A critical review on the production and application of graphene and graphene-based materials in anti-corrosion coatings

Bohdan Kulyk, Maria A. Freitas, Nuno F. Santos, Farzin Mohseni, Alexandre F. Carvalho, Kiryl Yasakau, António J. S. Fernandes, Adriana Bernardes, Bruno Figueiredo, Rui Silva, João Tedim & Florinda M. Costa

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Department of Physics, i3N, University of Aveiro, Aveiro, Portugal; CICECO, Department of Materials and Ceramic Engineering, University of Aveiro, Aveiro, Portugal; Graphenest, Aveiro, Portugal

ABSTRACT
Among the many potential applications of graphene and graphene-based materials, their use as protective films or as additives in coatings for corrosion protection has seen an increased level of interest in the last decade. Much of this interest is motivated by the need to implement additional functionalities, to enhance anti-corrosion performance and to ultimately extend the service life of metallic structures. Pristine graphene films, with their impermeable nature allied to flexibility and mechanical strength, appear as particularly attractive candidates for barrier films against corrosive agents, while graphene-based materials such as graphene oxide and reduced graphene oxide offer a wide range of opportunities for their dispersion in polymeric matrices for composite anti-corrosive coatings. Simultaneously, considerable progress in the development of scalable graphene and graphene-based materials production techniques has been made during the last several years. Currently, a broad range of graphene materials with different morphologies and properties is available, making the need for an adequate fit between the production method and the desired application even more evident. This review article aims to give the reader a general overview of the recent trends in both the production of graphene and graphene-based materials, and their implementation in different anti-corrosion solutions. Moreover, the present work provides a critical look on this subject, highlighting the areas in need of further exploration.

Abbreviations: 2D: two-dimensional; CVD: chemical vapor deposition; DMF: dimethylformamide; DMSO: dimethylsulfoxide; EIS: electrochemical impedance spectroscopy; EPD: electrophoretic deposition; FLG: few-layer graphene; G-NP: graphene-nanoparticle composite; GO: graphene oxide; HAp: hydroxyapatite; HOPG: highly-oriented pyrolithic graphite; LIG: laser induced graphene; NP: nanoparticle; NMP: N-methylpyrrolidone; PANI: polyaniline; PMMA: poly(methyl methacrylate); rGO: reduced graphene oxide; SEM: scanning electron microscopy; SLG: single-layer graphene; XPS: X-ray photoelectron spectroscopy; ZRC: zinc rich coating

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CONTACT Bohdan Kulyka bkulyk@ua.pt
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1. Introduction

With the discovery of graphene in 2004 by Geim and Novoselov\cite{1} a whole new field in materials science, chemistry and physics has opened. Seventeen years later we are approaching a time when the industry can provide this material in quality and quantity required for its application. Graphene is known for its outstanding electron mobility,\cite{2,3} highest thermal conductivity of all materials, on par with diamond and carbon nanotubes (CNTs) and 12 times that of copper,\cite{4-6} a mechanical breaking strength of 42 N/m (311 times higher than that of steel), and a Young’s modulus of 1.0 ± 0.1 TPa.\cite{7}

The application of graphene as a functional layer or as an additive in coatings for corrosion protection has been attracting considerable interest in the last decade. The main driving force behind this interest relies on the need to implement additional functionalities, to enhance coating performance and thus extend the service life of metallic structures under extreme conditions.\cite{8,9} Graphene’s impermeability to gases\cite{10} provided the first indication of it being capable of acting as a physical barrier to corrosive species. Moreover, its chemical inertness with respect to oxidative etching up to temperatures of 400 °C,\cite{11} along with the promise of an extremely lightweight coating that does not significantly affect the underlying material’s optical properties, have further motivated the research efforts into graphene’s application in corrosion protection.

Several recent reviews have described the breadth of research works in this area, thereby supporting the fact that graphene and graphene-based materials can and will play a central role in the protection of metallic substrates.\cite{12-14} However, it must be said that the remarkable figures ascribed to graphene are commonly (and incorrectly) presented in the works related to any graphene-based material, when in reality they are only applicable to the pristine single-layer graphene (SLG). As a result, the main aim of the present review is to render a critical look on the implementation of graphene and graphene-based materials in the field of protective coatings, combining a description of the different production methods of such materials with an overview of the recent trends in their implementation in corrosion protection, something which, to the best of our knowledge, has not yet been done.

As it will be made clear throughout this review, the range of properties of the graphene derivatives obtained by different processes is so wide that their grouping under the single word “graphene” would be scientifically inaccurate. Therefore, we will opt to follow the nomenclature for the classification of these materials according to the ISO/TS 80004-13:2017, namely:

1. Graphene, graphene layer, single-layer graphene (SLG), monolayer graphene: single layer of carbon atoms with each atom bound to three neighbors in a honeycomb structure.
2. Bilayer graphene (2LG): two-dimensional material consisting of two well-defined stacked graphene layers.
3. Few-layer graphene (FLG): two-dimensional material consisting of three to ten well-defined stacked graphene layers.
4. Graphene oxide (GO): chemically modified graphene prepared by oxidation and exfoliation of graphite, causing extensive oxidative modification of the basal plane. Graphene oxide is a single-layer material with high oxygen content, typically characterized by C/O atomic ratios of approximately 2.0, depending on the method of synthesis.
5. Reduced graphene oxide (rGO): reduced oxygen content form of graphene oxide.

Additionally, based on these definitions, single-layer, bilayer or few-layer graphene films or flakes will be generally grouped under the term “graphene,” while any other two-dimensional carbon materials, such as GO or rGO, will be referred to as “graphene-based materials.”

This review, outlined in Figure 1, is divided into four main sections. Firstly (section 2), different production methods are presented, giving particular
attention to methods to obtain high-quality graphene, high-throughput processes to produce graphene and methods to obtain graphene oxide and reduced graphene oxide. The following two sections are devoted to anti-corrosion applications, with works associated with graphene and graphene-based films (section 3) and composites using graphene (section 4) being discussed separately. The review ends with a critical overview of the field (section 5), describing not only the advantages but also challenges associated with implementation of graphene and graphene-based materials for corrosion protection.

2. Production methods of graphene and graphene-based materials

2.1. Conventional high crystalline quality graphene production methods

The first successful method to isolate graphene was through mechanical exfoliation of graphite by successive peelings using scotch tape. This method produces high-quality, single crystal, one atom thick graphene flakes. However, its throughput is very low, and its upscaling is unreasonable.

One of the earliest alternative synthesis methods that emerged was that of epitaxial graphene obtained by graphitization of silicon carbide (SiC). By heating up single-crystal SiC in ultrahigh vacuum conditions (or even at atmospheric pressure), the Si atoms are sublimated, with the remaining C atoms undergoing graphitization. This process can occur on both the Si-terminated face and on the C-terminated one. In the former case, a C-rich buffer layer covalently bonded to the substrate is present underneath the formed graphene, while in the latter case the interaction with the substrate is much weaker.

With the emergence of this synthesis approach, it was quickly established that the obtained graphene is of high quality, with charge carrier mobilities up to 27,000 cm² V⁻¹ s⁻¹ reported as early as 2006, for graphene obtained on the C-terminated face. However, the main disadvantage of graphene synthesis from SiC remains the high price of the substrates, as well as the associated limitations in terms of sizes and difficulties in micromachining of the obtained material.

Alternatively, chemical vapor deposition (CVD) has been one of the most prominent graphene synthesis techniques since the early reports showing surface segregation of carbon atoms into graphene domains and continuous layers from transition metals. Yu et al., for example, were able to form high-quality graphene layers by exposing a Ni foil to a mixture of CH₄, H₂, and Ar at 1000 °C (at atmospheric pressure). The process was explained on the basis of hydrocarbon gas decomposition, followed by carbon atom diffusion into the metal foil. A controlled cooling of the substrate then leads to carbon segregation, forming graphene layers at the surface. Moreover, the authors showed that this graphene can be transferred onto insulating substrates using a support layer.
rubber) applied to the grown film, followed by the etching of the metal and the application of the graphene/support stack onto the desired substrate. Shortly after, copper came forth as the preferred growth substrate, thanks to its low carbon solubility and catalytic effect on the hydrocarbon precursor breakdown, allowing to grow continuous SLG films (Figure 2(a)). Typical CVD graphene synthesis on copper is conducted at temperatures close to the latter’s melting point. However, low-temperature (300–400 °C) growth has also been demonstrated, using microwave plasma to assist in the decomposition of the hydrocarbon precursors. Currently, the CVD method allows to obtain graphene domains exceeding millimetric sizes (Figure 2(b)) and arbitrarily large single-crystal films, as well as graphene with charge carrier mobilities as high as 350,000 cm² V⁻¹ s⁻¹, with growth at both low and atmospheric pressures possible.

Here, it should be pointed out that, in many cases, graphene grown epitaxially or using a catalyst must be transferred onto a different substrate where it can be processed or incorporated directly for the desired application. An example of this is the transfer of graphene onto insulating substrates for lithographic fabrication of electronic devices, or onto transparent substrates for optical applications. In the case of CVD-grown graphene on copper, the copper can either be etched away, or the graphene can be separated from the copper. The latter can be done electrochemically, by forming hydrogen gas bubbles, through hydrolysis, between graphene and copper, exfoliating the former from the latter (Figure 2(c)). Alternatively, water molecules can be intercalated between the graphene and the substrate, weakening their interaction and allowing to pick up the sample using a stamp. Either way, in most cases, a supporting material must be used in order to avoid the tearing or any other damage to the graphene film after its separation from the copper. The most commonly used supporting material is poly(methyl methacrylate) (PMMA), which is usually spin-coated on the as-grown graphene at the beginning of the transfer process. However, the removal of the PMMA support layer after the transfer is complete, which is typically done by placing the transferred sample in acetone,
leaves behind PMMA residues which are known to be difficult to eliminate completely.\textsuperscript{[33,34]} These residues influence graphene’s properties, mainly by giving rise to a p-type doping. Some solutions to this problem have been reported in the literature, namely by means of high-temperature annealings\textsuperscript{[34]} or plasma treatments.\textsuperscript{[35]} Other approaches seek to avoid PMMA altogether by exploring other supporting materials that can be removed without leaving any residues.\textsuperscript{[36–39]}

Independently of the production method, high crystalline quality single-layer graphene can be characterized by narrow 2D and G characteristic Raman peaks, with a high ratio of the respective intensities, as well as by the absence (or very low intensity) of the characteristic D peak associated with structural defects (Figure 2(d)).\textsuperscript{[40]}

2.2. High-throughput graphene production methods

For many commercial applications, a high production throughput and economic viability are still the main requirements in terms of the preferred synthesis approach. After the original mechanical exfoliation of graphite using scotch tape, other methods of exfoliation were developed trying to follow a similar approach. The main ones are briefly reviewed below.

2.2.1. Ball milling exfoliation

Exfoliation of graphite by ball milling can be accomplished making use of either planetary ball mills\textsuperscript{[41–48]} (in dry or wet conditions) or stirred ball mills.\textsuperscript{[49,50]}

In a ball milling process, the precursor material is loaded, together with the milling balls, inside the milling chamber, followed by a continuous rolling of the balls along the chamber’s walls. This rolling action transfers energy to the precursor both through shear stresses due to the friction effect of the balls rolling along the chamber’s walls and through direct impact of the balls on the material to be milled (Figure 3(a)). Different types of equipment can exploit this general principle, such as planetary mills, vibratory mills, and attritor mills, among others.\textsuperscript{[52]} The motion of the grinding balls is typically caused by inertial forces due to the movement of the milling chamber, or, in the case of the attritor mill, due to the rotation of the impeller arms inside the chamber. The main parameters that can be controlled are thus milling speed, time, grinding medium, ball-to-precursor ratio, temperature, and milling atmosphere.\textsuperscript{[52]}

For the ball milling production of graphene, the most commonly used precursor is graphite, although
helical ribbon carbon nanofibers, for instance, have also been exfoliated by this technique. With graphite, the milling itself may not be sufficient for successful graphene exfoliation, as it has been shown to result in an amorphization of graphite’s crystalline structure, due to the damage caused by the milling balls, which distorts the aromatic rings of the graphite. This damage has been observed by both X-ray diffraction and Raman spectroscopy after 15 to 20 h of milling.

In one of the earliest examples of graphene production by ball milling, Zhao et al. milled 30 to 80 nm thick graphite nanoplatelets for 30 h in dimethylformamide (DMF), using a planetary mill. The shear forces induced by the milling process, along with the DMF-graphene interaction, allowed to exfoliate the graphite nanoplatelets into graphene sheets with up to three layers, as reported by the authors, while the low rotation speed of the rotating tray allowed to have less in-plane stress capable of damaging the graphite flakes. Other organic solvents have also been successfully employed for graphite exfoliation into graphene by ball milling, with the importance of a match between the surface tension of the solvent and that of graphene having been highlighted. Sodium dodecyl sulfate has also been used as a surfactant for ball milled graphene flakes, as demonstrated by Knieke et al. using a stirred media mill. As an alternative, naphthol polyoxyethylene ether (NPE), a more environmentally-friendly and more easily removable surfactant, has also been used, resulting in graphene flakes with fewer than 10 layers.

A systematic study of the influence of the process conditions and parameters in wet ball milling of graphite has been performed by Damm et al. The authors, besides stating that stirred media mills allow to exfoliate graphite in a relatively shorter time (few hours instead of few tens of hours) and with better temperature control than planetary mills, point out that the Hansen solubility parameters of the used solvents are some of the main factors determining their suitability for graphite exfoliation into graphene, as previously shown by Coleman’s group. More specifically, the parameter for the disperse interaction, \( \delta_D \), should have a value of around 18 MPa\(^1\) [N/A]\(^2\) (Figure 3(b)), as is the case for N-methylpyrrolidone (NMP) and cyclohexanone (CHO). The viscosity of the solvent also plays an important role, with the more viscous ones facilitating the energy transfer from the grinding balls to the graphite while attenuating the direct impact of the balls, in order to prevent in-plane damage. In summary, wet ball milling can be attractive for graphene production. However, it should be pointed out that, unfortunately, wet ball milling often requires additional purification steps to remove the solvents.

As a way to avoid the need to remove surfactants and solvents from the obtained graphene dispersions, as well as to avoid the functionalization of the graphene sheets, which can be detrimental to its properties, Liu et al. proposed a dry ball milling graphene production method. In a process taking up to 4 h, graphite was milled in the presence of ammonia borane \((\text{NH}_3\text{BH}_3)\). The authors claimed that ammonia borane weakens the Van de Waals forces between the graphene layers in graphite, by interacting strongly with the latter. This results in graphene flakes of 5 layers or fewer. However, a significant degree of oxidation was present, as shown via X-ray photoelectron spectroscopy (XPS), and the presented Raman and TEM analysis is insufficient to support such claims. Triazine exfoliating agents have also shown great promise in the context of graphite exfoliation by ball milling. Here, melamine was identified as having the best performance, due to its aromatic nucleus capable of interfacing with graphene sheets by \( \pi-\pi \) interactions, as well as due to its ability to form extended two-dimensional (2D) networks through hydrogen bonding, which lead to multipoint interactions with graphene sheets, promoting exfoliation and stabilization. Dry ball milling of graphite in the presence of certain gases and dry ice has been shown to permit the production of edge-functionalized graphene flakes that can be dispersed in several solvents, including pure water. In other work, a planetary mill was used for 24 h to produce edge-carboxylated graphene from graphite precursor using dry ice. The production of FLG was accomplished, but the structural quality of the products is questionable, and a significant degree of oxidation was observed. After thermal reduction, the Raman spectra show severe defect concentration, broadened vibrational modes and a near absence of 2D band, so that the product can hardly be denominated as graphene. In summary, the main advantage of the dry-milling technique resides in its simplicity and in the possibility of obtaining graphene without contaminations. However, prolonged process times (sometimes as long as 30 h), limit its throughput.

Despite the demonstrated capability of producing graphene flakes, ball milling presents some obvious disadvantages. It is a scalable process only limited by the size of the milling vessel, but it is apparent that the high degree of structural defects, including oxidation, as well as the inherent functionalization, in
many cases, restricts the spectrum of application of these graphene products. The bare impact of the balls fragments the graphene flakes into smaller flakes and induces severe structural defects to the graphene lattice, especially in basal planes.\cite{41,42,60} This can be minimized by slowing the rotation speed at the expense of the process time, which can extend to more than one day.\cite{45,56} Ball milling is, thus, time consuming and many thick flakes are present in the final products, despite some single- and few-layer graphene.\cite{52,53,54,55}

2.2.2. Sonication

In sonication, ultrasonic waves produce localized compressions and rarefactions in the liquid medium, forming vacuum cavities which then collapse, generating high pressure jets. These jets are capable of peeling off graphene layers from graphite, as well as weakening the Van der Waals interaction between the layers. This facilitates the intercalation of solvent molecules between graphitic planes, allowing for further exfoliation and consequent stabilization of the dispersion. This is exemplified by 1-pyrenemethylamine (Py-NH$_2$) and 1,3,6,8-pyrenetetrasulfonic acid (Py-SO$_3$),\cite{70} as well as 1-pyrenecarboxylic acid (PCA)\cite{71} and 9-anthracene carboxylic acid (9-ACA).\cite{72} Low boiling-point solvents, such as chloroform, acetone and isopropanol, have been explored by O’Neill and coworkers.\cite{73} Isopropanol, in particular, showed good results, both in terms of the number of layers obtained (less than 5), and in terms of stability (with 96% of graphene remaining stably dispersed for as long as 200 h after the initial dispersion). The authors claim that the Hansen solubility parameters of the solvents, particularly the dispersive parameter, $\delta_D$, dictate the quality of the success of the dispersion, having to match those of graphene ($\delta_D = 18 \text{ MPa}^{1/2}$, $\delta_p = 9.3 \text{ MPa}^{1/2}$, and $\delta_H = 7.7 \text{ MPa}^{1/2}$) for better results.

Despite not being suitable for graphene dispersion on its own, water has been employed for sonication exfoliation in conjunction with surfactants such as sodium dodecylbenzene sulfonate (SDBS).\cite{74} In this example, as much as 40% of the obtained dispersion have been reported as having less than 5 layers of graphene, with 3% of obtained flakes being single-layer.
| Concentration or yield | Thickness or number of layers | Precursor | Reagents | Process time and setup | Additional steps | Ref. | Year |
|------------------------|-------------------------------|-----------|----------|------------------------|-----------------|------|------|
| >0.1 mg/mL (after solvent dispersion), edge-functionalized | — | Natural graphite | H₂/dry ice/SO₃ (each for different edge functionalization) | 48 h, planetary mill, 500 rpm | Soxhlet extraction, freeze drying, dispersion in polar solvents | [42] | 2013 |
| ~0.9 mg/mL (and up to 6 mg/mL by recycling the sediment) | As low as 1 layer | Graphite flakes | SDS, water | Up to 12 h, planetary mill, 100 rpm | Centrifugation, sonication (2 h, 80 W), only the supernatant used | [43] | 2012 |
| 2.9 x 10⁻⁴ mg/mL | Multilayer | Graphite powder | Kerosene/2-ethylhexanol | 60 h, planetary mill, 400 rpm | Centrifugation, heat-treatment | [45] | 2017 |
| 35.16 g (starting from 25 g of graphite and 20 g of dry ice, edge-functionalized) | 4–5 layers | Graphite | Dry ice | 24 h, planetary mill, 150 rpm | Centrifugation, drying | [46] | 2019 |
| ≤5 layers | ~25% 1 layer | Graphite powder | NH₃BH₃ | 4 h, planetary mill, 150 rpm, in Ar atmosphere | Dispersion in ethanol, centrifugation | [48] | 2013 |
| 0.43 mg/mL | Mostly 3–4 layers | Isostatic graphite | N-methylpyrrolidone (NMP)/cyclohexanone (CHO)/i-propanol (IPA) | 1 h, stirred media mill, 1000 rpm | Centrifugation | [49] | 2015 |
| 25 mg/mL | Mostly ≤5 layers | Synthetic graphite | Water, sodium dodecyl sulfate (SDS) | 3–5 h, stirred media mill, 1500 rpm | Centrifugation | [50] | 2010 |
| ≤3 layers | Graphite nanosheets (30–80 nm thick) | N,N-dimethylformamide (DMF) | 30 h, planetary mill, 300 rpm | Centrifugation, ethanol wash, only supernatant used | Sedimentation, centrifugation | [55] | 2010 |
| 0.097 mg/mL (for DMF) | As low as 1 layer | Graphite nanosheets (30–80 nm thick, from expanded graphite) | Ethanol/formamide/acetonate/tetrahydrofuran (THF)/tetramethylurea (TMU)/DMF/NMP | 30 h, planetary mill, 300 rpm | | [56] | 2010 |
| 0.5–1.2 mg/mL (~10 wt% of the initial graphite) | Mostly <10 layers (as low as 1 layer) | Graphite powder | Naphthol polyoxyethylene ether (NPE) | 12 h, planetary mill, 400 rpm | Transfer to water, centrifugation, vacuum filtering, ethanol rinsing, water redispersion | [57] | 2010 |
| — | — | Graphite powder | NaCl | 2 h or 5 h, planetary ball mill, 350 rpm, in Ar atmosphere (0.4 MPa) | Ultrasonication in water, centrifugation, vacuum drying | [61] | 2017 |
| 0.37 mg/mL (for melamine) | — | Graphite powder | 2,4,6-triamino-1,3,5-triazine (melamine), 2,4,6-tri(N,N-diethylamino)-1,3,5-triazine (TNE-T), 2,4,6-trihydroxy-1,3,5-triazine (TOH-T), 2,4,6-trimethoxy-1,3,5-triazine (TOC-T), 2,4,6-trihydroxybenzene (TOH-B) | 30 min, planetary mill, 100 rpm, in air atmosphere | Redispersion in water or DMF, sonication (1 min), sedimentation | [59] | 2014 |
| — | As low as 2 layers | Graphite powder | Na₂SO₄ | 24 h, 150 rpm | Water wash, filtering, drying (90°C, 3 h) | [62] | 2014 |
graphene. As an alternative to SDBS, sodium deoxycholate (SDC) has also been used.[75]

Guardia and coworkers compared a wide range of different aqueous surfactants for bath sonication exfoliation of graphite.[76] Among these, the commercial nonionic surfactant Pluronic P-123 was identified as being particularly effective for the preparation of graphene dispersions, allowing to achieve concentrations as high as 1.5 mg/mL after 5 h of sonication. Tween 80 gave similarly good results. Dispersions obtained using P-123 were identified, by AFM profiles, to mostly have no more than 5 graphene layers, with 10% to 15% of flakes being single-layer graphene.

Nuvoli et al.[77] dispersed graphite, by bath sonication, in an ionic liquid (1-hexyl-3-methylimidazolium hexafluorophosphate, HMIH). This approach yielded graphene flake concentrations of 5.33 mg/mL, after 24 h of sonication and using an optimized concentration of the starting material (5.0 wt% of graphite flakes in HMIH).

As an alternative to bath sonication, horn ultrasonication has also been employed to exfoliate naturally occurring graphite flakes in the presence of a sodium cholate amphiphilic surfactant.[78] Supplemented by density gradient ultracentrifugation, this approach yielded graphene flake concentrations of 5.33 mg/mL, after 24 h of sonication and using an optimized concentration of the starting material (5.0 wt% of graphite flakes in HMIH).

In summary, sonication is an attractive technique for liquid-phase exfoliation, but, much like the other liquid-phase exfoliation approaches, it relies on the choice of an appropriate dispersant or stabilizing agent. The obtained yields are relatively low, and the long processing times, with continuous energy input, are a drawback of sonication exfoliation. Table 2 presents the conditions employed in a number of works on graphene production by sonication.

2.2.3. Shear stress exfoliation
Shear stress exfoliation of graphite can be accomplished in liquids using different types of apparatus, relying purely on fluid dynamics considerations in laminar or turbulent flow regimes. In particular, some authors[91,92] employed a rotating tube at sufficiently high rotating speeds to produce vortex fluidic films (e.g., 7000 rpm),[91] which are capable of producing enough shear forces to overcome Van der Waals interactions. The graphite precursors are mixed with organic or aqueous media and exfoliation takes place at the tube wall via combined lifting and slippage of graphene sheets.[58,91] The inclination of the tube was found to play a vital role in exfoliation, inducing
turbulence via combined centrifugal and gravitational interactions, which are crucial for the lifting/slippage process. Centrifugal forces hold the graphite precursors against the tube wall and allow for the slippage of graphene sheets. Despite the principle being successfully demonstrated for the production of good quality graphene, the shear stresses capable of graphene exfoliation are localized only at the tube surface, severely limiting the yield rate. This means that very high surface area fast rotating tubes are needed to improve the yield to satisfactory levels, which poses many technological challenges at increased costs. It was thus apparent that higher energy shear stress systems were needed to obtain graphene in a faster way and to increase the production yield. In this sense, it was found that pressurized fluids containing graphite precursors inserted in constriction channels also lead to the successful exfoliation of graphene and other 2D materials via fluid dynamics mechanisms (Figure 4(a)).[93–96] This type of exfoliation was the subject of theoretical simulations showing several mechanisms contributing to exfoliation besides shear stresses, namely collision, cavitation, and pressure release, the latter two mechanisms exerting normal forces aiding exfoliation. Among those producing shear forces are the turbulence-induced (Reynolds) stresses, velocity gradient stresses, and inter-flake collisions. Remarkable yields of 15% of mainly FLG with little oxidation have been demonstrated using high pressure pumps and closed circulation loops.[96] Interestingly, although constriction tubes are usually constructed in a straight geometry, recent theoretical simulations[95] have shown that helical tubes allow higher shear stresses, and thus more efficient exfoliation, despite turbulence being present in both cases.

Whilst most of the previous strategies use compressed fluids to obtain the desired shear rates, an alternative approach using compressed air-driven graphite suspensions through a tube was proposed in order to decrease the defects in the produced graphene because of the milder conditions compared to pressurized fluid approaches. An extreme shear rate of up to $3.3 \times 10^7 \text{ s}^{-1}$ was inferred at working pressures as low as 0.5 MPa.[97] Nevertheless, the calculated yield is still low, about 1.5%. The low yield is likely due to the fact that the extreme shear rates were found to be concentrated at the edges of the tube at the outlet. Indeed, by smoothing those edges the shear rates decreased by 3 orders of magnitude, barely above $\sim 10^4 \text{ s}^{-1}$. Authors claimed the production of unoxidized graphene, and that up to 62% and 35% of the flakes were monolayer and bilayer, respectively, via statistical analysis of AFM measurements. However, Raman measurements show a spectrum more attributable to FLG.

The most common and efficient approaches using fluid dynamics rely on rotor-stator[60,98–100] or rotating blades[101,102] immersed in a fluid containing the graphite precursor and suitable surfactants, commonly denominated by high-shear exfoliation (Figure 4(b)). The authors introducing the rotor-stator technique successfully produced highly concentrated graphene flake dispersions in sodium cholate and NMP, having no oxidation and low density of basal defects, judging by the low $I_{2D}/I_G$ Raman band ratio.[98] Exfoliation rates exceeding 5 g/hwere demonstrated using a 300 L system and exfoliating rates of 100 g/hwere extrapolated to $10^4 \text{ L high shear apparatus}$. Through several combinations of diameters and rotor speeds, it was noted that graphene flakes could be produced at relevant concentrations/yields if a minimum of $10^4 \text{ s}^{-1}$ shear rate is attained, independently if the regime is laminar or turbulent (discriminated by a Reynolds number of $10^3$). Hence, the exfoliation mechanism should be similar in both cases; nevertheless, turbulent regimes characterized by higher Reynolds numbers should in principle produce higher magnitude shear stresses, thus in principle allowing for enhanced yields.

An apparent disadvantage of high-shear exfoliation using rotor/stator is that the desired high shear rates are not uniformly distributed in all regions of the exfoliating media, being rather concentrated near the stator orifices and rotor-stator gap.[60,103] In this sense, high Reynolds numbers within the fluid are desirable because the developed turbulence homogenizes shear stresses within the exfoliation container. An obvious and easily accessible choice for this purpose are kitchen blenders equipped with rotating blades, as explored by some authors.[101,102,104] Commonly to ball stirring, shear force is not the only mechanism responsible for exfoliation in rotor-stator and rotating blade apparatus; it is aided by inter-flake collisions, as well as interactions with the rotor/blade such as flame edge collisions and cavitation[103] with a possible contribution from pressure differences induced by turbulence.[101] The high-speed rotating blade apparatus can produce enough shear stress for graphene exfoliation in all volume of the container, despite an obvious decrease of these stresses as the distance from the blade increases. Hence, using rotating blades seemingly ensures that shear stresses will dominate exfoliation at larger regions of the system.
Table 2. Summary of different sonication exfoliation approaches reported in the literature and the corresponding results.

| Concentration or yield | Thickness or number of layers | Precursor | Solvent | Surfactant | Sonication time | Sonication setup | Additional steps | Ref. Year |
|------------------------|--------------------------------|-----------|---------|------------|----------------|-----------------|------------------|-----------|
| 1.2 mg/mL              | Mostly 2–4 layers             | Natural graphite powder | NMP     | —          | 270 h          | 23 W, bath      | Centrifugation   | [64] 2010 |
| 3.18 mg/mL             | 28% monolayers                | Graphite powder | NMP     | —          | 30 min         | Bath            | Centrifugation   | [63] 2008 |
| 0.5 mg/mL              |                                | Graphite flakes | Benzylamine | —          | 2–4 h          | Bath            | Centrifugation   | [65] 2010 |
| 0.04 mg/mL             | Mostly <10 nm                 | Graphite flakes | NMP, progressively replaced with ethanol | —          | >2 h           | Bath            | Several centrifugation, solvent exchange and sonication steps | [66] 2010 |
| 0.02 mg/mL             | Mostly <5 layers              | Synthetic graphite | ODCB   | —          | 30 min         | Horn            | Prior homogenization in a high-shear mixer, centrifugation | [67] 2009 |
| 0.18 mg/mL             | 26% monolayers, 22% bilayers  | Graphite flakes | Water   | SDC        | 3 h            | Bath            | Centrifugation   | [68] 2010 |
| 0.1 mg/mL              | 28% monolayers, 19% bilayers  | NMP        | —       | —          | 6–9 h          | —               | —                | [69] 2009 |
| 0.06 mg/mL             | —                              | Graphite powder | Hexafluorobenzene, octafluorotoluene, perfluorobenzonitrile, perfluoropyridine, or pyridine | —          | >2 h          | Bath            | Several centrifugation, solvent exchange and sonication steps | [70] 2010 |
| 0.1 mg/mL (for pentafluorobenzonitrile) | 0.5–1 nm on average | Graphite powder | Water | SDBS | 30 min | Bath | Centrifugation | [71] 2011 |
| 0.05 mg/mL             | ~3% monolayers, 40% with <5 layers | Graphite powder | Water | SDBS | 30 min | Bath | Centrifugation | [74] 2009 |
| 0.5 mg/mL              | ~60% monolayers               | Kish graphite | Water | SDC | 90 min | Bath | Centrifugation | [75] 2010 |
| 1 mg/mL                | ~10%–15% of monolayers        | Natural graphite powder | Water | —          | 2 h            | Bath            | Centrifugation   | [76] 2011 |
| 5.33 mg/mL             | 2 nm on average               | Graphite flakes | 1-hexyl-3-methylimidazolium hexafluorophosphate | —          | 24 h          | 0.55 kW, bath   | Centrifugation   | [77] 2011 |
| 0.09 mg/mL             | 80% with a thickness of ≤1.2 nm | Natural graphite flakes | Water | Sodium cholate | —          | 51–52 W, horn | Centrifugation, density gradient centrifugation | [78] 2009 |
| 1.92 mg/mL             | 2 nm on average               | Graphite powder | Water | Silk nanofiber | 1 h       | 195 W, 5 s/2 s working pulse, probe | Centrifugation, 20 kHz, 1 kW, pressurized reactor, horn | [79] 2012 |
| ~0.24 mg/mL            |                                | Natural graphite | Water | Ethylene glycol | 50 min | —                | —                | [80] 2014 |
|                         | ~1.4 nm on average            | Graphite powder | Deionized water | TPA | 2 h          | 70 W, probe    | —                | [81] 2009 |
when compared to stator-rotor arrangements. This means that high yield rates are feasible using these simple systems that are readily scalable to mass production.

A more recent approach makes use of Taylor vortex flows to promote high-shear stress exfoliation of graphene,[105] denominated as Taylor–Couette flow. The apparatus comprises two concentric cylinders that are able to rotate independently and a graphite dispersion in an appropriate fluid contained between them (Figure 4(c)). If the inner cylinder rotation speed reaches a critical value whilst the outer one is held at rest, this special type of flow pattern can form, resulting in steady state axisymmetric toroidal vortices. These vortices pose strong shear forces and pressure gradients to the graphite precursors, leading to FLG production at yields of up to 5%. The authors have claimed a low oxidation degree and a low defect concentration of graphene via XPS and Raman spectroscopy. Nevertheless, the method used for inferring oxidation is not unequivocal. More studies are needed on this type of exfoliation, namely on the quality and characteristics of the produced graphene and on the possible limitations of scaling up to industrial production.

Advantageously, no compound intercalating steps are needed for high shear exfoliation, such as in electrochemical (described below) or some ball milling exfoliation routes, despite some attempts to combine both, using electrochemical intercalation of tetrabutylammonium tetrafluoroborate, resulting in stable graphene dispersions in NMP with yields of up to 16%.[106] In addition, this type of shear exfoliation provides higher yields compared to ball milling because the graphite precursors move within the exfoliating media and can be continuously exposed to the shear forces, thus increasing exfoliation efficiency.[60]

As a result, exfoliation rates are enhanced and processing times are significantly decreased, typically down to a few hours or less. Additionally, final product concentrations of up to about 1 g/L were reported, e.g., in aqueous solutions of PVP and sodium cholate surfactant[107] or even in household dishwashing liquids at a rate of up to 0.15 g/h.[102] It is also significant that, for similar exfoliation energy density, the exfoliation yield of shear stress using fluid dynamics can be about one order of magnitude larger when compared to purely sonication routes.[58,60,91,98] Finally, an increase of the processing time in shear stress leads to a linearly proportional increase in graphene product formation,[98,108] whilst in sonication routes a saturation tendency is observed.[64,109] Unsurprisingly, high-shear
exfoliation is reported to provide the same amount of FLG as obtained via sonication routes in about one tenth to one hundredth of the time.\textsuperscript{107}

On the other hand, similarly to other liquid-phase exfoliation approaches, the graphene products by shear exfoliation generally suffer from size and thickness polydispersity,\textsuperscript{107} and require centrifugation for separating larger flakes. Additionally, only very recently the effect of the viscosity of the exfoliating media on the produced graphene characteristics and yield was more systematically addressed,\textsuperscript{110} despite some older studies using solvents/surfactants with different viscosities.\textsuperscript{103} This is an example demonstrating that many features of the process are still lacking optimization. Furthermore, it is important to check if high yields, quality, and fast production times of graphene flakes are maintained in very large containers. Moreover, similarly to other liquid-phase exfoliation techniques, it matters to demonstrate the production of stable graphene dispersions using nontoxic and cheaper solvents at similar figures of merit compared to strong organic solvents (e.g., NMP, DMSO, and DMF, among others), ionic liquids or common surfactant aqueous solutions (e.g., SDS, SDBS and PVP).\textsuperscript{99} Pure water would be the optimal solution, due to its nontoxicity and low cost, but this is hindered by the hydrophobic nature of graphene. In this sense,\textsuperscript{103} the use of IPA aqueous solutions yielding graphene dispersions of up to 0.27 g/Lin one hour high-shear processing has been reported, which tended to be stable for up to 25 days after initial dispersion. Nevertheless, the stability is found to be highly dependent on the size and thickness of the graphene flakes. Other types of nontoxic dispersants are being unveiled for liquid-phase exfoliation of graphene and other 2D materials, such as nontoxic, cost-effective nonionic polymers,\textsuperscript{107}

Figure 4. (a) Schematic illustration of the high shear stress exfoliation process in a constriction channel. (b) Schematic illustration of the high shear stress exfoliation process in a rotating blade system. (c) High shear stress exfoliation relying on Taylor–Couette flow.
or the so-called green dispersants (e.g., black tea, alkaline lignin, black liquor). The latter, for example, resulted in FLG dispersions of up to 10 g/L, via shear stress mixing, providing a yield of about 18% after 10 h of process time. More studies within these approaches are needed to pave the way toward the environmental-friendly and cost-effective industrial production of graphene flakes using shear stress exfoliation. Table 3 summarizes some of the shear stress exfoliation approaches reported in the literature.

### 2.2.4. Electrochemical exfoliation

The production of graphene via electrochemical exfoliation is based on the physical deformation of a graphitic working electrode immersed in an appropriate electrolyte, via the exchange of an electric current. The experimental setup also comprises a counter (auxiliary) electrode against which the potential of the working electrode is applied, ideally providing a large surface area and low electrical resistance. The counter electrode should also be chemically inert (e.g., platinum) in order to avoid the production of species that could interfere with the graphene exfoliation process, but some recent approaches rely on the exfoliation of both electrodes in order to improve the yield.

In terms of industrial production, electrochemical exfoliation of graphene provides several advantages over other exfoliation types such as sonication or purely chemical procedures. In general, it encompasses simple, economically viable processes requiring no extreme temperature/pressure conditions. Furthermore, it permits obtaining graphene with lateral sizes of tens of micrometers, yields larger than 50 wt% and quantities up to a few grams in relatively short production times (e.g., a few hours versus days in chemical exfoliation and sonication). It has been shown that he applied potential is a critical parameter, through which it is possible to tune not only the quantity of the produced graphene, but also its quality. For instance, it has a profound impact on the degree of oxidation of the products. Hence, a third (reference or pseudo-reference) electrode is often employed (Figure 5(a)), through which negligible current is withdrawn, providing a stable potential to accurately and stably control the potential applied to the working electrode. Finally, it is a versatile technique that can be combined with other approaches such as sonication, microwave processing, and shear stress to increase the yield, although this poses some restrictions in terms of production scale-up and industrialization. The main disadvantages are common to sonication and high shear stress exfoliation, such as the lack of widely available, cheap, and “green” solvents needed to efficiently perform exfoliation and properly disperse the products. In this sense, NaOH/H2O2/H2O solutions are promising in terms of exfoliation efficiency, but the products still need to be posteriorly stabilized in surfactants. Finally, electrochemical exfoliation also suffers from a high degree of size polydispersity, requiring posterior purification or separation.

The general exfoliation principle is that the applied voltage promotes the intercalation of ionic species present in the electrolyte in between the graphite layers, leading to physical deformation and an increase in the interlayer distance, thus weakening the Van der Walls forces and culminating with the separation of graphene layers (Figure 5(a)). Two main types of electrochemical exfoliation of graphene exist depending on whether the electrode is anodically or cathodically driven. In anodic (cathodic) exfoliation the graphite electrode is positively (negatively) charged, promoting the intercalation of anions (cations). Anodic exfoliation provides faster and higher-yield graphene exfoliation at lower applied potentials when compared to cathodic exfoliation. In general, anodic exfoliation makes use of a variety of aqueous or organic solutions or ionic liquids. In particular, the usage of both organic and inorganic salts as intercalating agents has been attracting great attention, and recent studies point out sodium pyrophosphate and sodium phosphate as better exfoliating agents than sodium citrate, the former greatly enhancing the yield and allowing to obtain graphene with a reduced number of layers, but also with increased defect content. However, organic salts such as sodium citrate are also promising due to better solubility in NMP compared to inorganic salts.

In many cases, intercalation and exfoliation is attained using a constant voltage, the magnitude of which varies greatly among the literature, mainly influenced by the electrolyte properties and the desired characteristics of the graphene (see Table 4). The intercalated ions can undergo a change into the gas state, which enhances the exfoliation process by increasing structural deformation, e.g., via the formation of O2/CO2 or propylene gas. In this sense, a commonly employed strategy is to use aqueous solutions of sulfates. Parvez et al. analyzed several sulfate compounds in the exfoliation process and found up to 85% of graphene having three or less layers. According to the authors,
Table 3. Summary of different shear stress exfoliation approaches reported in the literature and the corresponding results.

| Yield or yield rate | Flake size | Thickness or number of layers | C/O ratio | Raman ID/IG ratio | Precursor | Media | Process time and setup | Additional steps or conditions | Ref. | Year |
|---------------------|------------|-------------------------------|-----------|------------------|-----------|-------|-----------------------|---------------------------------|------|------|
| <1 wt%              | ~1 μm      | 0.7 to 1.3 nm                 | —         | —                | Graphite flakes (0.1 mg/mL) | NMP    | Vortex fluidic film, 7000 rpm, 30 min | —                               | [91] | 2012 |
| 5 wt%               | <~2 μm     | <1.5 nm                       | ~20 to ~27 | ~0.4             | Graphite flakes (1 mg/mL) | DI Water/IPA (1:1) | Pressurized fluid, 2 h | Centrifugation                  | [93] | 2014 |
| 594 g/h             | Mostly <1 μm<sup>2</sup> | <100 nm                       | —         | —                | Thermally expanded graphite flakes (20 mg/mL) | 2 wt% polyacrylic acid aqueous solution | Pressurized fluid, 10 passes | —                                | [95] | 2019 |
| 15 ±0.3 wt%         | <~2 μm     | <12 layers                    | ~25       | ~0.5 to ~0.6     | Graphite flakes (10 mg/mL) | NMP    | —                               | Centrifugation                  | [96] | 2018 |
| 1 wt%               | 0.5–2.6 μm | <1.5 nm                       | ~24       | 0.13             | Graphite powder (40 mg/mL) | NMP    | Pressurized gas-driven, 90 min | Centrifugation                  | [97] | 2019 |
| 5.3 g/h             | Mostly <1 μm | Mostly <10 layers             | —         | 0.17 to 0.37     | Graphite flakes (50 mg/mL) | NMP, sodium cholate solution | Rotor-stator, 20 min | Centrifugation                  | [98] | 2014 |
| 7.3 wt%             | ~0.1 μm<sup>2</sup> | ~1.5 nm (~20% < 1 nm)         | ~21       | ~0.12            | Graphite flakes (3 mg/mL) | DMF    | Kitchen blender, 8 h          | Centrifugation                  | [101] | 2016 |
| 1 wt%, ~0.15 g/h    | Mostly <1 μm | <10 layers                    | —         | 0.3–0.7          | Graphite flakes (50 mg/mL) | Household dishwashing | Kitchen blender, 4 h | Centrifugation                  | [102] | 2014 |
| ~1 to 5 wt%         | Mostly <2 μm | Mostly <3 nm                  | —         | ~0.14            | Graphite powder (1 to 5 mg/mL) | NMP    | Taylor–Couette flow, 1 h | Centrifugation                  | [105] | 2016 |
| 16.0 ±0.2 wt%       | <1.5 μm    | Mostly 4–6 layers             | Residual O content | 0.24 ± 0.10     | Graphite flakes (1 mg/mL) | NMP    | Electrochemical intercalation; Centrifugation | —                                | [106] | 2017 |
| —                   | Mostly <1 μm | Mostly <2 nm                  | —         | 0.14–0.18        | Graphite powder (10 mg/mL) | 40 vol% IPA aqueous solution | Rotor-stator (variable sizes), 1 h | Centrifugation                  | [103] | 2014 |
| —                   | 200 nm to 1 μm | 2 to 8 layers, average of 4 | ~4        | 0.17             | Graphite flakes, (~17 mg/mL) | Black tea solution | Kitchen blender, 15 min | Centrifugation                  | [104] | 2017 |
| —                   | Mostly <1 μm | Mostly <20 nm                 | —         | ~0.2 to ~0.3     | Graphite flakes (40 mg/mL) | PVP and sodium cholate aqueous solutions | Rotor-stator, 2 h | Centrifugation                  | [107] | 2017 |
| Up to 18 wt%        | <3.5 μm    | Mostly <5 nm                  | —         | —                | Graphite powder | Black liquor/water solution (2:1) | Rotor-stator, 0.5 to 10 h | Sonication; Centrifugation | [108] | 2018 |
| —                   | ~1 to 5 μm | 1 to 5 layers (qualitative measurement) | —         | 0.1              | Graphite flakes (50 mg/mL) | NMP with zeolite particles | Cappuccino mixer, 6 h | Centrifugation                  | [111] | 2019 |
| Up to 65 wt%        | <~10 μm    | <~3 nm                        | —         | —                | Mildly oxidized graphite (1 mg/mL) | Aqueous solution | Millstone, 45 min | Centrifugation                  | [112] | 2018 |
Table 4. Summary of different electrochemical exfoliation approaches reported in the literature and the corresponding results.

| Yield or yield rate | Flake size | Thickness or number of layers | C/O ratio | Raman I_D/I_G ratio | Electrode | Electrolyte | Applied signal | Complementary steps or conditions | Ref. | Year |
|--------------------|------------|--------------------------------|-----------|---------------------|-----------|-------------|---------------|----------------------------------|------|------|
| ~15 wt%            | <1 μm      | Mono and few layer             | <1        | Graphite rod        | 0.05 M and 0.1 M SDS 10 mg/mL copper phthalocyanine tetrasulfonic acid tetrasodium salt             | 5 V DC (anodic) | —                     |                                   | [130] | 2009 |
| 5 to 8 wt%         | <3 nm      | —                              | <1        | Graphite rod        | 0.1 M Poly(sodium-4-styrenesulfonate) 2.4 g of 98% H2SO4 in 100 mL of DI water plus 11 mL of 30% KOH Alternating 10 V (anodic) and ~10 V (cathodic) LSV0 or sinusoidal, 7 to 10 V (anodic) | Preliminary intercalation step at 2.5 V (anodic) | Posterior deintercalation at ~0.5 V (cathodic); Sonication | [138] | 2011 |
| 60 to 77 wt%       | <5 layers  | ~21                            | ~0.3 to ~0.4 | Graphite rod or HOPG | 0.1 M (NH4)2SO4 | 10 V DC (anodic) | —                     |                                   | [137] | 2014 |
| 65 wt%             | >30 μm     | 2–7 nm                         | ~5        | Graphite flakes     | 0.1 M (NH4)2SO4 | 10 V DC (anodic) | Physical compression of graphite; Shear mixing | [116] | 2018 |
| 2.2 g L⁻¹ h⁻¹     | 2–5 layers | ~8                             | 0.7       | Graphite rod        | Glycine in HSO4/H2O | 1 V and 3 V DC (anodic) | —                     | Sonication at 8000 rpm               | [139] | 2015 |
| 1.6 mg cm⁻² h⁻¹   | <10 μm     | 1–3 layers                     | ~1        | HOPG                | 0.1 M H2SO4 | 1 to 10 V DC (Anodic) | Shear stress field (~74,000 s⁻¹) | Several transition metal compound additives 1 M NaOH pretreatment | [122] | 2016 |
| ~15 wt%            | Mono and few layer | ~27                        | 0.05      | Graphite foil      | 0.5 M Na2SO4 | 20 V DC (anodic) | —                     |                                   | [140] | 2018 |
| 56 wt%             | ~2 μm      | 2–4 layers                     | 11        | Graphite foil       | 0.5 M H2SO4 | 10 V DC (Anodic) | 1 M NaOH pretreatment | Sonication or shear mixing | [141] | 2015 |
| ~15 wt%            | 0.2–0.6 μm | 1–2 nm                         | ~0.4      | Graphite foil, powder and flakes, HOPG | 0.1 M K2SO4 | 10 V DC (Anodic) | —                     | Sonication | [142] | 2015 |
| ~15 wt%            | 0.6 μm     | 2–3 nm                         | ~17       | Graphite foil       | Sodium halides (0.05 M NaCl, 0.1 M NaBr, 0.1 M NaI) | 10 V DC (anodic) | —                     | Sonication | [143] | 2017 |
| 42 ± 7 wt%         | Average of 12 μm | Average of 10 nm | ~14 | HOPG | 0.1 M (NH4)2SO4 | 12 V DC (Anodic) | Thermal treatments up to 800 °C | Preliminary 1 V DC intercalation step | [144] | 2016 |
| 90 wt%             | 10 to 30 μm | 85% <7 layers                  | 7.5       | Pre-expanded graphite | 10 M H2SO4 | 2 V DC (Anodic) | —                     | Preliminary 1 V DC intercalation step | [145] | 2014 |
| ~77 wt%            | 5–15 nm    | —                              | ~0.3      | Graphite foil      | 0.25 M H2SO4 plus 11 mM KOH (30%) 10 V DC (anodic) | 80 °C | —                     |                                   | [146] | 2015 |
| ~30 wt%            | 0.5–1 μm   | Mostly 2 layers                | ~1.8      | Graphite rod       | 0.2M (NH4)2SO4 | 10 V DC (Anodic) | 2 V DC preliminary step; sonication; thermal reduction | — | [147] | 2017 |
| 75 wt%             | Mostly 1–5 μm | Mostly 1–3                  | ~21       | Graphite sheet     | 0.1M TBA-HSO4, NaOH (pH adjustment) | 10 V AC, 0.1 Hz (both anodic and cathodic) | —                     |                                   | [114] | 2017 |
| wt% and g h⁻¹ | μm | Layer Count | Current Density | Solution/Reagents | Processing Conditions | Additional Info |
|----------------|-----|-------------|-----------------|-------------------|----------------------|-----------------|
| 25, 1.5 g h⁻¹ | 18 | Mostly 1 to 3 | ~26 | 0.2–0.54 | Graphite powder | 65 g/L H₂SO₄, varying amounts of CH₄N₂S, alternating pulses | [148] 2015 |
| 63 ± 4% (cathodic), 45 ± 3% (anodic) | ~5 layers | ~9 (anodic), ~23.5 (cathodic) | — | Graphite rod | 0.1 M of ionic liquids in acetonitrile | — |
| — | ~25 μm | Mono and bilayer | ~25 | 0.16 | Graphite foil | 1.0 M H₂SO₄ | [150] 2017 |
| 13 wt% | ~33–50 μm | 1–5 layers | ~30–50 | ~0.2 | Graphite foil | 0.1 M Na₂SO₄, 0.1 M NaCl in DMF | — |
| 25 wt% | Mostly 2–5 layers | ~36 | <0.05 | Graphite foil | Lithium chloride and/or triethylamine hydrochloride in dimethyl sulfoxide | Preliminary step at −1.7 V DC | [152] 2014 |
| 0.5–2 g h⁻¹ | Mostly 1–50 μm | <5 nm | ~12 | 90% <0.5 | Graphite/polyvinylidene fluoride pellets or rods | LSV, −1.7 to −5 V DC, plus step at −5 V DC (cathodic) | [118] 2011 |
| Up to 70 wt% | Mostly <5 μm | 70% <5 layers | — | <0.1 | HOPG and graphite powder | 30 mg/mL LiClO₄ in propylene carbonate | Thermal treatment via MW radiation; sonication | [115] 2019 |
| — | Few μm | Mono, few and multilayer | — | — | Graphite foil | 1.0 M HClO₄ | Thermal treatment at 800°C in Ar; Sonication | [115] 2019 |
| 85 wt% (cathodic) and 48 wt% (anodic) | 25 g h⁻¹ | 70% of 1–5 μm | ~18 (anodic), ~21 (cathodic) | 0.1 M TBAClO₄ in propylene carbonate | 5 V DC (both anodic and cathodic) | Thermal treatment at 800°C in Ar; Sonication | [115] 2019 |
| 80 wt% | 2–4 nm | ~8.6 | ~0.3 | Graphite rod | Molten LiOH | Constant 15 A current | [154] 2012 |
| — | 100–200 nm | ~2 nm | 11.5 | — | HOPG and graphite rod | 0.1 M TMA ClO₄, TEA BF₄ or TBA BF₄, NaCl, DMSO, and thionin acetate salt | Sonication (for HOPG) | [120] 2014 |
| — | ~3 nm | 12.5 | ~0.1 | Graphite rod | — | — | [121] 2013 |

a Similarly to other production routes, centrifugation is also used in almost all the works listed in this table and is excluded from this entry for practicality.

b Linear sweep voltammetry.
hydroxyl ions from water reduction attack preferentially the graphite edges and grain boundaries, where expansion starts promoting the intercalation of sulfate ions and H$_2$O. Afterwards, sulfate ions and water molecules undergo reduction and oxidation, leading to the production of SO$_2$ and O$_2$ that assist the exfoliation process.

The major obstacle often observed using bulky graphite electrodes lies in the inherent mechanism of exfoliation. Since the electrode is physically deformed during intercalation, it may start disintegrating, creating thick graphite flakes that are no longer electrochemically active by lack of electrical contact with the electrode. This is the main reason justifying the need for posterior thinning of the graphitic products via sonication.[116,129,137,155,156] A potential breakthrough to solve this problem whilst maintaining good scalability prospects is the use of physical compression of the graphite,[116] taking place within a permeable and compressible container. Authors have shown appreciable yields of 38%, which could theoretically be improved to 65%, but the large-scale production with this figure of merit is yet to be demonstrated.

The formation of thick flakes occurs especially when employing acid electrolytes, because of the higher anion intercalation rate in such conditions.[156] Moreover, the products often denote an important degree of overoxidation.[117,156,157] In this sense, the usage of inorganic salts at neutral pH[137] has been also proposed in order to control anion intercalation, seemingly allowing the production of good quality and thin exfoliated graphene. The addition of antioxidant species to the electrolyte also enhances carbon to oxygen content ratio.[140,158] Nevertheless, the major issue is that, in general, anodic-based processes using aqueous solutions at the potentials required for successful anion intercalation also lead to water hydrolysis. This culminates with low yields, severe oxidation

![Figure 5.](image-url)

(a) Schematic illustration of the electrochemical exfoliation process. WE, RE and CE labels correspond to the working, reference and counter electrodes. Photograph of a graphite rod (b) before and (c) after electrochemical exfoliation by tetraethylammonium intercalation. Adapted from ref. [120] under the Creative Commons CC-BY license. (d) Photograph and (e) SEM image of HOPG expanded after tetrabutylammonium intercalation. Adapted from ref. [120] under the Creative Commons CC-BY license.
defects and/or other undesired functionalization, the extent of which depends mainly on the applied potential and on the pH of the electrolyte.

An alternative approach relies on the application of multi-potential programs, as opposed to the traditional constant potential process. A better control in the intercalation process leading to the formation of thinner graphene flakes (e.g., <3 nm) was reported. Recently, Chen et al. employed different potential sweep profiles between 7 and 10 V, followed by deintercalation steps at −0.5 V, in 0.1 M (NH₄)₂SO₄ solutions. A sinusoidal profile more than doubled the average area of the products, and an increase of 9% up to 96% of graphene with less than 5 layers was also obtained. Without the −0.5 V step, that percentage decreased to about 60%. Authors claimed that the potential sweeps, especially sinusoidal ones, greatly reduced the oxidative breaking of the graphite basal C = C bonds and promoted SO₄²⁻ intercalation in comparison with fixed potential approaches, by equilibrating these two kinetically dissimilar processes. On the other hand, at −0.5 V the basal plane disruption and anion intercalation is hindered, and anion release from in between the graphite layers is promoted. This allows a more controlled interlayer expansion and exfoliation, yielding higher surface area graphene with fewer layers and less defects. Despite the merits, the final optimized products still denote a considerable degree of oxidation as shown via XPS.

Despite the above-mentioned efforts to control the anodic exfoliation, cathodically-driven exfoliation is conceptually preferable to obtain higher structural quality graphene, mainly because cathodic processes avoid the formation of graphene oxide. Cathodic exfoliation is, in general, achieved using organic solvents or ionic liquids, because the insertion of aqueous cations into graphite is a slow and inefficient process, limited at higher potentials by the hydrogen evolution reaction. Ionic liquids are an interesting approach because of their tunable miscibility and viscosity, recyclability, high electrical conductivity, large electrochemical windows and surface energies compatible with graphene.

On the other hand, Li ions have been known for decades to intercalate graphite, and this fact was used to promote cathodic exfoliation of graphene. Nevertheless, Li ion insertion is a slow, inhomogeneous process that limits the yield, in most cases also requiring post-sonication to obtain a satisfactory exfoliation of graphene. The sluggish kinetics can be sped up by increasing the applied voltage magnitude, such as −30 V versus 1 to 10 V in anodic exfoliation (although reports exist using higher potentials, see Table 4). Nevertheless, this comes at the expense of energy efficiency via competing decomposition of solvent cations, lack of control over the characteristics of the produced graphene, with increased defect content and unwanted contaminations. There are, however, other compounds that can be cathodically intercalated into graphite. Na⁺/DMSO complexes with the aid of thionin acetate salt in aqueous solutions were shown to be viable intercalants, but still post-sonication was necessary. Cathodic exfoliation at low potentials (−4 V to −5 V) using tetraalkylammonium salts has been shown to yield few layer graphene products with minimal oxygen content and without the need for any subsequent exfoliation steps, which is one of the most impressive results in the literature on cathodic exfoliation (Figure 5(b–e)). Interestingly, little to none graphite expansion is seen using Li-based electrolyte as a comparison.

Some authors have put efforts to combine both anodic and cathodic exfoliation of graphite electrodes in the same run, either through application of alternating positive/negative DC pulses or AC pulses, or employing electrolytes allowing both anionic and cationic intercalation, including ionic liquids. Contrarily to purely anodic or cathodic exfoliation, these strategies allow for exfoliation of both the working and counter (graphite) electrodes. Using the latter approach, graphene was produced in tetrabutylammonium perchlorate (TBAClO₄) in propylene carbonate, which provides a potential window large enough for electrochemical anion (ClO₄⁻) and cation (TBA⁺) intercalation at the graphite anode and cathode, respectively. A posterior thermal treatment enhances interlayer expansion via transition of the intercalated ions to gas state. The obtained yield is admirable, but extensive temperature treatments, post sonication and repeated wash by vacuum filtration and centrifugation were necessary, so that the process is hardly two-step. In addition, the influence of the sonication on the final product quality and yield is unclear.

AC signals (±10 V, 0.1 Hz) in 0.1 M tetra-n-butylammonium bisulfate (TBA-HSO₄) have been proposed to achieve simultaneous cathodic and anodic exfoliation. The authors claimed outstanding yields and the low defect density of the products was attributed to both the higher intrinsic quality of cathodic exfoliation and the reduction of anodic graphene via the AC signal. The mechanism involves the initial oxygen radicals formation during the anodic process, which
attack graphite grain boundaries and defects preferentially, opening gaps for the consequent intercalation of sulfate ions. The sulfate ions are reduced at the cathodic counterpart forming SO$_2$ gas bubbles, which expand the graphite and allow the bigger TBA$^+$ ions to intercalate, expanding the graphite even further. Interestingly, little exfoliation at the cathode was observed when using ammonium sulfate. However, in other work, exfoliation could be accomplished simultaneously at both graphite electrodes using alternating DC pulses of ± 10 V in ammonium sulfate-based aqueous solutions.[113] The pulses were shown to promote anion (SO$_4^{2-}$) and cation (NH$_4^+$) intercalation and bubbling, thereby increasing the exfoliation efficiency.

Despite the recent progress, further studies are needed in order to establish electrochemical exfoliation as a definite alternative for scalable graphene production, preferentially without the aid of post-sonication or other co-assisting routes. Such should be accomplished using cheap, environmental-friendly, and widely available chemicals, which still remains a challenge.

### 2.3. Graphene oxide and reduced graphene oxide production methods

Graphene Oxide (GO) is another important member of the graphene-graphite family, which is a single layer of graphite oxide discovered in mid-19th century.[162]

There are multiple routes to prepare GO, most commonly by oxidizing graphite, followed by an exfoliation process. As a result of these chemical methods, a stable colloidal suspension of GO is achieved. The first example of oxidizing graphite came when Brodie employed a combination of potassium chlorate (KClO$_3$) and nitric acid (HNO$_3$) in his investigations to study the chemistry of graphite.[162]

As a result of these studies, a new material was developed, dispersible in pure or regular water, which he named “graphite acid.” In the late 19th century, Staudenmaier improved Brodie’s process by adding H$_2$SO$_4$ to the combination and by adding KClO$_3$ throughout the reaction rather than in a single addition.[163] Nearly 60 years later, Hummers used a combination of potassium permanganate (KMnO$_4$), sodium nitrate (NaNO$_3$) and H$_2$SO$_4$ to oxidize graphite,[164] while Hofmann’s method involved oxidation of graphite with KClO$_3$ and HNO$_3$.[165] Tour’s group used a combination of KMnO$_4$, H$_2$SO$_4$ and phosphoric acid (H$_3$PO$_4$) as the oxidants.[166] As a result of oxidizing with different combinations of compounds and reaction conditions, the carbon to oxygen ratio would vary from 1.17 to 0.74 in Staudenmaier to Tour methods, respectively.[167]

The most common source of graphite used for these chemical methods is flake graphite. The localized defects in the π-structure may serve as starting sites of the oxidation process.[168] Depending on the oxidation process, different oxygen-containing functional groups, such as carboxyl (C＝O), hydroxyl (C－OH) and epoxide (C－O), form on the surface and around the edges of the graphene layers, making them hydrophilic and reducing the interlayer forces. Depending on the relative humidity within the stacked GO sheets, the interlayer spacing varies from 0.6 nm to 1.2 nm.[169] These interlayer spacings weaken the interactions between the GO sheets and facilitate the exfoliation process. The GO is then exfoliated in the aqueous media via mechanical stirring, rapid heating or sonification to separate the layers of GO.[170] Currently, the Hummers method and its modified versions[166,171,172] are the most commonly employed methods to synthesize GO.

The less common methods of preparing GO involve direct oxidation of graphene layers via atomic oxygen in ultrahigh vacuum conditions,[173] photochemical reaction in ultraviolet light and oxygen,[174] or oxidation in nitric acid under potentiostatic conditions.[175]

Due to its disrupted sp$^2$ bonding network, GO is an electrical insulator (with electrical conductivity <1 μS/m). This disruption is also reflected in the Raman spectra of GO, which present broadened characteristic peaks, low I$_{2D}$/I$_G$ intensity ratios and a prominent D peak (Figure 6(a)). In order to increase the electrical conductivity of these 2D nanomaterials, GO can be reduced by a strong chemical reductant[176] or thermal reduction,[177] which results in the formation of rGO (Figure 6(b)). As a result of the deoxygenation process, the π-network is partially restored and recovers most of the electrical conductivity. The thermal properties are also improved as a result of the reduction process.[178] However, due to the residual oxygen functional groups, rGO does not exhibit the same electrical properties as those of pure graphene. Although the reduction process can bring GO’s optical and electrical properties closer to those of pristine graphene, it is still not enough to heal the vacancy defects.[179] One should note, however, that certain application can benefit from the presence of oxygen-containing functional groups. In the context of water purification, a prominent example of a potential practical application of GO, the oxygen-
containing functional groups ensure sufficient interlayer spacing in GO membranes, allowing capillary permeation of water through two-dimensional channels formed between the layers while maintaining impermeability toward gases such as He. Such behavior has led to an increased interest in the use of GO membranes not only in water purification, but also in gas separation applications.

2.4. Other production methods

A method with potential to produce graphene with medium to high quality in moderate quantities relies on the atmospheric synthesis by microwave-sustained plasmas. This method was reported for the first time by Dato et al. In this approach, the use of a substrate or a catalyst is not necessary, as the nucleation and growth of the graphene flakes take place within the gas phase. Complicated vacuum fixtures are also avoided in this open-air graphene synthesis alternative. The principle of the synthesis consists in the decomposition of the carbon precursor organic compounds (ex. ethanol) in a dense microwave argon plasma and the subsequent radical rearrangement into sp² coordinated carbon, along with CO and H₂ gaseous by-products. The material thus obtained is a ‘fluffy’ dark powder constituted by graphene sheets that show good structural quality when characterized by Raman, XPS and electron microscopy techniques, matching or even exceeding the best grade quality of liquid-phase exfoliated graphene.

Although the yield of the process is pointed out as its major disadvantage, growth rates as high as 2 mg/min have been reported in a surface wave induced based reactor (surfatron) assisted with IR/UV radiation excitation applied at the “cold” outlet gas flow section. In fact, this does not scale as well as much higher throughput methods like liquid-phase exfoliation, but considering the higher structural quality of the atmospheric plasma graphene deposits, this technique may become an option.

Alternatively to the standard atmospheric microwave plasma torch set up in the seminal work by Dato et al., there are at least three types of reactors capable of depositing graphene, mainly differing in terms of the configuration through which the microwave plasma is sustained.

i. Surfatron: a surfatron-based setup consists in creating a surface wave induced microwave configuration that enables maintaining a stable plasma in atmospheric pressure conditions.

ii. TIAGO: a device constituted by a waveguide, whose central section is reduced in height, and a cylindrical hollow metallic rod placed perpendicularly, producing a jet conical flame plasma surrounded by the air around it.

iii. Microwave slot antenna: a simple set up that allows initiating and maintaining atmospheric plasmas in H₂/Ar mixtures.

Since this approach to produce graphene is relatively new, in the last years more groups have been devoting their workforce to the development of this technique. Therefore, its full potential is not yet disclosed, and it is expected that graphene produced by microwave atmospheric plasmas will be soon reported embodying different applications, ranging from nanocomposites and energy storage devices to water purification or drug delivery systems.
Laser processing is a powerful tool explored in different processes, since it is a rapid and efficient method to produce complex structures at different scales. Thus, nowadays, laser technology is being used to produce graphene-based materials, since there is a strong demand for cost-effective fabrication of graphene patterns with low energy consumption in a short fabrication time (Figure 7(a)). This novel one-step process was developed, in 2014, by Tour’s group, and allows to obtain porous graphene-based structures through the incidence of a laser beam on a polymeric sheet at atmospheric conditions \([188]\) (Figure 7(b)). The laser-induced graphene (LIG) technique enables to transform sp\(^3\) carbon present in the polymeric substrate into sp\(^2\) carbon \([188]\) (Figure 7(c)). Laser irradiation can cause the increase of local temperature to produce LIG due to local decomposition and rearrangement of the precursor’s structure \([189]\). Moreover, gas release during the reaction creates a 3D porous graphitic material. This photothermal effect was firstly produced by the incidence of an infrared CO\(_2\) laser beam on a commercial polymer polyimide sheet\([188]\) but was further pursued by other authors using different laser sources (UV laser\([190]\) visible laser)\([191]\) and carbon precursor materials (wood, food, cork, paper, among others)\([192,193]\).

Figure 7(b) shows that LIG has the appearance of a foam with a percolation network of porous multi-layer graphenes, which is associated with the localized high temperature and pressure produced by laser irradiation. The image puts in evidence the very high surface area of this material, comparable to that of the wet-chemistry derived 3D graphene\([192]\).

This direct laser writing process has emerged as a facile method enabling the synthesis of flexible, patternable and scalable graphene-based structures, and has many advantages, such as nontoxicity and controllability, envisioning large-scale catalyst free non-contact production and maskless patterning. In fact, LIG has become a multifunctional graphene foam, with many applications having been explored since it was discovered in 2014, such as: supercapacitors for energy storage devices\([188]\) electrochemical biosensors\([194,195]\) piezoresistive sensors\([190,196]\) water treatment...
devices, touch screens and antennas, among others.

In another example of graphene production by laser, graphene films were obtained by coating a Ni substrate with graphite nanoparticles, followed by laser irradiation of the coated surface. Carbon atoms from the graphite particles dissolve into Ni bulk under the action of laser-induced heating, and, subsequently, precipitate onto the surface, where the graphene film is formed. The obtained material presents Raman spectra characteristic of high crystalline quality graphene, with SLG accounting for 25% of the overall area of the film.

Novel and innovative ways of producing graphene are still arising in the literature. One outstanding example is that of Tour’s group, who used a technique similar to flash sintering to produce graphene from organic waste in a fast and inexpensive way, with yields of 1 g per batch.

3. Anti-corrosion coatings using graphene and graphene-based films

Corrosion, the gradual degradation of metals and alloys as a result of chemical and/or electrochemical reactions with the surrounding environment, has an undeniable impact in terms of ecological health and economic loss. Material deteriorates, appearance is significantly altered, mechanical properties are lost and eventually the affected metallic parts end up losing their functionality. Protecting such structures is, therefore, of utmost importance. Coating technology is among the plethora of approaches developed to provide protection to metallic surfaces and improve their quality. In addition to passive barrier properties and esthetic appearance, coating formulations can serve other functional purposes, such as an active self-healing action triggered by specific stimuli. When it comes to corrosion prevention, the key requirements of an ideal barrier coating system include an inherent resistance to degradation in hostile conditions, an effective resistance toward the permeation of corrosive agents and mechanical integrity over the desired life of the coated components. Conventional anti-corrosion coating technologies that involve, for instance, the use of hexavalent chromium connote a negative environmental impact given the carcinogenic risk of such compounds. Furthermore, traditional approaches reveal several limitations, such as susceptibility to damage by heat, limited chemical stability, potentially high cost and the possibility of altering the physical properties of the metals being protected. Hence, there is a need for a disruptive eco-friendly alternative in order to overcome the aforementioned roadblocks and achieve long-lasting corrosion mitigation.

Graphene has emerged as a promising candidate due to its remarkable set of properties arisen from its unique structure. Being the barrier effect one of the most relevant characteristics in a multilayer system, much attention has been devoted to such material by virtue of its impermeability to fluids and gases. Apart from preventing the transport of corrosive or oxidative species toward the underlying substrate without notably altering its thermal conductivity and optical properties, graphene-based films enable a reduction of the applied layer in terms of quantity and volume. Besides, because of its chemical inertness, particularly regarding oxidation, graphene is highly resistant to corrosion in conditions under which other substances would undergo chemical transformations. However, despite this exceptional combination of features, the application of graphene in corrosion science appeared inconceivable, at first, since it was exclusively obtained via mechanical exfoliation, a low throughput, non-scalable method that renders graphene flakes smaller than 0.1 mm. Nonetheless, the discovery that not even the smallest atoms of helium can penetrate the dense lattice structure of graphene, along with the first large-area syntheses of such material on copper substrates through CVD, significantly altered this scenario. The seminal research conducted by Chen et al., in turn, can be considered the major turning point in the application of graphene as an anti-corrosive coating. Chen and coworkers unequivocally demonstrated, for the first time, the ability of CVD-grown graphene layers to protect refined metals in hostile environments over relatively short periods. Even though the early studies on the utilization of graphene in the field of corrosion prevention emerged later than those in other engineering fields, the number of citations is increasing at an accelerating pace, reflecting the quickly growing interest of corrosion scientists and engineers in this multipurpose material. Research on graphene-based anti-corrosion technologies can be split into two generalized groups, pure graphene anti-corrosive coatings and composite graphene anti-corrosive coatings. The achievements regarding both types were reviewed by Tong et al., for example.

3.1. Preparation methods and corrosion resistance performance

The studies of Chen et al. in 2011, on the ability of graphene films to protect metals from reactive
environments, attracted considerable attention to applying both graphene and graphene oxide films as a protective layer due to their barrier effect. Several studies showed the benefit of applying graphene layers on the surface of metals, such as nickel alloys, copper alloys and steel. The protective graphene layer could be prepared directly on the metal surface via methods such as CVD, electrophoretic deposition (EPD) and laser fabrication methods, or could be transferred to the surface using a mechanical transfer technology. Among all the preparation techniques, CVD is the most commonly used method due to its simplicity, quality of the film and the ability to coat large areas. The distance between the graphene film and the metal substrate largely affects the bonding strength between the graphene and the metal, and the graphene films prepared on different metals show different structures. It has been reported that the strength between the carbon atoms and copper, nickel and cobalt is weaker compared to gold, silver, and palladium. Therefore, other techniques such as mechanical transfer have been reported to be applied to coat any metal materials with high-quality graphene films.

Anisur et al. prepared graphene films on the surface of nickel via a CVD method and investigated the role of hydrogen flow and the cooling rate during graphene growth on the barrier properties and defect density of the film. Electrochemical impedance spectroscopy (EIS) and potentiodynamic tests on the samples immersed in a 0.1 M NaCl solution showed that graphene coatings prepared on Ni in an environment without H₂ flow offer nearly one order of magnitude superior corrosion resistance than that of uncoated Ni. SEM revealed that the hydrogen flow during the graphene growth facilitated wrinkle formation, resulting in a similar corrosion resistance to the uncoated Ni. Ye et al. prepared graphene on Ni substrates, using a laser fabrication method at room temperature, showing the possibility of fabrication of large-area graphene films with superior anti-corrosion performance compared to the bare substrate.

In the study of Pu et al., a CVD method was used to coat stainless steel and nickel-plated stainless steel with graphene films. The SEM analysis showed 100% graphitization of the nickel-plated stainless steel, while for the stainless steel sample the SEM images showed a poor coverage of the graphene film on the substrate surface. The polarization curves also showed that the corrosion currents of the graphene coated nickel-plated stainless steel sample were one fifth of the bare nickel-plated stainless steel. In order to eliminate the complications of transferring grown graphene films to another substrate, different preparation techniques were applied in other studies. Ye et al. employed a laser alloying process to form a Ni/Fe layer on the surface of carbon steel, followed by the growth of graphene films via laser irradiation. Anti-corrosion performance, including the results of polarization and EIS in 3.5% NaCl (aq) solutions, showed a significant improvement in the corrosion resistance of carbon steel, which is even superior to stainless steel. In another study, Quezada-Renteria et al. applied the EPD technique to grow rGO films as an anti-corrosive coating on carbon steel, which showed a reduction of up to three times in the corrosion rate compared to the bared carbon steel.

Tiwari et al. prepared graphene films on a copper surface using the CVD method. In order to increase the corrosion protection ability of the graphene films, a multilayer graphene coating was prepared. In this study a significant corrosion resistance in a 0.1 M NaCl solution was observed and the corrosion resistance was sustained for longer durations. This study showed that multilayer graphene could be effective in blocking the pathways of corrosive species to the metal surface. Raza et al. studied...
et al. \cite{213} prepared GO films on copper substrates via an EPD method. The results showed control of coating thickness by varying the GO/water suspension concentrations and the voltage in the EPD process. EIS results also showed that the corrosion rate of GO film covered copper was six times lower than that of bare copper in 0.6 M NaCl solution. Hsieh et al. \cite{214} applied the atomic layer deposition (ALD) technique to grow graphene films on copper to eliminate the structural defects in CVD-grown graphene, which resulted in an inhibition efficiency of more than 99%.

The main mechanism which allows graphene and graphene-based films and coatings to significantly reduce the corrosion rate of copper, nickel, and steel in aggressive solutions is the barrier effect (Figure 8(a)). The contact between the graphene film and the metal substrate makes it difficult for reactive agents such as oxygen and water to reach the metal surface. Thus, in order to have a long-lasting corrosion resistance and a high-quality film with a complete barrier effect, the film needs to fully cover the metal surface. There should also be a good adhesion between the graphene film and the substrate, with high resistance to wear and friction.

Nonetheless, although there are several reports showing promising results on the corrosion resistance of graphene-coated metals, recent studies have shown that graphene-coated metals could corrode at even greater rates than bare metals. \cite{215} These considerable variations in results arise due to the lack of complete surface coverage, the presence of defects including wrinkles and cracks. \cite{216,217} These defects are also the main reason for poor corrosion resistance in long-lasting applications (Figure 8(b)). Therefore, many researchers proposed that it was not practical to consider graphene films as a corrosion protective layer for long-term applications due to the difficulties in avoiding defects in the preparation process. \cite{218} There are other studies that tried to improve the protective ability of the graphene films by repairing the defects via an atomic layer precipitation technology. \cite{214} Some researchers have also shown that due to the high electrical conductivity of graphene it promotes electrochemical corrosion of metals. \cite{215} It has been shown that in the damaged regions, graphene and metal form corrosion microcells. In such cases, metals such as copper, nickel, aluminum, magnesium, iron, or steel, act as the anode, promoting galvanic corrosion. In addition, the as-grown graphene layers are characterized by low adherence to the subjacent metal as a consequence of weak interaction. \cite{200,202} Although some surface treatments and graphene functionalization routes can improve the quality of coating adhesion, such measures may not be considered good enough for long term durability.

4. Anti-corrosion composites using graphene and graphene-based materials

When performance is the absolute priority, the development of graphene-based nanocomposites is a promising route to trigger the commercialization of graphene-based coatings. The first studies of graphene-based composite coatings date from around 2012, \cite{219} and since then such films have been applied to a broad range of substrates, such as aluminum, steel, brass and copper. The preparation methodology and coating procedure of such materials can be established based on the traditional coating production processes with controllable workability. The most relevant coating methodologies include lamination, \cite{220} physical deposition or dispersion and subsequent curing, \cite{221,222} self-assembly, layer-by-layer (LbL) deposition and the sol-gel method. \cite{200}

4.1. Dispersion in polymeric matrices

In coating applications, graphene and its derivatives have stood out as efficient fillers in polymer matrices to form hybrid films. Polymer nanocomposites provide superior features at a relatively low cost and the processing procedure is less complex when compared to multilayer coating systems. Moreover, polymer/graphene hybrids address other problematic issues, such as galvanic corrosion. \cite{14,204} By utilizing graphene or its derivatives as nanometric fillers in the polymeric matrix, it is possible to lengthen the diffusion pathway of the corrosion medium in the coatings by taking advantage of its chemical inertness and barrier properties. \cite{13} Graphene has heretofore been combined with a wide variety of polymeric materials, such as epoxy (EP), \cite{223–253} polyurethane (PU), \cite{254–259} polyaniline (PANI), \cite{227,233,260–262} alkyd, \cite{263} polystyrene (PS), \cite{264} PMMA, \cite{265,266} polyvinyl butyral (PVB) and \cite{267,268} polydimethylsiloxane (PDMS). \cite{269}

The lack of functional groups on the surface of pure graphene, its high surface area (and, consequently, high surface energy) and the Van der Waals interactions pose major challenges for the dispersion of graphene as filler particles in coating matrices, resulting in poor compatibility. Such factors contribute to the agglomeration of graphene sheets in aqueous media and organic matter, which in turn compromises the overall coating performance and
undermines its corrosion resistance. The inclusion of graphene fillers in polymer matrices aims at hampering the diffusion of corrosive species through the coating, thus enhancing its barrier effect. However, their agglomeration has quite the opposite effect by providing direct diffusion pathways through the matrix. Thus, improving the dispersibility of graphene must be an utmost priority when developing graphene/polymer nanocomposites. And even though some researchers have simply incorporated unmodified graphene in polymeric matrices by way of physical means\cite{224} or in situ polymerization,\cite{270} the results were not considered ideal and such methods have proven suitable only for GO and rGO, due to the presence of epoxy and carboxyl groups on its surface.\cite{271} To achieve improved dispersibility, two main (often complementary) approaches can be identified (Figure 9): mechanical dispersion and dispersion through chemical modification. The latter aims to affect the interaction between graphene or graphene-based materials and the polymeric matrix. These interactions can be either covalent or non-covalent. Moreover, decoration with nanoparticles is also common, often taking advantage of the modified interactions. In the context of mechanical dispersion, Chang et al.,\cite{266} for example, prepared thermally reduced graphene oxides with varying carboxylic-group content, and found that higher amounts of carboxyl groups appeared to enhance the compatibility of these rGOs with the acrylic resin, thus improving their dispersion in the PMMA matrix. On the other hand, the simple mechanical dispersion of unmodified single-layered GO in epoxy resin required vigorous mechanical mixing and intense sonication processes in order to effectively deagglomerate the GO sheets.\cite{235}

The structural and chemical modification of graphene and graphene-based materials is among the most commonly used approaches to stabilize suspensions of such material in complex environments without agglomeration taking place. Surface functionalization can play an important role in achieving good interfacial bonding between graphene sheets and the surrounding matrices. Graphane, the fully hydrogenated analogue of graphene, was one of the first chemically modified graphene derivatives.\cite{272,273} Aside from hydrogenation, the controlled oxidation of graphite flakes to generate hydrophilic GO and its selective reduction into hydrophobic rGO are also among the simplest routes to introduce surface functional groups onto graphene layers.\cite{200} The chemical modification of graphene usually involves the use of coupling agents, such as silanes\cite{239,240,244,246,252,256,257} and other common dispersants.\cite{219,225–231,241,242,253,255,258,260,264,269} Generally, such additives contain both hydrophilic and organophilic functional groups, which can be combined with inorganic filler particles and organic coating matrices respectively. As a result, these compounds are able to improve the quality of the matrix/filler interactions and, consequently, enhance the overall performance of graphene/polymer composites.\cite{12} The anti-corrosion properties of protective hybrid systems benefit from the incorporation of functionalized graphene and graphene-based nanostructures, given their ability to retard

Figure 9. Different approaches for the dispersion of graphene and graphene-related materials in polymeric matrices.
corrosion processes by restraining the diffusion of corrosive species. Table 5 compiles some of the most relevant research studies on the surface modification of graphene and its derivatives, either by coupling agents or other chemical reagents, over the past few years.

The synthesis of graphene-nanoparticle composites (G-NPs) is another route to improve the dispersibility of graphene-based materials in polymeric matrices and ensure the integrity of their structure. Several nanoparticles have been utilized for such purpose, as summarized in Table 6. In addition to their availability, mechanical properties and photo and thermal stability—among other appealing characteristics—these nanostructures increase the interlayer spacing between GO sheets, preventing their aggregation,[248,250] and may be employed to impart new functionalities to graphene-based nanocomposites, as will be covered in the following section. By taking advantage of the synergistic effects arising from the combination of both materials, it is possible to develop high-quality coatings with excellent properties. The three main strategies to fabricate G-NPs nanohybrids are the following: pre-graphenization, post-graphenization, and syn-graphenization[204] Pre-graphenization involves mixing the pre-synthesized graphene or its derivatives with the desired nanoparticles. In post-graphenization, the composite (graphene precursor and nanoparticles) is pre-prepared and the precursor is then converted to chemically reduced graphene. In syn-graphenization, the two components are synthesized simultaneously in a one-pot system.

### Table 5. Chemical modification of graphene and graphene-based materials dispersed in polymer matrices.

| Filler                          | Dispersing agent                                           | Polymer matrix | Substrate | Ref. Year |
|---------------------------------|------------------------------------------------------------|----------------|-----------|-----------|
| Graphene oxide                  | (3-Aminopropyl) triethoxysilane (APTES)                   | Epoxy resin    | Steel     | [246] 2017 |
| Graphene oxide                  | γ-(3-epoxypropoxy) propytrimethoxysilane (KHS60)          | Epoxy resin    | Steel     | [240] 2018 |
| Graphene oxide                  | APTES                                                      | Epoxy resin    | Steel     | [240] 2018 |
| Graphene oxide                  | Poly(2-butanilene) (P2BA)                                 | Epoxy resin    | Steel     | [240] 2018 |
| Graphene oxide                  | Poly(3-phenylenediamine) (PPDA)                            | Epoxy resin    | Steel     | [240] 2018 |
| Reduced graphene oxide          | Titanate coupling agent (TGO)                             | Epoxy resin    | Steel     | [240] 2018 |
| Reduced graphene oxide          | Perigraniline                                              | Epoxy resin    | Steel     | [240] 2018 |
| Reduced graphene oxide          | PDA                                                        | Epoxy resin    | Steel     | [240] 2018 |
| Reduced graphene oxide          | Polyisocyanate (PI)                                        | Epoxy resin    | Brass     | [240] 2018 |

#### 4.2. Functional incorporation in anti-corrosion composites

Nearly all the above-mentioned works have reached a unanimous conclusion: the superior performance of the obtained graphene-based composite coatings is unquestionably associated with the barrier effect provided by graphene or its derivatives. As previously noted, the unique impervious 2D structure of graphene imparts it with exceptional barrier properties toward all fluids and gases. The extremely small pores of its hexagonal lattice ensure minimal permeability even for the smallest atoms, i.e., He. Furthermore, the dense and delocalized electron cloud of the Π-conjugated carbon network in graphene blocks the gap within its close packed aromatic rings and poses a repelling field to reactive atoms or molecules, thus providing a physical separation between the refined metal surface and environmental reactants. Hence, graphene can be understood as an impenetrable nanometric shield which can effectively isolate subjacent metal surfaces from invasive agents without altering their thermal conductivity and optical properties. By prolonging the infiltration path of corrosive media through the coating, graphene can greatly enhance its corrosion resistance and extend its service life.[12,200]

Let us take as an example the investigation carried out by Yu and coworkers[275] who exploited the impermeable nature of graphene to create a high-performance polymer/graphene sandwich-like anti-corrosive system. Given the paucity of reports on CVD-
grown graphene-based coatings able to provide long-term protection of metals, the authors aimed at developing a hybrid structure consisting of alternating SLG and PVB films on a commercial aerospace aluminum alloy (AA2024-T3). After comparing the effect of immersion in simulated seawater on uncoated AA2024, pure PVB-coated AA2024 and graphene/PVB-coated AA2024, the best electrochemical performance was assigned to the hybrid coating made up of two graphene layers sandwiched by three polymeric films (PVB/SLG/PVB/SLG/PVB). Such system entirely suppressed the deterioration of the aluminum substrate, the surface of which remained unaltered for up to 120 days of immersion in 3.5 wt% NaCl solution. In contrast, the graphene-free variants and the PVB/SLG/PVB system were able to provide corrosion protection only in the short-term (30 days), which clearly emphasized the dramatic improvement of the barrier properties of PVB coatings by inclusion of atomically thin graphene layers. Thus, Yu and his team managed to combine the adhesive and insulating properties of polymer films with the imperviousness of CVD-grown graphene sheets, while avoiding the latter’s intrinsic drawbacks, such as poor adhesion and galvanic corrosion.

Fabricating impeccable, defect-free coatings is considered virtually impossible to achieve, since the formation of cracks is often unavoidable during their preparation. Such fissures pose menacing entry points for corrosive media to invade the metal, and once the aggressive species penetrate the metal, its corrosion resistance and structural integrity are jeopardized. If crack extension occurs, more vulnerable areas will be exposed, and severe localized corrosion may occur. In addition to its ability to hinder the penetration of corrosive agents through coatings due to its impermeable nature, graphene can effectively increase their fracture toughness, preventing crack propagation and the consequential ingress of threatening substances. The mechanisms underpinning the effectiveness of graphene in hampering the expansion of existing cracks mainly involve the consumption of the crack growth energy. When the crack reaches the graphene/matrix interface, the crack tip stress will cause interfacial debonding. Microcracks will be formed either around or in the graphene itself and will consume a significant amount of the original fracture energy. If external forces continue to intensify, crack deflection may occur through the deteriorated graphene. As the crack deviates from the original expansion direction and the propagation path is lengthened, the crack propagation resistance is increased. Moreover, the addition of graphene to crystalline and semicrystalline polymers can also affect their crystallization behavior, as shown by Zhang et al. They prepared a transparent poly(lactic acid) (PLA) nanocomposite film reinforced with fully exfoliated octadecylamine-functionalized graphene (ODAG) as an effective nanofiller, by solution blending and casting method. Given the good compatibility between the two materials, ODAG could be uniformly dispersed in the PLA matrix and improved interfacial adhesion was achieved. In comparison with neat PLA, the obtained PLA/ODAG nanohybrid exhibited a 34% increase in tensile strength, 44% increase in Young’s modulus, and 300% improvement in tensile toughness with the addition of only 0.4 wt% ODAG. The simultaneous improvement in mechanical strength and toughness could be attributed not only to the homogeneous distribution of ODA-modified graphene across the PLA matrix, but also to the enhanced crystallization behavior of the thermoplastic polymer. The incorporation of ODAG facilitated the cold crystallization process of PLA and notably improved its crystallinity.

The effectiveness of corrosive-protective schemes relies heavily on the adhesion quality between the substrate and the deposited films and, therefore, it is imperative to ensure proper adhesion strength through the chemical bonds established between them. Apart from providing an effective barrier against the diffusion of water, oxygen and ions, functionalized graphene oxide can concurrently improve the adhesion strength between the organic coating

### Table 6. Nanoparticle-decorated graphene oxide dispersed in polymer matrices.

| Nps/graphene-based nanofiller | Polymer matrix | Substrate | Ref     | Year |
|-------------------------------|----------------|-----------|---------|------|
| SiO2/graphene oxide           | Epoxy resin    | Steel     | [234]   | 2016 |
| SiO2/graphene oxide           | Epoxy resin    | Steel     | [237]   | 2016 |
| SiO2/graphene oxide           | Epoxy resin    | Steel     | [247]   | 2017 |
| Al2O3/graphene oxide          | Epoxy resin    | Steel     | [274]   | 2015 |
| Fe3O4/graphene oxide          | Epoxy resin    | Steel     | [251]   | 2018 |
| CaCO3/graphene oxide          | Epoxy resin    | Steel     | [248]   | 2016 |
| TiO2/graphene oxide           | Epoxy resin    | Steel     | [249]   | 2015 |
| TiO2/graphene oxide           | Epoxy resin    | Aluminum  | [250]   | 2018 |
| TiO2/graphene oxide           | Epoxy resin    | Steel     | [238]   | 2016 |
| -(2,3-epoxypropoxypropytrimethoxysilane (GPTMS)-modified TiO2/graphene oxide | Epoxy resin    | Steel     | [238]   | 2016 |
and the subjacent metal and, consequently, increase the degree of corrosion protection. Parhizkar et al. [279] used APTES to chemically alter the surface of graphene oxide and improve the adhesion and corrosion protection properties of an epoxy coating. Both were effectively enhanced by the deposition of a functionalized GO (fGO) film on the surface of steel prior to the application of the subsequent organic coating. Moreover, the cathodic delamination rate of the latter was decreased. The same group also studied the effect of 3-(Triethoxysilyl)propyl isocyanate (TEPI)- and APTES-functionalized GO nanofillers (IGO and AGO, respectively) embedded in sol-gel based silane films on the adhesion strength, corrosion protection and cathodic disbanding of the outermost epoxy coating applied on the silane-treated steel substrate. [280] While the incorporation of non-functionalized GO displayed no significant effect, the well-dispersed IGO and AGO inclusions significantly improved the adhesion quality and corrosion resistance of the topmost layer, due to their excellent barrier properties, good compatibility with the silane matrix and covalent bonds established with the epoxy coating.

In an example of the use of graphene-based materials to provide enhanced functionality to the anti-corrosion coating, Fan et al. [281] fabricated GO/Al2O3 hybrids without the assistance of surfactants via two titration methods, wherein GO and Al2O3 colloids were utilized as tiritants for composites with low (1.23 to 1.92 wt.%) and high GO content (2.88 to 11.5 wt.%), respectively. After sintering by spark plasma sintering, FLG/Al2O3 nanocomposites were obtained and GO was simultaneously reduced to rGO. A percolation threshold as low as 0.38 vol.% was achieved with a graphene content of 2.35 vol.%. Additionally, the electrical conductivity surpassed 103 S m−1. The most captivating finding, however, was related to the charge carrier type, which changed from p- to n-type with the increase in graphene content. Such conversion was attributed to the doping effect induced by the alumina matrix, which is thickness-dependent with respect to rGO.

The synergistic advantage of combining graphene derivatives and metal oxides was also explored by Song [282] and Jiao. [283] The former prepared Co3O4/rGO nanosheet composites through a facile hydrothermal route, by exploiting the electrostatic and coordination interaction between the individual negatively charged GO nanosheets and Co2+ ions. The as-synthesized sheet-on-sheet nano hybrids exhibited good cyclic performance and coulomb efficiency with a specific capacitance over 402 F g−1 at a current density of 0.5-2.0 A g−1, mainly as a result of the high porosity of the composite structure. In the latter case, a highly-ordered Fe3O4-graphite nanosheets/epoxy composite was obtained by tethering the magnetic nanoparticles onto the surface of graphite nanoplatelets, by wet-chemical precipitation method, and aligning the latter in the epoxy resin under a low magnetic field. The Fe3O4-modified graphite nanoplatelets were embedded in the EP matrix in such a highly oriented manner that its permeability dropped considerably and the gas barrier properties increased one order of magnitude compared to the blank EP and more than 65% compared to the randomly arranged graphite nanoplatelets/EP specimens.

### 4.3. Composites for active corrosion protection

The addition of chemically treated graphene to organic coatings can go beyond the mere improvement in adhesion and passive barrier properties. By intentionally altering graphene’s structure with, for instance, corrosion inhibitors, highly efficient protective coatings can be developed. In order to overcome the presence of defects in epoxy coatings, metronidazole (MET)-modified graphene oxide nanocomposites (GME) were synthesized via a two-step method that involved grafting maleic anhydride (MA) on GO surface (GM), [232] followed by the covalent attachment of MET by the esterification mechanism. GME materials were shown to enhance epoxy films’ corrosion protection performance at a low weight fraction of 0.2 wt%. Given that MET could be released from the GME hybrids and form an adsorption film, the scratched specimens were able to self-repair the damaged areas as a means to prolong the lifetime of the subjacent metallic structure.

The adsorptive behavior of *Urtica Dioica* (U.D) leaf extract on the graphene oxide nanosheets was examined by Nikpour et al., [284] who investigated its influence on the corrosion inhibiting properties of an epoxy film. The achieved results revealed that the addition of GO-U.D and GO-U.D + Zn2+ to the epoxy layer with an artificial defect improved the coating resistance to corrosion and decreased its delamination. It was also observed that the inhibition effect of GO-U.D sheets became more prominent in the presence of zinc cations. The chelation between the U.D molecules and Zn species led to the deposition of a protective film on the active anodic and cathodic sites of steel, and both electrochemical reactions were suppressed. The corrosion-inhibiting effect of green corrosion inhibitors (GIs) obtained from U.D leaves was...
also explored by Ramezanzadeh and colleagues,\cite{261} who utilized three-dimensional GO nanosheets as a platform to fabricate a protective system comprised of PANI nanofibers and the sustainable inhibiting agents. The GO-PANI-GI nanostructures were synthesized through a layer-by-layer assembly technique, and their anti-corrosive properties were analyzed in saline solutions on mild steel panels. The investigation confirmed that positively charged PANI strongly anchored to GO surface via physisorption mechanism, and GIs stabilized near protonated PANI through intermolecular H-bond interactions. This multilayered graphene-based nanocomposite demonstrated effective corrosion inhibition performance due to the adsorption of GI molecules on the anodic and cathodic sites and the passivation of steel surface in the presence of PANI fibers.

The corrosion resistance of aminoazobenzene- and diaminobenzene-functionalized graphene oxide composites (AAB-GO and DAB-GO, respectively) has also been examined, and it has been shown that both composites act as efficient corrosion-inhibiting agents by adsorbing on the mild steel surface and that their performance boosts with concentration.\cite{285} It was also found that AAB-GO and DAB-GO act as mixed-type inhibitors, behaving predominantly in a cathodic manner. More recently, Sun and his team developed a facile CVD method to modify graphene nanosheets with molecular-sized polydimethylsiloxane.\cite{269} The synthesized nanostructures were used to reinforce epoxy films, which displayed a coating resistance one order of magnitude higher than that of neat epoxy. PDMS-modified graphene nanosheets were able to enhance its performance at a loading as high as 5 wt\%.

Qui et al.\cite{243} synthesized FLG functionalized by poly (2-aminothiazole) (PAT) with free basal plane defects in organic solvent based on the $\pi-\pi$ interaction between them. In contrast to neat epoxy and PAT-based films, the coatings containing PAT/graphene hybrids displayed excellent corrosion performance and lower wear rate by dint of multifunctional graphene and passive effect of PAT, which acted both as a corrosion inhibitor and a lubricating agent.

Another strategy combining the passive barrier property of graphene and active inhibition functionality has been recently developed by Hou,\cite{245} who proposed graphene oxide nanocontainers as a promising alternative to conventional capsule-based technologies, whose barrier effect is limited due to their spherical shape. The layer-by-layer self-assembled GO-based nanostructures, encapsulated with polymeric ionic liquid (PIL), an environmentally friendly corrosion inhibitor, endowed the resulting container-impregnated epoxy composites with effective active inhibition function and superior barrier properties. While the embedded PIL could exert its corrosion-inhibiting effect at defected areas, the impermeable GO nanosheets provided passive protection by preventing the penetration of corrosive species through the coating.

Other functional integration approaches involve the modification of graphene to produce compounds able to react with the underlying metals to form chemical conversion layers. The investigations carried out by Taheri\cite{262} and Ramezanzadeh\cite{233} are cases in point. The authors modified negatively charged graphene oxide sheets with PANI nanofibers with zinc cations (GO-PANI-Zn) and cerium oxide (GO-PANI-CeO$_2$), respectively, through a layer-by-layer assembly method. In the former case, zinc ions demonstrated electrostatic and cation-$\pi$ interactions with emeraldine base and salt forms of PANI. GO-PANI-Zn hybrids displayed improved thermal stability and active inhibition properties through adsorption of polyaniline as anodic and zinc cations as cathodic inhibitors, and consequent formation of a dense metal oxide passive layer. In addition, the results confirmed the high cation exchange capability of the synthesized nanostructures. Analogously, the excellent cation exchange ability of multifunctional GO-PANI-CeO$_2$ nanocomposites led to the creation of an adsorptive protective film on the metal surface and enhancement of barrier properties and corrosion inhibition performance of epoxy coatings.

The biomedical applications of graphene-based materials have also attracted ever-increasing interest, particularly the corrosion protection of metallic implants, such as Ti and its alloys, which can be deteriorated through wear and exposure to corrosive environments.\cite{286} In this context, graphene-reinforced hydroxyapatite (HAp) biocomposites appear as a potential solution.\cite{287,288} The incorporation of such materials as additives in HAp hybrids has been obtained through different approaches, namely biomimetic mineralization,\cite{289,290} in situ synthesis,\cite{291,292} electrospinning,\cite{293} spark plasma sintering\cite{294} and radio-frequency chemical vapor deposition.\cite{295} Recent reports have investigated the potential of electrodeposited graphene-based HAp composites as improved protective schemes for Ti substrates.\cite{287,288,296} These novel coatings exhibit not only enhanced mechanical performance and corrosion stability when compared to pure HAp films, but also superior bioactivity.
Taking into consideration the fact that highly conductive graphene can aggravate localized corrosion at exposed metal-coating interfaces, several efforts have been made to suppress its undesired corrosion-promotion effect.\cite{226,259,260,268} Contrarily, other authors have taken advantage of such feature to mitigate corrosion processes on metal surfaces by incorporating graphene-based conductive additives in organic coatings. Since the 1930s, zinc-rich coatings (ZRCs) have been extensively utilized for a multiplicity of industrial purposes under a wide range of hostile conditions. Such systems are able to protect metal substrates in the long term, even when slight mechanical damage occurs. Whilst the polymer matrix serves as a barrier toward corrosive agents, zinc particles provide cathodic protection by acting as sacrificial anodes. Given the electron pathway established between those particles and the underlying metal, a galvanic cell is formed, wherein the substrate behaves as cathode. Hence, the electron migration is suppressed, and corrosion processes are inhibited or retarded. Epoxy binders stand out amongst the most commonly used binders for ZRCs by virtue of their remarkable mechanical and chemical resistance and high adhesive strength. Since such film-forming elements possess low electrical conductivity, a considerable load of zinc particles is required to ensure electric continuity and the occurrence of preferential galvanic corrosion of zinc. However, an overload of zinc powder may adversely affect the coating flexibility and adhesion to the substrate, and eventually compromise its performance. For this reason, the protective mechanisms of ZRCs strongly rely on the value of pigment volume concentration. On the other hand, when this value is lower than the critical pigment volume concentration, the barrier effect prevails over cathodic protection. In this regard, efforts have been made to enhance the electrical conductivity of ZRCs without it being necessary to increase the amount of zinc particles. An attractive option involves the utilization of carbon-based conductive fillers. The uniform distribution of graphene additives can greatly enhance both

\[\text{Figure 10. Schematic illustration of the enhancement of the sacrificial anode protection in zinc-rich coatings by graphene through increased percolation and the barrier effect.}\]
percolation and barrier properties of ZRCs, when added in the proper amount. Several authors have focused on the impact of graphene and its derivatives on the corrosion resistance of ZRCs applied on steel\cite{297–302} and iron\cite{303} surfaces. The incorporation of graphene nanosheets into zinc-rich coatings can significantly enhance the role of the sacrificial anode (Figure 10). On one hand, the presence of graphene provides electrical continuity between zinc particles and the metallic substrate, thereby improving the percolation process with a lower amount of zinc pigments as compared with traditional ZRCs. On the other hand, graphene acts as a cathode and prevents the electrolyte from further permeating through the coating, seeking a cathodic counterpart for zinc powder particles. The resulting insoluble, non-conductive corrosion products effectively fill the permeation channel in the short term and, consequently, alleviate the diffusion of aggressive agents. Thus, the protective mechanism of zinc-rich epoxy coatings lies in the combination of barrier properties and cathodic protection.\cite{12}

5. Critical analysis

In the previous sections different methods for production of graphene and graphene-based materials were presented and their application in the field of corrosion protection were described. In this final section, additional considerations are presented considering two main points: the availability and quality of graphene-based materials for the target application and what can be done further in terms of characterization to obtain additional information on the developed systems.

5.1. Production method vs. application

The development of practical graphene-based anti-corrosion solutions depends heavily on the ability to produce this material in an economically viable fashion, while possessing the required properties. For anti-corrosion coatings based on graphene films, CVD presents itself as the best option. Building on its ability to provide large area single- and few-layer graphene, the recent progress in regard to CVD has led to arbitrarily large films with a very low surface density of grain boundaries. This allows to address one of the main shortcomings of graphene film anti-corrosion coatings, which are the structural defects which make the underlaying metal surface accessible to corrosive species from the surrounding environment. However, CVD growth is mostly limited to transition metal substrates, most often in the form of a foil or a thin sheet. This limits its direct growth on other metals. A number of approaches to transfer graphene from the original transition metal to the desired substrate exist, but these tend to be rather laborious and incompatible with target substrates possessing a non-planar geometry, not to mention the associated concern of introducing cracks, tears and wrinkles in the graphene during the process. These would defeat the purpose of graphene as an impermeable barrier to the corrosive species.

As composite coatings with graphene are concerned, the most viable options seem to be the different types of high-throughput exfoliation. High shear stress exfoliation, for example, benefits from dispensing the use of intercalating agents, while requiring considerably less time to achieve the same amount of FLG as through sonication, also avoiding the introduction of basal plane defects often seen after ball milling. Electrochemical exfoliation has also seen a lot of progress and is perhaps one of the more easily scalable exfoliation techniques. However, all these approaches share some common shortcomings. The thickness (number of graphene layers) and the lateral dimensions of the obtained flakes are often hard to control. Besides, the use of appropriate solvents is usually required, either during or after the process, to aid in the exfoliation and to avoid re-aggregation, with the best suited ones being environmentally hazardous. Environmentally friendly alternatives are available, but further research is needed. One should also note that these exfoliation approaches are also adequate for anti-corrosion solutions based on non-composite coatings formed from graphene flake suspensions.

Composite coatings containing graphene-based materials such as GO or rGO, on the other hand, continue to rely on the Hummers method and its variations as the most commonly employed production approach. This type of material is particularly relevant when functionalization is required, as happens in the cases where the dispersibility of the graphene-based material in a polymeric matrix needs to be improved. There, GOs and rGOs benefit from the presence of oxygen containing groups on their surface. And even then, vigorous mechanical dispersion strategies need to be implemented.

This underlines one of the main issues concerning composite anti-corrosion coatings with graphene and graphene-based materials: the need for functionalization, particularly with the aim of improving its dispersibility in polymeric matrices. In this context, both covalent and non-covalent modification approaches have delivered some promising results, as have the
different strategies revolving around nanoparticle immobilization. It is worth pointing out, however, that most of these approaches use graphene oxide as the starting point, leaving plenty of room for further work in the development of reliable functionalization techniques of pristine graphene flakes.

One should also be aware that, independently of the type of the anti-corrosion coating, it is important that any future work in this field matches the employed graphene or graphene-based material to the role and characteristics that are expected from it, as there is a number of works which ascribe the properties of pristine graphene to other graphene-based materials, such as GO. Both categories of materials have a place in the field of anti-corrosion technology, but the requirements for each one of them can vary greatly. While certain applications may require, for example, an easily attainable functionalization of the graphene-based material, and, thus, the presence of oxygen containing groups can be beneficial, others can demand sufficient electrical conductivity, as in the case of zinc-rich coatings, where pristine graphene flakes may result in better anti-corrosion performance.

5.2. Challenges and guidelines for characterization of anti-corrosion coatings using graphene

As described in sections 3 and 4, the contact between the graphene film and the metal substrate is of utmost importance for a long-lasting corrosion resistance. To achieve a maximal barrier effect under relevant service life conditions, a high-quality graphene film should fully cover the metal surface, despite their rough surfaces or curved shapes, while making it difficult for reactive agents such as oxygen and water to reach the metal surface. Therefore, a good adhesion between the graphene or graphene-based coating and the substrate, with high resistivity of this coating to wear and friction, is highly desirable.

The industry employs standardized methods to evaluate adhesion between coatings and substrates. As an example, the cross-cut tape test ASTM D3359 is widely used for ranking adhesion properties of coatings. Some works available in the literature have used this method to prove that organic coatings do not lose adhesion to the substrate upon modification by graphene-based materials.\textsuperscript{304,305} Furthermore, evaluation of adhesion in laboratories is often carried out using pull-off tests following the ASTM D4541 standard. Parhizkar et al.\textsuperscript{279,280} applied a pull-off method for testing adhesion properties of epoxy coatings deposited on the samples coated with a functionalized GO film, with sol-gel coatings containing GO additives or functionalized GO having been used for adhesion promotion between the metal and the epoxy coating. The results demonstrated an improvement in adhesion strength of the coating systems containing GO modified with 3-(triethoxysilyl) propyl isocyanate and APTES silanes. These systems demonstrated cohesion failure at the modified GO-coating/epoxy interface, unlike the simple unmodified GO/epoxy system. This effect was attributed to NH\textsubscript{2} and NCO functional groups present in the modified GO that can create strong covalent bonding with the top epoxy coating.

Overall, in the circumstances where adhesion strength is in demand or may be compromised by the additives used in coating systems, an appropriate testing should be performed. Adhesion tests can reveal at which interface the adhesive failure occurs, thus indicating a potentially weak spot in the coating system. Proper measures can be developed afterwards in order to curb the weakness.

Another aspect lacking extensive coverage in the literature so far is the detailed analysis of diffusion in coatings modified with graphene and graphene-derived materials. Most of the works on coatings modified with graphene and its derivatives claim that the aforementioned additives offer protection against diffusion of corrosive species through the coating. In fact, methods such as EIS have been used for characterization of coated metals for decades. Impedance spectra taken from samples under immersion in different electrolytes contain valuable information concerning the properties of metal/polymer and metal/solution interfaces, as well as other properties of the coatings. Using an appropriate equivalent circuit model allows to extract different parameters and assess evolution of parameters in time. The capacitance of a coating can be easily extracted from impedance spectra\textsuperscript{306} and it depends on surface area, thickness and the dielectric constant of the coating. Moreover, the swelling of the coating can have an impact on the capacitance of the coating during immersion, as the latter will depend on the amount of water absorbed, due to it having a much larger dielectric constant than an organic coating.\textsuperscript{307} A quantitative representation of water uptake by a coating was provided by the Brasher and Kingsbury equation,\textsuperscript{308} which correlates water uptake to the coating’s capacitance. Considering diffusion to be the main driving force for the water absorption, a particular solution of Fick’s second law allows to obtain the diffusion coefficient.\textsuperscript{309} Upon combining the Brasher and Kingsbury equation with the solution of Fick’s second law a new equation is derived. It displays a linear dependence between the capacitance and the square root
of time during a short time of immersion when water uptake follows the fickian diffusion. Diffusion coefficient of water is obtained from the slope of that dependence. Such an approach may be used to characterize organic coating systems containing graphene and its derivatives. However, this analysis is very demanding and requires careful impedance measurements and selection of correct equivalent circuit models for interpretation of EIS spectra.

In conclusion, like with any other type of material under scrutiny for industrial implementation, it is evident that graphene-like materials are not the only existing solution to address all the technological challenges in the field of protective coatings, especially in cases where more cost-effective additives exist and give similar performances.

However, it is undeniable that the studies available in the literature so far point to unique properties rendered by graphene and its derivatives which, together with the implementation of industrial processes to produce sufficient amounts of these nanostructured materials, will contribute to the generation of new solutions for corrosion protection in the forthcoming years.

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Disclosure statement

The authors declare no conflict of interest.

ORCID

Bohdan Kulyk http://orcid.org/0000-0002-0497-2216

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