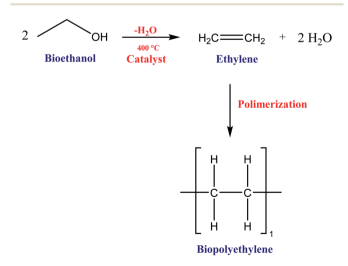
Superhydrophobic cellulose-based materials are on the micrometric scale and have low surface energy. These materials are in evidence because they are abundant on Earth. In addition to being biodegradable, non-toxic, and renewable, they have chemical, physical and mechanical properties superior to nonrenewable materials. In other words, they are a more sustainable and environmentally friendly alternative to polymers based on fossil fuels. Traditional hydrophobicity treatments do not insert the functionalities found in superhydrophobic cellulose materials. One of the methods used to manufacture cellulosebased superhydrophobic materials is wet chemistry, which produces more durable coatings (important for materials with abrasion resistance and wash cycles). The other method is the dry method, which has the main advantage of simplicity by using one-step processes without using organic solvents. Its promising features have potential application in self-cleaning, self-curing, oil and water separation, shielding against electromagnetic interference, etc. The perspective is that the subsequent studies aim to seek these materials for more practical and suitable applications in different areas.<sup>62,63</sup>

The most known cellulose-derived materials are carboxymethyl cellulose, hydroxyethyl cellulose, hydroxypropyl cellulose, ethylcellulose, and cellulose acetate. Their uses are being investigated for numerous purposes in food packaging, such as fruit, vegetables, fast food, bread, and others.<sup>54,56,64</sup>

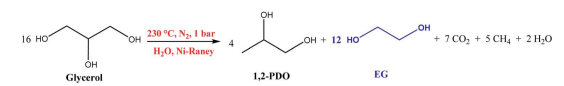
2.3.2. Starch. Starch (Fig. 3(b)) is a compound used by vegetables to store their energy, being present in plants' roots, seeds, and tubers. It is produced from wheat, rice, potatoes, cassava, and others. This polymer is comprised of two types of polysaccharides: amylose and amylopectin. The former is a linear polymer with  $\alpha$ -1-4 bonds among D-glucose monomers, and the latter is formed by branched α-1-6 linkages.<sup>54</sup> Moreover, it is a promising alternative raw material in the manufacture of food packaging due to its high availability, low cost, and sustainability, which can be added to the formulation of fossil polymers, increasing the biodegradability of the copolymer formed.<sup>64</sup> The properties of starch polymers can be varied and often unstable in the presence of water, as it is a hydrophobic material; also, microbial action might create pores through their structure and accelerate its disintegration, conferring to them a biodegradable feature. Furthermore, the expansion of starch carried out by steam consists of a promising alternative material and finds applications in the manufacture of cups, plates, and food packaging as green alternatives to replace polystyrene, a non-renewable material.34,35

**2.3.3. Biopolyethylene.** As afore mentioned, PE is mainly produced from petrochemicals, *i.e.*, non-renewable feedstock. However, biopolyethylene (bioPE) can also be made through the dehydration reaction of bioethanol that produces ethylene, as shown in Scheme 2. The ethylene produced by bioethanol dehydration is polymerized to produce biopolyethylene,<sup>65</sup> which has the same characteristics and applications for food packaging as polyethylene from fossil sources, as discussed in Section 2.2.1.<sup>54,56,65</sup>

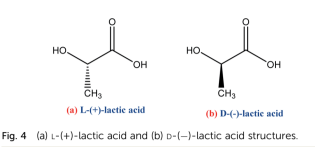
**2.3.4. Biopolyethylene terephthalate**. Biopolyethylene terephthalate (bio-PET) can be produced partially or totally from renewable sources. Considering a partially renewable protocol, one non-renewable precursor, TA or EG, is derived from renewable sources. In a fully renewable approach, both molecules are produced from renewable sources. In this case, TA is usually made by the oxidation of *p*-xylene – from the fermentation of lignocellulosic materials – and then condensed with



Scheme 2 Biopolyethylene production by bioethanol dehydration route.



Scheme 3 General route glycerol APR-hydrogenolysis reactions to produce ethylene glycol in a single process.



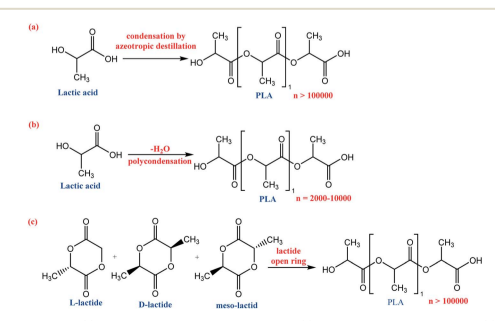
EG to form monomers.<sup>54,61</sup> Furthermore, EG can be produced through the hydrogenolysis reaction of glycerol<sup>66,67</sup> and by reforming the aqueous phase of glycerol (APR),<sup>68,69</sup> as shown in Scheme 3. In this way, it is worth mentioning that bio-PET possesses the same properties and applications for food packaging purposes as those mentioned in non-renewable PET Section 2.2.4.<sup>54,61</sup>

**2.3.5. Poly(vinyl alcohol).** Poly(vinyl alcohol) (PVA), Fig. 3(c), is produced by the polymerization of vinyl acetate and then hydrolyzed to PVA. The –OH groups in the PVA have intermolecular forces such as hydrogen bonds, leading to a high melting point,  $T_{\rm m} = 200$  °C, in which its structure is generally organized in a zigzag and presents a semi-crystalline feature. PVA is widely used due to its biocompatibility, biodegradability, water solubility, and low toxicity. Despite its hydrophilicity, PVA

has good gas barrier properties and tear resistance. Also, as a result of this feature, it is essential in its manufacture to add crosslinking agents to block its hydrophobicity and hence be able to apply in food packaging. In this way, PVA is generally used as a packaging film for fruits and vegetables.<sup>33,58,61</sup>

**2.3.6.** Poly(lactic acid). Poly(lactic acid) (PLA) is a polymer belonging to the group of aliphatic polyesters and synthesized from lactic acid monomers, obtained from biological sources mediated by biocatalysis and chemocatalysis (homo- and heterogeneous). In this context, lactic acid (LA) production is mainly prepared using biocatalysis, which occurs through the fermentation of sugar (sucrose, glucose, fructose, and others) that produces two stereoisomers of lactic acid, L(+), and D(-), Fig. 4.<sup>35,54,59,61</sup> Chemocatalysis can also furnish LA through catalysis to convert sugars derived from non-edible lignocellulosic biomass and glycerol. In the fermentation process, a homo or heterofermentative bacteria is used as an enzymatic catalyst.<sup>35,54,59,61</sup>

PLA can be synthesized by different polymerization routes,<sup>70</sup> Scheme 4, (a) LA condensation by azeotropic distillation; (b) LA polycondensation and dehydration; (c) lactide ring-opening polymerization. The main properties of PLA are good protection barrier, high mechanical and chemical resistance, low toxicity, transparency, and insolubility in water. However, it presents lower thermal stability compared to PET. Besides, the PLA barrier property to gases such as O<sub>2</sub> and CO<sub>2</sub> is lower than



Scheme 4 PLA production routes: (a) LA condensation by azeotropic distillation; (a) LA polycondensation, dehydration; (c) lactide ring-opening polymerization.

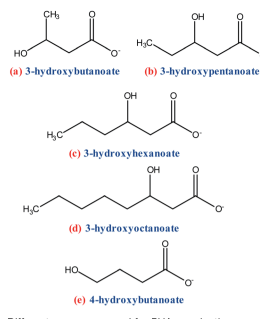


Fig. 5 Different monomers used for PHAs production.

PS and approximately equal to PET polymers. PLA mechanical properties are superior to those observed in PP, HDPE, and PS due to their chemical structure and hardness, and  $T_{\rm m}$  between 173–178 °C. The commercial PLA consists of a copolymer formed between the stereoisomers D(-) and L(+) of the lactic acid, as shown in Scheme 4c. Considering its applications, PLA is used to manufacture food packaging such as meat and soft drinks, among other purposes. Furthermore, packages composed of PLA/cellulose and PLA/natural fiber copolymers gain strength in several packaging and material sectors as they are biodegradable.<sup>35,54,59,61</sup>

**2.3.7. Poly(hydroxyalkanoates).** Poly(hydroxyalkanoates) (PHAs) are ester polymers with optically active carbon atoms. PHAs are thermoplastics and elastomers possessing a melting point ranging from 40 to 180 °C according to the monomer (substrate or reagent) used in the polymerization reaction. These polymers are usually produced by bacterial fermentation of large amounts of carbon, such as sugars and lipids. These bacteria can also consume the polymer when there are few sources of carbon to feed them.<sup>35,54,59,61</sup>

Fig. 5 shows the different monomers used in PHAs production.<sup>71</sup> The polymers shown in Fig. 5 also be chemically synthesized with biodegradability similar to those produced by a bacterial pathway. The most common PHAs is polyhydroxybutanoate (PHB), the polymerization product of 3-

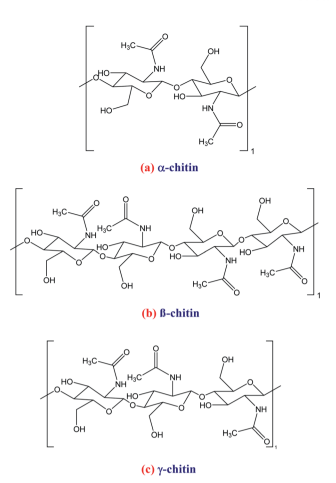
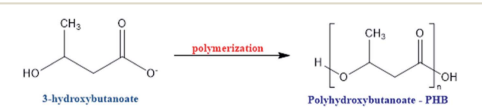
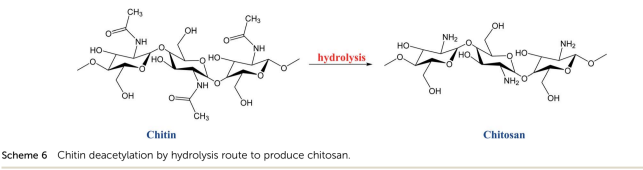


Fig. 6 Different chitin molecular structures, (a)  $\alpha-$  (b)  $\beta-$  and (c)  $\gamma-$  chitin.

hydroxybutanoate monomers,<sup>35,61,71,72</sup> Scheme 5. PHB has an orthorhombic crystalline structure, with a flat amorphous phase between the lamellae; it is a thermoplastic with high crystallinity (80%) and possesses higher rigidity and fragility compared to PP. It has good gas barrier properties, high hydrophobicity, resistance to organic solvents, and is highly biodegradable. The disadvantages of industrial PHB processing are low deformability and thermal stability, the difficulty of being processed industrially, and being expensive. However, some studies have been reported to modify PHAs, such as the use of their combination with plasticizers, acrylic acid, epoxidized soybean oil, and cellulose acetate to improve properties and facilitate the processing for future applications in food packaging, paper coating, and other purposes.<sup>35,54,59,61,64,71,73,74</sup>



Scheme 5 Different monomers used for PHB production polymerization of 3-hydroxybutanoate monomers.



2.3.8. Chitin and chitosan. Chitin (CT) or *N*-acetyl-D-glucosamine is a polysaccharide found in three different structures  $\alpha$ ,  $\beta$  and  $\gamma$ ,<sup>54,73,74</sup> Fig. 6.

The  $\alpha$ -chitin structure has higher physical and chemical resistance because it has an antiparallel arrangement between its oligomers, causing higher strength in the hydrogen bonds of its structure.  $\beta$ -Chitin is characterized by the parallel arrangement of chitin oligomers for the formation of its structure. The *N*-acetyl groups are structural spacers, facilitating the hydration of this macromolecule. The  $\gamma$ -chitin structure is formed by two parallel and one antiparallel chitin sheets, and the *N*-acetyl groups play the same role as in  $\beta$ -chitin. The strength and resistance of  $\beta$ - and  $\gamma$ -chitin structures have also been reported and were found in fungi.<sup>54,73,74</sup>

Nature produces billions of tons of crystalline chitin present as the exoskeleton of some insects, arthropods, and mollusks, and in the cell walls of fungi, yeasts, and plants. Chitin is highly hydrophobic, non-toxic, biodegradable, and easily compacted. It is generally soluble only in mixtures of hexafluoroacetone, chloroalcohols, and hexafluoroalcohols with aqueous solutions of strong inorganic acids. Due to its poor chemical reactivity, cellulose is industrially preferred to chitin. However, its derivative chitosan (CS) possesses higher solubility in a wide range of solvents and has gained considerable attention from researchers and industries, resurging the interest in nature chitin. Amines' presence in the chitin and chitosan structure can be chemically modified to improve their properties and hence furnishing several applications in technological and industrial fields.<sup>54,74</sup> In this sense, CS is produced by the hot leaching process of chitin, which causes the removal of the acetate group from the chitin molecule by hydrolysis,54,74 Scheme 6.

After this process, chitosan is washed, dried, ground, and ready to be commercialized.<sup>34,35,61</sup> The chitosan application for food packaging formulation tests showed excellent mechanical and barrier ( $O_2$  and moisture) properties, suitable antimicrobial and antioxidant activities, and good biodegradability, making chitosan a promising polymer in the packaging industry.<sup>41-44,75-78</sup>

Although biopolymers based on renewable sources consist a step forward for food packaging to achieve a more sustainable society, we are still highly dependent on polymers derived from non-renewable sources for such purposes. In addition, the current packaging technologies lack several points, such as higher resistance barrier properties, hence the research of new technologies for food packaging. In this way, nanocomposites (mixture of two different nanomaterials with distinct properties) endure a promising pathway for high-tech packaging. They might improve mechanical, thermal, barrier properties, cytotoxicity, microbiology, and biodegradable properties.<sup>41–44,75</sup>

#### 2.4. Nanocomposites-based packaging

The first example of composites for food packaging was reported at the beginning of this century (21st). Metal cans were covered with a multilayer plastic barrier preventing the contact of the food with the metal and thus increasing its shelf life. Also, other technological changes in food packaging were reported, such as the change in the internal atmosphere, the resistance of the packaging to the microwave oven, trays for gas and electric ovens, and retractable film labels, among other examples.<sup>51</sup>

Due to the several combinations of polymer matrix/ dispersed nanomaterials, polymer-based nanocomposites can provide different properties for a wide range of applications.<sup>33,79</sup>

Graphene-based nanocomposites have been utilized to improve UV resistance as a barrier against gases and still have good thermal, mechanical, and electrical properties compared to their polymeric matrices.<sup>80-83</sup> Furthermore, graphene-based nanomaterials combined with biodegradable polymers offer high potentials to be used as antimicrobial and antioxidant in active packaging technology resulting in more quality, safety, extended shelf-life, and added value.<sup>84-86</sup>

Some examples of nanocomposites based on polymer matrix/nanomaterials can be found in the literature. For example, polyvinyl alcohol (PVA)/graphene-based nanocomposites are just one example of polymeric nanocomposites

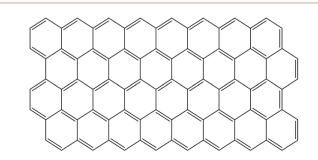
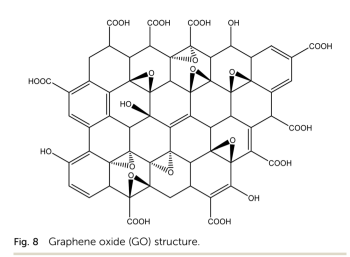


Fig. 7 Graphene structure (Gr).



with considerable potential in the food packaging industry. They have excellent gas and moisture barrier properties and good thermal resistance. Besides protecting the product, the new concept of packaging technologies brings technological innovations to adding novel features for barrier performance. An example is the active packaging technologies and modified atmosphere packaging for fresh vegetables and fruits.<sup>87</sup>

The interaction between the packaging, the product, and the environment through partial gas permeability increases shelf-life providing oxygen to cellular respiration.<sup>33</sup>

Biobased polymer packaging generally are films made from biopolymers and biodegradable polymers, such as starch,<sup>88</sup> pullulan,<sup>81</sup> PVA,<sup>85,86</sup> and PLA.<sup>89</sup> These nanocomposites can be used in food packaging as nanocoating films to control moisture transfer and gas exchange, provide security, and preserve nutrition and sensory features.<sup>79</sup>

Furthermore, in smart packaging based on polymer nanocomposites, graphene can be used as a sensor for biochemical or microbial changes in the product packed to detect specific food-borne pathogens or gases. Thus, this type of packaging can serve as oxygen and spoilage indicators for food safety and quality monitoring.<sup>90</sup> Due to the advanced features required for the food packaging industry, we highlight in this review graphene-based nanocomposites. Specially graphene oxide is a promising packaging material due to its unique characteristics, such as high barrier properties; good mechanical, electric, and thermal properties; large specific surface area; high electrical conductivity; carrier mobility; antibacterial, antifungal antioxidant, and biocompatibility.<sup>84,87,89,99,92-101</sup>

The electrical properties mean that the food in the package can be quickly heated and provide a package that "speaks to the consumer". The gas barrier means that the food can be preserved for longer, preventing oxidation and external contamination by microorganisms.<sup>80,83,91-95</sup>

Since all physicochemical properties of food packaging materials can be improved using graphene-based composites, a closer look at the graphene family exerts a pivotal role in better understanding the final product. The following section will present a detailed discussion regarding the graphene family and their nanocomposites.

# 3. Graphene-based nanocomposites applied to food packaging

#### 3.1. Graphene, graphene oxide, and reduced graphene oxide

Graphene (Gr) was introduced in 1986 by Bohen's group, aiming to understand a two-dimensional atomic sheet of graphite. However, in the 21st century, discoveries concerning graphene were made, and it was observed that graphene was thermodynamically unstable under ambient conditions. In 2004, Novoselov's group isolated and characterized a graphene monolayer obtained by mechanical exfoliation under ambient conditions. IUPAC defines graphene as a single layer of aromatic polycyclic carbon of almost infinite size, a single flat layer of carbon atoms with sp<sup>2</sup> hybridization, compacted together in a dense shapeshaped grid in two ordered dimensions (2D),<sup>11,12,39,40,99,100</sup> Fig. 7.

The graphene unit cell has a hexagonal shape and comprises two equivalent sub-networks of carbon atoms, connected by sigma ( $\sigma$ ) bonds, where the length of this C–C bond is 0.142 nm. Each carbon has a pi ( $\pi$ ) orbital in this network that generates a delocalized electronic effect. These bonds and structures provide excellent properties that are highly superior compared to other nanomaterials such as clays, zeolites, zinc oxide, and silicate-based.<sup>11,12,39,40,99,100</sup>

Graphene presents several interesting electronic properties due to its two-dimensional framework.<sup>101-103</sup> High carrier mobility is observed, reaching 15 000 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup> at room temperature (or 200 000 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup> when impurity scattering is minimized). The high carrier is explained by promoting electrons in the valence band to the conduction band, forming a positively charged space that allows the current formation under an external electric field.<sup>101-103</sup>

The structure of graphene can be modified to generate different structures derived from graphene (graphene-based materials), thus presenting other physicochemical properties.<sup>7–13,19–23</sup>

GO (Fig. 8) possesses reduced thermal and electrical properties than Gr. However, the formation of polar groups in the GO structure favors its application in composites and polymeric nanocomposites.<sup>11</sup> The structure of the GO is still the target of several discussions in the literature,<sup>7–13,19–23</sup> in which physicochemical studies indicate that epoxy and hydroxyl groups are created – usually in the basal planes of the GO leaves – and carboxyl groups commonly in the leaf edges or the pores of the

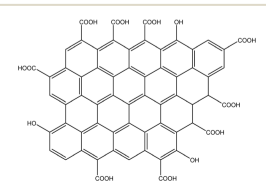


Fig. 9 Structure of reduced graphene oxide (rGO).

#### Review

GO structure. Also, it a variation in the amount, location, and functional available in the GO structure is observed, which can be controlled by the synthesis conditions and methods as a function of the specific application and thus reflected the desired properties.<sup>11</sup>

Graphene oxide may present insulating or semiconducting behavior, possessing sp<sup>2</sup> and sp<sup>3</sup>-bonded carbon atoms in its honeycomb structure. Oxygenated groups such as epoxy and hydroxyl groups are usually covalently bonded with sp<sup>3</sup> hybridized carbons, hindering their conductivity and leading to higher hydrophilicity.<sup>103-106</sup> Recent works stated that depending on the oxygen content, GO might present a bandgap from 1.7 eV (semiconductor) to 2.1 eV (insulator), having a higher electrical resistance than graphene.<sup>107</sup>

Generally, graphene oxide (GO) synthesis uses well-known methods described by Staudenmaier,<sup>108</sup> Brudie,<sup>109</sup> and Hummers.<sup>110</sup> In this manner, Staudenmaier,<sup>108</sup> and Brodie,<sup>109</sup> in a pivotal study, oxidized graphite with a combination of HNO<sub>3</sub> and KClO<sub>3</sub>. Furthermore, H<sub>2</sub>SO<sub>4</sub>, HNO<sub>3</sub>, and HClO<sub>4</sub> have also been used to promote graphite oxidation, as described by Staudenmaier,<sup>108</sup> Brudie,<sup>109</sup> and Hummers' method.<sup>110,111</sup>

However, the most used GO preparation methods considering biomedical and food packing purposes are the Hummers' method<sup>110</sup> and the modified Hummers' method.<sup>108,112</sup> Classical Hummers' process involves the treatment of graphite with H<sub>2</sub>SO<sub>4</sub>, NaNO<sub>3</sub>, KMnO<sub>4</sub> and H<sub>2</sub>O<sub>2</sub>.<sup>110</sup> The modified Hummers' method uses H<sub>2</sub>SO<sub>4</sub>, KMnO<sub>4</sub>, and H<sub>2</sub>O<sub>2</sub> without the addition of NaNO<sub>3</sub>, which generates less toxic gases during the procedure.<sup>112</sup> KMnO<sub>4</sub> is one of the most potent oxidizing agents in an acidic medium. In concentrated H<sub>2</sub>SO<sub>4</sub>, this oxidizing agent provides complete graphite intercalation, forming graphite bisulfate intercalated in all the graphene layers. This intercalation favors the diffusion of the KMnO<sub>4</sub> and thus leads to effective graphene oxidation to graphene oxide without the need for NaNO3.113,114 Furthermore, other variations of the Hummers' method are reported, by the replacement of H<sub>2</sub>SO<sub>4</sub> with a mixture of H<sub>2</sub>SO<sub>4</sub>/H<sub>3</sub>PO<sub>4</sub>, causing a less exothermic process without the formation of hazardous gases and also leading to more considerable oxidation in graphite, higher hydrophilicity and a more regular structure due to the cyclic phosphates creation.115

From a sustainable point of view, greener methods for synthesizing graphene and graphene oxide are reported, such as the synthesis using pyrolysis biochar from the lignocellulosic biomass leaves of *Cinnamomum Camphora*, in which the  $\pi$ - $\pi$  interaction of the biochar suspension with D-tyrosine solution promotes the separation of Gr layers by centrifugation.<sup>22</sup> Other graphene-like carbons from biomass pyrolysis methods found in the literature are the salt-based method, chemical blowing technique, model-based confinement, coupling to hydro-thermal carbonization treatment, and post-exfoliation.<sup>23</sup>

Notwithstanding, changes in GO structure can also be carried out by a reduction reaction and generating rGO. This reduction consists of partially removing oxygenated groups, thus conferring excellent structural stability compared to GO,<sup>10</sup> Fig. 9.

GO reduction to rGO can be carried out by chemical<sup>17,18</sup> or thermal<sup>19,116</sup> routes. Several reducing agents are used in the

chemical reduction, such as hydrazine, sulfur compounds, sodium borohydride, ascorbic acid, tyrosine, dimethylhydrazine, ethylenediamine, and others reducing agents and reducing atmosphere.<sup>17,18</sup> The first publication regarding GO reduction by chemical route was in 2006, in which it was used hydrazine and dimethylhydrazine as reducing agents. The nitrogen atom from hydrazine possesses a nucleophilic character, essential to reducing GO. In addition, other examples of reducing nucleophilic compounds are amines, hydrazine derivatives, and sodium borohydride. The reducing agents can also be electrophilic, such as borane in THF solution, one of the most common electrophiles for GO reduction.<sup>17</sup> Furthermore, another way to reduce GO is to treat the suspension in an acid solution under reflux. The number of oxygenated species to be reduced can be easily controlled by the reflux time.<sup>17</sup> Still, considering the chemical methods for rGO preparation, most of them use a lot of solvents and hazardous reductants, thus making the process expensive and non-sustainable due to the large amount of waste produced. In this way, other processes of chemical reduction of GO that use more sustainable reducers, such as tryptophan and ascorbic acid, have been reported. However, in these reduction processes, the agglomeration of rGO particles can occur, reducing their surface area.19 The thermal reduction process involves submitting GO to rapid heating under an inert atmosphere (Ar, N<sub>2</sub>). Firstly, GO the is dried, placed in a quartz tube and subjected to high temperatures (in the range of 700 to 1000 °C) for a short time, usually less than 1 h. During this thermal decomposition, H<sub>2</sub>O, CO<sub>2</sub> and CO are released, which increases the internal pressure, causing the separation of the leaves, forming the rGO.<sup>11</sup> The amount of oxygenated groups remaining in the rGO depends on the initial oxygen concentration in the GO, the proportion of epoxy and hydroxyl groups in the GO, and the temperature used to reduce the GO.117

In this way, chemical and thermal reduction methods generate rGO with similar electronic and structural properties. However, the thermal protocol does not require a purification step or solvents since it only uses temperature as a reducing agent, making it a faster and cheaper protocol.<sup>17,18,116</sup> Notwithstanding, decomposition gases (CO<sub>2</sub> and CO) can be collected and used as reagents in several green chemical processes.<sup>118-121</sup>

Thus, graphene-based materials have a high potential for application in the production of new energy storage batteries, magnetic sensors, materials, superconductors, medication administration, ultra-sensitive biosensors, tissue engineering, catalysis, nanocomposites and polymeric nanocomposites, and food packing.<sup>21</sup> Considering the encouraging features of GO and rGO, such as chemical, mechanical and antimicrobiological properties, these materials are highly promising to be used in food packing composites.

#### 3.2. Graphene-based nanocomposites - properties

It is essential to select the most suitable food packaging for storage and transportation to maintain food quality, *i.e.*, from the moment they leave the industry until reaching consumer usage. Also, proper packaging provides food with safety, lifetime extension, and protection against insects, microorganisms, light, and chemical contaminants. Furthermore, appropriate packaging can control the humidity, oxidation, diffusion of gases, and taste-odor maintenance.<sup>57,70</sup>

The molecular structure of the polymeric materials used for food packaging directly influences the polarity, rigidity, malleability, crystallinity, and intermolecular forces present in the polymeric chain, controlling the diffusion of the external molecules into the packaging. The chemical polymer composition can alter the barrier properties, impacting the characteristics of resistance, rigidity, cost, efficiency, ease of processing, and increasing its useful life.<sup>57,70</sup>

Polar polymers have good gas barrier properties. However, they are hydrophilic, and they allow water vapor permeation. On the other hand, nonpolar polymers do not allow the water vapor passage but have weak barrier properties, facilitating gas permeation independently of the polymer thickness. Copolymerization or additives incorporation in the polymeric matrix for food packaging formulations may affect these barrier properties as they fill and reinforce the polymeric matrix-free spaces. However, this will depend on the degree of adhesion and polymer and additive compatibility.<sup>57,70</sup>

Knowing the polymers' thermal properties is essential to understanding the physical and chemical transformation in nanocomposites structure with the temperature variation. The glass transition temperature, crystalline melting temperature, and crystallization temperature are the most critical transformations in these materials.<sup>70</sup>

# 3.3. Graphene-based preparation for nanocomposite production

In this way, the encouraging features of GO and rGO, such as chemical, mechanical and antimicrobiological properties, these materials are highly promising to be used in food packaging composites.<sup>21</sup> Generally, before producing graphene-based nanocomposites, the Gr should be in the GO or exfoliated forms (eGO). Therefore, incorporating the polymeric matrix into the structure of the eGO can be carried out, generating nanocomposites of commercial interest. GO exfoliation can be performed by two main routes: (i) thermal route; (ii) using polar solvents (water, ethanol, dimethylformamide - DFM, methyl pyrrolidone - NMP, propylene carbonate - PC) under stirring or ultrasonication.19,37 In the thermal process, the powdered GO is subjected to high temperatures (400-1000 °C) and a high heating rate for a few seconds. Then, the decomposition of the -OH and -COOH groups on the GO surface into CO, CO<sub>2</sub>, and H<sub>2</sub>O occurs. This process is advantageous since it produces a thermally exfoliated GO (TeGO) with a high surface area (700-1500 m<sup>2</sup> g<sup>-1</sup>), low apparent density, and platelets similar to the eGO platelets produced using the sonication process.<sup>19</sup>

The suspension is electrostatically stabilized in the solvent and ultrasound-assisted method by the carboxylated groups present in the GO lamellae peripheries.<sup>19</sup> The GO exfoliation performed by the sonication process in aqueous suspension furnishes GO monolayer platelet with high yield and hence forming eGO monolayers. However, the exfoliation process with agitation formed eGO monolayers with larger platelets, and this process is time-consuming and presents a low yield.<sup>19,37</sup> Fig. 10 shows the graphene oxide (GO) and exfoliated graphene oxide (eGO) structures, layers, and platelets.<sup>19</sup>

The exfoliated platelets by stirring or ultrasound can be later reduced by chemical processes. These processes involve the formation of a suspension between eGO and a reducing agent, such as sodium borohydride, hydrazine monohydrate, or dimethylhydrazine, in a polar solvent (water, NMP, DMF, DMSO) or a mixture of these solvents.<sup>19</sup>

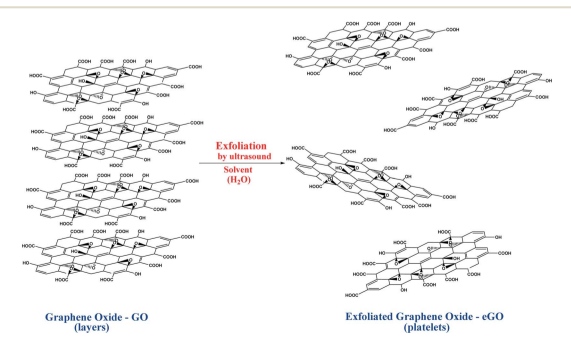


Fig. 10 Exfoliation of GO layers to eGO platelets using solvent and ultrasound.

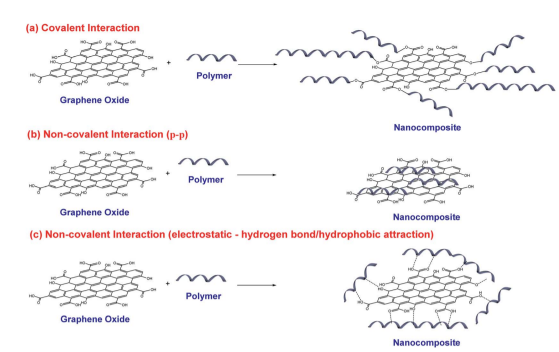


Fig. 11 Types of functionalization of GO with polymers for nanocomposites synthesis: (a) covalent functionalization; (b)  $\pi - \pi$  non-covalent functionalization and (c) electrostatic non-covalent functionalization.

Generally, chemically reduced eGO platelets result in materials (erGO) with a C : O ratio = 10 : 1, in which the oxygen content is related to the –OH and –COOH non-reduced groups in the eGO structure.<sup>19,37</sup> Biochemical reducing agents such as vitamin C (ascorbic acid) and tryptophan have also been tested but have resulted in the aggregation and reduction of the eGO platelet surface area.<sup>19,122,123</sup>

GO, rGO, eGO, and erGO structures have interesting functional groups, such as hydroxyl and epoxy, carboxyl, carbonyl, ketone, phenol, and lactone groups. These moieties allow the insertion of new atoms, molecules, or macromolecules (thermoplastic, thermosetting and biodegradable polymers) and thus creating the graphene-based nanocomposites.<sup>19,37,124</sup>

The functionalization of graphene-based nanomaterials can be performed by covalent or non-covalent pathways. The covalent functionalization occurs from creating the covalent bonds between functional groups on the eGO platelets surface with the polymeric matrix,<sup>16,19,37,38</sup> (Fig. 11(a)). This pathway can be carried out by nucleophilic substitution reactions, electrophilic substitution, condensation, cycloaddition, esterification, and hydrolytic condensation. In this manner, covalent functionalization is a propitious procedure for adjusting the charge transfer interface but impairs the electrical properties of the formed nanocomposite. On the other hand, non-covalent functionalization involves  $\pi$ - $\pi$  interactions (Fig. 11(b)), hydrogen bonding (Fig. 11(c)), and hydrophobic attraction.<sup>16,19,37,38</sup>

Covalent and non-covalent functionalization generates stable dispersions in organic solvents and can improve the compatibility of GO platelets with the polymers to be inserted in the nanocomposite structure.<sup>16,19,37,38</sup>

Graphene-based/polymer nanocomposites can be prepared by different methods such as: mixing (MX), melt-mixing (MM), *in situ* polymerization (ISP), and layer-by-layer (LBL) assembling. Here, it is worth mentioning that the chosen method will influence the properties of the final nanocomposite, such as surface area, molecular interaction, dispersion, and other properties.<sup>16,19,37,38</sup>

The MX method consists of adding a polymeric matrix solution to a graphene-based suspension. The resulting mixture is subjected to sonication, mechanical stirring and solvent evaporation. This method effectively obtains a fine dispersion of the polymeric material, but it uses non-benign solvents, and removing the solvent is difficult. As an example, the synthesis of graphene-based/poly(vinyl)alcohol nanocomposite.<sup>16,19,37,38</sup>

The MM technique is the most efficient and environmentally benign since solvents are not used. This method is commonly applied when *in situ* polymerization is not indicated, generally for thermoplastic polymers. The synthesis consists of melting the polymer and dispersing the graphene-based material through an extrusion and injection molding. However, the polymer can undergo thermal degradation, and the structure of the graphene-based nanocomposite can be disrupted by the high shear force involved during synthesis. For example, MM is applied to synthesize the graphene-based/polypropylene, graphene-based/poly(lactic acid), and graphene-based/ polyethylene terephthalate nanocomposites.<sup>16,19,37,38</sup>

ISP protocol is used to disperse the graphene-based material homogeneously in the polymeric matrix. In this technique, the graphene-based material is dispersed in the monomer followed by the polymerization reaction, with subsequent polymer grafting over the graphene-based surface. *In situ* polymerization uses heat or microwave radiation to form nanocomposites. Using ISP, a more homogeneous and fine dispersion occurs between graphene-based and the polymeric matrix, being very

types	Matrix : graphene-based ratio and others	Synthesis method	Polymers properties modifications and improvements	Important comments and applications	Ref.
PP-Gr	-Gr-PP 3 vol%; extrusion at 180 °C for 3 min; injection molding at $T_{\rm barrel} = 180$ °C and $T_{\rm mold} = 80$ °C, injection processes of 1 MDs	MI/MM	-↑Thermal conductivity	-Viscoelastic properties allow the fabrication of composites with desired properties at high	Kalaitizidou and co- workers (2007) <sup>24</sup>
	injection pressure at 1.1 ML a		-†O2 barrier	-Improvement of physical, -Improvement of physical, thermal, and mechanical properties	
PET-Gr	-Gr 1.5 wt%, sandwiched between two PET films, hot pressed, and chopped to granules	MM	-†Thermal stability	-can be applied to tood packaging -Gr uniformly dispersed in the matrix	Harel and co-workers (2012) <sup>26</sup>
	-Melt compound at 260 °C for 5 min		-↑Young's modulus, 1.16 to 1.40 GPa	-Enhanced crystallinity	
	-Molded into thin films (0.17 mm thick)		-↓Elongation at break, 40%	-The formed films present Gr exfoliated morphology	
	-Dried under vacuum at 120 $^\circ \mathrm{C}$ for 4 h		-↓Tensile strength, 56%	-Gr addition enhanced brittleness	
			-†02 barrier	-Can be applied to food packaging	
PS-GO	-2 g of PS, GO 20 wt%, and 4-vinyl benzyl chloride (VBC)	ISPs	- J Crystallinity	The nanocomposite showed high barrier and antimicrobial properties and could be utilized in yogurt containers and bottles for medicine capsules	Ghanem and co- workers (2020) <sup>25</sup>
	-Stirring -Dried for 24 h on a glass plate		<ul> <li>† Hydrophobicity</li> <li>† Thermic stability</li> <li>† Tg</li> <li>† Young modulus, 60%</li> <li>† Impact resistance</li> <li>↓ Water vapor transmission rate, 50 to 16.6 wt</li> </ul>	-Applied to food packaging	

" Gr: graphene; GO: graphene oxide; PP: polypropylene; PS: polystyrene; PET: polyethylene terephthalate; MM: melt-mixing; ISPs: in situ polymerization with solvent; IM: injection molding. increase; 1: decrease.

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Table 2 Renewable polymeric matrix/graphene-based group<sup>a</sup>

Nanocomposite types	Matrix : graphene-based ratio and others	Synthesis method	Polymers properties modifications and improvements	Important comments and applications	Ref
CS-GO	-GO + water, sonication for 45 min -CS + acetic acid solution 0.5% v/v (water)	XX	-†Young's modulus, 64% -†Tensile strength, 88%	-Good GO dispersion on CS matrix -The CS-GO nanocomposite formation occurred by H bonding between CS molecules with GO oxygenated groups and by electrostatic interaction between polycationic CS and the negative	Li and co- workers (2010) <sup>129</sup>
	-GO : CS ratio = 1 : 1, GO suspension was added on CS solution and stirring for 24 h -Dried at 40 °C, in glass plate until weight equilibration		-† $T_{ m gr}$ 175.4 to 180.4 °C	-Can be applied to food packaging	
PVA/GO and PVA/rGO	-PVA/GO and PVA/rGO: 0.3 wt%	MX	-↓O₂ permeability, 17–87 times bigger	-Promising for the development of transparent high-seas-barrier films	Lee and co- workers (2011) <sup>27</sup>
	-PET: a substrate		-73% light transmittance at 550 nm	-The H-bonding interaction between GO and PVA is not very large to change the PVA thermal	
	-Hybrids solutions were cast onto PET film at 90 °C			Can be applied to food packaging and bottles	
PLA/GO	-GO 0.4 wt%, dispersed in	SC	-†Young's modulus	-Environmentally benign	Magalhães and
	-GO + PLA/chloroform (plasticizer) solution. sonicated for 15 min		-↑Tensile strength	-Possible biomedical application	$(2013)^{130}$
	-PLA/GO films formed in polytetrafluoroethylene, coated nlare		$^{-1}$ O <sub>2</sub> and N <sub>2</sub> barrier	-Formation of transparent films	
	-Dried at room temperature for 7 days			-No detectable effects in the elongation at break -Can be annlied to food packaeine	
PVA-rGO	-GOr 0.8 wt%	MM	-↓Elongation at break, 451.9 to 229.1%	GO reduction process result in a good dispersion in PVA matrix	Hu and co- workers
	-PVA aqueous solution (1.0 g mL <sup><math>^{-1}</math></sup> ), 90.2 wt% -Glycerol (plasticizer), 9 wt% -Stirred at 95 °C for 24 h -Dried 2 days, 45 °C, in a glass plate, after 80 °C for 24 h		-†Tensile strength, 35.7 to 39.9 Mpa -†Thermal stability -†O2 barrier	-Can be applied to food packaging	(2013) <sup>131</sup>

Table 2 (Contd.)					
Nanocomposite types	Matrix : graphene-based ratio and others	Synthesis method	Polymers properties modifications and improvements	Important comments and applications	Ref
PVA-CS-rGO	-rGO 0.8 wt%	MM	-↓Elongation at break, 102.3 to 76.5%	-GO reduction process result in a good dispersion in CS-PVA matrix	Hu and co- workers (2013) <sup>131</sup>
	-PVA 63.1 wt%		-↑Tensile strength, 46.2 to 64.5 Mpa	-PVA/CS/rGO showed improvement in mechanical and thermal monorties	
	-CS 27.1 wt% -Glycerol (plasticizer), 9 wt% -Stirred at 95 °C for 24 h -Dried 2 days at 45 °C, in glass		-↑Thermal stability -↑O₂ barrier	-Can be applied to food packaging	
PLA/PEI-GO	PLA: 99.84 wt%	LBL	- $\downarrow$ O <sub>2</sub> permeability, $\approx$ 99%	-Use on biodegradable packaging materials	He and co- workers
	-PEI: 0.1 wt% in deionized water		-↑Tensile strength 93.2 to 120.2 MPa	-Increasing GO concentration in the nanocomposite structure can reduce oxygen nermeation	$(2014)^{132}$
	-GO: 0.06 wt%		-↑Elongation at break, 57.5 to 63.3%	-Can be applied to food packaging	
PVA-XGO	-XGO is GO functionalized with oxygen groups	MM	-†Young's modulus, 27.6 to 37.8 MPa	-The PVA-XGO mechanical and barrier properties depend on the dispersion and the alignment of oraphone-based in PVA matrix	Loryuenyog and co-workers (2015) <sup>133</sup>
	-XGO 0.3 wt%		-†Tensile strength, 25.4 to 37 9 MPa	Can be applied to food packaging	
	-PVA solution aqueous 10wt%, 20 g		- 15.0 Partial - 15.0 Partial - 10.2 barrier - 10.2 barrie		
CH-PAA-rGO	Solvent: deionized water	LBL	28.0% - $10_2$ permeability: 3.9 × $10^{-20}$ cm <sup>3</sup> cm <sup>-2</sup> Pa <sup>-1</sup> s <sup>-1</sup>	-Different pH's may affect the thickness of the film and also the	Grunlan and co- workers
	-CS: 0.1 wt% (pH = $3.5$ ; 4.5; 5.5)		${\rm cm^{-2}~Pa^{-1}~s^{-1}}$	permeaning or O <sub>2</sub> -In comparison with PET, the O <sub>2</sub> barrier of this nanocomposite is 20 rimes hicher	(6102)
	-PAA: $0.2 \text{ wt\%} \text{ (pH} = 3, 4 \text{ and } 5)$		↓O₂ transmission rate: 0,34 cm³ per m² per day per atm	The authors performed a thermal reduction of GO to increase the barrier and gas selectivity of this material by increasing the hydrophobicity of the film -Can be applied to food packaging	

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Table 2 (Contd.)					
Nanocomposite types	Matrix : graphene-based ratio and others	Synthesis method	Polymers properties modifications and improvements	Important comments and applications	Ref
	-Expholiated GO: 0.1 wt% (by sonication) -A support was used to deposite a thin layer of each polymer and GO.		-↑Gas selectivity (H2 and CO2)		
PLA-rGO	-Thermal reduction was performed at 175 °C for 90 min -The assembling process comprised a freestanding rGO film between two pieces of PLA film with heat pressing compression: 2000 pounds-force	LBL	-1 Hydrophobicity -1 Rresistance towards moisture penetration of 87.6% -1 O_2 barrier, 99%	-Promising solution to food packaging with outstanding environmental sustainability	Chen and co- workers (2016) <sup>135</sup>
GO-CS	at 65 °C for 30 min -50 mL GO solution (by sonication): obtain a homogeneous and stable dispersion solution -CS 1 wt%: dissolving CS in 0.5% (v/v) aqueous acetic acid solution -GO was dropwise into the CS solution	WX	-The spectrum of GO-CS nanocomposites exhibited neither a peak at 1596 cm <sup><math>-1</math></sup> related to $-NH_2$ absorbance vibration nor a peak at 1730 cm <sup><math>-1</math></sup> related to the C=O stretch of the carboxylic group	-Can be applied to food packaging	Xu & Liu (2017) <sup>93</sup>
GO-CS-TiO <sub>2</sub>	-Stirring for 24 h -Ratio GO : CS : TiO <sub>2</sub> (1 : 20 : 4): NPs2 -50 mL GO solution (by sonication, 60 W): obtain a homogeneous and stable dispersion solution -CS 1 wt%: dissolving CS in 0.5% (v/v) aqueous acetic acid solution (v/v) aqueous acetic acid solution GO was dropwise into the CS solution -GO was dropwise into the CS solution -Centrifuged at 6000 rpm for 1 h	XW	-Absorption bands of oxygen-containing functional groups were dramatically reduced -C-OH and carbonyl C=O bands were observed at 1200 and 1729 cm <sup>-1</sup> , respectively -Strong absorption bands at 450 and 670 cm <sup>-1</sup> , indicating the presence of the Ti-O-Ti bond in TiO <sub>2</sub>	-Non-toxic Exhibited significant antimicrobial effects against <i>B</i> . <i>subtilis</i> and <i>A. niger</i> biofilm formation -Can be employed as a cling film that effectively delays the loss of moisture in fruits and vegetables -Can be applied to food packaging	Xu & Liu (2017) <sup>93</sup>

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Table 2 (Contd.)

uposite         Matrix : graphene-based ratio and others         Synthesis method           -rfbe supernatant was discarded, and the deposit was freezedried for 12 h in freeze dryer for 12 h in freeze dryer for 12 h in freeze dryer (or 12 h in freeze dryer for 0.04 ± 1 wt%)         MX           e-GO         -callulose/16 wt% GO and cellulose/18 wt% GO and cellulose/18 wt% GO and cellulose/18 wt% GO         MX           Gr         -2 g CS was suspending in acetic         MX           -1mmersion in thermostatic bath:						
The reprenent we discription for 13 hin tree deposition with the rest of 13 hin tree deposition with the fiber nutrix clutonic polyaerytimide (FMA)     MX     1.0, permethility, 90.85%     CPM prevents the C0 effi- regregation in the fiber nutrix (1.84% GO)       Cutionic polyaerytimide (FMA)     MX     -1.0, permethility, 90.85%     CPM prevents the C0 effi- regregation in the fiber nutrix (1.84% GO)       Cutionic polyaerytimide (FMA)     MX     -1.0, permethility, 90.85%     CPM prevents the C0 effi- regregation in the fiber nutrix (1.84% GO)       0     -3 g CS was suspending in acetic     MX     -1.1 means in the monstatic (1.91%)     -1.0, permethility, 90.85%       0     -3 g CS was suspending in acetic     MX     -1.1 means in the monstatic (1.91%)     -1.0, permethility, 90.85%       0     -3 g CS was suspending in acetic     MX     -1.1 means in the monstatic (1.91%)     -1.0, permethility, 90.85%       0     -3 g CS was suspending in acetic     MX     -1.1 means in the monstatic (1.91%)     -1.0 means in the monstatic (2.91%)       0     -3 g CS was suspending in acetic     MX     -1.1 means in the monstatic (2.91%)     -1.0 means and acetic (2.91%)       0     -3 g CS was suspending in a terminal striter: 2.1     -1.0 means and acetic (2.91%)     -1.0 means and acetic (2.91%)       0     -3 g CS was suspending in a terminal striter: 2.1     -1.0 means and acetic (2.91%)     -1.0 means and acetic (2.91%)       0     -3 g CS was suspending in a termin	Nanocomposite types	Matrix : graphene-based ratio and others	Synthesis method	Polymers properties modifications and improvements	Important comments and applications	Ref
eC0     ccllubser/16 wr% C0 and ccllubser/16 wr% C0 and Cutonic polyacrylamide (ZAM)     -(Jo, permeablity, 99,45%     cCllubser/16 matrix (4 wr% C0)       Cationic polyacrylamide (ZAM)     -(Jo, permeablity, 99,45%     -(Do, permeablity, 99,45%       Cationic polyacrylamide (ZAM)     -(Jo, permeablity, 99,45%     -(Do, permeablity, 99,45%       Cationic polyacrylamide (ZAM)     -(Jo, permeablity, 99,45%     -(Do the large grapher for lar		-The supernatant was discarded, and the deposit was freeze-dried for 12 h in freeze dryer				
Cationic polyacylamide (CPAM)     - Thurst strength 81.4%     - Contonic polyacylamide (CPAM)       CrAMGGr. 0.04 : 1 w/%     - Cationic polyacylamide (CPAM)     - Thurst strength 81.4%     - Pennic polyacylamide (CPAM)       CPAMGGr. 0.04 : 1 w/%     - CAMGGr. 0.04 : 1 w/%     - Thurst strength 81.4%     - Pennic polyacylamide (CPAM)       CPAMGGr. 0.04 : 1 w/%     - CAMGGr. 0.04 : 1 w/%     - Thurst strength 81.4%     - Pennic polyacylamide (CPAM)       CPAMGGr. 0.04 : 1 w/%     - 2 g CS was suspending in acetic     MX     - 1 Flastic modulus, 2.5 to     - Pennic polyacylamide (CPAM)       CPAMGGr. 0.04 : 1 w/%     - 2 g CS was suspending in acetic     MX     - 1 Flastic modulus, 2.5 to     - Pennic polyacylamide (CPAM)       Solution 1 w/%     - 2 g CS was suspending in acetic     MX     - 1 Flastic polyacylamide (CPAM)     - Pennic polyacylamide (CPAM)       Gr. 6 w/%	ose-GO	-Cellulose/16 wt% GO and cellulose/4 wt% GO	MX	-↓O₂ permeability, 99.85%	-CPAM prevents the GO self- aggregation in the fiber matrix	Huang and co- workers (2016) <sup>32</sup>
(GO: 0.04 : 1 w <sup>6</sup> /s.     Permit the large scale preparation of GO(cellulose page with high doping amount of GO       8 vis usspending in acetic     MX     -1 Elastic modulus, 2.5 to doping amount of GO       9 substruction (0.1 M)     -3.8 GPa     Presence of copended graphice doping amount of GO       1 sit of the mostatic bath:     -3.8 GPa     Presence of copended graphice led to higher values of mechanical increase in chain tempolements of motion and the fracture strength, 50 to 3.8 GPa     Presence of copended graphice presence of motion amile or the increase in chain tempolements of motion and the fracture strength, 51 to 3.8 GPa       anical stirrer: 2 h     -1 Elongation at break, 15 3.8 GPa     -1 Elongation at break, 15 0.9%.       w <sup>6</sup> /s.     -1 Elongation at break, 15 3.8 v <sup>6</sup> /s.     -1 Elongation at break, 15 0.9%.       w <sup>6</sup> /s.     -1 Elficiency against mold controlled temperature: 25 3.8 v <sup>6</sup> /s.     -1 Elficiency against mold can be applied to food packaging controlled temperature: 25       w <sup>6</sup> /s.     -1 Therial startify, 32.4 to 9.9%.     -1 Therial startify, 32.4 to 9.9%.       w <sup>6</sup> /s.     -1 Therial startify, 32.4 to 9.9%.     -2 con be applied to food packaging con that active fracted antibucterial ation at 60 °C for 1 h, dried       6.0, 1 w <sup>6</sup> /s.     -1 Therial startify, 32.4 to 9.0, 1 w <sup>6</sup> /s.     -2 con be applied to food packaging the natural properties than CS       6.0, 1 w <sup>6</sup> /s.     -1 Therial startify, 32.4 to 9.0, 1 w <sup>6</sup> /s.     -2 con be applied to food packaging the natural properties than CS       6.0, 1 w <sup>6</sup> /s.     -1 The		-Cationic polyacrylamide (CPAM)		-↑Burst strength 81.4% (4 wt% GO)	-Potential packaging applications	
Gr     3.g Gs was suspending in acetic     MX     -1 Flastic modulus, 2.5 to acid solution (0.1 M)     Presence of expanded graphic treases in chain entragements       Immersion in thermostatic bath:     3.8 Gra     Presence of expanded graphic increase in chain entragements       Immersion in thermostatic bath:     -1 Flacture strength, 50 to 3.6 Gra     Presence of expanded graphic properties, likely due to the complex strength, 50 to 3.6 Gra       Achain canagements     -1 Florgation at break, 15 0.0 0.6 wt%     -1 Elongation at break, 15 0.0 0.6 wt%     -1 Elongation at break, 15 0.0 0.6 wt%       -Cra to 5 wt%     -1 Efficiency against mold -1 Efficiency against mold -1 Efficiency against mold -2 stors wt%     -1 Finsile strength, 32.4 to 0.0 0.6 wt%       -Cra to 5 wt%     -1 Finsile strength, 32.4 to wt%     -1 Finsile strength, 32.4 to 0.0 0.6 wt%       -Cra to 5 wt%     -1 Therale strength, 32.4 to wt%     -1 Therale strength, 32.4 to 0.0 0.6 wt%       -Cra to 5 wt%     -1 Therale strength, 32.4 to 0.0 0.6 wt%     -1 Therale strength, 32.4 to 0.0 0.6 wt%       -Cra to 1 wt%     -1 Therale strength, 32.4 to 0.0 0.6 wt%     -1 Therale strength, 32.4 to 0.0 0.6 Wt%       -Cra to 2 wt%     -1 Wt%     -1 Therale strength, 32.4 to 0.0 0.6 wt%       -Cro to 1 wt%     -1 Therale strength, 32.4 to 0.0 0.6 Wt%     -1 Therale strength, 32.4 to 0.0 0.6 Wt%       -Cro to 1 wt%     -1 Therale strength, 32.4 to 0.0 0.6 Wt%     -1 Therale strength, 0.0 of packaging 0.0 0.6 Wt%       -Cro to nut PLA mixed		-CPAM/GO: 0.04 : 1 wt%			-Permit the large scale preparation of GO/cellulose paper with high doping amount of GO -Can be applied to food packaging	
Immersion in thermostatic bath:Thracture strength, 50 toSmall concentrations of CA are sufficient to functionalize of mode $25  ^{\circ}$ C.0 MPa.0 MPaSmall concentrations of CA are sufficient to functionalize controlled temperature:.1 Engration at break, 15Small concentrations of CA are sufficient to functionalize chroan and are effective against the natural proliferation of mode $0.0 MPa$ .0 Mehanical stirrer: 2 h.1 Engration at break, 15.1 Engration at break, 15Small concentrations of CA are sufficient of mode $0.0 M$ .0.5 wt%s .0.5 wt%s .0.1 Efficiency against mold $0.0 M$ .1 Efficiency against mold $0.0 M$ .1 Efficiency against mold $0.0 M$ .0.6 mode .0.5 wt%s .0.6 wt%s.1 Efficiency against mold $0.0 M$ .1 Efficiency against mold $0.0 M$ .1 Efficiency against mold $0.0 M$ .0.6 mted milds $0.0 M M$ .1 Promys modulus, 22.7 to $0.0 M M$ .1 Efficiency against mold $0.0 M$ .2 mole $0.0 M M$ .1.7 mised with acetic acid solution $0.0 M M$ .1 Promised strength, 32.4 to $0.0 M M$ .2 mole $0.0 M M$ .2 mole $0.0 M M$ .1.1.2 °C complet $0.0 M M$ .1 mole $0.0 M M$ .1 mole $0.0 M M$ .2 mole $0.0 M M$ .1.2.3 M M.1.2 mole $0.0 M M$ .1 mole $0.0 M M$ .2 mole $0.0 M M$ .1.2.4 mole $0.0 6 M M$ .1 mole $0.0 M M$ .2 mole $0.0 M M$ .1.2.4 mole $0.0 6 M M$ .1 mole $0.0 M M$ .2 mole $0.0 M M$ .1.2.4 mole $0.0 0 M M$ .1 mole $0.0 M M$ .2 mole $0.0 M M$ .1.2.4 mole <b< td=""><td>A-Gr 6)</td><td>-2 g CS was suspending in acetic acid solution (0.1 M)</td><td>XW</td><td>-†Elastic modulus, 2.5 to 3.8 GPa</td><td>-Presence of expanded graphite led to higher values of mechanical properties, likely due to the increase in chain entanglements</td><td>Demitri and co- workers (2016)<sup>136</sup></td></b<>	A-Gr 6)	-2 g CS was suspending in acetic acid solution (0.1 M)	XW	-†Elastic modulus, 2.5 to 3.8 GPa	-Presence of expanded graphite led to higher values of mechanical properties, likely due to the increase in chain entanglements	Demitri and co- workers (2016) <sup>136</sup>
Mechanical stirter: 2 h. [Elongation at break, 15Can be applied to food packaging to 9%Gr. 6 wt% Gr. 6 wt% Or 5 wt% Or 5 wt% Concession (CS): water solution 1 v/ $C2A: 0.5 wt%$ . [Enficiency against mold to 9% Or 9% Or 9% Or 5 wt% Or 6 wt%. [Enficiency against mold to 9% Or 9% Or 9%. [Enficiency against mold to 9% Or 9%Gr. 6 wt% Or 6 wt% Or 6 wt%. [Tonicssi (CS): water solution 1 v/ S43.7 MPa. [Tonicssi (CS): water solution 1 v/ S43.7 MPa. [Tonics mold packaging0 0.6 wt% Or 6 wt% Or 6 wt%. [Tonics mold ms, 22.7 to S43.7 MPa. [Tonics mold ms, 22.7 to packaging. [Tonics mold ms, 22.7 to packaging12.0 °C owen's of 0.6 wt%. [Tonics mold ms, 22.7 to S43.7 MPa. [Tonics mold ms, 22.7 to packaging. [Tonics mold ms, 22.7 to packaging0. [Tonics mold ms, 22.7 MPa. [Tonics mold ms, 22.7 to packaging. [Tonics mold ms, 22.7 to packaging. [Tonics mold ms, 22.7 to packaging12.0 °C owen's at 12.0 °C owen's. [Tonics mold ms, 22.4 to packaging. [Tonics mold ms, 22.4 to packaging. [Tonics mold ms, 22.4 to packaging0. [Tonics mold ms, 22.7 to at 12.0 °C owen's. [Tonics mold ms, 22.4 to packaging. [Tonics mold ms, 22.4 to packaging12.0 °C owen's at 12.0 °C owen's. [Tonics mold ms, 22.4 to at 12.0 °C owen's. [Tonics mold ms, 22.4 to packaging. [Tonics mold ms, 22.4 to at 12.0 °C oshowed higher at 12.0 °C owen's0. [Tonics mold ms, 22.7 too at 12.0 °C owen's. [Tonics mold ms, 22.4 to at 12.0 °C owen's. [Tonics m		-Immersion in thermostatic bath: 25 °C		-† Fracture strength, 50 to 90 MPa	-Small concentrations of CA are sufficient to functionalize chitosan and are effective against the natural proliferation of mold	
Gr. 6 wt% C.1. 5 wt% C.1. 5 wt% C.1. 1 5 wt% C.2. 1. 5 wt% C.2. 1. 5 wt% C.1. 1 5 wt% C.1. 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		-Mechanical stirrer: 2 h		-↓Elongation at break, 15 to 9%	-Can be applied to food packaging	
Chitosan (CS): water solution 1 v/ wised with acetic acid solution $\neg \uparrow Young's modulus, 22.7 to5843.7 MPa\neg \uparrow roung's modulus, 22.7 topackaging\neg \bullet otential application for foodpackaging2 wt%\neg wt%\neg Troung's modulus, 22.7 to5843.7 MPa\neg \uparrow roung's modulus, 22.7 topackaging\neg \bullet otential application for foodpackaging2 wt%\neg 0.6 wt%\neg \uparrow Tensile strength, 32.4 toat 120 °C overnight\neg \uparrow Tensile strength, 32.4 toat 120 °C overnight\neg GO showed higherantimicrobial properties than CS\neg \uparrow Tensile strength, 32.4 topacteria\neg GS showed higherantimicrobial properties than CS\neg T = 27 MPa0\neg TA + GO, 1 wt%\neg T = 327 MPa\neg 120 °C overnight\neg GS showed higherantimicrobial properties than CS\neg T = 327 MPa10\neg 0.6 wt%\neg T = 32.7 MPa\neg T = 32.7 MPa\neg GS showed higher\neg T = 32.7 MPa10\neg 0.6 wt%\neg T = 32.7 MPa\neg T = 32.7 MPa\neg GS showed higher\neg T = 32.7 MPa10\neg 0.6 wt%\neg T = 32.7 MPa\neg T = 32.7 MPa\neg GS showed higher\neg GS showed higher\neg T = 32.7 MPa10\neg 0.6 wt%\neg T = 32.7 MPa\neg T = 32.7 MPa\neg GS showed higher\neg GS showed higher$		-Gr: 6 wt% -CA: 0.5 wt% -Under controlled temperature: 25 °C		-† Efficiency against mold		
-GO 0.6 wt%-fTensile strength, 32.4 to-CS-GO showed higher-GO 0.6 wt%43.27 MPa-CS-GO showed higher-Sonication at 60 °C for 1 h, dried-17hermal stabilityCS does not show toxicity for-FIA + GO, 1 wt%-7Thermal stability-CS does not show toxicity for-PLA + GO, 1 wt%MX-1 Tensile strength 32.4 to-Demonstrated antibacterial-GO and PLA mixed and sonicated-1 Elongation at break, 41-Can be applied to food packaging-GO and PLA mixed and sonicated-1 Elongation at break, 41-Can be applied to food packaging-fTg, 91.6 wt-7Tg, 91.6 wt-0.6 Mpa-fTg, 81.5 to 84.8 °C-7Tg, 81.5 to 84.8 °C-0.6 break at 600 packaging-fTg, 81.5 to 84.8 °C-0.0 brenrier, 40%-0.0 brenrier, 40%		-Chitosan (CS): water solution 1 v/ v% mixed with acetic acid solution 2 wt%	MM	-†Young's modulus, 22.7 to 5843.7 MPa	-Potential application for food packaging	Advincula and co-workers (2017) <sup>137</sup>
-FThermal stability $-7$ does not show toxicity for at 120 °C overnight at 120 °C overnight $-7$ Thermal stability $-3$ does not show toxicity for bacteria at 120 °C overnight $-7$ more than the strength $-7$ and $-3$ does not show toxicity for bacteria $-7$ and $-3$ does not show toxicity for bacteria $-7$ and $-6$ does not show toxicity for bacteria $-7$ and $-6$ does not show toxicity for bacteria $-7$ and $-6$ does not show toxicity for bacteria $-7$ and $-6$ does not show toxicity for bacteria $-7$ and $-6$ does not show toxicity for bacteria $-7$ for $-7$ and $-6$ does not show toxicity for bacteria $-7$ for $-7$ and $-6$ does not show toxicity for bacteria $-7$ for $-7$ for $-6$ does not show toxicity for the applied to food packaging to $-7$ for $-7$ for $-6$ does not show toxicity for $-7$ does not show toxicity for the applied to food packaging to $-7$ for $-7$ for $-6$ does not show toxicity for $-7$ does not show toxicity for $-7$ does not show toxicity for the applied to food packaging to $-7$ for $-7$ for $-7$ does not show toxicity $-7$ d		-GO 0.6 wt%		-↑Tensile strength, 32.4 to 43.27 MPa	-CS-GO showed higher antimicrobial properties than CS	
-PLA + GO, 1 wt%MX $\uparrow$ Tensile strength 32.4 toDemonstrated antibacterial-GO and PLA mixed and sonicated $\downarrow$ 60.6 Mpa-f can be applied to food packaging-GO and PLA mixed and sonicated $\downarrow$ Elongation at break, 41can be applied to food packagingfor 30 min (was added polyto 31.6% $\uparrow$ $T_{c}^{s}$ +5 °C $\uparrow$ $T_{c}^{s}$ 81.5 to 84.8 °C $\uparrow$ $T_{c}^{s}$ 94.8 °C $\downarrow$ O2 barrier, 40%		-Sonication at 60 °C for 1 h, dried at 120 °C overnight		-†Thermal stability	-CS does not show toxicity for bacteria -Can be applied to food packaging	
- J Elongation at break, 41 -Can be applied to food packaging to $31.6\%$ - $\uparrow T_{2^{\circ}} + 5 \degree C$ - $\uparrow T_{7^{\circ}} 81.5$ to $84.8 \degree C$ - $\downarrow O_2$ barrier, $40\%$	0	-PLA + GO, 1 wt%	MX	-↑Tensile strength 32.4 to 40.6 Mpa	-Demonstrated antibacterial activity in food packaging	Ahmed and co- workers (2017) <sup>89</sup>
		-GO and PLA mixed and sonicated for 30 min (was added poly ethylene glycol as plasticizer)		-JElongation at break, 41 to 31.6%	-Can be applied to food packaging	
				-1 T <sub>g</sub> , +5 °C -1 T <sub>c</sub> , 81.5 to 84.8 °C -J O <sub>2</sub> barrier, 40%		

Table 2 (Contd.)					
Nanocomposite types	Matrix : graphene-based ratio and others	Synthesis method	Polymers properties modifications and improvements	Important comments and applications	Ref
Gly- nanocellulose- CO (Channel)	- GO 0.5% wt	MM	-†Tensile strength, 18.4 to 25.0 MPa	Glycerol and GO in the nanocellulose matrix led to	Arcot and co- workers
GO (Grycerol) Nanocellulose/ GO = GGN)	- GO aqueous solution sonicated for 3 h		-↓Thermal stability	synergiscue eneces -GGN films show potential to be used in the food package since they can bear heavier products	(/102)
	-Glycerol 40 wt% (plasticizer) -GO, glycerol, and cellulose were		-↑Elasticity -↑Elongation at break, 1.9 +0 0.402	than a zip-lock bag -Can be applied to food packaging	
	-Films formed in Petri dishes; dried at 40 °C for 2 days		+↑ Contact angle of the films		
			-↑Moisture sorption -↑Water vapor permeability		
OS-rGO (OS: oxidized starch)	-rGO: 1.0 wt%	MM	- 1.02 better - 1 Tensile strength 58.5 to 17.2 MPa	The nanocomposite mechanical property can be controlled by the rGO reduction time	Jiang and co- workers (2017) <sup>139</sup>
	-Plasticizing the OS: mixture and stirring (OS- rGO) at 90 °C, 300 rpm for 60 h		-↓O₂ permeability	-Can be applied to food packaging	
PLA-CNC/Gr	-PLA/CNC/Gr: 95/0.5/0.5 wt%,	MC	-†Young's modulus, 8%	-Good GO/matrix dispersion	Montes and co-
	prepared by ment compounding -Stirring 70 rpm at 185 °C for 10 min		-†Tensile strength, 11%	-Significant improvement in the antifungal activity by Gr	$(2018)^{140}$
			-↑O₂ barrier, 23% -↑Hydrophobicity	<ul> <li>Transparency of the materials</li> <li>Potential applications for food</li> <li>nackaoine</li> </ul>	
			- $\uparrow$ Thermic stability - $\uparrow$ $T_{g}$ : 51.8 to 53.5 °C - $\uparrow$ $T_{m}$ : 175.7 to 178.2 °C - $\uparrow$ Crystallinity: 24.8 to 28.0%	6 0	
EVOH-GO	-GO: 0.5 wt%	ММ	-†Electrical conductivity	-Excellent GO dispersion on	Lagaron and co-
	-EVOH: 99.5 wt%		$\uparrow T_{ m g}$	polynieric maurix -Can be used in smart food packaring	workers (2018) <sup>141</sup>
	-GO/EVOH sonication 15 min -Flectrosninning process		- $\uparrow T_{ m c}$	00	
GO-CS	-50 mL GO solution (by sonication): obtain a homogeneous and stable dispersion solution	МХ	-The spectrum of GO-CS nanocomposites exhibited neither a peak at 1596 cm <sup><math>-1</math></sup> related to $-NH_2$ absorbance	-Can be applied to food packaging	Xu & Liu (2017) <sup>93</sup>

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Table 2 (Contd.)					
Nanocomposite types	Matrix : graphene-based ratio and others	Synthesis method	Polymers properties modifications and improvements	Important comments and applications	Ref
	-CS 1 wt%: dissolving CS in 0.5% (v/v) aqueous acetic acid solution -GO was dropwise into the CS solution		vibration nor a peak at $1730 \text{ cm}^{-1}$ associated with the C=O stretch of the carboxylic group		
PLA/starch-Gr	-Stirring for 24 h -PLA/starch-Gr, 0.1 wt% of Gr -Vacuum oven overnight -The extrusion was pelletized at 50 °C for 4 h	ММ	-↑Elongation at break, 103.4% -↑Toughness, 500–900% -↓Young's modulus, 1.2 to 0.8 GPa	-Suitable for food packaging application -Can be applied to food packaging	Auras and co- workers (2018) <sup>142</sup>
Starch/gelatina- GO	-Starch/gelatin-GO matrix, 99.15 wt% -Starch/gelatin ratio 1 : 3 -GO 0.85 wt%	ММ	- $\downarrow$ O <sub>2</sub> permeability, 50% - $\uparrow$ Tensile strength 57.97 to 76.09 MPa - $\uparrow$ Young's modulus, 20.59 to 35.91 MPa - $\downarrow$ Elongation at break, 6.6 to 3.13%	Good candidate for biodegradable food packaging production	Baniasadi and co-workers (2018) <sup>31</sup>
PVA/Cu2O/TiO2- rGO	<ul> <li>-Sonication for 30 min</li> <li>-Cu<sub>2</sub>O-TiO<sub>2</sub>/rGO</li> <li>-Synthesized by ultrasonic</li> <li>-Synthesized by ultrasonic</li> <li>reduction and wet impregnation</li> <li>method using TiO<sub>2</sub> NPs, Cu(NO<sub>3</sub>)<sub>2</sub></li> <li>3H<sub>2</sub>O and GO</li> </ul>	sc	<ul> <li>↑ Thermal stability</li> <li>↓ Water vapor permeability</li> <li>↑ Zone of inhibition</li> <li>↑ Antibacterial activity under visible light</li> </ul>	-Environmentally benign -Uniform distribution of the particles in the PVA films	Venkatapras and co-workers (2018) <sup>143</sup>
	-PLA-Cu <sub>2</sub> O-TiO <sub>2</sub> /rGO -PLA-Cu <sub>2</sub> O-TiO <sub>2</sub> /rGO -Cu <sub>2</sub> O-TiO <sub>2</sub> /rGO 12.5 mg mL <sup>-1</sup> -PLA + Cu <sub>2</sub> O-TiO <sub>2</sub> /rGO, solutions were stirred -Dried in Petri plates at room		-↑Antimicrobial activity	-Effective antimicrobial activity under visible light against four different microorganisms -Potential for ambient light food packaging	
PLA-GO PLA-ST- <i>f</i> GO	temperature for 48 h -PLA/0.5 wt% GO (10 : 90) -fGO: GO functionalized with maleic anhydride and dodecyl	MM SC	-↓Water vapor permeability -↑T <sub>m</sub> , 165.9 to 170.2 °C -↑Young's modulus, 30% -↑Thermal stability	-Lightweight and strong packaging materials for food and industrial applications -Homogeneous dispersion of fGO in PLA-ST matrix	Peijs and co- workers (2018) <sup>28</sup> Sheng & Xiong (2019) <sup>144</sup>
	amine -4 g of PLA + 0.45 g of starch dissolved in chloroform -5 wt% tributyl citrate		-↑Crystallization capacity	-Potential versatile nanohybrids for food packaging and pharmaceutical industries	

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Table 2 (Contd.)					
Nanocomposite types	Matrix : graphene-based ratio and others	Synthesis method	Polymers properties modifications and improvements	Important comments and applications	Ref
	-Stirred for 1 h -0.4 wt% fGO, added and stirred for 1 h -Dried at 40 °C for 4 days in a polytetrafluorethylene frame		<ul> <li>↓ Plasticizer migration rates</li> <li>↑ UV shielding capacity</li> <li>↑ Hydrophobicity</li> <li>↑ Aging resistance</li> </ul>		
CS-rGO and CS- GO	-GO or rGO + acetic acid solution 1% v/v (water), 20 mL sonication for 9 min	МХ	-↑Storage modulus -↓Electric conductivity, graphite 1200 to GO 3.1 S cm <sup>-1</sup> and to rGO 60 S cm <sup>-1</sup>	-The addition of rGO did not affect the CS matrix hydrophilicity	Malmonge & Basso (2019) <sup>30</sup>
	<ul> <li>-CS (0.4 mg) + acetic ácid solution 1% v/v (water), 20 mL</li> <li>-GO or rGO suspension was added to CS solution, stirring for 5 min and sonication for 9 min</li> <li>-Solvent remotion and dried</li> </ul>		-↓GO and rGO dispersion on CS matrix	-The GO and rGO addition on the CS matrix did not accelerate the composite degradation process -Can be applied to food packaging	
Starch/PVA-GO	-Starch/5 wt%, PVA-GO -GO: 2 mg mL -All films should be adjusted before being tested for temperature $(T \approx 24 ^{\circ}\text{C})$ and relative humidity $(\approx 50\%)$	WX	-↑Tensile strength 18.40 to 25.28 MPa -↓Elongation at break, 184.03 to 143.78% -↑ thermal stability	-Starch/PVA-GO composite film can resist the water molecules migration -Can be applied to food packaging	Lin & Pu (2019) <sup>145</sup>
PVA-GA-GO	-22.5 mg GO in water, 2.71 mL, sonication 1 h -GO/water suspension was added in PVA-GA matrix and stirred at 6000 rpm 15 min -Dried overnight, at room temperature, on a glass plate	W	-↑Young's modulus 0.66 to 1.55 MPa -↑Tensile strength, 0.66 to 1.51 MPa -↑Thermic stability -↓Water vapor transmission rate, 38.49 to 32.13%	-PVA-GA-GO demonstrated antibacterial activity and can be applied to food packaging	Chowdhury & Mah (2020) <sup>146</sup>
PHB-Gr	-0.7 wt% Gr on PHB -Solvent: chloroform -Gr nanoparticles were dispersed in chloroform (1 mL) by	ХИ	-↓O <sub>2</sub> permeability, 1.53 to 0.4 mm per m <sup>2</sup> per day <sup>2</sup> per atm	-Was found to be environmentally safe and highly biodegradable -Applications in packaging of light-sensitive food products -High strength is desired for the packaging of heavy food items	Pakshirajan & Pugazhenthi (2020) <sup>29</sup>

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Table 2 (Contd.)					
Nanocomposite types	Matrix : graphene-based ratio and others	Synthesis method	Polymers properties modifications and improvements	Important comments and applications Re	Ref
	sonication for 45 min (15 s on cycle: 45 s off/cycle) -Stirring at 100 rpm for 60 min		-↓Water vapor permeability, 9.26 to 4 mm per m² per day² per atm ↑ Thermal stability ↑ Tm, 172.8 to 182.5 °C ↑ Tensile strength, 4.5 to 9.0 MPa -↓ Elongation at break, 15 to 12.2% -↓ Transparency to UV and visible light		
<sup><i>a</i></sup> Gr: graphene; GO: gr: cellulose nanocrystals; PEP: pickering emulsio ↓: decrease.	<sup>a</sup> Gr: graphene; GO: graphene oxide, rGO: reduced graphene oxid cellulose nanocrystals; EVOH: poly(ethylene-co-vinyl alcohol); PAA PEP: pickering emulsion polymerization; ISPs: <i>in situ</i> polymerizat ↓: decrease.	e; CS: chitosan; PLA: po : poly(acrylic acid); OS: o ion with solvent; LBL: lay	ly lactic acid; PVA: poly(vinyl alcohol) xidized starch; PHB: polyhydroxybuta ver-by-layer; PS: plasticized-starch; SP:	<sup>a</sup> Gr. graphene; GO. graphene oxide, rGO. reduced graphene oxide; CS. chitosan; PLA: poly lactic acid; PVA: poly(vinyl alcohol); PEI: polyethyleneimine; ST. starch; GA. glutaraldehyde; CN cellulose nanocrystals; EVOH: poly(ethylene- <i>co</i> -vinyl alcohol); PAA: poly(acrylic acid); OS: oxidized starch; PHB: polyhydroxybutanoate; MX: mixing; MM: melt-mixing; MC: melt compoundi PEP: pickering emulsion polymerization; ISPs: <i>in situ</i> polymerization with solvent; LBL: layer-by-layer; PS: plasticized-starch; SP: solution processing; SC: solution casting method. 7: increa ↓: decrease.	raldehyde; CN t compoundii 10d. ↑: increa

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effective for both heating techniques. For example, this protocol is used to synthesize the graphene-based/polystyrene nanocomposite.<sup>16,19,37,38</sup>

The LBL technique offers the possibility of building nanocomposite alternating layers between the graphene-based material and polymeric matrix. This synthesis method creates a nanomaterial with good dispersion and graphene-based orientation. Also, it allows the possibility of controlling the amount of graphene added between the polymer layers and the morphology of the generated nanocomposite. For example, the GO/poly(vinyl) alcohol nanocomposite construction can be mentioned.<sup>16,19,37,38</sup>

Graphene-based nanocomposites have several applications in biosensors, energy-related devices, food packaging, and biomedical, among others. However, many challenges are still encountered, especially for materials used in health and food. However, many challenges are still faced, especially for health and food packaging materials. There are already successes in using graphene nanocomposites with polymeric matrices of renewable sources in drug delivery. The low solubility of graphene in an aqueous solution makes its implementation difficult. The aqueous stability and the solubility of graphene can be modified by the induction of  $\pi$ - $\pi$  interactions with drugs, aromatic molecules, or other organic molecules. However, it is possible to circumvent it by replacing graphene with graphene oxide in the nanocomposite assembly. The advantages of using GO and rGO to construct polymeric nanocomposites consist of oxygenated groups (hydroxyl, epoxide, carboxyl, and carbonyl) on both GO and rGO nanosheets that favor the assembly of new nanometric structures and the new polymeric nanocomposite is formed. This assembly occurs through the stacking system between the nanomaterial and the polymeric matrix and between the hydrogen bonds and  $\pi$ - $\pi$  interactions that will significantly improve the properties of the polymeric nanocomposite formed as biocompatibility, low toxicity, thermal detachment, and mechanical stability.125,126

#### 3.4 Graphene-based nanocomposites - types

Gr, GO, and rGO are used as polymeric matrices additives for several nanocomposites' formulations. In this sense, these polymeric nanocomposites formulations often present improved properties for food packaging.<sup>127,128</sup>

Several research groups have been developing new materials for the manufacture of packaging based on nanocomposites, such as clays, tourmaline, zeolites, oxides, and carbon-based materials, such as graphene-based, to improve and control food packaging properties.<sup>25,33–36,51,61</sup> In this sense, no polymeric matrix has all the necessary/desired properties in food packaging production. However, with the nanotechnology advent, polymeric nanocomposites (polymer matrix + nanomaterial) arise, improving their thermal, mechanical, electrical, optical, and barrier properties of food packaging.<sup>70</sup>

The matrix/graphene-based nanocomposites will be divided into two groups in the next section: (a) non-renewable polymeric matrix/graphene-based group (petroleum-derived polymers)

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and (b) renewable polymeric matrix/graphene-based group (biomass-derived polymers).

**3.4.1.** Non-renewable polymeric matrix/graphene-based group. Table 1 shows the main nanocomposites types comprised of the non-renewable polymeric matrix and graphene-based material, the experimental conditions, synthesis methods, and properties.

Table 1 shows that melt-mixing is the most used synthesis method for constructing polymeric nanocomposites (nonrenewable origin). The addition of graphene-based nanomaterials increases the nanocomposite's thermal, barrier, and other properties.

**3.4.2.** Polymeric matrix renewable/graphene-based group. Table 2 shows the main nanocomposites based on renewable polymeric matrices, experimental conditions, synthesis methods, and properties modifications.

In this sense, Table 2 shows that MX and MM are the methods for synthesizing polymeric nanocomposites (from renewable sources) based on graphene most used by most research groups. It is due to their ease and simplicity of handling on a small scale. It is indisputable that the addition of graphene-based nanomaterials (Gr, GO, and rGO) generates a significant improvement in the various properties of these nanomaterials when compared to their pure polymeric matrices.

Improvements in polymeric nanocomposites occurred in their thermal, mechanical (high tensile strength, low elongation at break, increase in Young's modulus), and barrier properties (low permeability to water vapor; low permeability to  $O_2$ ). Generally, there is also an excellent dispersion of Gr, GO, or rGO in the polymer matrix, generating environmentally safe and biodegradable nanocomposites. However, the addition of GO or rGO in the polymeric matrices did not accelerate the degradation process of the composite. Many graphene-based polymeric nanocomposites have demonstrated antimicrobial activity and can apply to food packaging. A comparison between Tables 1 and 2 shows that several studies have been devoted to using renewable polymeric matrix/ graphene compared to non-renewable-based composites. This perspective shows the great scientific concern for environmental causes. The creation of new packaging from nano-composites is one of the most impressive and promising packaging technology results for the near future challenges.<sup>124</sup>

# 4. Microbiological properties of graphene-based nanocomposites

Fungal and bacterial transmission in contaminated objects and surfaces is one of the vectors of disease transmission by these microorganisms, although it is not the main transmission route. They adhere to surfaces and colonize the plants, insects, animals, fabrics, food, medical apparatus, packaging, and hard materials. This transmission by adhesion to solid organic and inorganic surfaces is a relevant challenge to public health, epidemiological significance, and economic losses.<sup>147</sup> Depending on the surface type and its humidity condition, the microorganisms can survive and grow for extended storage periods, sometimes even months. Removing them is necessary to disinfect and destroy microorganisms by using organic substances and chemical disinfectants that can effectively combat these pathogenic microorganisms.<sup>148</sup>

Another strategy is to use organic and inorganic compounds with fungal and bactericidal activities incorporated in polymeric matrices (fabrics, paper, paints, *etc.*) to eliminate microorganisms by contact. In this way, graphene and graphenebased materials emerge as a practical pathway to minimize the proliferation of these microorganisms.

Graphene is a biocompatible material with low or no cytotoxicity and promising for food packaging composites, and its oxidized form has different properties and antimicrobial effects in solution and when deposited on surfaces. In that perspective, several studies reported in the literature involving the graphene family and reporting its antibacterial activity<sup>148-157</sup> (*e.g.*, fungi

Table 3	Selected recent applications of graphene-based nanocomposites as antimicrobials	

Material	Microorganism	Ref
Reduced graphene oxide/Ag/Ag <sub>2</sub> S	E. coli	165
Reduced graphene oxide/ZnO	E. coli	166
Graphene oxide	S. aureus	159
Graphene oxide-catechol	E. coli/S. aureus	165
Reduced graphene oxide/CuO	E. coli/S. aureus	166
Graphene oxide cellulose/CuO	S. aureus, B. subtilis, E. coli/P. aeruginosa	167
Graphene oxides NiS-MoO <sub>3</sub>	E. coli/S. pyogenes	168
Reduced graphene oxide/Ag	E. coli/K. pneumoniae	169
Reduced graphene oxide/Ag	S. aureus/E. coli/P. aeruginosa	170
Graphene oxide	E. coli K12	171
Graphene oxide/p-aminophenol	E. coli/S. aureus	172
Graphene oxide/modified sodium anthraquinone-2-sulfonate	E. coli	173
Graphene oxide chloramine bromosuccinimide/FeCl <sub>3</sub> /KIO <sub>3</sub>	P. pneumonia/S. aureus	174
Graphene Oxide-chitosan/Ag	Pseudomonas sp	175
Graphene oxide/Ag	E. coli/S. aureus	176
Graphene oxide/ampicillin, chloramphenicol, or tetracycline	P. aeruginosa	177
Graphene oxide/NiS–In <sub>2</sub> O <sub>3</sub>	E. coli/S. aureus	178

and bacteria) – making these materials highly promising for food packaging nanocomposites.<sup>158-161</sup>

Regarding graphene-family composites, their antibacterial activity can be related to several intrinsic properties emerging from geometric and electronic structures.<sup>162</sup> The most common mechanism accepted for the antibacterial activities of graphene is oxidative stress caused in the microorganism by transferring electrons to their membrane and deactivating its protein functions and lipids through the reactive oxygen species (ROS). Furthermore, also can be found studies in the literature reporting that antibacterial activity can occur when phospholipids are extracted by supramolecular interactions with graphene and damage in the permeability of the bacterial cell wall and/or membrane.<sup>163,164</sup>

Moreover, the antimicrobial properties of the graphene family might be improved by the insertion of molecules by covalent bonds and the deposition of metal ions on its surface. These graphene-based nanocomposites materials may contain nanoparticles of metal ions/oxides/sulfides (manganese disulfide, cadmium sulfide), titanium dioxide, zinc oxide, copper oxide, polymers, antibiotics, among others.<sup>163,164</sup>

Nanotechnology involving the graphene family and its composites can be a solution for coping with bacteria in various situations. Table 3 shows some selected most recent examples of graphene-based-nanomaterials with high efficiency against bacteria, such as *Escherichia coli*, *Klebsiella pneumoniae*, Pseudomonas pneumonia, and *P. aeruginosa*, *Staphylococcus aureus*, *Staphylococcus pyogenes*, and *Bacillus subtilis*.

Despite the importance of graphene-based nanocomposites for safer packaging with resistance against microorganisms, it is noteworthy to mention that the specialized literature lacks in filling the gaps regarding the risks of nanomaterials used. In this way, information regarding their interaction with cellular components, migration to the food, tolerance dosage levels, toxicity to the human organism, long-term exposure, and toxicity to ecologically organisms when disposed of incorrectly.

# 5. Graphene-based nanocomposites– toxicity and (bio)degradability

Graphene and its derivatives (GD) have chemical, mechanical, electrical, and optical properties of interest for different applications such as biotechnology, biosensors, biomedicine, nanocomposites, catalysis, and drug delivery systems, metal detection, and removal, among others. Because of this, many studies are being developed to assess the effects of GD toxicity on the environment and the human body.<sup>29</sup> GD toxicity is influenced by the shape, size, concentration, synthesis methods, and type of functional groups. For example, the PHB/ graphene nanocomposite showed promising results in improving the food packaging life and high biodegradability. However, graphene in polymer nanocomposites can be released into the air and groundwater depending on size, shape, and concentration.<sup>155,179,180</sup>

In the formulation of nanocomposite PVA/gelatin was added to cellulose nanocrystals to make food packaging. The addition of cellulose nanocrystals in the nanocomposite showed improved properties (mechanical, thermal, and barrier). The results indicate that PVA-gelatin films reinforced with cellulose nanocrystals can be considered a potential biodegradable packaging material, especially for food packaging.<sup>63,181</sup> They can be tested in the construction of graphene-based polymeric nanocomposites.

Although the excellent adsorption capacity of graphene allows the capture of organic contaminants in aqueous media, DGs can promote the spread of these contaminants in groundwater, where DGs are more unstable and have greater mobility. Because of this, some research has been developed to evaluate the effects of these materials on aquatic organisms. It has been observed that the cell division of photosynthetic microorganisms is inhibited as the concentration of GD is high. On the other hand, invertebrates present several changes: changes in biochemical performances, the survival rate and inhibition of the swimming behavior of crustaceans, and

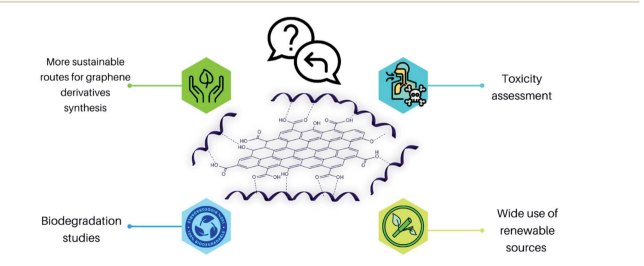


Fig. 12 Challenges in the field of graphene-based polymeric nanocomposites.

alterations in membranes. The impact observed in zebrafish was from inhibition of embryo growth to cardiac development.<sup>182</sup>

Graphene and its derivatives have excellent properties such as good dispersibility and stability in human physiological environments, making them promising for application in biomedicine, drug delivery systems, and food packaging. However, studies of GD toxicity in human biological organisms are not conclusive. Yang's group (2013) performed biodistribution and toxicology nanographene oxide analyses in mice, by oral and intraperitoneal administration. The researchers noted that the GDs were not adsorbed by the organs but were easily excreted. Despite this, they emphasize that toxicity will depend on the surface coating, size, and routes of administration.<sup>183</sup> Liu et al. (2012) conducted a study of size and dose on the biodistribution of GO in mice. The results showed that sizes between 1 and 5 µm accumulate in the lungs and between 110 and 500 nm in the liver. Therefore, they concluded that GO is not suitable for human use.184,185

On the other hand, some research shows that GD effectively fights cancer cells and is safe for healthy cells. The administered dose influences the cytotoxicity and apoptosis of human cells; that is, the use of small doses ( $<50 \text{ mg L}^{-1}$ ) of these materials proved to be safe.<sup>186–188</sup> In addition, Manikandan *et al.* (2020) considered that PHB/graphene nanocomposites have a negligible cytotoxic effect.<sup>1</sup> Thus, the toxicity of GD in human cells still needs thorough research, as there are few publications on genotoxicity tests and lots of contradictory information.

One of the biggest challenges in this area is producing Gr, GO and rGO by greener processes and on an industrial scale to formulate polymeric nanocomposites that are increasingly less toxic and aggressive to the environment through greener and more sustainable routes, Fig. 12.

### **Final considerations**

The development of novel packaging from polymeric nanocomposites based on graphene opens several possibilities for the food packaging industry and reduces production costs. It also brings lightness and higher resistance to these novel packaging resulting in numerous applications. The Gr, GO, and rGO application in nanocomposites significantly improves the packaging's thermal, mechanical, and barrier properties, reducing waste and increasing food safety. The more significant interest in using renewable polymers in the novel polymeric nanocomposites creation with the use of origin natural and biodegradable polymeric matrices and a smaller number of investigations using polymers is also noticeable of fossil origin, showing the concern with the environment. In general, studies are still lacking on the application of graphene nanocomposite packaging in wet foods and also on its toxicity and biodegradability, suggesting that further research in this regard should be carried out to ensure the safety and efficiency application of graphene and graphene-based in the formulation of new food packaging.

## Author contributions

The authors confirm contribution to the paper as follows: conceptualization: Vinicius Rossa and Thiago M. Lima. investigation: Vinicius Rossa, Luanne E. Monteiro, Sancler C. Vasconcelos and Vinicius G. Madriaga. Data curation: Vinicius Rossa, Luanne E. Monteiro, Sancler da Costa Vasconcelos, Vinicius G. Madriaga, Eric Thomas Tai Shimabukuro, Anna Paula Carvalho, Vitor Francisco Ferreira and Thiago M. Lima. Writing - original draft preparation: Vinicius Rossa, Luanne E. Monteiro, Sancler C. Vasconcelos, Anna Paula Carvalho, Sibele Berenice Castella Pergher and Thiago M. Lima. Visualization: Thiago M. Lima, Vinicius Rossa, Vinicius G. Madriaga, Sancler C. Vasconcelos, Sibele Berenice Castellã Pergher and Thiago M. Lima. Writing - review & editing: Thiago M. Lima, Vinicius Rossa, Sibele Berenice Castellã Pergher, Fernando de Carvalho da Silva, Vitor Francisco Ferreira, Anna Paula Carvalho and Carlos Adam Conte Junior. Supervision: Thiago M. Lima and Vinicius Rossa.

# Conflicts of interest

There are no conflicts to declare.

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